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# **DELIRIUM: diagnosis, prevention and management**

**Draft for consultation, November 2009**

Produced by the National Clinical Guideline Centre

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**FOREWORD (To be written before publication)**

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## 11 Guideline Review Panel

12 The Guideline Review Panel is an independent panel that oversees the development of the  
13 guideline and takes responsibility for monitoring its quality. The members of the Guideline  
14 Review Panel will be added when available.

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1 **Stakeholder Involvement**

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3 **To be added after consultation.**

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## 1 Abbreviations

<b>ACS</b>	Acute confusional state (delirium)
<b>ADL</b>	Activities of Daily Living
<b>AGU</b>	Acute Geriatric Unit
<b>AMT</b>	Abbreviated Mental Test
<b>ANOVA</b>	Analysis of variance
<b>APACHE</b>	Acute Physiology and Chronic Health Evaluation (severity of illness classification system)
<b>ARDS</b>	Acute respiratory distress syndrome
<b>ASA</b>	American Society of Anesthesiologists (score for illness severity)
<b>ASE</b>	Attention Screening Examination
<b>BEHAVE-AD</b>	Behavioural Pathology in Alzheimer's Disease Rating
<b>BNF</b>	British National Formulary
<b>CABG</b>	Coronary artery bypass grafting
<b>CAM</b>	Confusion Assessment Method
<b>CAM-ICU</b>	Confusion Assessment Method for the ICU
<b>CCA</b>	Cost-consequences analysis
<b>CD</b>	Compact disc
<b>CDR</b>	Clinical Dementia Rating scale
<b>CDT</b>	Clock Drawing Test
<b>CEA</b>	Cost-effectiveness analysis
<b>c.f.</b>	Confer (refer to)
<b>CGBRS</b>	Crichton Geriatric Behavioural Rating Scale
<b>CGI</b>	Clinical global impression scale
<b>CGI-GI</b>	Clinical global impression scale: global improvement item
<b>CGI-SI</b>	Clinical global impression scale: severity of illness item
<b>CHF</b>	Chronic heart failure
<b>CI / 95% CI</b>	Confidence interval / 95% confidence interval
<b>CIPFA</b>	Chartered Institute of Public Finance and Accountancy
<b>CNS</b>	Central nervous system
<b>COPD</b>	Chronic obstructive pulmonary disease
<b>CT</b>	Computed tomography
<b>CUA</b>	Cost-utility analysis
<b>DH</b>	Department of Health
<b>DI</b>	Delirium Index
<b>DRS / DRS-98 or</b>	Delirium Rating Scale / DRS-revised-98

<b>DRS-R-98</b>	
<b>DSA</b>	Deterministic Sensitivity Analysis
<b>DSI</b>	Delirium Symptom Interview
<b>DSM (DSM III, III-R or IV)</b>	Diagnostic and Statistical Manual of Mental Disorders (edition III, III-R or IV)
<b>ED</b>	Emergency Department
<b>EQ-5D</b>	EuroQol-5D
<b>FCEs</b>	Finished Consultant Episodes
<b>FIM</b>	Functional Independence Measure
<b>GA</b>	General anaesthesia
<b>GDG</b>	Guideline Development Group
<b>GI</b>	Gastrointestinal
<b>GP</b>	General Practitioner
<b>GRADE</b>	Grading of Recommendations Assessment, Development and Evaluation
<b>HES</b>	Hospital Episode Statistics
<b>HR</b>	Hazard Ratio
<b>HRQoL</b>	Health-related quality of life
<b>HT / 5-HT / 5-HT<sub>3</sub></b>	5-hydroxytryptamine / 5-hydroxytryptamine 3
<b>HTA</b>	Health technology assessment
<b>Hx</b>	History (in appendices)
<b>ICD-10</b>	International Classification of Diseases, 10 <sup>th</sup> edition
<b>ICDSC</b>	Intensive Care Delirium Screening Checklist
<b>ICER</b>	Incremental cost-effectiveness ratio
<b>ICU</b>	Intensive Care Unit
<b>IQR</b>	Interquartile range
<b>INMB</b>	Incremental Net Monetary Benefit
<b>IQCODE</b>	Informant Questionnaire on Cognitive Decline in the Elderly
<b>IRR</b>	Inter-rater reliability
<b>K</b>	Cohen's kappa
<b>ITT</b>	Intention to treat
<b>LOS</b>	Length of Stay
<b>LR<sup>+</sup></b>	Positive likelihood ratio
<b>LTC</b>	Long-term care
<b>LY</b>	Life-year
<b>MD</b>	Mean difference
<b>MDAS</b>	Memorial Delirium Assessment Scale
<b>MDC</b>	Major diagnostic category
<b>MI</b>	Myocardial infraction
<b>MMSE</b>	Mini-Mental State Examination

<b>MRI</b>	Magnetic resonance imaging
<b>MTI</b>	Multi-component Targeted Interventions
<b>NCGC</b>	National Clinical Guidelines Centre
<b>NH</b>	Nursing Home
<b>NHS</b>	National Health Service
<b>NHSEED</b>	The NHS Economic Evaluation Database
<b>NICE</b>	National Institute for Health and Clinical Excellence
<b>NINDS-AIREN</b>	National Institute of Neurological Disorders and Stroke and Association Internationale pour la Recherché et l'Enseignement en Neurosciences
<b>NNT</b>	Number needed to treat
<b>NPV</b>	Negative predictive value
<b>NSAID</b>	Non-steroidal anti-inflammatory drug
<b>OBS</b>	Organic Brain Syndrome
<b>OECD</b>	Organisation for Economic Co-operation and Development
<b>OR</b>	Odds ratio
<b>PASA</b>	NHS Purchasing and Supply Agency
<b>PCA</b>	Patient controlled analgesia
<b>PICO</b>	Framework incorporating patients, interventions, comparison and outcome
<b>POPS</b>	Proactive care of older people undergoing surgery
<b>PPP</b>	Purchasing Power Parity
<b>PPV</b>	Positive predictive value
<b>p.r.n</b>	Pro re nata
<b>PSA</b>	Probabilistic sensitivity analysis
<b>PSS</b>	Personal Social Services
<b>PSSRU</b>	Personal Social Services Research Unit
<b>QALY</b>	Quality-adjusted life year
<b>QUADAS</b>	Quality assessment tool for diagnostic accuracy studies
<b>RASS</b>	Richmond Agitation Sedation Scale
<b>RCT</b>	Randomised controlled trial
<b>RFs</b>	Risk factors
<b>ROC</b>	Receiver operating characteristic
<b>RR</b>	Relative risk
<b>SD</b>	Standard deviation
<b>SDC</b>	Saskatoon Delirium Checklist
<b>SE</b>	Standard error
<b>SICU</b>	Surgical Intensive Care Unit
<b>SPC</b>	Summary of product characteristics
<b>SPMSQ</b>	Short Portable Mental Status Questionnaire
<b>SR</b>	Systematic review

<b>TICS</b>	Telephone interview for cognitive status
<b>VAS</b>	Visual analogue scale

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## 1 Glossary of Terms

<b>Abstract</b>	Summary of a study, which may be published alone or as an introduction to a full scientific paper.
<b>Acute confusional state (ACS)</b>	A synonymous term for delirium.
<b>Algorithm (in guidelines)</b>	A flow chart of the clinical decision pathway described in the guideline, where decision points are represented with boxes, linked with arrows.
<b>Allocation concealment</b>	The process used to prevent advance knowledge of group assignment in a RCT. The allocation process should be impervious to any influence by the individual making the allocation, by being administered by someone who is not responsible for recruiting participants.
<b>AMT (Abbreviated Mental Test)</b>	A quick and easy to use screening test to detect cognitive impairment.
<b>Anticholinergic</b>	A group of drugs which inhibit the transmission of parasympathetic nerve impulses and inhibit the brain neurotransmitter acetylcholine.
<b>Antipsychotic</b>	Also known as neuroleptic drugs, these are a class of psychoactive drugs.
<b>Applicability</b>	The degree to which the results of an observation, study or review are likely to hold true in a particular clinical practice setting.
<b>Arm (of a clinical study)</b>	Sub-section of individuals within a study who receive one particular intervention, for example placebo arm
<b>Association</b>	Statistical relationship between two or more events, characteristics or other variables. The relationship may or may not be causal.
<b>Atypical antipsychotic</b>	These are the second-generation antipsychotics. They are chemically different from and have different side effects than the older 'typical' antipsychotic medications.
<b>Baseline</b>	The initial set of measurements at the beginning of a study (after run-in period where applicable), with which subsequent results are compared.
<b>Before-and-after study</b>	A study that investigates the effects of an intervention by measuring particular characteristics of a population both before and after taking the intervention, and assessing any change that occurs.
<b>Bias</b>	Systematic (as opposed to random) deviation of the results of a study from the 'true' results that is caused by the way the study is designed or conducted.
<b>Blinding</b>	Keeping the study participants, caregivers, researchers and outcome assessors unaware about the interventions to which the participants

	have been allocated in a study.
<b>Cardio-aspirin</b>	Lower dose treatment with aspirin to reduce the occurrence of vascular disease.
<b>Carer (caregiver)</b>	Someone other than a health professional who is involved in caring for a person with a medical condition.
<b>Case-control study</b>	Comparative observational study in which the investigator selects individuals who have experienced an event (For example, developed a disease) and others who have not (controls), and then collects data to determine previous exposure to a possible cause.
<b>Case-series</b>	Report of a number of cases of a given disease, usually covering the course of the disease and the response to treatment. There is no comparison (control) group of patients.
<b>Clinical efficacy</b>	The extent to which an intervention is active when studied under controlled research conditions.
<b>Clinical effectiveness</b>	The extent to which an intervention produces an overall health benefit in routine clinical practice.
<b>Clinical question</b>	In guideline development, this term refers to the questions about treatment and care that are formulated to guide the development of evidence-based recommendations.
<b>Clinician</b>	A healthcare professional providing direct patient care, for example doctor, nurse or physiotherapist.
<b>Cochrane Review</b>	The Cochrane Library consists of a regularly updated collection of evidence-based medicine databases including the Cochrane Database of Systematic Reviews (reviews of randomised controlled trials prepared by the Cochrane Collaboration).
<b>Cognitive impairment</b>	A brain disorder in which various thinking abilities such as memory or attention are impaired.
<b>Cohort study</b>	A retrospective or prospective follow-up study. Groups of individuals to be followed up are defined on the basis of presence or absence of exposure to a suspected risk factor or intervention. A cohort study can be comparative, in which case two or more groups are selected on the basis of differences in their exposure to the agent of interest.
<b>Comorbidity</b>	Co-existence of more than one disease or an additional disease (other than that being studied or treated) in an individual.
<b>Comparability</b>	Similarity of the groups in characteristics likely to affect the study results (such as health status or age).
<b>Concordance</b>	This is a recent term whose meaning has changed. It was initially applied to the consultation process in which doctor and patient agree therapeutic decisions that incorporate their respective views, but now includes patient support in medicine taking as well as prescribing communication. Concordance reflects social values but does not address medicine-taking and may not lead to improved adherence.

<b>Confidence interval (CI)</b>	A range of values for an unknown population parameter with a stated 'confidence' (conventionally 95%) that it contains the true value. The interval is calculated from sample data, and generally straddles the sample estimate. The 'confidence' value means that if the method used to calculate the interval is repeated many times, then that proportion of intervals will actually contain the true value.
<b>Confounding</b>	In a study, confounding occurs when the effect of an intervention on an outcome is distorted as a result of an association between the population or intervention or outcome and another factor (the 'confounding variable') that can influence the outcome independently of the intervention under study.
<b>Confusion Assessment Method (CAM)</b>	An assessment tool that has been validated to help detect delirium that is carried out by means of a clinical interview.
<b>Control group</b>	A group of patients recruited into a study that receives no treatment, a treatment of known effect, or a placebo (dummy treatment) - in order to provide a comparison for a group receiving an experimental treatment, such as a new drug.
<b>Cost benefit analysis</b>	A type of economic evaluation where both costs and benefits of healthcare treatment are measured in the same monetary units. If benefits exceed costs, the evaluation would recommend providing the treatment.
<b>Cost-consequences analysis (CCA)</b>	A type of economic evaluation where various health outcomes are reported in addition to cost for each intervention, but there is no overall measure of health gain.
<b>Cost-effectiveness analysis (CEA)</b>	An economic study design in which consequences of different interventions are measured using a single outcome, usually in 'natural' units (For example, life-years gained, deaths avoided, heart attacks avoided, cases detected). Alternative interventions are then compared in terms of cost per unit of effectiveness.
<b>Cost-effectiveness model</b>	An explicit mathematical framework, which is used to represent clinical decision problems and incorporate evidence from a variety of sources in order to estimate the costs and health outcomes.
<b>Cost-utility analysis (CUA)</b>	A form of cost-effectiveness analysis in which the units of effectiveness are quality-adjusted life-years (QALYs).
<b>Data synthesis</b>	A generic term to describe methods used for summarising (comparing and contrasting) evidence into a clinically meaningful conclusion in order to answer a defined clinical question. This can include systematic review (with or without meta-analysis), other quantitative methods or qualitative and narrative summaries.

<b>Decision analysis</b>	An explicit quantitative approach to decision making under uncertainty, based on evidence from research. This evidence is translated into probabilities, and then into diagrams or decision trees which direct the clinician through a succession of possible scenarios, actions and outcomes.
<b>Decision problem</b>	A clear specification of the interventions, patient populations and outcome measures and perspective adopted in an evaluation, with an explicit justification, relating these to the decision which the analysis is to inform.
<b>Discounting</b>	Costs and perhaps benefits incurred today have a higher value than costs and benefits occurring in the future. Discounting health benefits reflects individual preference for benefits to be experienced in the present rather than the future. Discounting costs reflects individual preference for costs to be experienced in the future rather than the present.
<b>Dominance</b>	An intervention is said to be dominated if there is an alternative intervention that is both less costly and more effective.
<b>Dosage</b>	The prescribed amount of a drug to be taken, including the size and timing of the doses.
<b>DSM III, III-R or IV</b>	Diagnostic and Statistical Manual of Mental Disorders (edition III, III-R or IV). Diagnostic test used to diagnose delirium.
<b>Economic evaluation</b>	Comparative analysis of alternative health strategies (interventions or programmes) in terms of both their costs and consequences.
<b>Effect (as in effect measure, treatment effect, estimate of effect, effect size)</b>	The observed association between interventions and outcomes or a statistic to summarise the strength of the observed association.
<b>Effectiveness</b>	See 'Clinical effectiveness'.
<b>Efficacy</b>	See 'Clinical efficacy'.
<b>Epidemiological study</b>	The study of a disease within a population, defining its incidence and prevalence and examining the roles of external influences (For example, infection, diet) and interventions.
<b>EQ-5D (EuroQoL-5D)</b>	A standardise instrument used to measure a health outcome. It provides a single index value for health status.
<b>Evidence</b>	Information on which a decision or guidance is based. Evidence is obtained from a range of sources including randomised controlled trials, observational studies, expert opinion (of clinical professionals and/or patients).
<b>Exclusion criteria (literature review)</b>	Explicit standards used to decide which studies should be excluded from consideration as potential sources of evidence.
<b>Exclusion criteria (clinical study)</b>	Criteria that define who is not eligible to participate in a clinical study.

<b>Extended dominance</b>	If Option A is both more clinically effective than Option B and has a lower cost per unit of effect, when both are compared with a do-nothing alternative then Option A is said to have extended dominance over Option B. Option A is therefore more efficient and should be preferred, other things remaining equal.
<b>Extrapolation</b>	In data analysis, predicting the value of a parameter outside the range of observed values.
<b>Extrapyramidal</b>	Pertaining to the tissues and structures outside the cerebrospinal pyramidal tracts of the brain that are associated with movement of the body, excluding motor neurons, the motor cortex, and the corticospinal and corticobulbar tracts.
<b>Follow-up</b>	Observation over a period of time of an individual, group or initially defined population whose appropriate characteristics have been assessed in order to observe changes in health status or health-related variables.
<b>Generalisability</b>	The extent to which the results of a study based on measurement in a particular patient population and/or a specific context hold true for another population and/or in a different context. In this instance, this is the degree to which the guideline recommendation is applicable across both geographical and contextual settings. For instance, guidelines that suggest substituting one form of labour for another should acknowledge that these costs might vary across the country.
<b>Gold standard</b>	See 'Reference standard'.
<b>GRADE / GRADE profile</b>	A system developed by the GRADE Working Group to address the shortcomings of present grading systems in healthcare. The GRADE system uses a common, sensible and transparent approach to grading the quality of evidence. The results of applying the GRADE system to clinical trial data are displayed in a table known as a GRADE profile.
<b>Harms</b>	Adverse effects of an intervention.
<b>Health economics</b>	The study of the allocation of scarce resources among alternative healthcare treatments. Health economists are concerned with both increasing the average level of health in the population and improving the distribution of health.
<b>Health-related quality of life (HRQoL)</b>	A combination of an individual's physical, mental and social well-being; not merely the absence of disease.

<b>Heterogeneity</b>	Or lack of homogeneity. The term is used in meta-analyses and systematic reviews when the results or estimates of effects of treatment from separate studies seem to be very different – in terms of the size of treatment effects or even to the extent that some indicate beneficial and others suggest adverse treatment effects. Such results may occur as a result of differences between studies in terms of the patient populations, outcome measures, definition of variables or duration of follow-up.
<b>Hypothesis</b>	A supposition made as a starting point for further investigation.
<b>Imprecision</b>	Results are imprecise when studies include relatively few patients and few events and thus have wide confidence intervals around the estimate of effect.
<b>Incident delirium</b>	Newly occurring case(s) of delirium
<b>Inclusion criteria (literature review)</b>	Explicit criteria used to decide which studies should be considered as potential sources of evidence.
<b>Incremental analysis</b>	The analysis of additional costs and additional clinical outcomes with different interventions.
<b>Incremental cost</b>	The mean cost per patient associated with an intervention minus the mean cost per patient associated with a comparator intervention.
<b>Incremental cost effectiveness ratio (ICER)</b>	The difference in the mean costs in the population of interest divided by the differences in the mean outcomes in the population of interest for one treatment compared with another.  $\text{ICER} = (\text{Cost}_A - \text{Cost}_B) / (\text{Effectiveness}_A - \text{Effectiveness}_B).$
<b>Incremental net benefit (INB)</b>	The value (usually in monetary terms) of an intervention net of its cost compared with a comparator intervention. The INB can be calculated for a given cost-effectiveness (willingness to pay) threshold. If the threshold is £20,000 per QALY gained then the INB is calculated as: (£20,000 x QALYs gained) – Incremental cost.
<b>Index</b>	In epidemiology and related sciences, this word usually means a rating scale, for example, a set of numbers derived from a series of observations of specified variables. Examples include the various health status indices, and scoring systems for severity or stage of cancer.
<b>Indirectness</b>	The available evidence is different to the clinical question being addressed, in terms of PICO (population, intervention, comparison and outcome).
<b>Intention to treat analysis (ITT)</b>	A strategy for analysing data from a randomised controlled trial. All participants are included in the arm to which they were allocated, whether or not they received (or completed) the intervention given to that arm. Intention-to-treat analysis prevents bias caused by the loss of participants, which may disrupt the baseline equivalence established by randomisation and which may reflect non-adherence to the protocol.

<b>Intervention</b>	Healthcare action intended to benefit the patient, for example, drug treatment, surgical procedure, psychological therapy.
<b>Intraoperative</b>	The period of time during a surgical procedure.
<b>Length of stay</b>	The total number of days a participant stays in hospital.
<b>Licence</b>	See 'Product licence'.
<b>Life-years gained</b>	Mean average years of life gained per person as a result of the intervention compared with an alternative intervention.
<b>Likelihood ratio</b>	The likelihood ratio combines information about the sensitivity and specificity. It tells you how much a positive or negative result changes the likelihood that a patient would have the disease. The likelihood ratio of a positive test result (LR+) is sensitivity divided by 1 - specificity.
<b>Literature review</b>	An article that summarises the evidence contained in a number of different individual studies and draws conclusions about their findings. It may or may not be systematically researched and developed.
<b>Long-term care</b>	Residential care within a facility that may include ongoing skilled nursing care and/or assistance with activities of daily living. Long-term care facilities include nursing homes, residential homes and EMI (elderly mentally infirm) homes.
<b>Loss to follow-up</b>	Also known as attrition. The loss of participants during the course of a study. Participants that are lost during the study are often called dropouts.
<b>Markov model</b>	A method for estimating long-term costs and effects for recurrent or chronic conditions, based on health states and the probability of transition between them within a given time period (cycle).
<b>Meta-analysis</b>	A statistical technique for combining (pooling) the results of a number of studies that address the same question and report on the same outcomes to produce a summary result. The aim is to derive more precise and clear information from a large data pool. It is generally more reliably likely to confirm or refute a hypothesis than the individual trials.
<b>Mini-Mental State Examination (MMSE)</b>	A commonly used instrument for screening cognitive function. It is not suitable for making a diagnosis but can be used to indicate the presence of cognitive impairment.
<b>Multidisciplinary team</b>	A team of healthcare professionals with the full spectrum of clinical skills needed to offer holistic care to patients with complex problems.
<b>Multivariate model</b>	A statistical model for analysis of the relationship between two or more predictor (independent) variables and the outcome (dependent) variable.

<b>Negative predictive value (NPV)</b>	[In screening/diagnostic tests:] A measure of the usefulness of a screening/diagnostic test. It is the proportion of those with a negative test result who do not have the disease, and can be interpreted as the probability that a negative test result is correct. It is calculated as follows: $NPV = \text{Number with a negative test who do not have disease} / \text{Number with a negative test}$ .
<b>Number needed to treat (NNT)</b>	The number of patients that who on average must be treated to prevent a single occurrence of the outcome of interest.
<b>Observational study</b>	Retrospective or prospective study in which the investigator observes the natural course of events with or without control groups; for example, cohort studies and case-control studies.
<b>Odds ratio</b>	A measure of treatment effectiveness. The odds of an event happening in the treatment group, expressed as a proportion of the odds of it happening in the control group. The 'odds' is the ratio of events to non-events.
<b>Outcome</b>	Measure of the possible results that may stem from exposure to a preventive or therapeutic intervention. Outcome measures may be intermediate endpoints or they can be final endpoints. See 'Intermediate outcome'.
<b>P-value</b>	The probability that an observed difference could have occurred by chance, assuming that there is in fact no underlying difference between the means of the observations. If the probability is less than 1 in 20, the P value is less than 0.05; a result with a P value of less than 0.05 is conventionally considered to be 'statistically significant'.
<b>Placebo</b>	An inactive and physically identical medication or procedure used as a comparator in controlled clinical trials.
<b>Polypharmacy</b>	The use or prescription of multiple medications.
<b>Positive predictive value (PPV)</b>	[In screening/diagnostic tests:] A measure of the usefulness of a screening/diagnostic test. It is the proportion of those with a positive test result who have the disease, and can be interpreted as the probability that a positive test result is correct. It is calculated as follows: $PPV = \text{Number with a positive test}$ .
<b>Postoperative</b>	Pertaining to the period after patients leave the operating theatre, following surgery.
<b>Post-test probability</b>	For diagnostic tests. The proportion of patients with that particular test result who have the target disorder ( $\text{post test odds} / [1 + \text{post-test odds}]$ ).
<b>Power (statistical)</b>	The ability to demonstrate an association when one exists. Power is related to sample size; the larger the sample size, the greater the power and the lower the risk that a possible association could be missed.
<b>Preoperative</b>	Pertaining to the period before surgery commences.
<b>Pre-test probability</b>	For diagnostic tests. The proportion of people with the target disorder in the population at risk at a specific time point or time

	interval. Prevalence may depend on how a disorder is diagnosed.
<b>Prevalent delirium</b>	Cases of delirium that are present at the first assessment of the person; it cannot be determined when the delirium began.
<b>Primary care</b>	Healthcare delivered to patients outside hospitals. Primary care covers a range of services provided by GPs, nurses and other healthcare professionals, dentists, pharmacists and opticians.
<b>Primary outcome</b>	The outcome of greatest importance, usually the one in a study that the power calculation is based on.
<b>Product licence</b>	An authorisation from the MHRA to market a medicinal product.
<b>Prognosis</b>	A probable course or outcome of a disease. Prognostic factors are patient or disease characteristics that influence the course. Good prognosis is associated with low rate of undesirable outcomes; poor prognosis is associated with a high rate of undesirable outcomes.
<b>Prospective study</b>	A study in which people are entered into the research and then followed up over a period of time with future events recorded as they happen. This contrasts with studies that are <i>retrospective</i> .
<b>Publication bias</b>	Also known as reporting bias. A bias caused by only a subset of all the relevant data being available. The publication of research can depend on the nature and direction of the study results. Studies in which an intervention is not found to be effective are sometimes not published. Because of this, systematic reviews that fail to include unpublished studies may overestimate the true effect of an intervention. In addition, a published report might present a biased set of results (e.g. only outcomes or sub-groups where a statistically significant difference was found).
<b>Quality of life</b>	See 'Health-related quality of life'.
<b>Quality-adjusted life year (QALY)</b>	An index of survival that is adjusted to account for the patient's quality of life during this time. QALYs have the advantage of incorporating changes in both quantity (longevity/mortality) and quality (morbidity, psychological, functional, social and other factors) of life. Used to measure benefits in cost-utility analysis. The QALYs gained are the mean QALYs associated with one treatment minus the mean QALYs associated with an alternative treatment.
<b>Quantitative research</b>	Research that generates numerical data or data that can be converted into numbers, for example clinical trials or the national Census which counts people and households.
<b>Quick Reference Guide</b>	An abridged version of NICE guidance, which presents the key priorities for implementation and summarises the recommendations for the core clinical audience.

<b>Randomisation</b>	Allocation of participants in a research study to two or more alternative groups using a chance procedure, such as computer-generated random numbers. This approach is used in an attempt to ensure there is an even distribution of participants with different characteristics between groups and thus reduce sources of bias.
<b>Randomised controlled trial (RCT)</b>	A comparative study in which participants are randomly allocated to intervention and control groups and followed up to examine differences in outcomes between the groups.
<b>RCT</b>	See 'Randomised controlled trial'.
<b>Receiver operated characteristic (ROC) curve</b>	A graphical method of assessing the accuracy of a diagnostic test. Sensitivity is plotted against 1-specificity. A perfect test will have a positive, vertical linear slope starting at the origin. A good test will be somewhere close to this ideal.
<b>Reference standard</b>	The test that is considered to be the best available method to establish the presence or absence of the outcome – this may not be the one that is routinely used in practice.
<b>Relative risk (RR)</b>	The number of times more likely or less likely an event is to happen in one group compared with another (calculated as the risk of the event in group A/the risk of the event in group B).
<b>Remit</b>	The brief given by the Department of Health and Welsh Assembly Government at the beginning of the guideline development process. This defines core areas of care that the guideline needs to address.
<b>Reporting bias</b>	See publication bias.
<b>Resource implication</b>	The likely impact in terms of finance, workforce or other NHS resources.
<b>Retrospective study</b>	A retrospective study deals with the present/ past and does not involve studying future events. This contrasts with studies that are <i>prospective</i> .
<b>Secondary outcome</b>	An outcome used to evaluate additional effects of the intervention deemed a priori as being less important than the primary outcomes.
<b>Selection bias</b>	A systematic bias in selecting participants for study groups, so that the groups have differences in prognosis and/or therapeutic sensitivities at baseline. Randomisation (with concealed allocation) of patients protects against this bias.
<b>Selection criteria</b>	Explicit standards used by guideline development groups to decide which studies should be included and excluded from consideration as potential sources of evidence.
<b>Sensitivity</b>	Sensitivity or recall rate is the proportion of true positives which are correctly identified as such. For example in diagnostic testing it is the proportion of true cases that the test detects.  See the related term 'Specificity'

<b>Sensitivity analysis</b>	<p>A means of representing uncertainty in the results of economic evaluations. Uncertainty may arise from missing data, imprecise estimates or methodological controversy. Sensitivity analysis also allows for exploring the generalisability of results to other settings. The analysis is repeated using different assumptions to examine the effect on the results.</p> <p>One-way simple sensitivity analysis (univariate analysis): each parameter is varied individually in order to isolate the consequences of each parameter on the results of the study.</p> <p>Multi-way simple sensitivity analysis (scenario analysis): two or more parameters are varied at the same time and the overall effect on the results is evaluated.</p> <p>Threshold sensitivity analysis: the critical value of parameters above or below which the conclusions of the study will change are identified.</p> <p>Probabilistic sensitivity analysis: probability distributions are assigned to the uncertain parameters and are incorporated into evaluation models based on decision analytical techniques (For example, Monte Carlo simulation).</p>
<b>Significance (statistical)</b>	<p>A result is deemed statistically significant if the probability of the result occurring by chance is less than 1 in 20 (<math>p &lt; 0.05</math>).</p>
<b>Specificity</b>	<p>The proportion of true negatives that are correctly identified as such. For example in diagnostic testing the specificity is the proportion of non-cases incorrectly diagnosed as cases.</p> <p>See related term 'Sensitivity'.</p> <p>In terms of literature searching a highly specific search is generally narrow and aimed at picking up the key papers in a field and avoiding a wide range of papers.</p>
<b>Stakeholder</b>	<p>Those with an interest in the use of the guideline. Stakeholders include manufacturers, sponsors, healthcare professionals, and patient and carer groups.</p>
<b>Subsyndromal delirium</b>	<p>A person who has some, but not all, the features of delirium.</p>
<b>Systematic review</b>	<p>Research that summarises the evidence on a clearly formulated question according to a pre-defined protocol using systematic and explicit methods to identify, select and appraise relevant studies, and to extract, collate and report their findings. It may or may not use statistical meta-analysis.</p>
<b>Treatment allocation</b>	<p>Assigning a participant to a particular arm of the trial.</p>
<b>Typical antipsychotic</b>	<p>These are sometimes referred to as first generation antipsychotics because they are the older medications used to treat psychotic symptoms. They were not called "typical" until the newer generation of these drugs (the 'atypical antipsychotics') were developed.</p>

**Univariate**

Analysis which separately explores each variable in a data set.

**Utility**

A measure of the strength of an individual's preference for a specific health state in relation to alternative health states. The utility scale assigns numerical values on a scale from 0 (death) to 1 (optimal or 'perfect' health). Health states can be considered worse than death and thus have a negative value.

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# 1 Introduction

## 2 1.1 What is a guideline

3 Our clinical guidelines are recommendations for the care of individuals in specific  
4 clinical conditions or circumstances within the National Health Service (NHS) –  
5 from prevention and self-care through primary and secondary care to more  
6 specialised services. We base our clinical guidelines on the best available  
7 research evidence, with the aim of improving the quality of healthcare. We use  
8 predetermined and systematic methods to identify and evaluate the evidence  
9 relating to specific clinical questions.

10 Clinical guidelines can:

- 11 • provide recommendations for the treatment and care of people by health  
12 professionals
- 13 • be used to develop standards to assess the clinical practice of individual  
14 health professionals
- 15 • be used in the education and training of health professionals
- 16 • help patients to make informed decisions
- 17 • improve communication between patient and health professional

18 While guidelines assist the practice of healthcare professionals, they do not  
19 replace their knowledge and skills.

20 We produce our guidelines using the following steps:

- 21 • Guideline topic is referred to the National Institute for Health and Clinical  
22 Excellence (NICE) from the Department of Health
- 23 • Stakeholders register an interest in the guideline and are consulted  
24 throughout the development process.
- 25 • The scope is prepared by the National Clinical Guideline Centre (NCGC)
- 26 • The NCGC establish a guideline development group
- 27 • A draft guideline is produced after the group assesses the available  
28 evidence and makes recommendations
- 29 • There is a consultation on the draft guideline.
- 30 • The final guideline is produced.

31

32 The NCGC and NICE produce a number of versions of this guideline:

- 33 • the **full guideline** contains all the recommendations, plus details of the  
34 methods used and the underpinning evidence

- 1 • the **NICE guideline** presents the recommendations from the full version in a
- 2 format suited to implementation by health professionals and NHS bodies
- 3 • the **quick reference guide** presents recommendations in a suitable format
- 4 for health professionals
- 5 • information for the public ('**understanding NICE guidance**') is written using
- 6 suitable language for people without specialist medical knowledge.

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8 This version is the full version. The other versions can be downloaded from the  
9 NCGC website at [ADD website](#) or are available from NICE [www.NICE.org.uk](http://www.NICE.org.uk).

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## 11 1.2 The need for this guideline

12 Delirium, sometimes called 'acute confusional state' (ACS) is characterised by a  
13 disturbance of consciousness and a change in cognition that develops over a  
14 short period of time.

15 Although the clinical presentation of delirium differs considerably from patient to  
16 patient, there are several characteristic features that help make the diagnosis.  
17 The standard criteria for delirium, are described in the 'Diagnostic and Statistical  
18 Manual of Mental Disorders' [DSM-IV] (1994):

- 19 • disturbance of consciousness (i.e., reduced clarity of awareness of the
- 20 environment) with reduced ability to focus, sustain, or shift attention.
- 21 • a change in cognition (such as memory deficit, disorientation, language
- 22 disturbance) or the development of a perceptual disturbance that is not
- 23 better accounted for by a pre-existing, established, or evolving
- 24 dementia.
- 25 • the disturbance develops over a short period of time (usually hours to days)
- 26 and tends to fluctuate during the course of the day.
- 27 • there is evidence from the history, physical examination, and laboratory
- 28 findings that: (1) the disturbance is caused by the direct physiological
- 29 consequences of a general medical condition, (2) the symptoms in criteria
- 30 (a) and (b) developed during substance intoxication, or during or shortly
- 31 after, a withdrawal syndrome, or (3) the delirium has more than one
- 32 aetiology".

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34 Features of delirium are recent onset of fluctuating awareness, impairment of  
35 memory and attention, and disorganised thinking. Additional features may  
36 include hallucinations and disturbance of sleep-wake cycle. There are three  
37 clinical subtypes of delirium: hyperactive (characterised by hallucinations,  
38 delusions, agitation, and disorientation), hypoactive (sleepy state, uninterested in  
39 activities of living, often unrecognised or labelled as dementia) or mixed  
40 (patients can move between the two subtypes). Delirium may be present when a  
41 person is admitted to hospital (prevalent delirium) or develop during an  
42 admission (incident delirium).

1 Delirium is a common but complex clinical syndrome that is known to be  
2 associated with poor outcomes.

3 There is a need for guidance to improve methods of appropriate identification,  
4 diagnosis, prevention and management of delirium. Failure to diagnose delirium,  
5 or misdiagnosis (mainly as dementia), can lead to inappropriate treatment being  
6 given. Delirium is often preventable and improvements in care practices and  
7 other treatments are needed. The improved management of delirium has the  
8 potential to generate cost savings.

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### 10 **1.3 The NCGC**

11 This guideline was commissioned by NICE and developed by the NCGC. The  
12 NCGC is one of four national collaborating centres (Cancer, Women and  
13 Children's Health, Mental Health and the NCGC) funded by NICE and comprises  
14 a partnership between a variety of academic, professional and patient-based  
15 organisations. As a multidisciplinary centre we draw upon the expertise of the  
16 healthcare professions and academics and ensure the involvement of patients in  
17 our work. Further information on the centre and our partner organisations can be  
18 found at our website [\(web address to be added before publication\)](#).

### 19 **1.4 Remit**

20 The following remit was received by the NCGC from the Department of Health  
21 in October 2007 as part of NICE's 17th wave programme of work.

22 The Department of Health asked the Institute:

23 *"Remit: To prepare a clinical guideline on the diagnosis, prevention and*  
24 *management of delirium"*

25

### 26 **1.5 What the guideline covers**

27 This guideline covers adult patients (18 years and older) in a hospital setting and  
28 adults (18 and older) in long-term residential care. The guideline addresses: risk  
29 factors to identify people at risk of developing delirium; diagnosis of delirium in  
30 acute, critical and long-term care; as well as pharmacological and non-  
31 pharmacological interventions for a) reducing the incidence of delirium and its  
32 consequences, and b) to reduce the severity, duration and consequences of  
33 delirium in people who develop the condition.

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35 Further details of the scope of the guideline can be found in Appendix A.

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### 37 **1.6 What the guideline does not cover**

38 This guideline does not cover children and young people (under the age of 18  
39 years), people receiving end-of-life care, people with intoxication and/or

1 withdrawing from drugs or alcohol, and people with delirium associated with  
2 these states.

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#### 4 **1.7 Who developed this guideline**

5 A multidisciplinary Guideline Development Group (GDG) comprising professional  
6 group members and consumer representatives of the main stakeholders  
7 developed this guideline (see section on Guideline Development Group  
8 Membership and acknowledgements).

9 NICE funds the NCGC and thus supported the development of this guideline. The  
10 GDG was convened by the NCGC and chaired by Professor John Young in  
11 accordance with guidance from NICE.

12 The group met every 6-8 weeks during the development of the guideline. At the  
13 start of the guideline development process, all GDG members declared interests  
14 including consultancies, fee-paid work, share-holdings, fellowships and support  
15 from the healthcare industry. At all subsequent GDG meetings, members  
16 declared arising conflicts of interest, which were also recorded (Appendix B).

17 Members are either required to withdraw completely or for part of the  
18 discussion if their declared interest makes it appropriate, however this was not  
19 deemed necessary for any group members on this guideline.

20 Staff from the NCGC provided methodological support and guidance for the  
21 development process. They undertook systematic searches, retrieval and  
22 appraisal of the evidence and drafted the guideline. The glossary to the  
23 guideline contains definitions of terms used by staff and the GDG.

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## 2 Methodology

### 2.1 Guideline methodology

This guideline was commissioned by NICE and developed in accordance with the guideline development process outlined in 'The guidelines manual' (NICE2009).

### 2.2 Developing the clinical questions

Clinical questions were developed to guide the literature searching process and to facilitate the development of recommendations by the GDG. They were drafted by the technical team and refined and validated by the GDG. The questions were based on the scope (Appendix A).

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#### 2.2.1 List of all clinical questions

The full list of clinical questions addressed by the guideline is summarised in table 2.1 below:

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Table 2.1: full list of clinical questions

Question wording
<b>Diagnosis</b>
Assessment methods for identifying people at risk of delirium
Identification of symptoms that indicate patients may have delirium
Practical diagnostic tests for identifying patients with delirium in different settings
Diagnostic criteria for identifying patients with delirium
<b>Prognosis</b>
Risk factors for delirium
Precipitating factors for delirium
Consequences of, and following, delirium
<b>Interventions</b>
Prevention of delirium in a hospital setting
Pharmacological interventions for the prevention of delirium in a hospital setting
Single component, non-pharmacological interventions for the prevention of delirium in a hospital setting
Multi-component interventions for the prevention of delirium in hospital setting
Prevention of delirium in a long-term care setting
Pharmacological interventions for the prevention of delirium in long-term care
Single component, non-pharmacological interventions for the prevention of delirium in a long-term care setting
Multi-component interventions for the prevention of delirium in long-term care
Treatment of delirium in a hospital setting

Pharmacological interventions for the treatment of delirium in a hospital setting
Single component, non-pharmacological interventions for the treatment of delirium in a hospital setting
Multi-component interventions for the treatment of delirium in a hospital setting
Treatment of delirium in a long-term care setting
Pharmacological interventions for the treatment of delirium in a long-term care setting
Single component, non-pharmacological interventions for the treatment of delirium in a long-term care setting
Multi-component interventions for the treatment of delirium in a long-term care setting
Patient information
Information for people with delirium or at risk of delirium, and their carers
Other
Prevalence of delirium in different settings

1

## 2 2.2.2 Review questions

3 From these clinical questions, the technical team produced review questions and  
 4 protocols to address these questions. The protocols were converted into the  
 5 methods section (see 2.4).

## 6 2.3 Literature search

### 7 2.3.1 Clinical literature search

8 The search strategies and the databases searched are presented in detail in  
 9 Appendix C. All searches were carried out on the following core databases:  
 10 Medline, Embase, Cinahl and The Cochrane Library. Additional databases were  
 11 searched for individual reviews as appropriate.

12 Databases were searched using relevant subject headings and free-text terms.  
 13 Where appropriate, study design filters were applied. Non-English language  
 14 studies and abstracts were not reviewed.

15 Searches were initially performed for articles published since 1994, the  
 16 publication date of the DSM-IV which is the reference standard for the diagnosis  
 17 of delirium. Following guidance from the GDG, a further search back to 1987  
 18 was carried out in order to retrieve studies using the earlier *Diagnostic and*  
 19 *Statistical Manual III (Revised)* (DSMIII-R) as the reference standard.

20 All searches were updated to 17<sup>th</sup> August 2009. Hand-searching was not  
 21 undertaken following NICE advice that exhaustive searching on every guideline  
 22 review topic is not practical or efficient (Mason 2002). Reference lists of articles  
 23 were checked for studies of potential relevance.

24

### 25 2.3.2 Sifting process

26 Once the search had been completed, the following sifting process took place:

- 27 • 1st sift: one reviewer sifted the title/abstract for articles that potentially  
 28 met the eligibility criteria.

- 1           • 2nd sift: full papers were ordered that appeared relevant and eligible or  
2           where relevance/eligibility was not clear from the abstract.
- 3           • 3rd sift: full papers were appraised that meet eligibility criteria.  
4           Generally, one reviewer appraised the papers using an inclusion criteria  
5           form, and this was checked where necessary by a second reviewer.

6  
7           Once individual papers were retrieved, the articles were checked for  
8           methodological rigour (see section 2.4.7), applicability to the UK and clinical  
9           significance. Assessment of study quality concentrated on dimensions of internal  
10          validity and external validity. At this stage, some studies were excluded if the  
11          interventions were not licensed for use in the UK or they were not regularly used  
12          in the UK. Studies in which the interventions were obsolete were also excluded.

### 13 14           **2.3.3       Economic literature search**

15          Economic evidence was obtained from systematic searches of the following  
16          databases in accordance with the NICE Guidelines Manual: Medline, Embase, the  
17          Health Technology Appraisals (HTA) database and the NHS Economic  
18          Evaluations Database (NHSEED). The latter two databases were searched via The  
19          Cochrane Library.

20          Detailed search strategies can be found in Appendix J.

## 21 22 23           **2.4   Clinical effectiveness review methods**

24          This section describes the methods of reviewing that are common to all reviews of  
25          intervention studies, to reviews of prognostic factors and to reviews of diagnostic  
26          test accuracy. Further specific details are given in the individual reviews.

### 27 28           **2.4.1 Selection criteria: general**

29          The following selection criteria were to be applied to studies to determine their  
30          suitability for inclusion in the reviews:

#### 31 32           **2.4.1.1 Types of studies**

33          For intervention studies, the randomised trial (RCT) and quasi randomised trial  
34          (for example, allocation by alternation, and date of birth) were to be the  
35          primary trial designs. Non-randomised studies could be included only if there  
36          was no other evidence, with preference given to large cohort studies and  
37          comparative non-randomised designs; case series or case reports were not

1 included and before-and-after studies were considered cautiously for prevention  
2 studies only.

3  
4 For prognostic factor reviews, RCTs comparing groups with different risk factors  
5 (e.g. types of surgery) and cohort studies (prospective and retrospective)  
6 investigating the incidence of delirium or the consequences of delirium were to  
7 be the main study designs. We note that, for some risk factors (e.g. age), the  
8 randomised trial cannot be used as the study design. If there were no cohort  
9 studies available, case-control studies and cross-sectional surveys could be  
10 considered, with allowance made for the fact that they have increased potential  
11 for bias.

12  
13 For reviews of diagnostic test accuracy, the cross sectional study was to be the  
14 primary study design. Studies were to be those in which diagnoses obtained  
15 using a new (index) test were compared with 'true' diagnoses obtained using a  
16 reference standard, with both tests being carried out in the same patients. Case  
17 control studies were to be considered only in the absence of cross sectional  
18 studies.

19  
20 Studies were to be excluded if there were fewer than 20 patients in each arm  
21 for comparative studies and if there were fewer than 20 patients overall for  
22 cohort studies. We did not restrict the size of the studies of diagnostic test  
23 accuracy.

24  
25 Studies were limited to the English language, initially, with the exception of  
26 studies translated for Cochrane reviews, but the GDG directed that a search was  
27 carried out for any RCT, regardless of the language.

28

#### 29 **2.4.1.2 Types of participants**

30 For intervention studies, reviews were to be carried out separately to address  
31 interventions for prevention and treatment of delirium. Separate reviews were  
32 also done in the two main population groups: patients in a hospital setting and  
33 people in long-term care.

34 For prognostic factor reviews, the populations were not to be treated  
35 separately, although it was noted which population was concerned.

36  
37 Reviews of diagnostic test accuracy are sensitive to the population, so long-term  
38 care, hospital setting and intensive care unit (ICU) were to be treated  
39 separately.

40  
41 For all reviews, participants were to be adults (18 years and older) who were:

- 42 • Patients in a hospital setting, including surgical, medical, ICU, Accident and  
43 Emergency departments, and those in mental health settings

- 1           • In long-term care settings

2  
3           Studies including children or young people were to be considered if the mean  
4           age was 18 years or older. Studies in the community could be included as  
5           indirect evidence for the long-term care population.

6  
7           Excluded populations were to be:

- 8           • Children and young people (younger than 18 years).  
9           • People receiving end-of-life care.  
10          • People with intoxication and/or those who are withdrawing from drugs or  
11            alcohol, and/or (**treatment intervention reviews**) people with delirium  
12            associated with these states

13  
14          For the treatment intervention reviews: participants were to have delirium.  
15          Delirium is defined according to criteria described in the DSM-IV (1994) (see  
16          Appendix I). Typically delirium is diagnosed by examining changes in cognitive  
17          function, and this is linked to the DSM-IV criteria. Validated instruments, based on  
18          the operational application of the DSM-IV or DSM-III-R diagnostic criteria, are  
19          given in Appendix I.

20

## 21   **2.4.2 Selection criteria: reviews of interventions**

### 22   **2.4.2.1 Types of intervention**

23          The interventions considered varied across reviews. Interventions could be  
24          pharmacological or non-pharmacological (e.g. haloperidol, music therapy).

25  
26          Pharmacological interventions were to be restricted to those licensed for use in  
27          the UK, but these drugs were not necessarily those indicated for delirium (there  
28          are no drugs for delirium in the British National Formulary (BNF)).  
29          Pharmacological reviews were to be carried out by class rather than by  
30          individual drug, but drugs within a class were to be reported as subgroups (e.g.  
31          atypical antipsychotics: olanzapine and risperidone).

32  
33          Different doses, regimens and routes of delivery were to be permitted and  
34          studies were to be initially combined in analyses, regardless of these features.

35

### 36   **2.4.2.2 Types of comparisons**

37          The following comparisons were to be included:

- 1 i. Delirium intervention (**A**) versus placebo
- 2 ii. **A** versus usual care/no intervention
- 3 iii. **A** plus second intervention (**X**) versus **X** alone
- 4 iv. Within a class of interventions, **A1.1** versus **A1.2**
- 5 v. Across classes of interventions: **A1** versus **A2**

6

7

In analyses, comparisons (i) and (ii) could be combined, but (iii) was to be treated separately because of possible drug interactions.

8

9

#### 10 **2.4.2.3 Types of outcome measures**

11 For studies of interventions for the prevention of delirium, the primary outcome  
12 was to be incidence of delirium. All included types and severities of delirium  
13 were to be combined. For reviews of patients in hospital, the primary outcome  
14 was to be measured during the hospital stay.

15

16 For the incidence of delirium, studies should report that the DSM-IV or the DSM-  
17 III-R and validated scales associated with them were used (see Appendix I).  
18 Other acceptable methods could include a structured clinical interview.

19

20

Secondary outcomes were to be:

21

- Duration of delirium

22

- Severity of delirium

23

- Length of stay in hospital

24

- Incidence of dementia or cognitive impairment

25

- Number of patients discharged to new long-term care placement (for  
26 studies in a hospital setting)

26

27

- Mortality

28

- Quality of life (patient)

29

- Quality of life (carer)

30

- Activities of daily living

31

- Use of psychotropic medication

32

- Incidence of post traumatic stress disorder

33

- Admission to hospital (for long-term care studies)

34

35

For studies of interventions for the treatment of delirium, the primary outcomes  
36 were to be:

36

- 1           • Duration of delirium
- 2           • Complete response (number recovered from delirium)
- 3
- 4       Secondary outcomes:
- 5           • Severity of delirium
- 6           • Length of stay
- 7           • Incidence of dementia / cognitive impairment
- 8           • Number of patients discharged to new long-term care placement (for those
- 9                 in hospital)
- 10          • Mortality
- 11          • Number of patients with persisting delirium
- 12          • Quality of life (patient)
- 13          • Quality of life (carer)

14

15       For all intervention reviews, other outcome measures to be recorded were:

- 16           • Adverse effects associated with the intervention (e.g. extrapyramidal
- 17                 symptoms)

18

### 19   **2.4.3 Selection criteria: reviews of prognostic factors**

20       Two types of prognostic factor reviews were carried out, investigating prognostic  
21       factors for delirium, and studying the consequences of delirium for people with  
22       delirium.

23

#### 24   **2.4.3.1 Prognostic (risk) factors**

25       The risk factors to be considered for delirium are listed at the start of that  
26       review (section 6.2.1).

27       For the consequences of delirium review, the risk factor was to be one of:

- 28           • Incident delirium (although prevalent delirium was also acceptable)
- 29           • Persistent delirium: this was defined after McAvay (2006) as 'delirium in
- 30                 patients who met the full criteria for delirium at the discharge interview,
- 31                 or who had full delirium during the hospitalisation and partial symptoms
- 32                 at discharge'.
- 33           • Severity of delirium

1

2 **2.4.3.2 Types of outcome measures**

3 For the risk factors review, the following outcomes were to be included:

- 4 • Incidence of delirium
- 5 • Incidence of persistent delirium
- 6 • Severity of delirium
- 7 • Duration of delirium

8

9 For the consequences review, the following outcomes were to be included:

- 10 • Dementia/Cognitive impairment
- 11 • Progression of dementia
- 12 • Discharge to care home (for people who were in hospital)
- 13 • Falls
- 14 • Hospital admission (for people who were in long-term care)
- 15 • Post discharge care
- 16 • Post traumatic stress disorder
- 17 • Pressure Ulcers
- 18 • Mortality
- 19 • Impact on carers
- 20 • Length of stay
- 21 • Quality of life for patients

22

23 **2.4.4 Selection criteria: reviews of diagnostic test accuracy**24 **2.4.4.1 Prior tests**

25 No prior tests were to have been undertaken

26

27 **2.4.4.2 The index test**28 The following index tests, including the people operating them, were to be  
29 examined, subdivided by setting:

- 30 • Hospital:
  - 31 ○ Abbreviated Mental test (AMT); anyone could do this test
  - 32 ○ Clock-drawing; could be used by untrained nurses or volunteers

- 1                   ○ Confusion Assessment Method [long version] (long CAM); should
- 2                   be carried out by trained healthcare professionals
- 3                   ○ Confusion Assessment Method [short version] (short CAM); should
- 4                   be carried out by trained healthcare professionals
- 5                   ○ Delirium Rating Scale (DRS-98); should be carried out by trained
- 6                   healthcare professionals
- 7                   ○ Mini Mental State Examination (MMSE) or other cognitive
- 8                   assessment instrument: trained healthcare professionals.
- 9                   • ICU:
- 10                  ○ CAM-ICU and Richmond Agitation Sedation Scale (RASS)
- 11                  (together); should be carried out by trained healthcare
- 12                  professionals

13

#### 14   **2.4.4.3 The reference standard**

15           The reference standard was to be DSM-IV or ICD-10; carried out by a trained

16           specialist.

17

#### 18   **2.4.4.4 The target condition**

19           The target condition was to be delirium; subsyndromal delirium was not to be

20           included.

21

#### 22   **2.4.5 Outcomes**

23           For studies of diagnostic test accuracy, the outcomes to be recorded were

24           sensitivity, specificity, positive predictive value, negative predictive value,

25           likelihood ratio, diagnostic odds ratio, pre- and post-test probabilities. These

26           were to be calculated from raw data, and occasionally raw data could be back-

27           calculated from test accuracy statistics.

28

#### 29   **2.4.6 Data extraction**

30           Data from included studies were extracted by one reviewer for each review,

31           and randomly checked by a second reviewer, and entered into a Microsoft

32           Access relational database that had been especially designed for the guideline.

33

#### 34   **2.4.7 Appraisal of methodological quality of intervention studies**

35           For randomised trials, the following factors were considered in assessing the

36           potential for bias:

- 1           • *A priori* sample size calculation:
- 2           • Method of generation of the randomisation sequence:
- 3           • Allocation concealment at randomisation:
  - 4                 ○ The means of preventing the treatment assignment being known
  - 5                 before the time of allocation
- 6           • Baseline comparability of treatment groups for relevant risk factors
- 7           • Patients stated to be blinded, especially for comparisons with placebo:
  - 8                 ○ Blinding involves hiding the nature of the intervention from
  - 9                 participants, clinicians and treatment evaluators after allocation
  - 10                has taken place
  - 11                ○ Blinding may be not be possible depending on the nature of the
  - 12                interventions
  - 13                ○ Blinding may be more important for some outcomes than others:
- 14          • Outcome assessor stated to be blinded
- 15          • No missing data for each outcome:
  - 16                ○ Studies with at least 20% of data missing from any group were
  - 17                to be considered to be potentially biased, more so if there is a
  - 18                differential drop out from any one group or if the missing data is
  - 19                known to be significantly different from the remaining data
  - 20                ○ Those with moderate loss to follow up (20 to 50%) were to be
  - 21                considered in sensitivity analyses
  - 22                ○ Those with 50% or more patients missing from any one group
  - 23                were to be regarded as flawed and not analysed further (but
  - 24                would be included in the review)
- 25          • Intention to treat analysis:
  - 26                ○ Trial participants should be analysed in the groups to which they
  - 27                were randomised regardless of which (or how much) treatment
  - 28                they actually received, and regardless of other protocol
  - 29                irregularities **and**
  - 30                ○ All participants should be included regardless of whether their
  - 31                outcomes were actually collected
- 32
- 33          **For non-randomised intervention studies**, the following factors were
- 34          considered in assessing the potential for bias; further details are given in The
- 35          Cochrane Handbook for Systematic Reviews of Interventions
- 36          (<http://www.cochrane-handbook.org/> : Box 13.1.a: Some types of non-
- 37          randomised study design used for evaluating the effects of interventions)
- 38          • Selection bias:
  - 39                ○ Account is taken of the confounding factors, either by design (e.g.
  - 40                matching or restriction to particular subgroups) or by methods of
  - 41                analysis (e.g. stratification or regression modelling with propensity
  - 42                scores or covariates)

- 1                   ○ Confounding factors for delirium intervention reviews that the  
2                   GDG believed should be taken into consideration were: age,  
3                   cognitive impairment, sensory impairment, polypharmacy
- 4                   ● Prospectiveness:
- 5                   ○ On the basis of identification of participants; baseline assessment  
6                   and treatment allocation; assessment of outcomes
- 7                   ● Blinding (see RCTs)
- 8                   ○ Of patients
- 9                   ○ Of outcome assessors
- 10                  ● No loss to follow up (see RCTs)
- 11                  ● Intention to treat (see RCTs)
- 12

#### 13   **2.4.8 Appraisal of methodological quality of studies of prognostic factors**

14                  Cohort studies were assessed using criteria based on the Newcastle-Ottawa  
15                  checklist and the NICE Guidelines Manual. The following criteria were  
16                  considered, with examples given for risk factors for the incidence of delirium –  
17                  similar arguments apply for the consequences review:

- 18                  ● Representativeness of the exposed cohort:
- 19                  ○ Truly representative of the community e.g. random sample from  
20                  the guideline's population\*
- 21                  ○ Somewhat representative of the community e.g. hospital patients  
22                  only\*
- 23                  ○ Selected group e.g. cardiac operations
- 24                  ○ No description of the derivation of the cohort or unclear.
- 25
- 26                  ● Selection of the non exposed cohort:
- 27                  ○ Drawn from the same community as the exposed cohort\*
- 28                  ○ Drawn from a different source – e.g. compared with general  
29                  population levels in epidemiological studies
- 30                  ○ No description of the derivation of the non exposed cohort or  
31                  unclear.
- 32
- 33                  ● Ascertainment of exposure:
- 34                  ○ Measurement of risk factor using an adequate method (e.g.  
35                  MMSE for dementia)\*
- 36                  ○ Measurement of risk factor using a partly adequate method\*

- 1                   ○ Measurement of risk factor using an inadequate method (e.g.  
2                   retrospective examination of chart records)
- 3                   ○ No description.
- 4
- 5                   ● Demonstration that the outcome of interest was not present at the start of  
6                   the study:
- 7                   ○ Yes (includes analyses that excluded patients with prevalent  
8                   delirium)\*
- 9                   ○ No.
- 10
- 11                  ● Prospectiveness:
- 12                  ○ Prospective study\*
- 13                  ○ Retrospective study
- 14                  ○ Unclear.
- 15
- 16                  ● Comparability of cohorts on the basis of the design or analysis:
- 17                  ○ Cohorts balanced at baseline for important factors (see below)\*
- 18                  ○ Adjusted for confounding factors in the analysis and has at least  
19                  10 events per factor in the analysis\*
- 20                  ○ Study has at least 8 to 10 events per factor and analysis is  
21                  adjusted for at least 3 of 4 relevant factors in the analysis\*
- 22                  ○ Study adjusts for some confounders (or keeps them constant): 2 of  
23                  4 included in the analysis
- 24                  ○ Study has fewer than 8 to 10 events per factor in the analysis
- 25                  ○ Study does not adjust for confounders.
- 26

27                  In cohort studies, the best way to adjust for confounders is to use regression  
28                  methods to adjust for all the factors at once in a multivariate analysis. For  
29                  validity, there should be at least ten patients for each factor in the regression  
30                  equation for continuous outcomes, or at least ten patients having the event (e.g.  
31                  delirium) per factor for dichotomous outcomes. However, if there are insufficient  
32                  relevant factors taken into account, the quality of the study should be  
33                  downgraded.

34

35                  The relevant factors that had to be included in the analysis were decided *a-*  
36                  *priori* by the GDG using consensus methods. For the non-pharmacological risk  
37                  factors review for the incidence of delirium, they were: age; sensory impairment,  
38                  dementia/cognitive impairment and polypharmacy. For the pharmacological risk  
39                  factors review, polypharmacy was excluded. The relevant factors for each  
40                  consequence of delirium are given in that review. To qualify as a well adjusted

1 study, the analysis should include at least 3 out of 4 of these factors (or they  
2 should be kept constant).

3 • Ascertainment of outcome:

- 4 ○ Measurement of delirium using an adequate method (e.g. DSMIV,  
5 CAM)\*
- 6 ○ Measurement of delirium using a partly adequate method (e.g.  
7 MMSE)
- 8 ○ Measurement of delirium using an inadequate method (e.g.  
9 retrospective examination of chart records)
- 10 ○ No description.

11

12 • Adequacy of follow up of cohorts:

- 13 ○ Complete follow-up: all participants accounted for\*
- 14 ○ Participants lost to follow-up unlikely to introduce bias: more than  
15 80% follow up\*
- 16 ○ Follow-up rate less than 80% and no description of those lost
- 17 ○ No statement.

18

19 Studies were considered to be of acceptable quality if the asterisked statements  
20 were met, otherwise their quality rating was downgraded. All these factors were  
21 taken into consideration to give an overall quality rating.

22

23 **2.4.9 Appraisal of methodological quality of studies of diagnostic test accuracy**

24 For studies of diagnostic test accuracy, the study quality was assessed using a  
25 modified version of the 'QUADAS' list, with each item scored as yes, no or  
26 unclear (Whiting 2003). The following factors were considered in assessing the  
27 potential for bias:

- 28 • Representative spectrum: whether or not the patients had delirium and  
29 were representative of the population of the review.
  - 30 ○ Studies that recruited a group of healthy controls and a group  
31 known to have the target disorder were coded as 'no' on this item
- 32 • Clear description of selection criteria
- 33 • Reference standard likely to classify the target condition correctly
- 34 • Acceptable delay between tests: period between the reference standard  
35 and the index test was short enough to be reasonably sure that the  
36 target condition did not change between the 2 tests; for delirium, the  
37 GDG considered this to be about half a day

1  
2 An overall assessment for each study was given of ++ (good), + (acceptable,  
3 with some reservations) and – (unacceptable)

4

#### 5 **2.4.10 Data synthesis for intervention trials**

6 Meta-analysis of similar trials, where appropriate, was carried out using *The*  
7 *Cochrane Collaboration's* analysis software, Review Manager (Version 5). Trials  
8 were pooled using a fixed effects model and plotted on forest plots. Where  
9 there was significant heterogeneity, a random effects model was used as a  
10 sensitivity analysis.

11

12 For dichotomous studies, intention to treat analyses (including all participants  
13 according to their assigned groups) were used, when reported by the study  
14 authors, and failing that, available case analyses (all those reporting an  
15 outcome) as reported by the authors. When there were incomplete data  
16 reported (more than 20% missing in any one group), we carried out sensitivity  
17 analyses, excluding these studies.

18

19 When it was possible to combine studies, outcomes were summarised for  
20 dichotomous data using relative risks. Numbers needed to treat, with their 95%  
21 confidence intervals (95% CI) and the control group rate (range of rates) to  
22 which they apply, were calculated from the risk difference where appropriate.  
23 The number needed to treat (NNT) is the number of patients who would have to  
24 be treated for one to have an improved outcome.

25

26 For continuous data, weighted mean differences were used to summarise the  
27 pooled data, and where the studies had different scales, standardised mean  
28 differences were used. Sometimes it may be necessary to invert scales (e.g. if  
29 one has the maximum value meaning poor outcome and in another it means a  
30 good outcome).

31

32 Studies, in which one or more reported final values and others reported change  
33 scores, were combined if the scales used were the same, otherwise they were  
34 reported separately. If both final values and change scores were reported in a  
35 single study, the former were used. Summary statistics and their 95% confidence  
36 intervals were reported where sufficient detail allowed their calculation,  
37 together with the control group range.

38

39 Where there were differences between studies in the way the results were  
40 reported, for example, summary statistics only or raw data, the summary statistic  
41 (e.g. RR) and its standard error was calculated from 95% Confidence intervals,  
42 and the studies combined using the generic inverse variance method in Review  
43 Manager. For continuous outcomes reporting the difference in means with a p-  
44 value, the standard error was also calculated.

1  
2 Where possible, account was taken of unit of randomisation errors (e.g. cluster  
3 trials).

4  
5 Results from RCTs and non-randomised studies were not combined, but were  
6 reported as subgroups. Generally non-randomised studies were not included if  
7 the RCT data were adequate, but if the RCTs were very small or of poor quality,  
8 non-randomised studies could be included to give supplementary information.

9  
10 Heterogeneity between trials was assessed by visual inspection of forest plots,  
11 noting where there was poor overlap of horizontal lines, and by using statistical  
12 measures: the  $\chi^2$  test for heterogeneity and the level of inconsistency,  $I^2$  ( $I^2 = [(\chi^2$   
13  $- df) / \chi^2] \times 100\%$ , where df is the degrees of freedom). We considered that  
14 there was heterogeneity if the heterogeneity p-value was less than 0.1 and/or  $I^2$   
15 was greater than 50%. Any heterogeneity was explored further (see subgroup  
16 analyses below) and unexplained heterogeneous results were not used as the  
17 basis for recommendations.

#### 18 19 **2.4.10.1 Stratifications**

20 Separate reviews were carried out for prevention and treatment, and for setting  
21 (hospital and long-term care).

#### 22 23 **2.4.10.2 Combining studies**

24 Studies were combined regardless of:

- 25 • medical or surgical patients
- 26 • ICU or not
- 27 • risk of delirium, including baseline levels of dementia (for prevention  
28 reviews)
- 29 • dose of intervention

30  
31 In pharmacological reviews, all the drugs in a particular class were considered in  
32 the same review, with individual drugs considered as subgroups in meta-analysis.

#### 33 34 **2.4.10.3 Subgroup analyses**

35 If there was heterogeneity, subgroup analyses were carried out to investigate it.

36 The following subgroups were considered:

Delirium: full guideline DRAFT (November 2009)

- 1           • For prevention reviews: people at high risk of delirium, such as those with
- 2            dementia, may be distinguished from lower risk groups.
- 3           • Patients in ICU
- 4           • Type of intervention
- 5           • Dose of intervention
- 6           • Illness severity
- 7

#### 8   **2.4.10.4 Sensitivity analyses**

9           Sensitivity analyses were carried out to investigate assumptions within the  
10          analyses. These included the following:

- 11          • Methodological quality
- 12          • Fixed effects model
- 13          • Other features specific to each review.

14  
15          In terms of methodological quality, we paid particular attention to allocation  
16          concealment and loss to follow-up (missing data). We did not include studies with  
17          more than 50% missing data in the analyses. Otherwise we carried out sensitivity  
18          analyses on studies that had between 20 and 50% missing data in any group.

19

#### 20   **2.4.11                   Data synthesis for prognostic factor reviews**

21          Odds ratios or relative risks, with their 95% confidence intervals, from  
22          multivariate analyses were extracted from the papers, and standard errors were  
23          calculated from the 95% CIs. The log (odds ratio) with its standard error was  
24          then entered into the generic inverse variance technique of Review Manager 5.  
25          Studies were not combined in a meta-analysis because they were observational  
26          studies. Sensitivity analyses were carried out on the basis of study quality, and  
27          the results were represented on forest plots and reported as ranges.

28

#### 29   **2.4.12                   Data synthesis for reviews of diagnostic test accuracy**

30          For diagnostic test accuracy studies, 2 by 2 tables were constructed from raw  
31          data, which allowed calculation of sensitivity, specificity, positive predictive  
32          value, negative predictive value, likelihood ratio, diagnostic odds ratio, pre- and  
33          post-test probabilities. Some of this was done using an Access database, and  
34          Review Manager (version 5) was also used for the calculation of sensitivity and  
35          specificity and the representation of these in both forest plots and the receiver  
36          operating characteristic (ROC) space.

37

### 1 2.4.13 Grading evidence

2 The GRADE<sup>‡</sup> scheme (GRADE working group 2004) was used to assess the  
3 quality of the evidence for each outcome using the approach described below,  
4 and evidence summaries across all outcomes were produced.

5  
6 According to the GRADE scheme, evidence is classified as high, moderate, low or  
7 very low:

- 8 • High: further research is very unlikely to change our confidence in the  
9 estimate of effect
- 10 • Moderate: further research is likely to have an important impact on our  
11 confidence in the estimate of effect and may change the estimate
- 12 • Low: further research is very likely to have an important impact on our  
13 confidence in the estimate of effect and is likely to change the estimate
- 14 • Very low: any estimate of effect is very uncertain.

15  
16 The procedure adopted when using GRADE was:

- 17 • A quality rating was assigned, based on the study design, for example,  
18 RCTs started as high and observational studies as low.
- 19 • This rating was up- or down-graded according to specified criteria: study  
20 quality, consistency, directness, preciseness and reporting bias. These  
21 criteria are detailed below. Criteria were given a downgrade mark of –  
22 1 or –2 depending on the severity of the limitations.
- 23 • The downgrade/upgrade marks were then summed and the quality rating  
24 revised. For example, a decrease of –2 points for an RCT would result in  
25 a rating of 'low'.
- 26 • Wherever possible, reasoning was explained for the downgrade marks.

27

#### 28 2.4.13.1 Risk of bias

29 Risk of bias is assessed against standard criteria, depending on the study design.  
30 For randomised trials, we took into account: the adequacy of allocation  
31 concealment; blinding of participants for comparisons and outcomes susceptible  
32 to bias; attrition (missing data) and baseline comparability. A downgrade mark  
33 of –1 was given for inadequate or unclear allocation concealment and for a loss  
34 to follow-up of more than 20% in any one group or overall. Studies with more  
35 than 50% missing data were excluded from the analysis unless they were the  
36 only study, in which case they were given a downgrade mark of –2. If the

---

<sup>‡</sup> GRADE – Grading of Recommendations Assessment, Development and Evaluation

1 evidence was a meta-analysis of several studies, we took into consideration the  
2 proportion and weighting of higher risk studies, and in some instances carried out  
3 sensitivity analyses disregarding these studies and giving a separate rating for  
4 the new meta-analysis.

5

#### 6 **2.4.13.2 Inconsistency**

7 When several RCTs have widely differing estimates of treatment effect  
8 (heterogeneity or variability in results), the results are regarded as inconsistent.  
9 We defined this as a p-value for heterogeneity less than 0.1 and/or an I<sup>2</sup> value  
10 greater than 50%. Where this was the case, we gave a downgrade mark of -1.  
11 If the p-value was less than 0.1 and the I<sup>2</sup> value was greater than 80%, we  
12 gave a downgrade mark of -2. Where possible, we carried out pre-defined  
13 subgroup analyses to investigate heterogeneity and reported these results  
14 separately.

15

#### 16 **2.4.13.3 Indirectness**

17 Directness refers to the extent to which the population, interventions, comparisons  
18 and outcome measures are similar to those defined in the inclusion criteria for the  
19 reviews. Indirectness is only relevant if there is a compelling reason to expect  
20 important differences in the size of the effect. For example, many interventions  
21 have more or less the same relative effects across patient groups, so  
22 extrapolation is possible and reasonable. There are various types of indirectness  
23 that can be found in studies, but most relevant to this guideline are:

- 24 • When the setting is different from those of the guideline, e.g. community  
25 setting, rather than long-term care
- 26 • When the method for assessment of delirium is partly adequate or  
27 inadequate

28

#### 29 **2.4.13.4 Imprecision**

30 This is a rather subjective, but nevertheless important category. Evidence is  
31 considered to be imprecise if:

- 32 • There are sparse data (only a few events and they are uninformative).
- 33 • The confidence interval for the effect estimate is consistent with different  
34 conclusions, for example, both a clinically important effect (benefit or  
35 harm) and no clinically important effect; or the CI is consistent with  
36 important harms, no clinically important effect and important benefits.  
37 Precision requires the GDG to decide what are clinically important harms  
38 and benefits for that outcome measure. For dichotomous outcomes we  
39 used a relative risk reduction of 25% (RR of 1.25 or 0.75) to indicate the  
40 clinically important threshold. For continuous outcomes the GDG  
41 determined that the clinically important threshold for a difference  
42 between intervention groups was 0.5 days for a stay in ICU, 1 day for a

1 stay in hospital, 1 day for duration of delirium, and a change of 20% on  
2 any of the scales used (linearity assumed).

- 3 • If the confidence interval did not cross either of the clinically important  
4 thresholds (i.e. precise rating), the sample size was taken into  
5 consideration. If there was a power calculation for that outcome and  
6 comparison, it was used to decide if a study was 'small', otherwise the  
7 optimal information size was calculated (or 300 events total was  
8 assumed).

#### 10 **2.4.13.5 Reporting bias**

11 Reporting bias occurs in two main ways:

12 Publication bias, in which papers are more likely to be published if their results  
13 are statistically significant. The existence of publication bias in the studies in a  
14 meta-analysis can be investigated in a limited way using funnel plots, in which  
15 the standard error is plotted against the log odds ratio, the log relative risk or  
16 the mean difference. Asymmetry is indicative of reporting bias. This method is  
17 usually only useful when there are at least five studies. The GDG decided that  
18 industry sponsored studies should not be regarded as potentially biased.

### 20 **2.5 Economic literature reviewing process**

21 Information on cost-effectiveness is important for guideline development as it  
22 aids decision making on the application of intervention options in the different  
23 population groups considered in the guideline. It provides evidence on the cost  
24 and health impact of different intervention options considered during the process  
25 of guideline development. At the initial stage of the delirium guideline  
26 development, the health economist in conjunction with the GDG identified priority  
27 areas for cost-effectiveness evidence. The use of delirium prevention and  
28 treatment interventions in hospital and long-term care settings were identified as  
29 high priority areas for cost-effectiveness evidence. They were classified as high  
30 priority as the prevention and treatment of delirium would save NHS and PSS  
31 (Personal Social Services) resources as well as improve patients' health related  
32 quality of life. Information on the additional benefit associated with different  
33 strategies was also required. It was therefore necessary to look for health  
34 economic information on the intervention strategies and we started by reviewing  
35 published economic evaluations.

36  
37 A systematic review was carried out to identify and appraise existing published  
38 economic evaluations that are relevant to the guideline's clinical questions. An  
39 article had to present a full or partial economic evaluation to be included in this  
40 review. A full economic evaluation compares all relevant cost and patient  
41 outcomes and uses these to estimate a single measure of incremental cost and  
42 benefits. The different forms of economic evaluation include cost-effectiveness,  
43 cost-utility, cost-benefit or cost-minimisation analysis. A partial economic

1 evaluation only reports some of the relevant cost and patient outcomes. Studies  
2 reporting data from non-OECD (Organisation for Economic Co-operation and  
3 Development) member countries were excluded as these were felt to be less  
4 applicable to current practice in the UK. Publications that dealt with palliative  
5 care were removed as these were outside the scope of the guideline. For trial  
6 based economic evaluations, studies were excluded if they did not meet the  
7 inclusion criteria for the clinical effectiveness review.

8  
9 We initially searched Medline, Embase, NHSEED and HTA databases starting  
10 from 1994 to June 2008. An economics filter was applied to the Medline and  
11 Embase searches to identify relevant economic literature. The search terms used  
12 in Medline are given in Appendix J. The economics and quality of life filter is as  
13 listed in Appendix J. The terms were suitably adapted for searches in Embase,  
14 NHSEED and HTA. A total of 755 publications were sifted by the Health  
15 Economist. Sifting was done by reading the title and abstract of the publications  
16 and full papers were ordered for any potential economic evaluations. We  
17 ordered 12 publications (Bracco et al 2007, Pitkala et al 2008, Rizzo et al  
18 2001, Robinson et al 2002, The Medical and Health Research Council of the  
19 Netherlands [ongoing], Beaupre et al 2006, Heyman & Lombardo 1995, Caplan  
20 & Harper 2007, Pandharipande et al 2007, Rubin et al 2006, Webster et al  
21 1999, Caplan et al 2006) and four of them were reviewed (Bracco et al 2007,  
22 Pitkala et al 2008, Rizzo et al 2001, Robinson et al 2002). The outcomes of  
23 interest were intervention and non-intervention costs, the incidence and severity  
24 of delirium, incidence of complete recovery from delirium, Quality-adjusted life  
25 year (QALY) measure, and delirium-attributable mortality rate. The four papers  
26 reviewed (Bracco et al 2007, Pitkala et al 2008, Rizzo et al 2001, Robinson et  
27 al 2002) are described under the relevant clinical questions (Appendix J).

28 None of the identified economic evaluations were directly applicable to the  
29 guideline population. None of the studies assessed costs from a UK NHS and PSS  
30 perspective and none measured health benefits in QALYs. None of the studies  
31 discounted future costs and outcomes appropriately and none carried out a  
32 robust sensitivity analysis on the results of the economic analysis. We carried out  
33 update searches up to August 2009 but did not identify further relevant  
34 economic evaluation studies. As there was a lack of high quality, relevant  
35 evidence on the cost-effectiveness of the interventions included in the guideline, it  
36 became necessary to develop an original economic evaluation model to  
37 determine the cost-effectiveness of strategies for the prevention and treatment  
38 of delirium in different care settings.

## 39 40 **2.6 Cost-effectiveness modelling**

41 The details of the economic model are described in Appendix J.

42 We developed original models for intervention strategies in hospital care  
43 settings but could not develop any models for prevention and treatment  
44 strategies in the long-term care setting. This was because there was a lack of  
45 evidence from the long-term care setting which could be used to construct a cost-  
46 effectiveness model. The evidence on the adverse consequences of delirium came  
47 from studies that were carried out in the hospital setting (section 8). The efficacy

1 estimates of the interventions that we modelled came from studies carried out in  
2 hospital settings. Furthermore, the costing of the multi-component interventions  
3 was based on the assumption that they were applied in the hospital. We were  
4 not confident that we could use this evidence to model the cost-effectiveness of  
5 these interventions in long-term care setting.

6  
7 The outcomes of interest for the model were incremental cost and QALY gained.  
8 Costs were assessed from an NHS and PSS perspective. These outcomes were  
9 used to estimate the incremental cost-effectiveness ratio and net monetary  
10 benefit. Incremental net monetary benefit is defined below. Future costs and  
11 QALYs were discounted at a rate of 3.5% per annum. This is in line with the  
12 reference case advocated by NICE (NICE 2008 [manual on TA]).

13  
14 In the base case analysis, the cost effectiveness of an intervention was  
15 determined using the threshold, £20,000 per QALY, and all interventions were  
16 compared to the usual care. If an intervention strategy costs less than the  
17 comparator and generates greater benefit it is described as being dominant  
18 and is unequivocally cost-effective. If the intervention is more effective but more  
19 costly, the incremental cost per QALY is estimated and compared to the cost-  
20 effectiveness threshold of £20,000 to £30,000 per QALY in line with the  
21 principles stated in the NICE Technology Appraisal Manual (NICE 2008 [manual  
22 on TA]). Another alternative to using incremental cost and QALYs to estimate  
23 cost-effectiveness is the use of the Incremental Net Monetary Benefit (INMB). The  
24 INMB is the monetary value of an intervention compared to an alternative for a  
25 specific cost-effectiveness threshold. It is calculated as

26  
27  $\text{Cost-effectiveness Threshold} * \text{incremental QALY} - \text{incremental cost.}$

28  
29 An intervention is cost-effective if it has an INMB that is greater than zero.

30  
31 We constructed our model using the best available evidence and according to  
32 the NICE reference case for economic evaluation (NICE 2008 [manual on TA]).  
33 We described explicitly the assumptions made in the model as well as the  
34 uncertainties in the model input parameters. The results of the model were  
35 interpreted by the GDG bearing the assumptions in mind. We used deterministic  
36 and probabilistic sensitivity analyses to explore the impact of the assumptions  
37 and uncertainties on the model results. We discussed the limitations of the model.  
38 Further details on the cost-effectiveness model are given in chapter 16. For those  
39 clinical questions which were not prioritised for an original economic evaluation  
40 the GDG considered the likely cost-effectiveness of the interventions by making  
41 a qualitative judgement on the likely costs, health benefits and potential harms  
42 of interventions.

1

## 2 **2.7 Development of the recommendations**

3 Over the course of the guideline development process, the GDG was presented  
4 with the following:

- 5 • The clinical and economic evidence reviews. All evidence tables are in  
6 Appendices D, E and G.
- 7 • Forest plots of results from studies, including meta-analyses where  
8 appropriate.
- 9 • A description of the methods and results of the cost-effectiveness analysis  
10 (chapter 16).

11 Recommendations were drafted on the basis of this evidence whenever it was  
12 available.

13 When clinical and economic evidence was poor or absent, the GDG proposed  
14 recommendations based on their expert opinion.

15 The GDG also developed a care pathway algorithm according to the  
16 recommendations (see section 3.2).

17

## 18 **2.8 Research recommendations**

19 When areas were identified for which good evidence was lacking, the guideline  
20 development group considered making recommendations for future research.  
21 Decisions about inclusion were based on factors such as:

- 22 • the importance to patients or the population
- 23 • national priorities
- 24 • potential impact on the NHS and future NICE guidance
- 25 • ethical and technical feasibility

26

27 The GDG identified five high priority research recommendations (after discussion  
28 and voting). The full list of recommendations for future research, as well as those  
29 chosen as high priority, can be found in Appendix H.

30

## 31 **2.9 Prioritisation of recommendations for implementation**

32 To assist users of the guideline in deciding the order in which to implement the  
33 recommendations, the GDG identified ten key priorities for implementation. The  
34 decision was made after discussion and independent voting by the GDG. They  
35 selected recommendations that would:

- 1           • have a high impact on outcomes that are important to patients
- 2           • have a high impact on reducing variation in care and outcomes
- 3           • lead to a more efficient use of NHS resources
- 4           • promote patient choice
- 5           • promote equalities

6

7           In doing this the GDG also considered which recommendations were particularly  
8           likely to benefit from implementation support. They considered whether a  
9           recommendation:

- 10           • relates to an intervention that is not part of routine care
- 11           • requires changes in service delivery
- 12           • requires retraining staff or the development of new skills and competencies
- 13           • highlights the need for practice to change
- 14           • affects and needs to be implemented across various agencies or settings  
15            (complex interactions)
- 16           • may be viewed as potentially contentious, or difficult to implement for  
17            other reasons

18

## 19   **2.10 Validation of the guideline**

20           The first draft of this guideline was posted on the NICE website for an 8-week  
21           consultation between 11th November 2009 and 6<sup>th</sup> January 2010 and  
22           registered stakeholders were invited to comment. The GDG responded to  
23           comments and an amended version of the guideline was produced.

24

## 25   **2.11 Related NICE guidance**

26           NICE has developed/is developing the following guidance (details available  
27           from [www.nice.org.uk](http://www.nice.org.uk)), some of which has been referred to in this guideline:

- 28           • Acutely ill patients in hospital: recognition of and response to acute illness in  
29            adults in hospital. NICE clinical guideline 50 (2007). Available from  
30            [www.nice.org.uk/CG050](http://www.nice.org.uk/CG050).
- 31           • Infection control: prevention of healthcare-associated infection in primary  
32            and community care NICE clinical guideline 2 (2003). Available from  
33            [www.nice.org.uk/CG2](http://www.nice.org.uk/CG2).

- 1 • Nutrition support in adults: Nutrition support in adults: oral nutrition support,  
2 enteral tube feeding and parenteral nutrition. NICE clinical guideline 32  
3 (2006). Available from [www.nice.org.uk/CG032](http://www.nice.org.uk/CG032).
- 4 • Dementia: supporting people with dementia and their carers in health and  
5 social care. NICE clinical guideline 42 (2006). Available from  
6 [www.nice.org.uk/CG042](http://www.nice.org.uk/CG042).
- 7 • Drug misuse: opioid detoxification. NICE clinical guideline 52 (2007).  
8 Available from [www.nice.org.uk/CG0452](http://www.nice.org.uk/CG0452).
- 9 • Surgical site infection – prevention and treatment of surgical site infection.  
10 NICE clinical guideline 74 (2008). Available from  
11 [www.nice.org.uk/CG074](http://www.nice.org.uk/CG074).
- 12 • Schizophrenia – core interventions in the treatment and management of  
13 schizophrenia in primary and secondary care (update). NICE clinical  
14 guideline 82 (2009). Available from [www.nice.org.uk/CG082](http://www.nice.org.uk/CG082).
- 15 • Alzheimer's disease - donepezil, galantamine, rivastigmine (review) and  
16 memantine for the treatment of Alzheimer's disease. NICE technology  
17 appraisal 111 (2007). Available from [www.nice.org.uk/TA111](http://www.nice.org.uk/TA111).
- 18 • Schizophrenia - the clinical effectiveness and cost effectiveness of newer  
19 atypical antipsychotic drugs for schizophrenia. NICE technology appraisal  
20 43 (2002). Available from [www.nice.org.uk/TA43](http://www.nice.org.uk/TA43).
- 21 • Parkinson's disease – national clinical guideline for diagnosis and  
22 management in primary and secondary care. NICE clinical guideline 35  
23 (2006). Available from [www.nice.org.uk/CG035](http://www.nice.org.uk/CG035).
- 24 • Violence – the short-term management of disturbed/violent behaviour in in-  
25 patient psychiatric settings and emergency departments. NICE clinical  
26 guideline 25 (2005). Available from [www.nice.org.uk/CG025](http://www.nice.org.uk/CG025).
- 27 • Medicines adherence – involving patients in decisions about prescribed  
28 medicines and supporting adherence. NICE clinical guideline 76 (2009).  
29 Available from [www.nice.org.uk/CG076](http://www.nice.org.uk/CG076).
- 30 • Alcohol use disorders in adults and young people: clinical management.  
31 NICE clinical guideline. Publication expected May 2010.
- 32 • Alcohol dependence and harmful alcohol use. NICE clinical guideline.  
33 Publication expected January 2011.

34  
35

## 36 2.12 Updating the guideline

37 This guideline will be updated when appropriate. The decision to update will  
38 balance the need to reflect changes in the evidence against the need for  
39 stability, as frequent changes to the recommendations would make  
40 implementation difficult. We check for new evidence 2 and 4 years after  
41 publication, to decide whether all or part of the guideline should be updated. In  
42 exceptional circumstances, if important new evidence is published at other times,

1 we may conduct a more rapid update of some recommendations. Any update  
2 will follow the methodology outlined in the NICE guidelines manual (NICE 2009).

3

## 3 Summary of recommendations

Below are the recommendations that the GDG selected as the key priorities for implementation followed by the algorithm. The full list of guideline recommendations can be found in chapter 4 and the full list of recommendations for future research can be found in Appendix H.

6

### 3.1 Key priorities for implementation

The GDG identified ten key priorities for implementation. The decision was made after discussion and voting by the GDG. The recommendations chosen by the GDG as key priorities for implementation are listed below. The numbering of the recommendations in parentheses is as per the NICE version of the guideline.

In addition the GDG wanted to highlight the importance of being aware of delirium and its consequences and so a prominent statement has been included below.

#### Awareness of delirium and its consequences

Be aware that people in hospital or long-term care may be at risk of delirium, which can have serious consequences (such as increased risk of dementia and/or death) and, for people in hospital, may increase their risk of new admission to long-term care.

20

#### 3.1.1 Risk factor assessment

• When people first present to hospital or long-term care, assess them for the following risk factors:

- Age 65 years or older.
- Cognitive impairment: a previous history of cognitive impairment or, if cognitive impairment is suspected, confirm it using a standardised and validated cognitive impairment measure.
- Current hip fracture.
- Severe illness (a clinical condition that is deteriorating or is at risk of deterioration)<sup>1</sup>.

If any of these risk factors is present, the person is considered at risk of delirium.  
[1.1.1]

---

<sup>1</sup> For further information on recognising and responding to acute illness in adults in hospital see 'Acutely ill patients in hospital' (NICE clinical guideline CG50).

1

2 **3.1.2 Indicators of prevalent delirium**

3 • At presentation, assess people at risk for indicators of delirium, which are  
4 sudden changes or fluctuations in usual behaviour. These may be  
5 reported by the person at risk, or a carer or relative. The changes may  
6 be in any of the following:

- 7 ○ cognitive function: for example, worsened concentration, slow  
8 responses, confusion
- 9 ○ perception: for example, visual or auditory hallucinations
- 10 ○ physical function: for example, reduced mobility, reduced  
11 movement, restlessness, agitation, changes in appetite, sleep  
12 disturbance
- 13 ○ social behaviour: for example, poor cooperation, withdrawal, or  
14 alterations in communication, mood and/or attitude.

15 If any of these indicators is present, a healthcare professional who is  
16 trained and competent in the diagnosis of delirium should carry out a  
17 clinical assessment to confirm the diagnosis. [1.2.1]

18

19 **3.1.3 Interventions to prevent delirium**

- 20 • Ensure that people at risk of delirium have a care environment that:
  - 21 ○ avoids unnecessary room changes
  - 22 ○ maintains a team of healthcare professionals who are familiar to  
23 the person at risk. [1.3.1]
- 24 • Within 24 hours of admission, assess people at risk for clinical indicators  
25 contributing to delirium (recommendations 1.3.3.1–1.3.3.9). Based on this  
26 assessment, provide a multicomponent intervention package tailored to  
27 the person's individual needs and care setting. [1.3.2]
- 28 • The tailored multicomponent intervention package should be delivered by a  
29 multidisciplinary team trained and competent in delirium prevention. The  
30 tailored package should address the clinical indicators in  
31 recommendations 1.3.3.1–1.3.3.9. [1.3.3]

32

33 **3.1.4 Diagnosis (specialist clinical assessment)**

- 34 • If indicators of delirium are identified, carry out a clinical assessment using  
35 the Diagnostic and Statistical Manual of Mental Disorders (DSM-IV)  
36 criteria or short Confusion Assessment Method (short CAM). In critical care  
37 or in the recovery room after surgery, CAM-ICU should be used. A  
38 healthcare professional who is trained and competent in the diagnosis of  
39 delirium should carry out the assessment. [1.5.1]

- 1                   • Ensure that the diagnosis of delirium is documented in the person's  
2                   healthcare record. [1.5.2]

3

#### 4   **3.1.5 Non-pharmacological interventions**

- 5                   • In people diagnosed with delirium, identify and manage the possible  
6                   underlying cause or combination of causes. [1.6.1]
- 7                   • Ensure effective communication and reorientation and provide reassurance  
8                   for people diagnosed with delirium. Family, friends and carers may be  
9                   able to help with this. [1.6.2]

10

#### 11   **3.1.6 Pharmacological interventions**

- 12                   • If non-pharmacological approaches are ineffective, consider giving short-  
13                   term (for 1 week or less) haloperidol<sup>2</sup> or olanzapine<sup>2</sup> if people with  
14                   delirium are distressed or a risk to themselves or others. [1.6.4]

15

16

17

18

19

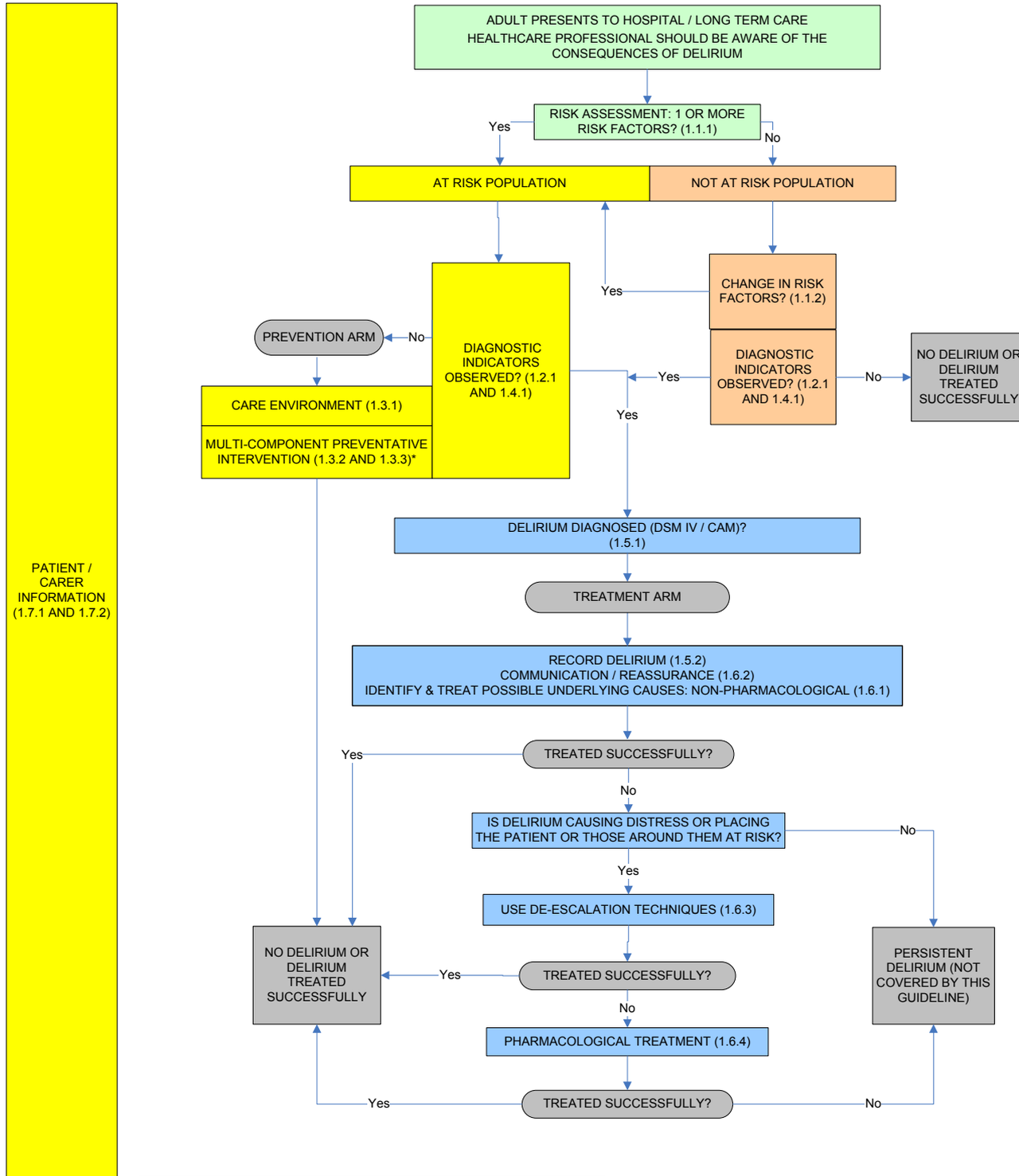
---

<sup>2</sup> Haloperidol and olanzapine do not have UK marketing authorisation for this indication.

1

2 3.2 Algorithm (link to full recommendations)

3



4 5

Clinical indicators that can contribute to delirium	Preventative interventions and actions
Disorientation	1.3.3.1 Address reorientation through the following actions: <ul style="list-style-type: none"> <li>• Provide clear signage, soft lighting, a 24-hour clock and a calendar, all easily visible to the person at risk.</li> <li>• Introduce cognitively stimulating activities (for example, structured reminiscence) and reorienting communication.</li> <li>• Facilitate regular visits from family and friends.</li> </ul>
Dehydration and/or constipation	1.3.3.2 Address dehydration and/or constipation through the following actions: <ul style="list-style-type: none"> <li>• Ensure adequate fluid intake to prevent dehydration by encouraging the person to drink. Consider offering subcutaneous or intravenous fluids if necessary.</li> <li>• Take advice where necessary when managing fluid balance in people with co-morbidities (for example heart failure or chronic kidney disease).</li> </ul>
Infection	1.3.2.2 Address problems with infection through the following actions: <ul style="list-style-type: none"> <li>• Look for and treat infection.</li> <li>• Avoid unnecessary catheterisation.</li> <li>• Implement good infection control procedures in line with 'Infection control' (NICE clinical guideline CG2).</li> </ul>
Pain	1.3.3.4 Address problems with pain through the following actions: <ul style="list-style-type: none"> <li>• Find out whether the person has pain.</li> <li>• Look for non-verbal signs of pain, particularly in those with communication difficulties (for example, people with learning difficulties or dementia, or people on a ventilator or who have a tracheotomy).</li> <li>• If people have been prescribed pain relief, ensure they receive it.</li> </ul>
Polypharmacy effects	1.3.3.5 Address problems with polypharmacy effects through the following actions: <ul style="list-style-type: none"> <li>• Carry out a drug review for people taking multiple medications in line with 'Medicines adherence' (NICE clinical guideline CG76).</li> </ul>
Poor nutrition and/or constipation	1.3.3.6 Address problems with poor nutrition and/or constipation through the following actions: <ul style="list-style-type: none"> <li>• Follow the advice given on nutrition in 'Nutrition support in adults' (NICE clinical guideline CG32).</li> <li>• If people have dentures, ensure they are well fitting.</li> </ul>
Restricted or limited mobility or immobility	1.3.3.7 Address problems with restricted or limited mobility or immobility through the following actions: <ul style="list-style-type: none"> <li>• Encourage people to:               <ul style="list-style-type: none"> <li>○ walk around</li> <li>○ carry out active range-of-motion exercises, <b>and</b></li> <li>○ mobilise early after surgery.</li> </ul> </li> </ul>
Sensory impairment	1.3.3.8 Address problems with sensory impairment through the following actions: <ul style="list-style-type: none"> <li>• Ensure hearing and visual aids are available to and used by people who need them, and that they are in good working order.</li> </ul>
Sleep disturbance	1.3.3.9 Address problems with sleep disturbance through the following actions: <ul style="list-style-type: none"> <li>• Promote good sleep patterns and sleep hygiene by:</li> <li>• scheduling medication rounds to avoid disturbing sleep, <b>and</b></li> <li>• reducing noise to a minimum during sleep periods.</li> </ul> For more information on good sleep hygiene, see also 'Parkinson's disease' (NICE clinical guideline CG35).

## 4 Recommendations and evidence to recommendations

### 4A. Full list of guideline recommendations

The numbering of the recommendations in parentheses is as per the NICE version of the guideline.

#### 4.1 Awareness of delirium and its consequences

Be aware that people in hospital or long-term care may be at risk of delirium, which can have serious consequences (such as increased risk of dementia and/or death) and, for people in hospital, may increase their risk of new admission to long-term care.

#### 4.2 Risk factor assessment

- When people first present to hospital or long-term care, assess them for the following risk factors:

- Age 65 years or older.
- Cognitive impairment: a previous history of cognitive impairment or, if cognitive impairment is suspected, confirm it using a standardised and validated cognitive impairment measure.
- Current hip fracture.
- Severe illness (a clinical condition that is deteriorating or is at risk of deterioration)<sup>3</sup>.

If any of these risk factors is present, the person is considered at risk of delirium. [1.1.1]

- Observe people at every opportunity for any changes in the risk factors for delirium. [1.1.2]

---

<sup>3</sup> For further information on recognising and responding to acute illness in adults in hospital see 'Acutely ill patients in hospital' (NICE clinical guideline CG50).

1

## 2 4.3 Indicators of prevalent delirium

3 • At presentation, assess people at risk for indicators of delirium, which are  
4 sudden changes or fluctuations in usual behaviour. These may be  
5 reported by the person at risk, or a carer or relative. The changes may  
6 be in any of the following:

- 7 ○ cognitive function: for example, worsened concentration, slow  
8 responses, confusion
- 9 ○ perception: for example, visual or auditory hallucinations
- 10 ○ physical function: for example, reduced mobility, reduced  
11 movement, restlessness, agitation, changes in appetite, sleep  
12 disturbance
- 13 ○ social behaviour: for example, poor cooperation, withdrawal, or  
14 alterations in communication, mood and/or attitude.

15 If any of these indicators is present, a healthcare professional who is  
16 trained and competent in the diagnosis of delirium should carry out a  
17 clinical assessment to confirm the diagnosis. [1.2.1]

18

## 19 4.4 Interventions to prevent delirium

20 • Ensure that people at risk of delirium have a care environment that:  
21 ○ avoids unnecessary room changes  
22 ○ maintains a team of healthcare professionals who are familiar to  
23 the person at risk. [1.3.1]

24

25 • Within 24 hours of admission, assess people at risk for clinical indicators  
26 contributing to delirium (recommendations 1.3.3.1–1.3.3.9). Based on this  
27 assessment, provide a multicomponent intervention package tailored to  
28 the person's individual needs and care setting. [1.3.2]

29

30 • The tailored multicomponent intervention package should be delivered by a  
31 multidisciplinary team trained and competent in delirium prevention. The  
32 tailored package should address the clinical indicators in  
33 recommendations 1.3.3.1–1.3.3.9. [1.3.3]

34

### 35 Disorientation

36 [1.3.3.1] Address reorientation through the following actions:

- 37 ○ Provide clear signage, soft lighting, a 24-hour clock and a  
38 calendar, all easily visible to the person at risk.

- 1                   ○ Introduce cognitively stimulating activities (for example, structured
- 2                   reminiscence) and reorienting communication.
- 3                   ○ Facilitate regular visits from family and friends.

#### 4 5                   Dehydration and/or constipation

6                   **[1.3.3.2]** Address dehydration and/or constipation through the following

7                   actions:

- 8                   ○ Ensure adequate fluid intake to prevent dehydration by
- 9                   encouraging the person to drink. Consider offering subcutaneous
- 10                  or intravenous fluids if necessary.
- 11                  ○ Take advice where necessary when managing fluid balance in
- 12                  people with comorbidities (for example heart failure or chronic
- 13                  kidney disease).

#### 14 15                  Infection

16                  **[1.3.3.3]** Address problems with infection through the following actions:

- 17                  ○ Look for and treat infection.
- 18                  ○ Avoid unnecessary catheterisation.
- 19                  ○ Implement good infection control procedures in line with 'Infection
- 20                  control' (NICE clinical guideline CG2).

#### 21 22                  Pain

23                  **[1.3.3.4]** Address problems with pain through the following actions:

- 24                  ○ Find out whether the person has pain.
- 25                  ○ Look for non-verbal signs of pain, particularly in those with
- 26                  communication difficulties (for example, people with learning
- 27                  difficulties or dementia, or people on a ventilator or who have a
- 28                  tracheotomy).
- 29                  ○ If people have been prescribed pain relief, ensure they receive it.

#### 30 31                  Polypharmacy effects

32                  **[1.3.3.5]** Address problems with polypharmacy effects through the

33                  following actions:

- 34                  ○ Carry out a drug review for people taking multiple drugs in line
- 35                  with 'Medicines adherence' (NICE clinical guideline CG76).

#### 36 37                  Poor nutrition and/or constipation

38                  **[1.3.3.6]** Address problems with poor nutrition and/or constipation

39                  through the following actions:

- Follow the advice given on nutrition in ‘Nutrition support in adults’ (NICE clinical guideline CG32).
- If people have dentures, ensure they are well fitting.

#### Restricted or limited mobility or immobility

[1.3.3.7] Address problems with restricted or limited mobility or immobility through the following actions:

- Encourage people to:
  - walk around
  - carry out active range-of-motion exercises, **and**
  - mobilise early after surgery.

#### Sensory impairment

[1.3.3.8] Address problems with sensory impairment through the following actions:

- Ensure hearing and visual aids are available to and used by people who need them, and that they are in good working order.

#### Sleep disturbance

[1.3.3.9] Address problems with sleep disturbance through the following actions:

- Promote good sleep patterns and sleep hygiene by:
  - scheduling medication rounds to avoid disturbing sleep, **and**
  - reducing noise to a minimum during sleep periods.

For more information on good sleep hygiene, see also ‘Parkinson’s disease’ (NICE clinical guideline CG35).

## **4.5 Indicators: daily observations (all people in hospital or long-term care)**

- Observe at least daily, all people in hospital or long-term care for indicators of delirium, which are sudden changes or fluctuations in usual behaviour (see recommendation 1.2.1).  
If any of these indicators is present, a healthcare professional who is trained and competent in the diagnosis of delirium should carry out a clinical assessment to confirm the diagnosis. [1.4.1]

## 1 **4.6 Diagnosis (specialist clinical assessment)**

2           • If indicators of delirium are identified, carry out a clinical assessment using  
3           the Diagnostic and Statistical Manual of Mental Disorders (DSM-IV)  
4           criteria or short Confusion Assessment Method (short CAM). In critical care  
5           or in the recovery room after surgery, CAM-ICU should be used. A  
6           healthcare professional who is trained and competent in the diagnosis of  
7           delirium should carry out the assessment. [1.5.1]

8

9           • Ensure that the diagnosis of delirium is documented in the person's  
10          healthcare record. [1.5.2]

11

## 12 **4.7 Treatment of delirium**

### 13 **4.7.1 Non-pharmacological interventions**

14           • In people diagnosed with delirium, identify and manage the possible  
15          underlying cause or combination of causes. [1.6.1]

16

17           • Ensure effective communication and reorientation and provide reassurance  
18          for people diagnosed with delirium. Family, friends and carers may be  
19          able to help with this. [1.6.2]

20

21           • If the person with delirium is distressed or a risk to themselves or others, first  
22          use verbal and non-verbal techniques to de-escalate the situation before  
23          considering pharmacological interventions. For more information on de-  
24          escalation techniques, see 'Violence' (NICE clinical guideline 25). [1.6.3]

25

### 26 **4.7.2 Pharmacological interventions**

27           • If non-pharmacological approaches are ineffective, consider giving short-  
28          term (for 1 week or less) haloperidol<sup>4</sup> or olanzapine<sup>4</sup> if people with  
29          delirium are distressed or a risk to themselves or others. [1.6.4]

30

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<sup>4</sup> Haloperidol and olanzapine do not have UK marketing authorisation for this indication.

## 1 4.8 Information giving and support

- 2 • Offer information to people who are at risk of delirium or who have  
3 delirium, and their family and/or carers, which:
  - 4 ○ describes people's experience of delirium
  - 5 ○ informs them that the experience of delirium is common and is  
6 usually temporary
  - 7 ○ encourages people at risk and their families and/or carers to tell  
8 their healthcare team about any sudden changes or fluctuations in  
9 usual behaviour
  - 10 ○ encourages the person with delirium to share their experiences  
11 during recovery with the healthcare professional. [1.7.1]
- 12 • Ensure that information provision meets the cultural, linguistic, cognitive and  
13 language needs of the person. [1.7.2]
- 14
- 15
- 16

## 17 4B. Evidence to recommendations

18

### 19 4.9 Risk factor assessment (recommendations 1.1.1 and 1.1.2)

20

#### 21 4.9.1 Quality of evidence

22 There was moderate or low quality evidence from the risk factors review for  
23 each of 20 risk factors for the incidence of delirium, and limited evidence for the  
24 duration, severity and persistence of delirium. The GDG separated the evidence  
25 into three categories: those risk factors for which the GDG had some confidence  
26 in the evidence, those for which it had slight confidence and those for which there  
27 was inconsistency or uncertainty. The risk factors in which the GDG had some  
28 confidence were:

- 29 • Age as a continuous variable
- 30 • Age over 65 years
- 31 • Age over 80 years
- 32 • Cognitive impairment
- 33 • Vision impairment
- 34 • Illness severity using the APACHE II as a continuous variable
- 35 • Fracture on admission
- 36 • Infection

1           • Physical restraint

2           These risk factors are of two types, those that can be modified (e.g. infection)  
3           and those that are not modifiable (e.g. age). The magnitude of the independent  
4           modifiable risk factors ranged from an odds ratio of around 1.7 (visual  
5           impairment) to around 3.0 (infection). The magnitude of the independent non-  
6           modifiable risk factors ranged from about 3.0 (age over 65 years) to about 6.6  
7           (fracture).

8

9           **4.9.2 GDG considerations**

10           The GDG wished to define an at-risk group of people, who would be targeted  
11           to receive the multicomponent preventative intervention (section 4.11).

12

13           The GDG recognised that the multicomponent interventions addressed modifiable  
14           risk factors only. There was no expectation that the incidence of delirium would  
15           be reduced for people who did not have any modifiable risk factors. In defining  
16           the at-risk group, the GDG considered which risk factors were important. People  
17           who had non-modifiable risk factors for delirium had a higher baseline risk, and  
18           the additional presence of a modifiable risk factor would raise the risk of  
19           developing delirium. For example, one person with no risk factors might have a  
20           baseline risk of 5%, and another person aged 75 years with a hip fracture  
21           might have a 50% risk of delirium. If the two people also had an infection (e.g.  
22           with a relative risk of 2), the risks of delirium would be 10% and 100% for the  
23           two cases.

24

25           Taking these factors into consideration, and that the clinical and cost-  
26           effectiveness evidence only applied to people at intermediate and higher risk of  
27           delirium, the GDG concluded that the intervention(s) should not be offered to  
28           everyone in hospital or long-term care, and that non-modifiable risk factors  
29           should be used to define the 'at-risk' group. The modifiable risk factors would  
30           then be addressed by the multicomponent intervention.

31

32           The GDG, therefore, decided not to include visual impairment, infection and  
33           physical restraint in the definition of the at-risk group; infection and visual  
34           impairment are covered by the multicomponent intervention. The evidence  
35           pertaining to physical restraint as a risk factor for the severity and persistent  
36           delirium was low and moderate quality. The GDG noted that restraint is  
37           sometimes used in patients with delirium to prevent them causing harm to  
38           themselves, for example, self-extubation in ICU. In addition, restraint can  
39           indirectly result from medical interventions, for example, by intravenous infusions  
40           reducing people's ability to mobilise. The GDG therefore decided against  
41           including restraint as a risk factor as part of the multicomponent intervention.

1

2 In formulating the recommendations, the GDG considered the following points for  
3 the non-modifiable risk factors:

4 • Age: a cut-off point of 65 years; this decision was based on the weight of  
5 evidence from the risk factors review, in particular the evidence from one  
6 moderate quality study (Pandharipande 2006), which demonstrated 65  
7 years as a clear cut off point. From the evidence on age as a continuous  
8 variable, the GDG were confident that increasing age (above age 65  
9 years) increases the risk of delirium.

10 • Cognitive impairment: the GDG emphasised that either a known history  
11 should be ascertained, or that suspected cognitive impairment should be  
12 confirmed with a validated measure.

13 • Current hip fracture: there was moderate quality evidence for 'fracture on  
14 admission' as a risk factor (fracture type unspecified) and low quality  
15 evidence for emergency hip fracture surgery in comparison with elective  
16 surgery for hip or knee arthritis. The GDG consensus was that the risk  
17 factor should be 'current hip fracture'

18 • Illness severity: the GDG debated the appropriate measure that should be  
19 used to measure illness severity. It was agreed to cross refer in the  
20 recommendation to the NICE guideline on acutely ill patients in hospital;  
21 and to state that, in a hospital setting.

22

23 The risk factor review evidence did not find any studies conducted solely in the  
24 long-term care settings, but the GDG agreed that the same risk factors would be  
25 applicable regardless of the setting.

26

27 The GDG discussed when people should be assessed for risk factors, and agreed  
28 that this should be conducted when the person presents to hospital or long-term  
29 care setting.

30

31 The GDG recognised that during the course of a hospital stay or long-term care,  
32 there might be a change in the risk factors for delirium in the group previously  
33 defined as not at risk, particularly in terms of illness severity. The GDG therefore  
34 added recommendation 1.1.2 covering risk factors developing subsequently to  
35 the initial presentation.

36

## 1 **4.10 Interventions to prevent delirium: care environment**

### 2 **(recommendation 1.3.1)**

3

#### 4 **4.10.1 Quality of evidence**

5 For environmental risk factors there was low quality evidence from the risk  
6 factors review pertaining to the severity of delirium and no evidence relating to  
7 the incidence of delirium. The GDG extrapolated the evidence to cover the  
8 incidence of delirium and added to it from their experience, referring to some of  
9 the multicomponent prevention studies. There was no economic evidence for this  
10 recommendation.

11

#### 12 **4.10.2 GDG considerations (strong agreement)**

13 Frequent changes in surroundings, of both room and people, may contribute to  
14 feelings of disorientation and delirium, and with frequent changes of staff,  
15 information may be lost. The GDG recognised that trying to reduce the number  
16 of room moves may conflict with service provision and operational factors for  
17 example assessment wards, single sex wards and that delirium in itself may  
18 trigger for a patient being moved to a side ward.

19 Factors related to reorientation can help towards minimising risk, this included use  
20 of a 24hour clock. This was included in the recommendation addressing  
21 disorientation as part of the multicomponent intervention package.

22

## 23 **4.11 Interventions to reduce the risk of delirium: non-pharmacological** 24 **intervention (recommendations 1.3.2, 1.3.3 and 1.3.3.1-1.3.3.9)**

25

#### 26 **4.11.1 Quality of evidence**

27 Recommendations 1.3.2–1.3.3 derive from moderate and high (Inouye 1999 and  
28 Marcantonio 2001) and low quality evidence from the multicomponent  
29 prevention review for patients in hospital (primary evidence source), supported  
30 by mixed quality evidence from the non- pharmacological risk factors review,  
31 low quality evidence from the hydration review, moderate quality evidence from  
32 the pharmacological risk factors review and GDG consensus. The latter was also  
33 informed by three other NICE guidelines.

34 Economic evidence for the hospital setting was obtained by modelling the  
35 preventative pathway and was informed by the evidence from the  
36 multicomponent prevention review, and the review on the consequences of

1 delirium. It was also informed by evidence on cost, quality of life and baseline  
2 risks.

3 There was no clinical or cost-effectiveness evidence for the long-term care  
4 population, and recommendations for this setting were based on indirect  
5 evidence from the hospital population.

6

#### 7 **4.11.2 GDG considerations: multicomponent interventions in a hospital setting** 8 **for the prevention of delirium (strong agreement)**

9 The evidence from two studies was of moderate and high quality (Inouye 1999  
10 and Marcantonio 2001). Each of the multicomponent interventions (and not each  
11 study) were incorporated into the economic model (using the same risk profiles as  
12 those described in the studies) and was found to be cost effective. There was a  
13 degree of uncertainty around the cost-effectiveness estimates, but this uncertainty  
14 was not judged by the GDG to be sufficient enough to affect the general  
15 conclusion.

16

17 One of the components in recommendation 1.3.3 is hydration (recommendation  
18 1.3.3.2), and the GDG considered the merits of developing a stand-alone good  
19 practice recommendation on hydration for all patients in hospital or long-term  
20 care. In addition to the evidence review, the GDG considered further information  
21 about which they were aware on hydration in the long-term care setting, which  
22 suggested an improvement in the well-being of the residents when a drinking-  
23 water regimen was implemented, although there was no control for comparison.  
24 On balance they decided that the evidence base was weak and a stand-alone  
25 recommendation might dilute the importance of other factors, for example  
26 infection. It was agreed that strategies for hydration would be captured in the  
27 multicomponent prevention intervention.

28

29 The GDG discussed whether the preventative intervention should be given to all  
30 patients, or only to those at risk of delirium, or whether to carry out sensitivity  
31 analyses to determine separately the cost effectiveness for intermediate and  
32 high risk groups. They concluded that the recommendation should be restricted to  
33 patients who are at-risk of delirium, but that healthcare professionals should be  
34 encouraged to give the intervention to all patients in that category. They defined  
35 the at-risk group according to the risk factors review (see section 4.9).

36

37 The GDG recognised that the initial stage of the multicomponent intervention was  
38 assessment of the patient's needs, and a recommendation was made for  
39 multicomponent intervention interventions that are tailored to individual needs.  
40 Both of the higher quality intervention studies (Inouye 1999 and Marcantonio  
41 2001) included this initial assessment stage, and the GDG agreed this was very  
42 important. The GDG also concurred with the evidence from the Marcantonio  
43 (2001) study, that this assessment should be made within 24 hours of admission.

1

2 In line with evidence from the Inouye (1999) study, the GDG agreed that a  
3 multidisciplinary team should carry out the multicomponent intervention, and  
4 considered it important that the healthcare team members concerned should be  
5 trained and competent in carrying out these tasks.

6

7 The GDG discussed whether to recommend one or both of the multicomponent  
8 intervention 'packages' (described by the two reviewed studies) or whether to  
9 produce a more general recommendation that selected individual elements from  
10 each package, together with evidence from the other reviews.

11

12 The GDG concluded that the latter course of action should be taken and that the  
13 two packages could be used to make a broad recommendation since the studies  
14 showed that when risk factors were addressed by providing better quality care,  
15 outcomes were improved. Hence the studies were deemed by the GDG to be  
16 'proof of concept' studies.

17

18 The GDG discussed which clinical indicators should be addressed by the  
19 multicomponent interventions, and the final list was based upon the available  
20 evidence and GDG clinical expertise. Each indicator that was included, and the  
21 evidence for them is listed below:

22

- 23 • Disorientation – evidence from the Inouye (1999) study and the non-  
24 pharmacological risk factors review
- 25 • Dehydration / constipation – evidence from the Inouye (1999) and  
26 Marcantonio (2001) studies, from the hydration review and from GDG  
27 expertise
- 28 • Infection – evidence from the Marcantonio (2001) study, the non-  
29 pharmacological risk factors review and GDG expertise; cross reference  
30 to the NICE Infection Control guideline. For catheterisation evidence  
31 came from the Marcantonio (2001) and Inouye (1999) studies and the  
32 non-pharmacological risk factors review, and GDG clinical expertise
- 33 • Pain – evidence from the Marcantonio (2001) study, indirect evidence from  
34 the pharmacological risk factors review and GDG expertise. The GDG  
35 emphasised that both verbal and non verbal signs of pain should be  
36 assessed, particularly in patients with dementia or learning difficulties.
- 37 • Polypharmacy effects - evidence from the Marcantonio (2001) study, from  
38 the non-pharmacological risk factors review and GDG expertise. The  
39 GDG advised recommending a drug review that addressed the type of  
40 drugs as well as the number; the GDG also supported the principle that if  
41 clinicians add a new drug, another should be taken away.

- 1 • Poor nutrition / constipation - some evidence from the Marcantonio (2001)  
2 study and from lower quality multicomponent prevention studies, and  
3 GDG expertise; cross reference to the NICE nutrition guideline
- 4 • Restricted / limited mobility or immobility – evidence from the Marcantonio  
5 (2001) and Inouye (1999) studies
- 6 • Sensory impairment – evidence from the Inouye (1999) and Marcantonio  
7 (1999) studies, and from the non-pharmacological risk factors review for  
8 visual impairment
- 9 • Sleep disturbance – evidence from the Inouye (1999) study and GDG  
10 clinical expertise; cross reference to the NICE Parkinson’s Disease  
11 guideline. Although the GDG considered it important that patients slept  
12 well in hospital, they decided to exclude the use of sleep enhancers  
13 (which was part of the Inouye (1999) study intervention) because  
14 evidence from the pharmacological risk factors review suggested that the  
15 drugs may also cause delirium

#### 17 **4.11.3 GDG considerations: multicomponent interventions in the long-term** 18 **care setting for the prevention of delirium**

19 There was no evidence for multicomponent preventative interventions in a long-  
20 term care setting, and very limited evidence for the consequences of delirium.  
21 Clinical effectiveness was therefore extrapolated from the hospital setting and  
22 GDG experience. Health economic modelling was not carried out because there  
23 was a lack of data for this setting and a large number of assumptions would  
24 have had to be made by the GDG, leading to serious uncertainty in outcomes.  
25 GDG consensus was that a multicomponent intervention for long-term care could  
26 have massive potential cost-savings for the NHS, was unlikely to do any harm to  
27 patients, and could probably be fairly easily accommodated within current care  
28 without incurring high costs. Therefore, they decided to recommend that the  
29 multicomponent intervention package should be tailored to the care setting, and  
30 that further research should be carried out. This led to writing a research  
31 recommendation (see Appendix H). The GDG considered it important that the  
32 care staff concerned should be trained and competent in carrying out these  
33 tasks.

34  
35 The GDG noted that some of the low quality multicomponent prevention studies  
36 examined the effectiveness of an educational intervention for staff. The GDG  
37 felt that this showed some potential, not least in the prevention of delirium  
38 resulting from increased staff awareness and this is reflected in a research  
39 recommendation (see Appendix H).

40

## 4.12 Diagnosis (recommendations 1.2.1, 1.4.1 and 1.5.1)

### 4.12.1 Quality of evidence

Two stages in the diagnostic process are identified: an initial screening stage and a confirmation stage. In the absence of evidence, the first stage comprises GDG consensus recommendations, with strong agreement, that were partly informed by the standard operational definition of delirium (the DSM criteria) and partly by GDG clinical experience. For the second stage, there was moderate to low quality evidence from the review of diagnostic test accuracy for different tests, comparing them with the reference standard of the DSM IV criteria. This review and the epidemiology review also compared different criteria over the years that have been developed as the standard operational definition for delirium.

### 4.12.2 GDG considerations – 1<sup>st</sup> stage (recommendations 1.2.1 and 1.4.1)

The initial screening stage is intended to alert any healthcare professional, including the non-specialist, to warning signs that the patient has, or is developing, delirium.

The GDG debated when would be an appropriate time to carry out the initial stage, and considered completing the initial assessment at the person's first presentation to hospital or long-term care. This would mean that all patients presenting to the accident and emergency department would have to undergo the test and the GDG considered this impractical in an accident and emergency setting. Therefore, they decided that only people who had already been determined to be at-risk of delirium (see recommendation 1.1.1) should be screened for prevalent delirium (recommendation 1.2.1), and that all people who were 'in hospital' (i.e. admitted) or in long-term care should subsequently be observed for signs of delirium (recommendation 1.4.1). This group included both those initially determined to be 'at-risk' and those determined to be not at-risk.

The GDG considered using a simple validated diagnostic tool such as the clock drawing test and MMSE, but noted from the evidence that these tools had low sensitivity. The GDG was keen that screening for delirium was based upon clinical signs and symptoms that could be easily identified by the non-specialist. The GDG noted that warning signs are sudden changes or fluctuations in usual behaviour of the hospital patient or person in long-term care, and compiled a list of clinical indicators based on their clinical experience. It was noted that it is often the case that the patient or their family or carer notice and report changes in behaviour, which would otherwise be unnoticed by the healthcare professional. The GDG decided to emphasise and include this in the recommendation.

1 **4.12.3 GDG considerations – 2nd stage (recommendation 1.5.1)**

2 The GDG considered whether to use the DSM IV diagnostic criteria for delirium,  
3 noting that this should be applied by a trained healthcare professional, or  
4 whether to recommend a diagnostic test. The GDG concluded that it was  
5 important to give people the option to use either DSM IV or a diagnostic test.  
6 The tests examined in the review of diagnostic test accuracy showed that both  
7 the long and short versions of the CAM, CAM-ICU and the AMT, all had  
8 acceptable sensitivity. The GDG noted that the long version of the CAM was not  
9 used in clinical practice and serial tests (such as AMT and MMSE) may be  
10 considered for those under elective care, but have limited clinical utility in  
11 relation to patients with a high risk of delirium. The GDG therefore decided the  
12 short version of CAM and CAM-ICU should be recommended as alternatives to  
13 DSM IV.

14

15 The GDG noted the evidence from one moderate quality study (Radtko 2008)  
16 that CAM had only 43% sensitivity for diagnosing delirium in a population that  
17 was in the recovery room following surgery. The GDG considered this to be an  
18 inappropriate test for this population and agreed to recommend using the CAM-  
19 ICU in critical care or in they recovery room following surgery.

20

21 **4.13 Recording delirium, awareness of and general consequences of**  
22 **delirium (recommendation 1.5.2)**

23

24 **4.13.1 Quality of evidence**

25 There was low and moderate quality evidence from the consequences of delirium  
26 review for patients in hospital, but no evidence for the consequences of delirium  
27 in long-term care. Reference was also made to the epidemiology review.

28

29 **4.13.2 GDG considerations**

30 The GDG noted from the epidemiological review, that there was widespread  
31 occurrence of delirium throughout the healthcare system but it was poorly  
32 reported. Moreover, the GDG observed that, in their experience, healthcare  
33 professionals were often unaware of the possibility that delirium might occur. The  
34 GDG thought that the slogan, “Think Delirium” summarised their rationale for this  
35 recommendation (1.5.2). The GDG wished to reinforce the importance of  
36 accurately recording delirium by making a recommendation on coding  
37 (recommendation 1.5.2).

38

39 The GDG considered the evidence review of the consequences of delirium, noting  
40 that dementia, death and new admission to long-term care were all significant

1 consequences of delirium. The GDG felt that awareness of this information was  
2 of significant importance, but acknowledged that a recommendation could not be  
3 made stating 'be aware of the consequences of delirium'. They recognised the  
4 difficulty of implementing and auditing a recommendation based on 'awareness'.  
5 However, in order not to lose the important message, the GDG agreed that a  
6 prominent statement conveying this message would appear at the start of the list  
7 of recommendations.

8 The GDG proposed a research recommendation (see Appendix H) to investigate  
9 the occurrence of delirium in the long-term care setting, and the consequences of  
10 delirium in that population.

11

## 12 **4.14 Treatment of delirium (recommendations 1.6.1–1.6.4)**

13

### 14 **4.14.1 Quality of evidence**

15 There was low quality evidence for the treatment of people with delirium from  
16 the multicomponent treatment review, and moderate quality evidence from the  
17 pharmacological treatment review and the adverse effects review. The GDG  
18 noted that the major adverse event considered (the incidence of stroke) came  
19 from indirect evidence, in people who would have received the drugs for long  
20 periods of time, unlike the short-term use in delirium.

21 Economic evidence was obtained by modelling the treatment pathway for two  
22 pharmacological interventions, and was informed by the pharmacological  
23 treatment review and the review on the consequences of delirium. It was also  
24 informed by evidence on cost, quality of life and baseline risks.

25 The GDG also considered evidence from the non-pharmacological risk factors  
26 review and the patient information review, and drew on their clinical experience.  
27 Their discussions were informed by the NICE guideline on Parkinson's Disease,  
28 and the recommendations cross refer to the NICE guideline on Violence.

29

### 30 **4.14.2 GDG considerations**

31 The multicomponent treatment review showed some indication of clinical  
32 effectiveness of the multicomponent intervention in one study (Pitkala 2006), but  
33 the GDG considered the measure of delirium to be too unreliable to support this  
34 in economic modelling. However, the GDG did draw on the components  
35 comprising the multicomponent interventions, and used them, together with  
36 information from the risk factors review to make a consensus recommendation on  
37 treating possible underlying causes of delirium (recommendation 1.6.1). The  
38 GDG recognised that sometimes there was more than one underlying factor.

1 The GDG recognised the importance of talking and listening to the person  
2 experiencing delirium. The GDG specifically took on board the messages  
3 conveyed by the patient representatives on the GDG describing how difficult it  
4 was for them to tell relatives and staff about their changes in cognition.

5 As a separate issue the GDG felt that evidence from the multicomponent  
6 treatment review and GDG experience underlined the importance of reinforcing  
7 and addressing orientation for example date, day, time and place. Hospital  
8 environments, artificial lighting and time loss through disturbed sleep patterns /  
9 unconsciousness can easily lead to disorientation with potential knock on  
10 implications to delirium. Familiar faces of family, friends and carers may also  
11 help with orientation. Recommendation 1.6.2 should be carried out for all  
12 people diagnosed with delirium.

13

14 The GDG referred to the NICE Violence guideline and how to calm down an  
15 escalating situation. The GDG considered that non-pharmacological de-  
16 escalation approaches should be tried before resorting to drug treatment. This  
17 was partly on the basis of their clinical experience and partly in view of their  
18 reservations about the evidence on drugs.

19

20 There was little evidence for the use of pharmacological agents for the treatment  
21 of delirium. The GDG observed that there was evidence from one moderate  
22 quality RCT, but did not wish to make a strong recommendation on the basis of a  
23 single study which had a risk of bias (Hu 2006).

24 The health economic analysis showed that haloperidol and olanzapine were cost  
25 effective compared with placebo for treating delirium, but the uncertainty  
26 around the cost effectiveness estimates precluded recommending one drug over  
27 another. The GDG took into consideration the possible harms of the drugs, for  
28 which the evidence was largely indirect. The GDG were uncertain whether there  
29 was a risk of stroke when using these drugs in the short-term treatment of  
30 delirium. Due to the limited evidence the GDG did not wish to consider a class  
31 effect and hence made recommendations for individual drugs (recommendation  
32 1.6.4).

33

34 On balance, weighing up the effects of reduced mortality and dementia, versus  
35 possible increased risk of stroke, and taking into account the cost effectiveness  
36 analysis, the GDG decided that the benefits outweighed the risks, and that they  
37 should recommend drug treatment after other treatment interventions had been  
38 tried. In the light of the adverse events associated with these drugs for longer  
39 term use, and their uncertainty about the evidence, the GDG did not want to  
40 recommend the routine use of these drugs for everyone with delirium. The GDG  
41 therefore decided to make a weak recommendation (as reflected by the  
42 recommendation wording) that healthcare professionals consider giving  
43 pharmacological treatment as short term treatment. Short-term treatment was  
44 defined as 1 week or less, based on the evidence from the Hu (2006) study and  
45 usual practice. The GDG considered that this treatment should only be given to  
46 patients who had severe or distressing symptoms and whose behaviour meant

1 their safety or the safety of those around them is compromised. This was in line  
2 with the summary of product characteristics (SPC) indications for these drugs for  
3 the treatment of symptoms: 'rapid control of agitation and disturbed behaviours  
4 in patients with schizophrenia or manic episode' for olanzapine and 'As an  
5 adjunct to short term management of moderate to severe psychomotor agitation,  
6 excitement, violent or dangerously impulsive behaviour' for haloperidol' (SPCs).

7

8 The GDG wished to investigate further the clinical and cost effectiveness of the  
9 range of pharmacological agents currently used for treating delirium and  
10 proposed a research recommendation (see Appendix H).

11

12

13

#### 14 **4.15 Information giving and support: recommendations 1.7.1 and 1.7.2**

15

##### 16 **4.15.1 Quality of evidence**

17 There was qualitative and quantitative evidence from the patient information  
18 review, which informed GDG discussions.

19

##### 20 **4.15.2 GDG considerations**

21 The GDG discussed who should be given information about delirium and at what  
22 stage(s) in the patient pathway. It was decided that it was not practical to give  
23 every person that presented in hospital or long-term care information about  
24 delirium and it was also not beneficial to unduly worry those who were not at  
25 risk. It was therefore decided that information would be most useful to people in  
26 hospital or long-term care at two stages in the pathway: to those who had been  
27 assessed and found to be at risk of delirium, and at a later stage to people  
28 diagnosed with delirium.

29 The GDG also noted from the evidence that it was important for information to  
30 be given to the relatives and carers of people at risk of delirium and to relatives  
31 and carers of people diagnosed with delirium, as well as the patients themselves.

32 The evidence review and experience of the patient representatives indicated the  
33 content of the patient information recommendations. The GDG considered that  
34 information about delirium could easily be incorporated into existing material for  
35 patients and relatives.

36 The GDG also decided to make a recommendation about patient information in  
37 accordance with equalities legislation and NICE's equality scheme. This was

1 because the information given should be accessible to people with additional  
2 needs such as physical, sensory or learning disabilities, and to people who do not  
3 speak or read English. Standard information delivery may not be applicable /  
4 beneficial to people with different cultural, linguistic, cognitive and literacy  
5 needs.

6

#### 7 **4.15.3 Single component non pharmacological prevention: music therapy**

8 The GDG considered the evidence which showed a significantly lower incidence  
9 of delirium in the group receiving music therapy compared with usual care. The  
10 GDG noted that the studies were at high risk of bias as an unvalidated method  
11 of assessing delirium incidence was used. The GDG did not want to make a  
12 recommendation based on this evidence and proposed music therapy should be  
13 considered in a future research recommendation (see Appendix H).

14

### 15 **4.16 Pharmacological prevention of delirium**

#### 16 **4.16.1 Quality of evidence**

17 There was limited low quality evidence described in the pharmacological  
18 prevention review. Each of the studies had quality issues (or were small sized):

- 19 • One study was not representative of the population (the donezepil study  
20 was investigating patients who were fit and healthy with no cognitive  
21 impairment)
- 22 • One study was not representative of the intervention or the population (the  
23 risperidone study used a dose that was very different from that used in  
24 clinical practice, and the study included a relatively young population  
25 (age range: 51 to 71 years) undergoing cardiac surgery
- 26 • One study was unrepresentative of the intervention because it combined  
27 benzodiazepines with meperidine.
- 28 • Two studies investigated haloperidol. One study had a high risk of bias and  
29 the other study assessed the adjunctive effect of haloperidol to a  
30 proactive geriatric consultation intervention.

31

#### 32 **4.16.2 GDG considerations**

33 The GDG agreed that typical antipsychotics, atypical antipsychotics,  
34 cholinesterase inhibitors and benzodiazepines should be considered as a  
35 research recommendation (see Appendix H). They noted that risperidone has  
36 been withdrawn for use in dementia because of the increased risk of stroke  
37 associated with its long-term use. For ethical reasons, the trial should only be  
38 carried out in a population at high risk of delirium.

## 5 The epidemiology of delirium: an assessment of need

### 5.1 Introduction

Delirium is a common clinical syndrome that can be found throughout the healthcare system. In order to understand more fully the clinical burden and associated health economic implications of delirium, it is necessary to first understand the epidemiology in terms of the occurrence of delirium within individual healthcare settings.

Operationalised diagnostic criteria for delirium have been formulated in the Diagnostic and Statistical Manual of Mental Disorders (American Psychiatric Association 1980; American Psychiatric Association 1987; American Psychiatric Association 1994) (DSM III, DSM III-R and DSM-IV), and in the International Classification of Diseases (10<sup>th</sup> Edition) (World Health Organisation 1992) (ICD-10). There is good diagnostic agreement between DSM-IV and its predecessors, with DSM-IV identifying all patients diagnosed with delirium by DSM III and DSM III-R in one prospective cohort study of elderly hospital patients and nursing home residents (Laurila 2004, and section 12.6).

There is a notable disparity between the DSM and ICD-10 criteria for the diagnosis of delirium. The DSM-IV criteria are more inclusive in terms of diagnosis of delirium, with ICD-10 being relatively restrictive. In a cohort of elderly medical hospital patients and nursing home residents (mean age 88.4 years), 24.9% met the diagnostic criteria of DSM-IV, whilst only 10.1% of the same cohort were diagnosed with delirium when the diagnostic criteria of ICD-10 were applied (Laurila 2004). A comparison of the DSM-IV and ICD-10 criteria (table 5.1) reveals the ICD-10 criteria to include additional requirements for the diagnosis of delirium. The stricter inclusion criteria and additional diagnostic requirements of ICD-10 have an associated impact on case detection and identifies a cohort of patients who are more frequently dependent for care needs and more likely to be resident in the long-term care setting (Laurila 2004).

In this guideline, we have identified the simplified, more inclusive, DSM-IV criteria as being the standard operational definition for delirium.

1

2 Table 5.1: DSM-IV and ICD-10 Diagnostic Criteria (American Psychiatric  
3 Association 1994; World Health Organisation 1992)

<b>DSM-IV Diagnostic Criteria (American Psychiatric Association, 1994)</b> <b>In order to be diagnosed with delirium, a patient must show all of the four features listed below:</b>	<b>ICD-10 Diagnostic Criteria (World Health Organisation, 1992)</b> <b>For a definite diagnosis, symptoms, mild or severe, should be present in each one of the following (five) areas:</b>
1. A disturbance of consciousness (i.e. reduced clarity of awareness of the environment) is evident, with reduced ability to focus, sustain or shift attention	a) Impairment of consciousness and attention (on a continuum from clouding to coma; reduced ability to direct, focus, sustain, and shift attention)
2. There is a change in cognition (such as memory deficit, disorientation, language disturbance) or the development of a perceptual disturbance that is not better accounted for by a pre-existing or evolving dementia.	b) Global disturbance of cognition (perceptual distortions, illusions and hallucinations – most often visual; impairment of abstract thinking and comprehension, with or without transient delusions, but typically with some degree of incoherence; impairment of immediate recall and of recent memory but with relatively intact remote memory; disorientation for time as well as, in more severe cases, for place and person)
3. The disturbance develops over a short period of time (usually hours to days) and tends to fluctuate during the course of the day.	
4. There is evidence from the history, physical examination, or laboratory findings that the disturbance is caused by the direct physiological consequences of a general medical condition, substance intoxication or substance withdrawal.	
	c) Psychomotor disturbances (hypo- or hyperactivity and unpredictable shifts from one to the other; increased reaction time; increased or decreased flow of speech; enhanced startle reaction)
	d) Disturbance of the sleep-wake cycle (insomnia or, in severe cases, total sleep loss or reversal of the sleep-wake cycle; daytime drowsiness; nocturnal worsening of symptoms; disturbing dreams or nightmares, which may continue as hallucinations after awakening)
	e) Emotional disturbances, e.g. depression, anxiety or fear, irritability, euphoria, apathy, or wondering perplexity.

4

### 5 5.1.1 Epidemiological terminology

6 Confusion can exist between the epidemiological terms **prevalence** and  
7 **incidence**. Prevalence represents the number of existing cases at a single point  
8 in time. Incidence represents the number of new cases that develop within a  
9 cohort over a defined period of time. The term 'occurrence rate' has been  
10 proposed as an alternative when there is ambiguity or overlap between the  
11 measurement of prevalence and incidence (Boyle 1998).

1 Prevalent delirium in hospital therefore defines the presence of delirium at the  
2 point of admission to hospital. Incident delirium in hospital represents the  
3 development of delirium after hospital admission.

4

5 This is an important distinction to make as incident (new) cases of delirium are  
6 more likely to be amenable to strategies aimed at preventing the onset of  
7 delirium. It is therefore of key importance to provide *a priori* definitions of  
8 prevalence, incidence and occurrence rates with regard to delirium. Where it is  
9 not possible to use these definitions because of healthcare setting, alternatives  
10 will be considered, for example in the surgical setting, in which the concept of  
11 pre- and post-operative delirium is likely to hold importance.

12 As the emergency department represents a healthcare setting in which patients  
13 spend a short period of time prior to admission to the hospital bed base or  
14 discharge home, the concept of point prevalence is most relevant in this setting  
15 and incidence/occurrence rates will not be measured.

16 Long-term care represents the permanent residence of an individual, rather than  
17 respite care on a temporary basis. The concepts of point prevalence  
18 (prevalence at a single point in time) and period incidence (cumulative incidence  
19 over a defined period of time) are likely to be relevant in the long-term care  
20 setting.

21

## 22 **5.1.2 *A priori* definitions**

23 The following *a priori* definitions form the basis for the review of study data and  
24 subsequent data categorisation:

25

### 26 **5.1.2.1 *Prevalent delirium***

27 The presence of delirium within the first 24 hours of admission to a healthcare  
28 setting (or the duration of the preoperative period within the surgical cohort).

29

### 30 **5.1.2.2 *Incident delirium***

31 The development of delirium subsequent to the first 24 hours of admission (24  
32 hours postoperatively in surgical cohorts), measured at least daily, until discharge  
33 from hospital or death.

34

### 35 **5.1.2.3 *Occurrence rate***

1 Where study data reveal overlap between the *a priori* definitions of prevalent  
2 and incident data, or where the *a priori* conditions are not met, the term  
3 'occurrence rate' will be used.

4

#### 5 **5.1.2.4 Total Delirium**

6 Where there is more than one measure of rate of delirium available (e.g. both  
7 prevalent and incident delirium), or where occurrence rate represents data  
8 collected from healthcare admission to discharge, a fourth term, total delirium,  
9 will be summated to reflect the occurrence of delirium throughout the duration of  
10 stay.

11

## 12 **5.2 Selection criteria for epidemiological studies**

13

### 14 **5.2.1 Types of study**

15 Prospective cohort and cross-sectional studies were to be included.  
16 Epidemiological data derived from the control arm of randomised clinical trials  
17 and case-control studies could be considered if there was evidence of  
18 reasonable representativeness of the sample. Retrospective studies were to be  
19 excluded.

20

### 21 **5.2.2 Patient population & healthcare setting**

22 Selection criteria for the patient population are defined in the methods section.  
23 Settings included are hospital and long-term care. In much of the guideline, the  
24 hospital patient population has been considered as a whole, but it is clear that  
25 this population is diverse and heterogeneous. For this epidemiological review,  
26 each healthcare setting was to be considered separately and data were to be  
27 grouped according to individual healthcare settings.

28 Studies were preferred if they were conducted in the UK. However, studies were  
29 to be included regardless of the country in which they were conducted, although  
30 the representativeness was to be taken into consideration in the analysis.

31 The DSM-IV criteria for delirium were to be the desired operational definition.  
32 As set out in the introduction, there is consistency between cases of delirium  
33 identified with DSM-IV versus DSM III-R and DSM III. Studies using a case  
34 definition based on the DSM-IV, DSM III-R or DSM III criteria [or a diagnostic tool  
35 validated against DSM-IV, DSM III-R or DSM III e.g. Confusion Assessment  
36 Method (CAM), DRS] were therefore to be included. As set out in the  
37 introduction, there is a notable disparity between cases of delirium that are  
38 identified with application of ICD-10 as compared with DSM-IV. Consequent to  
39 this, studies using the ICD-10 criteria for delirium were excluded from the  
40 epidemiological review.

1

### 2 **5.3 Hospital Episode Statistics (HES) data**

3 Locally generated clinical coding data is collated nationally in the Hospital  
4 Episode Statistics (HES) database, the national statistical data warehouse for the  
5 NHS. Clinical coding of data is used for clinical research, epidemiological  
6 mapping and health resource allocation. A bespoke HES dataset was generated  
7 in order to assess the agreement between the epidemiological profile of delirium  
8 as determined by prospective cohort data and clinical coding data collated by  
9 the HES database.

10

### 11 **5.4 Characteristics of included studies**

12 The initial search produced 1,767 citations of potential relevance and, following  
13 examination of all titles and abstracts, 199 full-text articles were retrieved for  
14 further consideration. 124 papers were excluded. Reasons for exclusion are  
15 reported in Appendix G.

16 We included 75 studies (Adamis 2005; Andrew 2006; Angles 2008; Balas  
17 2007; Benoit 2005; Bickel 2008; Brauer 2000; Breitbart 1996; Caeiro 2004;  
18 Cole 1994; Contin 2005; Dubois 2001; Edelstein 2004; Edlund 1999; Edlund  
19 2001; Edlund 2006; Elie 2000; Ely 2001; Faezah 2008; Franco 2001;  
20 Furlaneto 2006; Galanakis 2001; Goldenberg 2006; Greene 2009; Hamann  
21 2005; Han 2009; Henon 1999; Holden 2008; Holmes 2000; Inouye 1998;  
22 Inouye 1998; Inouye 1999; Jones 2006; Kagansky 2004; Kakuma 2003;  
23 Kawaguchi 2006; Koebrugge 2009; Koster 2008; Leslie 2005; Lewis 1995; Lin  
24 2004; Marcantonio 1994; Martin 2000; McAlpine 2008; McCusker 2003;  
25 McNicoll 2003; Milbrandt 2004; Milisen 2001; Morrison 2003; Naughton 1995;  
26 Naughton 2005; O'Keefe 1999; Ouimet 2007; Pandharipande 2008; Patten  
27 1997; Peterson 2006; Pisani 2006; Pitkala 2005; Ramirez-Bermudez 2006;  
28 Roberts 2005; Robinson 2008; Robinson 2009; Rockwood 1999; Rolfson 1999;  
29 Rudolph 2005; Rudolph 2006; Rudolph 2007; Santana Santos 2005; Santos  
30 2004; Sasajima 2000; Thomason 2005; Uldall 2000; van der Mast 1999; Van  
31 Rompaey 2009; Yoshimura 2004) and these are summarised in Appendix D. In  
32 four studies (Bickel 2008; Galanakis 2001; Inouye 1998; Pitkala 2005), more  
33 than one distinct cohort was examined and reported separately, thus giving data  
34 for 79 cohorts reported in 75 studies.

35

#### 36 **5.4.1 Study design**

37 Sixty-five studies had a prospective cohort design (Adamis 2005; Angles 2008;  
38 Balas 2007; Benoit 2005; Bickel 2008; Brauer 2000; Caeiro 2004; Contin  
39 2005; Dubois 2001; Edlund 1999; Edlund 2001; Edlund 2006; Ely 2001;  
40 Faezah 2008; Franco 2001; Furlaneto 2006; Galanakis 2001; Goldenberg  
41 2006; Greene 2009; Hamann 2005; Henon 1999; Holden 2008; Holmes 2000;

1 Inouye 1998; Inouye 1998; Inouye 1999; Jones 2006; Kagansky; Kawaguchi  
 2 2006; Koebrugge 2009; Koster 2008; Leslie 2005; Lin 2004; Marcantonio  
 3 1994; Martin 2000; McAlpine 2008; McCusker 2003; McNicoll 2003; Milbrandt  
 4 2004; Milisen 2001; Morrison 2003; Naughton 1995; Naughton 2005; O'Keefe  
 5 1999; Ouimet 2007; Pandharipande 2008; Patten 1997; Peterson 2006; Pisani  
 6 2006; Ramirez-Bermudez 2006; Roberts 2005; Robinson 2008; Robinson 2009;  
 7 Rockwood 1999; Rolfson 1999; Rudolph 2005; Rudolph 2006; Rudolph 2007;  
 8 Santana Santos 2005; Santos 2004; Sasajima 2000; Thomason 2005; Uldall  
 9 2000; van der Mast 1999; Van Rompaey 2009; Yoshimura 2004), five were  
 10 cross sectional studies (Elie 2000; Han 2009; Lewis 1995; Naughton 1995;  
 11 Pitkala) and two studies were randomised trials (Breitbart 1996; Cole 1994).

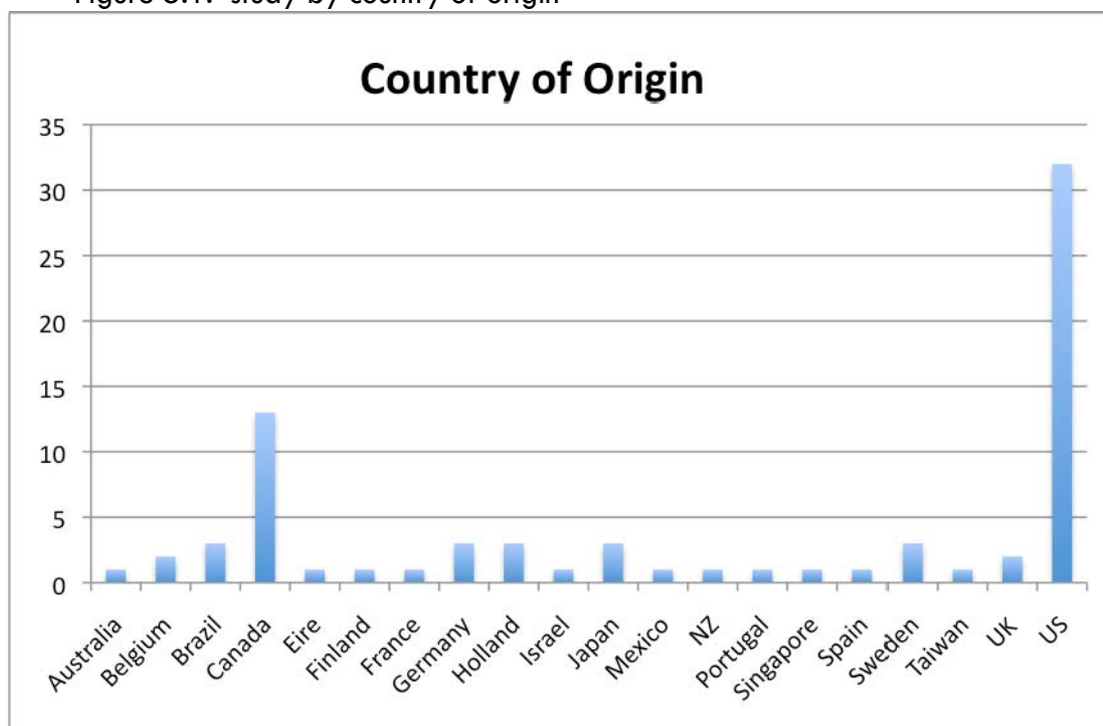
12 Eleven studies had fewer than 100 participants (Adamis 2005; Angles 2008;  
 13 Edlund 2009; Goldenberg 2006; Koebrugge 2009; Milisen 2001; Robinson  
 14 2008; Rolfson 1999; Rudolph 2005; Rudolph 2006; Santana Santos 2005); 11  
 15 studies had more than 500 participants (Brauer 2000; Holmes 2000; Inouye  
 16 2008; Leslie 2005; Marcantonio 1994; McCusker 2003; Morrison 2003; Ouimet  
 17 2007; Peterson 2006; Rudolph 2007; Van Rompaey 2009) and the remaining  
 18 50 studies had between 100 and 500 participants.

19 The majority of included studies were of North American origin (figure 5.1), with  
 20 only two studies based in the UK health service setting (Adamis 2005; Holmes  
 21 2000).

22

23

Figure 5.1: study by country of origin



24

25

26

27 Thirty-eight studies selected adult patients with age cut-off points (Adamis 2005;  
 28 Balas 2007; Bickel 2008; Brauer 2000; Breitbart 1996; Cole 1994; Edlund

1 2001; Edlund 2006; Elie 2000; Faezah 2008; Franco 2001; Furlaneto 2006;  
 2 Galanakis 2001; Goldenberg 2006; Greene 2009; Han 2009; Henon 1999;  
 3 Holden 2008; Holmes 2000; Inouye 1998; Inouye 1998; Inouye 1999; Jones  
 4 2006; Kagansky 2004; Koebrugge 2009; Leslie 2005; Lewis 1995;  
 5 Marcantonio 1994; Martin 2000; McAlpine 2008; McNicoll 2003; Naughton  
 6 1995; Naughton 2005; Pisani 2006; Pitkala 2005; Rockwood 1999; Santos  
 7 2004; Santana Santos 2005). One study selected patients above the age of 40  
 8 years, three those above the 50 years, six selected patients above 60 years, 17  
 9 above 65 years, eight above 70 years and three studies selected patients  
 10 above the age of 75 years.

11 Mean patient age varied between healthcare settings, with a higher mean age  
 12 of study participants noted in the general medicine and long-term care cohorts  
 13 (see Appendix D). A younger mean age of study participants was notable in the  
 14 ICU, HIV/AIDS medicine and psychiatry settings.

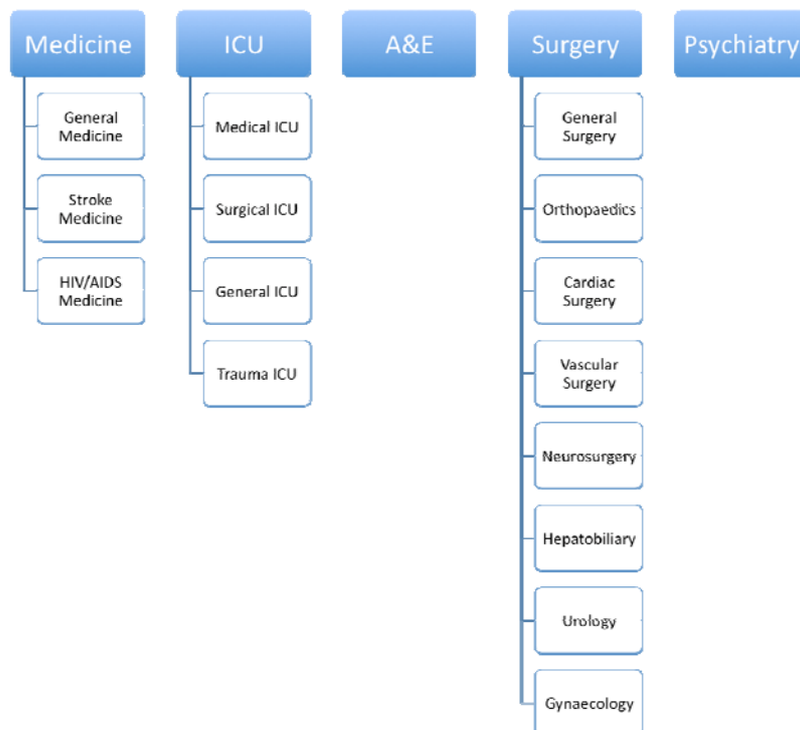
15

#### 16 5.4.2 Healthcare Setting

17 Studies were first assessed and grouped according to healthcare setting (Figure  
 18 5.2).

19

20 Figure 5.2. Hospital study populations grouped by healthcare setting



21

22

23

1 Where applicable, study populations were further categorised into, for  
2 example, acute and elective surgical patient groups. The long-term care setting  
3 was considered separately.

4 Both the ICU and acute stroke unit settings are frequently a form of enhanced  
5 specialist care within standard/usual care pathways. Thus, patients with ongoing  
6 delirium episodes may be admitted from the inpatient bed base to the  
7 ICU/acute stroke unit and therefore the occurrence rate can be a useful record  
8 of delirium rate for these specific healthcare settings. This model of ICU/acute  
9 stroke unit care is commonplace within the UK healthcare system.

10

## 11 **5.5 Methodological quality of studies**

12 The study cohort as a whole was assessed for representativeness on the grounds  
13 of the inclusion and exclusion criteria defined in each individual study. Inclusion  
14 and exclusion criteria were broadly similar between studies in each healthcare  
15 setting. Three studies (Andrew 2006; Edelstein 2004; Kakuma 2003) stated  
16 exclusion criteria showing that the study cohort was not representative of the  
17 population for that setting (see Appendix E). This is an important consideration  
18 for this epidemiology review, and these studies were therefore not analysed  
19 further.

- 20 • One study (Andrew 2006) was in a long-term care setting whereby people  
21 with dementia were excluded from the cohort.
- 22 • One study (Edelstein 2004) was in a hip fracture setting whereby only  
23 ambulatory home dwelling people were included in the cohort.
- 24 • One study (Kakuma 2003) was in an emergency department setting  
25 whereby people presenting from long-term care residents were excluded  
26 from the participant cohort.

27

28 Fourteen studies listed dementia as an exclusion criterion (Andrew 2006; Bickel  
29 2008; Contin 2005; Koebrugge 2009; Lin 2004; Roberts 2005; Rudolph 2007)  
30 or severe dementia (Franco 2001; Galanakis 2001; Han 2009; Kagansky  
31 2004; Leslie 2005; Martin 2000; McNicoll 2003). However, as many of these  
32 studies were in the surgical and ICU setting, it was felt that the exclusion of  
33 people with dementia in these studies would not necessarily affect the  
34 representativeness of the study cohort.

35 As set out earlier, studies using the DSM-IV, DSM III-R or DSM III criteria (or a  
36 diagnostic tool validated against DSM-IV, DSM III-R or DSM III) were considered  
37 for inclusion. As delirium may often be present at admission and may be present  
38 for a short period of time with a tendency to fluctuate, included studies were  
39 appraised for quality on the basis of (1) an initial assessment for delirium within  
40 the first 24 hours of admission (post admission, preoperative period in the  
41 surgical studies) and (2) the frequency of subsequent assessments for delirium.  
42 Included studies were also appraised on the basis of sample size. These three  
43 criteria form the overall basis of the methodological quality assessment  
44 (Appendix E).

1 The relative importance of each quality criterion varies according to the type of  
2 epidemiological measurement. For example, prevalent delirium represents  
3 delirium within the first 24 hours of admission (preoperative period in the  
4 surgical cohort). With regard to this measure, the study size is therefore the key  
5 index. With regard to occurrence rate, the frequency of measurement of  
6 delirium and the study duration are potentially of greater importance.

7 Therefore, where studies recorded more than one measure of delirium (e.g. both  
8 prevalent delirium and occurrence rates), these were given separate quality  
9 assessments (Appendix E).

10 The studies were pragmatically and qualitatively grouped into high, medium and  
11 low quality on the basis of the quality criteria (Appendix E). Studies in which the  
12 sample size was small, in which the assessment of delirium was notably infrequent  
13 and/or the overall study length was short compared to the expected length of  
14 healthcare stay were considered to be at high risk of bias if a combination of  
15 these factors were present. Studies in which the methodology was unclear were  
16 also considered to lead to risk of bias. There was significant heterogeneity  
17 noted in frequency of assessment of delirium across all studies.

18 On the basis of these factors, four studies (Edlund 1999; Rudolph 2005; Santana  
19 Santos 2005; Van Rompaey 2009) were excluded from the overall results  
20 summary as they were felt to give potential for bias. These studies are  
21 highlighted in blue and given in italics in the study summary tables (Appendix D).

22

## 23 5.6 Results

24 Full data are given in Appendix D. Sixteen studies reported incidence or  
25 prevalence in different healthcare settings. Three studies reported data for more  
26 than one setting:

- 27 • Pitkala 2005: General medicine (prevalence 32.6%); long-term care  
28 (15.9%)
- 29 • Bickel 2008: Orthopaedics acute hip fracture (occurrence 41%);  
30 orthopaedics elective surgery (12.5%)
- 31 • Galanakis 2001: Orthopaedics acute hip fracture (occurrence 40.5%);  
32 orthopaedics elective surgery (14.7%)

33

34 Summary data are reported by healthcare setting (table 5.2); in many  
35 healthcare settings the number of studies available for inclusion was limited, and  
36 the number ranged from 1 to 17 across all settings. Where more than one study  
37 is included, the median and range are given.

38

### 1 5.6.1 Sensitivity analysis

2 A sensitivity analysis was performed whereby the studies qualitatively graded  
 3 as low quality were excluded from the dataset (table 5.5 – end of chapter).  
 4 Removal of low quality studies led to significant change in a small number of  
 5 cumulative results. Where this was the case, the sensitivity analysis results are  
 6 preferred and these are shown in table 5.2 with the full results in square  
 7 brackets. Exclusion of one low quality study with a low occurrence rate in the  
 8 medical ICU setting led to a significant increase in the median (range) values for  
 9 the occurrence of delirium, from 70.9 (22.4 – 83.3) to 80 (48 – 83.3). Following  
 10 the sensitivity analysis, there was a decrease in the median (range) occurrence  
 11 rate of delirium in the cardiac surgery setting, from 32 (13.5 – 50) to 21 (13.5 –  
 12 33.6), and an increase for the acute hip fracture setting. There was no apparent  
 13 change in the rates of delirium in other healthcare settings when low quality  
 14 studies were excluded. Where the only studies in a particular healthcare setting  
 15 were low quality, this is indicated in the table.

16  
 17 Table 5.2: Summary data by healthcare setting. Full results are shown in the red  
 18 text

Healthcare setting	No. of studies	Prevalence % (Median, Range)	Incidence % (Median, Range)	Occurrence Rate % (Median, Range)	Total delirium % (median, range)
General Medicine	16	21.4 (18 – 32.6)	15.2 (12.5 – 17.9)	22 (5.7 – 42) [22 (3– 42)]	25 (15 – 42) [23.7 (15 – 42)]
Stroke Medicine	2	12	No data available	24.3	24.3
HIV/AIDS Medicine	2	No data available	No data available	12 (12 – 12)	12 (12 – 12)
Medical ICU	7	36.6	24.4	80 (48 – 83.3) [70.9 ( 22.4 – 83.3)]	70.9 (48 – 83.3)
Surgical ICU	4	No data available	No data available	43.5 (29.8 – 70)	36.9 (29.8 - 44) [43 (29.8 – 44)]
Trauma ICU	1	No data available	No data available	59 (low quality)	No data available
General ICU	3	No data available	No data available	31.8 (19 – 45)	38.4 (31.8 – 45)
Emergency Department	4	9.8 (9.6 – 11.1)	No data available	No data available	9.8 (9.6 – 11.1)
General Surgery	5	No data available	No data available	11.4 (9 – 24)	No data available
Orthopaedics (Acute Hip Fracture)	10	22 (16.5 – 29.7)	30.3 (12.5 – 48.1)	28.3 (9.5 – 41) [17.4 (9.5 – 41)]	35 (29 – 68.1) [44.8 (29 – 41.1)]
Orthopaedics (Elective)	3	No data available	No data available	13.6 (12.5 – 14.7) [14.7 (12.5 – 22)]	No data available
Orthopaedics (Spinal Surgery)	1	No data available	No data available	3.8	No data available
Cardiac Surgery	5	No data available	No data available	21 (13.5 – 33.6) [32 (13.5 – 50)]	No data available
Vascular Surgery	2	No data available	No data available	31.1 (29.1 – 33)	No data available
Neurosurgery	1	No data available	No data available	14.9	14.9

Hepatobiliary	1	No data available	No data available	17	No data available
Urology	1	No data available	No data available	7	No data available
Gynaecology	1	No data available	No data available	17.5 (low quality)	No data available
Psychiatry	1	No data available	No data available	2.8	No data available
Long-term care	1	No data available	No data available	15.9 (low quality)	No data available

1

2

### 3 5.6.2 UK Data

4 Two included studies gave data on rates of delirium in the UK healthcare setting.  
5 The first, a prospective cohort study in a general medical setting with a sample  
6 size of 940 (Adamis 2005), recorded an occurrence rate of delirium of 37.3%.  
7 The second, a larger prospective cohort study in an orthopaedic setting with a  
8 sample size of 731 (Holmes 2000), recorded an occurrence rate of delirium of  
9 14.8% (this study was considered to be of low quality). The limited number of  
10 studies available in UK healthcare settings leaves significant uncertainty as to the  
11 actual rates of delirium within the UK healthcare system.

12

### 13 5.6.3 Hospital Episode Statistics (HES) Data

14 In order to compare the epidemiological data with national clinical coding data,  
15 a bespoke dataset was requested from HES. The dataset provided information  
16 on the 2006 – 2007 total number of Finished Consultant Episodes (FCEs) of  
17 delirium (ICD code F05, delirium not induced by alcohol and other psychoactive)  
18 thus reflecting the scope of the guideline. The data were subcategorised by  
19 specialty and age (table 5.3).

20 Primary diagnoses represent the first of up to 14 diagnoses in the HES dataset  
21 and provide the main reason as to why the patient was in hospital. Subsequent  
22 to the primary diagnosis are up to 13 secondary diagnoses that record other  
23 diagnoses related to the episode. The bespoke delirium F05 dataset included  
24 both primary and secondary coded diagnoses of delirium, hence capturing all  
25 episodes of delirium in the UK healthcare setting in 2006 – 2007. It is likely that  
26 one episode of delirium corresponds to one patient having delirium. In order to  
27 calculate incidence of delirium as a percentage, the total number of FCEs in  
28 2006 – 2007 (again split by specialty) was also requested. The latter is the  
29 record of the primary diagnoses only, which approximates to the number of  
30 admissions to each specialty. Therefore the HES delirium percentage is a  
31 reasonable reflection of the total delirium rate.

32 The dataset was split by age. The HES dataset captures episodes between the  
33 ages of 15 – 44 years followed by age 45 – 64 years. In order to provide a  
34 dataset that was representative of the mean age and inclusion criteria of the

1 study cohort populations and in order that non-adult data was not introduced  
2 into the dataset, data were extracted from the HES dataset with a lower age  
3 limit of 45 years.

4

5 Table 5.3: Delirium Finished Consultant Episodes and Total Episodes by Specialty  
6 (Copyright © 2009, Re-used with the permission of The Health and Social Care  
7 Information Centre. All rights reserved)

Main Specialty	Delirium FCEs	Total Specialty FCEs	Total Delirium Episode Rate %
General Medicine	4706	2034768	0.23
Geriatric Medicine	3474	583506	0.59
Critical Care	15	102040	0.14
A & E	262	267476	0.01
Trauma & orthopaedics	204	652304	0.03
General Surgery	179	1041513	0.02
Adult Mental Illness	121	39839	0.30

8

9

#### 10 5.6.4 Epidemiology data compared with coded HES data

11 HES data are generated over the course of the hospital admission. As discussed  
12 above, the proportion of episodes of delirium is very similar to the total rate of  
13 delirium in the study summary tables (Appendix D). In order to assess the  
14 reliability of the HES data, table 5.4 shows both the HES data and the  
15 appropriate median total delirium rate (from the sensitivity analyses) as  
16 reported by the epidemiological research studies and where total delirium rate  
17 was available.

18

19 Table 5.4: Comparison of Median Total Delirium Rates with HES Total Delirium  
20 Episode Rates (Copyright © 2009, Re-used with the permission of The Health  
21 and Social Care Information Centre. All rights reserved)

Main Specialty	Median (Range) Total Delirium Rate (Epidemiology Data) %	Total Delirium Episode Rate (HES data) %
General Medicine	25 (15 – 42)	0.31
Critical Care	31.8 (19 – 45)	0.23
A & E	9.8 (9.6 – 11.1)	0.14
Trauma & orthopaedics	28.3 (9.5 – 41)	0.06

22

1

2 There is a clear and significant disparity between the expected total delirium  
3 rates from epidemiology data and the rates of delirium extracted from HES  
4 coding data. Less than one percent of the expected cases of delirium are  
5 identified by the coding process. There are also differences in the relative  
6 numbers of patients in the various healthcare settings, e.g. trauma & orthopaedic  
7 surgery has a similar level of delirium compared with general medicine in the  
8 studies, but the HES data show a much lower level for orthopaedic surgery.

9

## 10 **5.7 Discussion**

11 Accurate coding of clinical data relies on all of the following taking place: the  
12 recognition of the underlying diagnosis, recording of the diagnosis by a clinician  
13 in the medical notes and subsequent extraction of the correct diagnosis /  
14 diagnoses from the medical notes by clinical coders. It is possible that there is an  
15 attrition of delirium diagnoses at each of these three stages. Clinicians often fail  
16 to identify delirium in the hospital setting, with up to two thirds of cases of  
17 delirium remaining unrecognised (Inouye 1998). The 'terminological chaos'  
18 (Lindesay 1999) of delirium creates a situation in which a variety of terms are  
19 used to describe the diagnosis of delirium. If the correct diagnostic terminology  
20 for delirium is not used, clinical coders will be unable to extract accurate  
21 diagnostic data from the clinical record and hence there is the potential for  
22 considerable under-reporting of delirium at a national healthcare level.

23 Delirium is ubiquitous throughout the healthcare system, being particularly  
24 common in the critical care, hip fracture, vascular surgery, cardiac surgery and  
25 general medical patient populations. Delirium also appears to be common in the  
26 long-term care setting, with a point prevalence estimate of 15.9% when  
27 residents with dementia are included within the prospective cohort (we note that  
28 this study was considered to be of low quality).

29 In many healthcare settings there are few studies and these studies are often of  
30 lower quality. There is therefore significant uncertainty present with regard to  
31 the true epidemiology of delirium in a significant proportion of healthcare  
32 settings. In these healthcare settings further large prospective cohort studies of  
33 high methodological quality would help provide rigorous data informing the true  
34 epidemiology of delirium.

35 There is a paucity of prospective cohort studies of delirium in the UK healthcare  
36 environment, with the majority of epidemiological data originating from North  
37 America. There are potential differences between the structure and organisation  
38 of healthcare in the UK compared to North America that may limit between-  
39 system comparisons and there is consequent uncertainty regarding the true rates  
40 of delirium within the UK healthcare system.

41 There is a significant disparity between the expected rates of delirium from  
42 prospective epidemiological studies and the rates of delirium as recorded in the  
43 HES data set. National clinical coding is systematically failing to accurately

1 record the considerable scale and consequent importance of delirium as a  
2 healthcare priority.

3

4

5 Table 5.5: Sensitivity analysis - low quality studies removed, amended data  
6 highlighted in bold with number of low quality studies removed

Healthcare setting	No. of studies	Prevalence % (median, Range)	Incidence % (median, Range)	Occurrence Rate % (median, Range)	Total delirium % (median, range)
General Medicine	16	21.4 (18 - 32.6)	15.2 (12.5 - 17.9)	<b>22 (5.7 - 42)</b> 4 removed	<b>25 (15 - 42)</b> 1 removed
Stroke Medicine	2	12	No data available	24.3	24.3
HIV/AIDS Medicine	2	No data available	No data available	<b>12 (1 removed)</b>	<b>12 (1 removed)</b>
Medical ICU	7	36.6	24.4	<b>80 (48 - 83.3)</b> 1 removed	<b>70.9 (48 - 83.3)</b>
Surgical ICU	4	No data available	No data available	<b>44 (29.8 - 70)</b> 1 removed	<b>36.9 (29.8 - 44)</b> 1 removed
Trauma ICU	1	No data available	No data available	<b>59 (low quality)</b>	No data available
General ICU	3	No data available	No data available	31.8(19 - 45)	38.4 (31.8 - 45)
Emergency Department	4	9.8 (9.6 - 11.1)	No data available	No data available	9.8 (9.6 - 11.1)
General Surgery	5	No data available	No data available	<b>9 (9 - 11.4)</b> 2 removed	No data available
Orthopaedics (Acute Hip Fracture)	10	<b>23.1 (16.5 - 29.7)</b> 1 removed	<b>12.5</b> 1 removed	<b>28.3 (9.5 - 41)</b> 4 removed	<b>35 (29 - 41)</b> 2 removed
Orthopaedics (Elective)	3	No data available	No data available	<b>13.6 (12.5 - 14.7)</b> 1 removed	No data available
Orthopaedics (Spinal Surgery)	1	No data available	No data available	3.8	No data available
Cardiac Surgery	5	No data available	No data available	<b>21 (13.5 - 33.6)</b> 2 removed	No data available
Vascular Surgery	2	No data available	No data available	<b>29.1 (1 removed)</b>	No data available
Neurosurgery	1	No data available	No data available	14.9	14.9
Hepatobiliary	1	No data available	No data available	<b>17</b>	No data available
Urology	1	No data available	No data available	7	No data available
Gynaecology	1	No data available	No data available	<b>17.5 (low quality)</b>	No data available
Psychiatry	1	No data available	No data available	2.8	No data available
Long-term care	1	<b>15.9 (low quality)</b>	No data available	<b>15.9 (low quality)</b>	No data available

7

## 6 Risk factors for delirium: non-pharmacological

### 6.1 Clinical introduction

Delirium is a complex syndrome and patients appear to differ in their susceptibility to the condition. For example, some patients develop delirium with a urinary infection, while others do not. Understanding the underlying risk factors for delirium helps to explain this clinical variation. It also provides an opportunity to identify people who are at higher risk of delirium and, importantly, consider modifying key risk factors such that delirium incidence might be reduced.

### 6.2 Selection criteria

Selection criteria were as outlined in the general methods section apart from the types of risk factor described below.

#### 6.2.1 Types of risk factor

Any variable reported to be a risk factor for delirium was to be considered, including the following *a-priori* ones predicted by the GDG:

##### 6.2.1.1 Patient Characteristics

- Age
- Sex
- Dementia
- Sensory impairment
- Severity of illness
- Depression
- Multiorgan failure
- Polypharmacy (having more than one drug)
- Dehydration
- Electrolyte disturbance
- Continence
- Constipation

- 1           • Hypoxia
- 2           • Immobility/ bedridden
- 3           • Infection
- 4           • Malnutrition
- 5           • Sleep deprivation

#### 6    **6.2.1.2 Environmental**

- 7           • Setting
- 8           • Lighting
- 9           • Orientation
- 10          • Sensory overload

#### 11   **6.2.1.3 Procedural**

- 12          • Type of anaesthesia
- 13          • Cardiac surgery
- 14          • Hip fractures
- 15          • Insertion of urinary catheter
- 16          • Any iatrogenic intervention
- 17          • Smoking cessation
- 18          • Physical restraint

### 20   **6.3 Description of studies**

21           Eighty-four papers were evaluated for inclusion. Eleven studies were excluded  
 22           because fewer than 20 patients developed delirium (Clayer 2000: n=9;  
 23           Duggleby 1994: n=16; Eriksson 2002: n=12; Hamann 2005: n=7; Kaneko  
 24           1997: n=6; Kawaguchi 2006: n=13; Koebrugge 2009: n=17; McAlpine 2008:  
 25           n=18; Milstein 2000: n=10; Naughton 1995: n=18; Wakefield 1996: n=16);  
 26           25 five other studies were excluded and are listed in Appendix G with reasons  
 27           for exclusion.

28  
 29           Eleven other studies that were identified in update searches were included in the  
 30           review, but not analysed in depth because they were considered to be of low or  
 31           biased quality or they did not add to the body of evidence (Angles 2008;  
 32           Chang 2008; Detroyer 2008; Galankis 2001; Gao 2008; Greene 2009;  
 33           McManus 2009; Oh 2008; Robinson 2008; Van Rompaey 2009; Yang 2008).  
 34           One additional study was identified from the pharmacological risk factors  
 35           review (Pandharipande 2006; chapter 7).

36

### 1 6.3.1 Study Design

2 The 38 included studies had different study designs:

- 3 • 32 were prospective cohort studies (Andersson 2001; Böhner 2003;  
4 Bucerius 2004; Caeiro 2004; Edlund 2001; Ely 2007; Furlaneto 2006;  
5 Goldenberg 2006; Hofsté 1997; Inouye 1993; Inouye 2007; Kazmierski  
6 2006; Korevaar 2005; Leung 2007; Levkoff 1992; Margiotta 2006;  
7 McCusker 2001; Olin 2005; Ouimet 2007; Pandharipande 2006; Pisani  
8 2007; Pompei 1994; Ranhoff 2006; Rolfson 1999; Rudolph 2007;  
9 Santos 2004; Schor 1992; Sheng 2006; Veliz-Reissmüller 2007; Weed  
10 1995; Zakriya 2002)
- 11 • 3 were retrospective cohort studies (Levkoff 1988; Redelmeier 2008;  
12 Yildizeli 2005)
- 13 • 3 had a cross-sectional design (Ramirez-Bermudez 2006; Sandberg 2001;  
14 van Munster 2007).

15 The latter three studies were not reported further, because this is a weak study  
16 design and other data were available from the cohort studies. Details of the  
17 additional study (Pandharipande 2006) are given in section 7.3, and only  
18 reported here exceptionally.

19  
20 None of the studies were carried out in the UK. The other studies were conducted  
21 in various other countries:

- 22 • Thirteen in the USA (Ely 2007; Goldenberg 2006; Inouye 1993; Inouye  
23 2007; Leung 2007; Levkoff 1988; Levkoff 1992; Pisani 2007; Pompei  
24 1994; Rudolph 2007; Schor 1992; Weed 1995; Zakriya 2002)
- 25 • Four in Sweden (Andersson 2001; Edlund 2001; Olin 2005; Veliz-  
26 Reissmüller 2007)
- 27 • Four in Canada (Ouimet 2007; McCusker 2001; Redelmeier 2008; Rolfson  
28 1999)
- 29 • Two in The Netherlands (Hofsté 1997; Korevaar 2005)
- 30 • Two in Germany (Böhner 2003; Bucerius 2004)
- 31 • Two in Brazil (Furlaneto 2006; Santos 2004)
- 32 • Two in Italy (Margiotta 2006; Ranhoff 2006)
- 33 • One in each of Turkey (Yildizeli 2005), Portugal (Caeiro 2004), Poland  
34 (Kazmierski 2006), Australia (Sheng 2006), and Taiwan (Lin 2008).

35  
36 Of the prospective cohort studies, sample sizes ranged from 53 (Ely 2007) to  
37 16,184 patients (Bucerius 2004). Four studies had fewer than 100 patients, ten  
38 studies had 100 or more patients, thirteen had more than 200 patients, and five  
39 studies were very large (table 6.1). Of the three retrospective cohort studies,  
40 samples sizes were 432 patients (Yildizeli 2005), 1,285 patients (Levkoff 1988)  
41 and 28,4158 (Redelmeier 2008).

1 Table 6.1: sample sizes of prospective and retrospective cohort studies

Studies with fewer than 100 patients	Studies with 100 or more patients	Studies with more than 200 patients	Large studies
Ely 2007: n=53	Böhner 2003: n=153	Andersson 2001: n=457	Bucerius 2004 n=16,184
Goldenberg 2006: n=77	Edlund 2001: n=101	Caeiro 2004: n=218	Levkoff 1988: n=1,285
Olin 2005: n=61	Furlaneto 2006: n=103	Hofsté 1997: n=321	Ouimet 2007 n=764
Rolfson 1999: n=75	Inouye 1993: n=107	Inouye 2007: n=491	Pompei 1994: n=755
	Korevaar 2005: n=126	Kazmierski 2006: n=260	Redelmeier 2008: n=28,4158
	Lin 2008: n=151	Leung 2007: n=203	Rudolph 2007 n=1,218
	Sheng 2006: n=156	Levkoff 1992: n=325	
	Veliz-Reissmüller 2007: n=107	Margiotta 2006: n=330	
	Weed 1995: n=138	McCusker 2001: N=444	
	Zakriya 2002: n=168	Pisani 2007: n=304	
		Pompei 1994: n=432 and n=323	
		Ranhoff 2006: n=401	
		Santos 2004: n=220	
		Schor 1992: n=291	
		Yildizeli 2005: n=432	

2

3

4

5 All of the studies included hospital patients. The study by Pompei (1994)  
6 analysed data separately from two studies: n=432 from Chicago Hospital and  
7 n=323 from New Haven Hospital (data were not combined).

7

8

9 The study by Levkoff (1992) reported data separately for patients who were  
10 admitted to hospital from institutional settings (n=114, 35%), and those who  
11 were admitted from community settings (n=211), as well as combining the  
12 samples (reported for some risk factors). Nine other studies reported the  
patients' pre-hospital setting:

13

14

- Goldenberg (2006) had 79% of patients from the community and 21% from skilled nursing facilities

15

16

- Inouye (1993) reported that 3% of patients had been living in a nursing home

17

- Pisani (2007) had 18% patients from a nursing home

18

- Schor (1992) had 30% of patients from an institutional setting

19

20

- Andersson (2001) had 53% of patients living alone and 11% in sheltered accommodation

21

22

- Pompei (1994) Chicago hospital had 31% patients living alone and Pompei (1994) New Haven hospital had 41% living alone

23

- Ranhoff (2006) had 25% patients living alone

- 1           • Sheng (2006) had 90% patients living alone  
 2           • McCusker (2001) had 71% living alone, 18% from a foster home/senior  
 3           residence, and 11% from a nursing home

4  
 5           Eighteen studies were carried out in patients admitted for surgery (Andersson  
 6           2001; Böhner 2003; Bucerius 2004; Edlund 2001; Furlaneto 2006; Goldenberg  
 7           2006; Hofsté 1997; Kazmierski 2006; Leung 2007; Olin 2005; Redelmeier  
 8           2008; Rolfson 1999; Rudolph 2007; Santos 2004; Veliz-Reissmüller 2007;  
 9           Weed 1995; Yildizeli 2005; Zakriya 2002):

- 10           • Seven studies were conducted in patients undergoing cardiac operations  
 11           generally (Veliz-Reissmüller 2007), with and without cardiopulmonary  
 12           bypass (CPB) (Bucerius 2004), or with CPB only (Hofsté 1997), or  
 13           undergoing coronary artery bypass graft (CABG) surgery (Rolfson 1999;  
 14           Santos 2004), or open heart surgery (Kazmierski 2006), or aortic,  
 15           carotid, and vascular surgery (Böhner 2003)
- 16           • Five studies were in patients who had surgery for hip fracture (Andersson  
 17           2001; Edlund 2001; Furlaneto 2006; Goldenberg 2006; Zakriya 2002)
- 18           • One study was in patients who had major elective or urgent thoracic  
 19           surgery (Yildizeli 2005)
- 20           • One study was in patients who had abdominal surgery (Olin 2005)
- 21           • One study was in patients who had head and neck cancer surgery (Weed  
 22           1995)
- 23           • Two studies were in patients undergoing non-cardiac surgery (Leung 2007;  
 24           Rudolph 2007)
- 25           • One study was in patients undergoing cardiac, thoracic, neurosurgical,  
 26           vascular, musculoskeletal, lower urologic and gynaecologic, breast and  
 27           skin, external head and neck, and ophthalmologic surgery (Redelmeier  
 28           2008).

29  
 30           Four studies evaluated patients from both surgical and medical wards (Levkoff  
 31           1988; 1992; Pompei 1994; Schor 1992): in the study by Levkoff (1992) the  
 32           principal diagnoses of patients admitted to hospital included circulatory,  
 33           digestive, respiratory or genitourinary system diseases; endocrine, nutritional  
 34           and metabolic diseases; fractures; cancer; diseases of the skin or other reasons  
 35           not stated. Reasons for admission were not stated in the study by Pompei (1994).  
 36           In the study by Schor (1992), 61% were admitted to medical wards, 21% to  
 37           general surgery, and 8% to orthopaedic surgery.

38  
 39           Seven studies evaluated patients in medical wards only (Caeiro 2004 – stroke  
 40           unit; Inouye 1993; Inouye 2007; Korevaar 2005; Margiotta 2006; McCusker  
 41           2001; Sheng 2006):

- 42           • Two studies included acute stroke patients (Caeiro 2004; Sheng 2006)
- 43           • One study included patients admitted to an internal medicine ward with  
 44           diagnoses including infectious disease, malignancy, gastrointestinal

1 bleeding, water and electrolyte disturbances and other reasons not  
2 stated (Korevaar 2005)

3 • Reasons for admission were not stated in four studies (Inouye 1993; Inouye  
4 2007; Margiotta 2006; McCusker 2001).

5  
6 Six studies evaluated patients in intensive care units (ICUs) (Ely 2007; Lin 2008;  
7 Ouimet 2007; Pandharipande 2006; Pisani 2007; Ranhoff 2006):

8 • Three studies included mechanically ventilated patients in ICU (Ely 2007; Lin  
9 2008; Pandharipande 2006;)

10 • One study was in patients with admission diagnoses of respiratory,  
11 gastrointestinal haemorrhage, sepsis, neurological or other causes (Pisani  
12 2007)

13 • One study included patients admitted to a sub-intensive care unit for older  
14 people; diagnoses included respiratory failure, cardiac diseases, stroke,  
15 gastrointestinal bleeding, cancer-related problems, acute renal failure or  
16 other diagnoses not stated (Ranhoff 2006)

17 • Reasons for admission were not stated in the study by Ouimet (2007)

18

### 19 6.3.2 Population

20 Details about the population are summarised in this section, focussing on the  
21 principal risk factors; further details are given in Appendix F.

22

23 The mean **age** ranged from 51.7 years (Yildizeli 2005) to 87.4 years (Levkoff  
24 institution 1992). Age ranges are given in table 6.2; two studies did not report  
25 on patient age (Böhner 2003; Levkoff 1988). The GDG concluded that two  
26 studies had a narrow age range that could be considered to be effectively  
27 constant (Olin 2005; Rolfson 1999).

28

29

30 Table 6.2: Patient ages (+/- indicates that the range was calculated from the  
31 mean +/- 1 standard deviation)

Study	Age range (years)	Study	Age range (years)
Andersson (2001)	65-96	Margiotta (2006)	65-100
Böhner (2003)	not stated	McCusker (2001)	76-90 (+/-)
Bucerius (2004)	54-75 (+/-)	Olin (2005)	70-80
Caeiro (2004)	24-86	Ouimet 2007)	48-78
Edlund (2001)	65-102	Pandharipande 2006	25-90
Ely (2007)	31-79	Pisani (2007)	66-83
Furlaneto (2006)	71-90	Pompei (1994) (Chicago)	68-83
Goldenberg 2006)	66-98	Pompei (1994) (Yale)	73-85 (+/-)
Hofsté (1997)	29-83	Ranhoff (2006)	60-94
Inouye (1993)	73-86 (+/-)	Redelmeier (2008)	67-80
Inouye (2007)	73-85 (+/-)	Rolfson (1999)	69-74
Kazmierski (2006)	25-81	Rudolph (2007)	63-75 (+/-)
Korevaar (2005)	71-87 (+/-)	Santos (2004)	66-78
Leung (2007)	66-78 (+/-)	Schor (1992)	73-88 (+/-)
Levkoff (1988)	not stated	Sheng (2006)	65-95

Levkoff (1992)	74-89 (+/-)	Veliz-Reissmüller (2007)	65-95
Levkoff institution (1992)	80-95 (+/-)	Weed (2005)	mean 64
Levkoff community (1992)	71-85 (+/-)	Yildizeli (2005)	18-86
Lin (2008)	64-86	Zakriya (2002)	50-98

The studies varied in the proportions of patients reported to have **cognitive impairment** at baseline. In addition, the GDG decided that, when this was not clearly stated, it was unlikely that patients undergoing elective cardiac surgery would have cognitive impairment at baseline. This gave the following subgroups:

- No studies were carried out in which all the patients had cognitive impairment
- Twenty-two studies reported that some patients had cognitive impairment or dementia at baseline (Caeiro 2004; Edlund 2001; Ely 2007; Furlaneto 2006; Goldenberg 2006; Hofsté 1997; Inouye 1993; Inouye 2007; Kazmierski 2006; Korevaar 2005; Leung 2007; Levkoff 1992; Margiotta 2006; McCusker 2001; Olin 2005; Pisani 2007; Pompei 1994; Rolfson 1999; Schor 1992; Sheng 2006; Veliz-Reissmüller 2007; Weed 2005)
- Inouye (1993) also excluded patients with severe underlying dementia
- Two studies stated that patients with cognitive impairment at baseline were excluded from their studies (Andersson 2001; Santos 2004) and four studies excluded patients with pre-existing dementia (Kazmierski 2006; Lin 2008; Rudolph 2007; Zakriya 2002).
- Rudolph (2007) included patients with mild cognitive impairment, but not dementia
- Kazmierski (2006) reported results for cognitive impairment as a risk factor
- One ICU study (Ranhoff 2006) reported scores on the MMSE at discharge from the hospital and used this together with measures of pre-admission activities of daily living (ADL) to determine pre-existing dementia (which the authors described as 'probably demented'). This is, at best, an indirect measure of pre-existing dementia, but it was used in the multivariate analysis
- It was not stated if the patients had cognitive impairment at baseline in five studies (Böhner 2003; Bucerius 2004; Levkoff 1988; Ouimet 2007; Redelmeier 2008).
  - Three of these studies were carried out in elective heart surgery patients who would be unlikely to have cognitive impairment (Böhner 2003; Bucerius 2004; Redelmeier 2008)
  - However, we note that three elective cardiac surgery studies stated that some patients had cognitive impairment at baseline (e.g. Rolfson 1999; Veliz-Reissmüller 2007)

Of the studies that assessed cognitive impairment and/or dementia, 18 used the Mini Mental State Examination (MMSE) score, two used DSM-IV; four used Informant Questionnaire on Cognitive Decline in the Elderly (IQCODE); and two

1 used the Blessed dementia questionnaire; four studies did not report what scale  
 2 was used (table 6.3). One study (Caeiro 2004) had less than 10% of patients  
 3 with cognitive impairment, so that any results for cognitive impairment in this  
 4 study were likely to be inaccurate. The GDG considered that the cut-off point of  
 5 28 on the MMSE scale, used in the Veliz-Reissmuller (2007) study, was unreliable  
 6 and this study was not included in the analyses for cognitive impairment.  
 7

8 Table 6.3: Cognitive impairment and/or dementia

Study	Cognitive impairment and/or dementia
Caeiro (2004)	Unstated scale: 3% had dementia/cognitive decline
Edlund (2001)	DSM-IV: 21 of 101 (21%) patients had dementia
Ely (2007)	IQCODE: 16% had a mean score of 4 or more
Furlaneto (2006)	MMSE: mean 12.07 (SD 9.04) in delirium group and 17.74 (SD 8.78) in control group; Blessed dementia questionnaire to caregiver: 45% had a score above 4
Goldenberg (2006)	MMSE: mean score 21.6 (range 2 to 30); DSM-IV: 53 of 77 (69%) had dementia
Hofsté (1997)	MMSE: 23% reported to have cognitive disorders
Inouye (1993)	MMSE: mean score 24.2 (5.0); 36% with a score below 24
Inouye (2007)	MMSE: mean 23.1 (SD 6.3); 39% with a score below 24; modified Blessed dementia questionnaire to family member: 20% had a score above 4
Kazmierski (2006)	MMSE: 53% in group with delirium and 16% in group without delirium (preoperatively) had a score equal to or below 24
Korevaar (2005)	MMSE: 53% had a score below 24; IQCODE: 43% had a mean score of 3.9 or more
Leung (2007)	MMSE: mean score 33 (SD 3.2)
Levkoff (1992)	Unstated scale: 24% had cognitive impairment
Margiotta (2006)	MMSE: mean score 16.9 (SD 6.8) in patients with delirium and 22.1 (SD 7.0) in patients without delirium
McCusker (2001)	IQCODE: 60% with a score of 3.5 or more
Olin (2005)	MMSE: mean score 28 (SD 3)
Pisani (2007)	IQCODE: 31% had a mean score of 3.3 or more
Pompei (1994)	MMSE: 37% had cognitive impairment
Ranhoff (2006)	MMSE on discharge: mean score was 19.1 (SD 11) prior to hospital admission; 30% had MMSE score less than 18 and/or Barthel Index less than 95 and/or IADL impairment on 1 or more tasks
Rolfson (1999)	MMSE: 9% in group with delirium and 12% in group without delirium using a cut-off of 24
Rudolph (2007)	MMSE: mean 27.8 (SD 1.6) at baseline
Santos (2004)	MMSE: no patients with cognitive impairment
Schor (1992)	Unstated scale: 42% had a history of cognitive impairment in delirium group and 10% in group without delirium
Sheng (2006)	MMSE: overall scores at one month were 23.4 (SD 6); 8% were reported to have dementia
Veliz-Reissmüller (2007)	MMSE: median score 29 (range 17-30) in group with delirium and 30 (range 27-30) in group without delirium; cut-off was 28
Weed (1995)	MMSE: mean score 26.3 in patients with delirium and 27.4 in patients without delirium
Zakriya 2002	Method of assessment not stated

9  
 10  
 11 Sensory impairment was reported in twelve studies (Andersson 2001; Böhner  
 12 2003; Edlund 2001; Inouye 1993; 2007; Margiotta 2006; McCusker 2001;  
 13 Pisani 2007; Ranhoff 2006; Schor 1992; Sheng 2006; Weed 2005). Four  
 14 studies excluded patients with severe visual and/or hearing impairment (Levkoff

1 1992; Olin 2005; Santos 2004; Schor 1992); Hofsté (1997) and Rolfson (1999)  
 2 excluded people who were blind or deaf, but the GDG did not consider this to  
 3 be a modifiable risk factor for sensory impairment and noted that there would  
 4 be other people who did have other degrees of sensory impairment. The studies  
 5 did not generally give much information on how sensory impairment was  
 6 assessed:

- 7 • Andersson (2001) and Pisani (2007): stated it was patient reported and  
 8 proxy reported respectively
- 9 • Ranhoff (2006): patient/close relative was asked if they had vision  
 10 problems affecting daily activity
- 11 • Inouye (1993) and Inouye (2007): Jaeger- and Snellen-type tests for  
 12 standard vision – visual impairment was defined as corrected vision  
 13 worse than 20/70 on both near and distant binocular tests. For hearing  
 14 impairment, the Inouye (2007) study used a whisper test and Inouye  
 15 (1993) used a Welch-Allyn audioscope and questions designed to screen  
 16 for hearing loss – hearing impairment was defined if the patient heard  
 17 fewer than three of eight tones on the audioscope (at 40 dB and  
 18 frequencies of 500, 1000, 2000 and 4000 Hz) and a score of 4 or less  
 19 (of 8) on the screening tests
- 20 • McCusker (2001): no details, but the study also included in the analysis  
 21 whether or not the patient was wearing reading glasses
- 22 • Sheng (2006) in stroke patients recorded ‘vision field loss’

23  
 24 Levels of sensory impairment are given in table 6.4.

25  
 26  
 27 Table 6.4: Sensory impairment

Study	Visual impairment	Hearing impairment
Andersson 2001	31%	39%
Böhner 2003	61%	24%
Edlund 2001	23%	30%
Inouye 1993	6%	54%
Inouye 2007	38%	Not reported
McCusker 2001	20% with visual/hearing impairment; the authors also reported that 48% patients were wearing glasses, and 8% used a hearing aid	
Margiotta 2006	Some patients with sensory impairment (details not reported)	
Pisani 2007	11%	17%
Ranhoff 2006	29%	Not reported
Schor 1992	33%	21%
Sheng 2006	18%	Not reported
Weed 2005	5%	11%

28  
 29  
 30 Eight studies reported on the number of drugs (**polypharmacy**) taken by patients  
 31 (Goldenberg 2006; Inouye 2007; Korevaar 2005; Olin 2005; Ranhoff 2006;

1 Rolfson 1999; Veliz-Reissmüller 2007; Weed 1995). Where reported, the mean  
2 number of drugs ranged from 1.4 (Rolfson 1999) to 8.5 (Ranhoff 2006).

- 3 • Goldenberg (2006) reported that 87% of the patients had more than  
4 three medications at baseline (means were not reported)
- 5 • Inouye (2007) reported that 56% of the patients had more than three  
6 hospital medications in one day, and 29% had more than three  
7 psychoactive medications in one day
- 8 • Korevaar (2005) reported that the mean number of drugs used before  
9 admission was 4.4 (SD 3.2) in patients with delirium and 4.9 (SD 3.6) in  
10 patients without delirium
- 11 • Olin (2005) reported that the mean number of drugs taken was 3.0 (SD 3)  
12 in patients with delirium and 2.1 (SD 2) in patients without delirium
- 13 • Ranhoff (2006) reported that the mean number of drugs used was 8.5 (SD  
14 3.4) in patients with prevalent delirium, 8.0 (SD 3.2) in patients with  
15 incident delirium, and 7.3 (SD 3.1) in patients without delirium
- 16 • Rolfson (1999) reported that mean number of *selective drugs* used  
17 (dimenhydrinate, meperidine, or any benzodiazepine) was 1.4 in patients  
18 with delirium and 1.6 in the patients without delirium
- 19 • Veliz-Reissmüller (2007) reported that the mean number of drugs taken  
20 was 6.2 (SD 3.4) in the group with delirium and 6 (SD 3) in the group  
21 without delirium
- 22 • Weed (1995) reported that the mean number of medications was 3.4 in  
23 patients with delirium and 3.0 in patients without delirium.

24  
25 The GDG considered a definition of polypharmacy and did not agree on a  
26 suitable cut-off point: either 3 or 5 drugs were suggested, depending on setting.  
27 The GDG ruled that, for studies in older patients undergoing cardiac surgery,  
28 polypharmacy was likely to be present in all patients (i.e., Böhner 2003; Bucerius  
29 2004; Rolfson 1999; Santos 2004; Veliz-Reissmüller 2007). Similarly the GDG  
30 regarded studies in ICU as having the majority of patients with polypharmacy  
31 (i.e., Ely 2007; Lin 2008; Ouimet 2007; Pisani 2007; Ranhoff 2006).

32  
33 Comorbidities were reported in most of the studies, with the exception of Inouye  
34 (1993); Inouye (2007) and Rolfson (1999). Generally, they included conditions  
35 related to heart disease (congestive heart failure, previous myocardial  
36 infarction, atrial fibrillation), angina, stroke, hypertension, diabetes, obesity,  
37 renal dysfunction, chronic obstructive pulmonary disease, asthma, hypothyroid,  
38 cancer, and depression. Two studies reported baseline Charlson Comorbidity  
39 Index data (Inouye 2007; McCusker 2001). In these studies, the mean scores  
40 were 2.7 (SD 2.1) and 2.7 (SD 2.0) respectively.  
41

## 42 **6.4 Methodological quality of included studies**

43 The methodological quality of studies was assessed according to the type of  
44 study design. In evaluating the literature, RCTs and cohort studies were selected  
45 to be the best available evidence source for this review. Cross-sectional and

1 case-control studies were not included in this review unless there was no other  
2 information.  
3

#### 4 **6.4.1 RCTs**

5 No RCTs met the inclusion criteria.  
6

#### 7 **6.4.2 Cohort studies**

##### 8 **6.4.2.1 Representativeness and prospectiveness**

9 None of the 35 cohort studies were considered to be truly representative of the  
10 population (i.e. adults in surgical and/or medical wards in hospital or people in  
11 long-term care). In all studies except the McCusker (2001) study, the non-  
12 exposed cohort was drawn from the same community as the exposed cohort. The  
13 McCusker (2001) was a secondary analysis of data from two related concurrent  
14 studies, an RCT in patients with delirium, and non-delirious patients were selected  
15 from patients screened for delirium but free of the condition.  
16

17 All studies were prospective apart from three (Levkoff 1988; Redelmeier 2008;  
18 Yildizeli 2005), which were retrospective.  
19

##### 20 **6.4.2.2 Missing data**

21 Eight studies reported less than 20% loss to follow-up (Caeiro 2004; Edlund  
22 2001; Inouye 2007; Leung 2007; Lin 2008; Rolfson 1999; Rudolph 2007; Veliz-  
23 Reissmüller 2007); the remaining studies reported that all the patients were  
24 followed up, with the exception of McCusker (2001) and Pandharipande 2006,  
25 in which it was not clearly reported.  
26

27 One study reported an *a priori* sample size calculation (Rolfson 1999). In this  
28 study, a sample size of 81 was estimated assuming  $\alpha=0.05$ ,  $\beta=0.20$ ,  
29 and a desired margin of error of 0.10, with an anticipated proportion of  
30 delirium of 30%. The sample size of this study was 75.  
31

##### 32 **6.4.2.3 Delirium at baseline**

33 The studies varied in the number of patients with prevalent delirium (delirium at  
34 baseline): further details are given in Appendix D.

- 35 • Sixteen studies reported that none of the patients had delirium at baseline  
36 (Andersson 2001; Böhner 2003; Goldenberg 2006; Inouye 1993; Inouye  
37 2007; Kazmierski 2006; Levkoff 1988; Lin 2008; Olin 2005; Rolfson  
38 1999; Rudolph 2007; Santos 2004; Schor 1992; Veliz-Reissmüller 2007;  
39 Yildizeli 2005; Zakriya 2002)

- 40 ○ eight of these studies excluded patients with delirium at baseline  
41 from their studies (Andersson 2001; Goldenberg 2006; Inouye

- 1 1993; Inouye 2007; Kazmierski 2006; Olin 2005; Rolfson 1999;  
2 Schor 1992; Zakriya 2002).
- 3 • Six studies reported that some patients had delirium at baseline (Edlund  
4 2001; Furlaneto 2006; Levkoff 1992; Margiotta 2006; Pompei 1994;  
5 Ranhoff 2006).
- 6 ○ Two studies excluded these patients from the analysis: (Edlund  
7 2001: 61% of all patients; Levkoff 1992:10%)
- 8 ○ Three studies (four cohorts) included these patients in the analysis  
9 together with patients with incident delirium:
- 10 - Furlaneto (2006): 17% (17/103) prevalent, 13%  
11 (13/103) incident; 57% of all delirium was prevalent  
12 (17/30)
- 13 - Pompei (1994) Chicago: 5% (21/463) prevalent, 9%  
14 (43/463) incident; 33% of all delirium was prevalent  
15 (21/64)
- 16 - Pompei (1994) Yale: 15% (48/323) prevalent, 12%  
17 (38/323) incident; 56% of all delirium was prevalent  
18 (48/86)
- 19 - Margiotta (2006): 9% (31/330) prevalent, 10%  
20 (32/330) incident; 49% was prevalent (31/63)
- 21 ○ One study (Ranhoff 2006) reported that 16% (62/401) of  
22 patients had prevalent delirium, and 14% (55/410) had incident  
23 delirium; 53% of all delirium was prevalent. This study was  
24 carried out in a sub-ICU and prevalent delirium was diagnosed  
25 within 24 hours of admission to ICU. The GDG did not believe that  
26 incident and prevalent delirium could be distinguished in this  
27 population (because patients had come from other parts of the  
28 hospital) and all delirium was assumed to be incident.
- 29 • For 11 studies, it was unclear if the patients had delirium at baseline  
30 (Bucerius 2004; Caeiro 2004; Ely 2007; Hofsté 1997; Korevaar 2005;  
31 Leung 2007; Margiotta 2006; Ouimet 2007; Pisani 2007; Redelmeier  
32 2008; Sheng 2006; Weed 1995).
- 33 ○ In all of these studies the authors evaluated patients who  
34 'developed' delirium, but they did not specifically state if any of  
35 the patients had existing delirium.
- 36 ○ Two of these studies (Bucerius 2004; Hofsté 1997) included  
37 patients undergoing elective cardiac surgery and the GDG  
38 decided that this type of operation was unlikely to be carried out  
39 in patients with preoperative delirium.
- 40 ○ Four studies (Ely 2007; Ouimet 2007; Pandharipande 2006;  
41 Pisani 2007) were carried out in ICU and the GDG considered  
42 that these patients were likely to have incident delirium only
- 43 • One study evaluated delirium severity (McCusker 2001); the authors  
44 reported that 73% of patients had prevalent delirium (although  
45 prevalent (versus incident) delirium was included as a risk factor in the  
46 multivariate analysis).

#### 1 **6.4.2.4 Method of delirium assessment**

2 A number of validated instruments were used to evaluate delirium incidence or  
 3 duration using DSM-IV or DSM-III-R criteria. The GDG considered that 27 studies  
 4 had an adequate method of assessment; two had a partially adequate method  
 5 (Levkoff 1992; Schor 1992); three had an inadequate method (Levkoff 1988;  
 6 Redelmeier 2008; Yildizeli 2005) and one did not state the method (Weed  
 7 1995).  
 8

##### 9 • Adequate method

- 10 ○ Fifteen studies used the CAM (Ely 2007; Furlaneto 2006;  
 11 Goldenberg 2006; Inouye 1993; Inouye 2007; Korevaar 2005;  
 12 Leung 2007; Lin 2008; Margiotta 2006; Olin 2005; Pisani 2007;  
 13 Ranhoff 2006; Rolfson 1999; Veliz-Reissmüller 2007; Zakriya  
 14 2002)
- 15 ○ Two studies used the Organic Brain Syndrome (OBS) scale  
 16 (Andersson 2001; Edlund 2001) (the study by Andersson 2001  
 17 used a modified version of this scale)
- 18 ○ Two studies used the DRS (Böhner 2003; Caeiro 2004)
- 19 ○ One study used the used the Intensive Care Delirium Screening  
 20 Checklist (ICDSC) (Ouimet 2007)
- 21 ○ One study used the CAM-ICU test with the Richmond Agitation  
 22 Sedation Scale (RASS) (Pandharipande 2006)
- 23 ○ One study used the Saskatoon Delirium Checklist (SDC) (Hofsté  
 24 1997)
- 25 ○ Six studies assessed delirium based on clinical observations using  
 26 DSM-IV, DSM-III-R or (Bucerius 2004; Kazmierski 2006; Pompei  
 27 1994; Rudolph 2007; Santos 2004; Sheng 2006).
- 28 ○ Two studies (Levkoff 1992; Schor 1992) used the Delirium  
 29 Symptom Interview (DSI) which assesses the domains of delirium  
 30 specified in DSM III. The GDG considered this to be an adequate  
 31 method.

32

##### 33 • Inadequate

- 34 ○ Three studies assessed delirium by retrospective chart review  
 35 (Levkoff 1988; Redelmeier 2008; Yildizeli 2005)
- 36 ○ The study by Weed (1995) did not report what diagnostic  
 37 criteria were used to assess delirium, or what instrument was  
 38 applied.

39

40 One study evaluated severity of delirium as an outcome measure (McCusker  
 41 2001). In this study, the authors developed in their group a Delirium Index (DI)  
 42 based on the CAM criteria, which ranged from 0 to 21 (maximum severity). This  
 43 was compared with the Delirium Rating Scale which showed reasonably good

1 correlation (Pearson correlation coefficient 0.84). However, the GDG regarded  
2 this as indirect evidence, and this was supported by the review of diagnostic test  
3 accuracy (chapter 12).

4  
5 The GDG considered the three retrospective studies (Levkoff 1988; Redelmeier  
6 2008; Yildizeli 2005) to be biased because the method of assessment was  
7 based on review of medical notes. The GDG agreed that the two studies  
8 (Levkoff 1992; Schor 1992), which used the DSM III (or methods based on DSM  
9 III) for assessment had an adequate method of assessment.

#### 11 **6.4.2.5 Confounders taken into account**

12 Of the 35 cohort studies, 32 conducted multivariate analyses (Andersson 2001;  
13 Böhner 2003; Bucerius 2004; Caeiro 2004; Edlund 2001; Ely 2007; Furlaneto  
14 2006; Goldenberg 2006; Hofsté 1997; Inouye 1993; Inouye 2007; Kazmierski  
15 2006; Korevaar 2005; Leung 2007; Levkoff 1988; Levkoff 1992; Lin 2008;  
16 McCusker 2001; Ouimet 2007; Pandharipande 2006; Pisani 2007; Pompei  
17 1994; Ranhoff 2006; Redelmeier 2008; Rolfson 1999; Rudolph 2007; Santos  
18 2004; Schor 1992; Sheng 2006; Veliz-Reissmüller 2007; Yildizeli 2005; Zakriya  
19 2002).

20 Three studies conducted only univariate analyses for the incidence of delirium:  
21 Margiotta 2006; Olin 2005; Weed 1995) and these studies were not  
22 considered further. Details of the factors included in the multivariate analysis are  
23 given in Appendix F.

24 We considered whether the cohort studies took account of particular  
25 confounders, either in the study design or the multivariate analysis. The GDG had  
26 identified, by consensus, four risk factors to be important: age, sensory  
27 impairment, polypharmacy and cognitive impairment. Following GDG discussion  
28 it was decided *post-hoc* to record whether the multivariate analyses included  
29 severity of illness or comorbidity, as well as polypharmacy.

30 Studies were summarised according to the number of key risk factors included in  
31 the multivariate analysis and the ratio of events to covariates (the GDG  
32 considered a ratio of 1 or less to be flawed and a ratio of 2 or 3 to be possibly  
33 confounded). We assumed that the key risk factors were the same for severity of  
34 delirium and duration of delirium. The following combinations were found:

- 35 • Confounders taken into account: all/most (4 or 3) of the important risk  
36 factors (RFs) taken into account in the multivariate analysis or held  
37 constant and a ratio of events to variables of 10 or more
  - 38 ○ Bucerius (2004) had a ratio of 39 (3 key RFs: age included in the  
39 analysis; cognitive impairment excluded because elective cardiac  
40 operations and polypharmacy constant because elective cardiac  
41 operations in older patients; missing key RF: sensory impairment)
  - 42 ○ Levkoff (1992) had a ratio of 23 (2-3 key RFs: age and cognitive  
43 impairment included in the analysis, and patients with severe  
44 sensory impairment were excluded; illness severity included. No  
45 systematic standardised method was used to detect cognitive  
46 impairment, with reliance on medical chart review)

- 1                   ○ McCusker (2001) had a ratio of 18 (3 key RFs: age, dementia,  
2                   and sensory impairment included in the analysis; missing key RF:  
3                   polypharmacy; comorbidity included)
- 4                   ○ Schor (1992) had a ratio of 10 (2-3 key RFs: age and cognitive  
5                   impairment included in the analysis and patients with severe  
6                   hearing or vision impairment excluded; missing key RF:  
7                   polypharmacy; unstated scale for cognitive impairment)
- 8                   ● Possibly confounded: all/most of the important risk factors taken into  
9                   account in the multivariate analysis but an insufficient ratio of events to  
10                  variables
- 11                  ○ Ranhoff (2006) had a ratio of 7 (all 4 key RFs included in the  
12                  analysis)
- 13                  ○ Böhner (2003) had a ratio of 7 (3 key RFs: age and cognitive  
14                  impairment included in the analysis and polypharmacy constant  
15                  because elective cardiac operations in older patients; missing key  
16                  RF: sensory impairment)
- 17                  ○ Goldenberg (2006) had a ratio of 6 (3 key RFs included in the  
18                  analysis – not sensory impairment)
- 19                  ○ Pandharipande (2006) had a ratio that ranged from 4 (66/17)  
20                  to 7 (118/17) (3 key RFs: age, dementia, visual impairment )
- 21                    - The study reported the number with delirium for two  
22                    subgroups: those who received antipsychotics (66/75 had  
23                    delirium) and those who received anticholinergics (52/63);  
24                    it is unclear if any patients had both drugs, therefore the  
25                    number with delirium was considered to range from 66 to  
26                    118.
- 27                  ○ Veliz-Reissmüller (2007) had a ratio of 4 (3 key RFs: age and  
28                  cognitive impairment included in the analysis and polypharmacy  
29                  constant because elective cardiac operations in older patients;  
30                  missing key RF: sensory impairment; inappropriate cut off point on  
31                  MMSE scale for cognitive impairment)
- 32                  ○ Sheng (2006) had a ratio of 3 (3 key RFs included in the analysis  
33                  – not polypharmacy)
- 34                  ○ 3 studies had ratio of 2:
- 35                    - Andersson (2001) (all 4 key RFs included in the analysis;  
36                    comorbidity was also included)
- 37                    - Santos (2004) (3-4 key RFs: age and cognitive impairment  
38                    included in the analysis; polypharmacy constant because  
39                    elective cardiac operations in older patients; patients with  
40                    severe sensory impairment excluded)
- 41                    - Inouye (1993) (3 key RFs included in the analysis; not  
42                    polypharmacy; illness severity included)
- 43

- 1 • Possibly confounded: not enough of important risk factors taken into account  
2 in the multivariate analysis (2/4) but a sufficient ratio of events to  
3 covariates
- 4 ○ Age and cognitive impairment
    - 5 - Rudolph (2007) had a ratio of 16 (1-2 RFs: age included
    - 6 in the analysis and patients with dementia (not mild
    - 7 cognitive impairment) were excluded)
  - 8 ○ Age and polypharmacy
    - 9 - Ouimet (2007) had a ratio of 19 (2 RFs: age included in
    - 10 the analysis and polypharmacy constant because patients
    - 11 in ICU; illness severity also included)
    - 12 - Redelmeier (2008) had a ratio of 200 (2 key RFs: age
    - 13 included in analysis and polypharmacy likely constant
    - 14 because surgical patients)
  - 15 ○ Cognitive impairment and polypharmacy
    - 16 - Lin (2008) had a ratio of 10 (2 RFs: patients with
    - 17 dementia excluded and polypharmacy constant because
    - 18 patients in ICU)
  - 19 ○ Cognitive impairment and sensory impairment
    - 20 - Inouye (2007) had a ratio of 10 (2 RFs: dementia and
    - 21 vision impairment included in analysis; illness severity also
    - 22 included)
- 23
- 24 • Possibly confounded: not enough of important risk factors taken into account  
25 in the multivariate analysis (2/4) and not high enough ratio of events to  
26 covariates
- 27 ○ Age and cognitive impairment
    - 28 - Hofsté (1997) had a ratio of 9 (2 key RFs: age included in
    - 29 analysis and cognitive impairment constant because
    - 30 elective cardiac operations)
    - 31 - Korevaar (2005) had a ratio of 4 (age and cognitive
    - 32 impairment included in the analysis)
    - 33 - Leung (2007) had a ratio of 3 (age and cognitive
    - 34 impairment included in the analysis)
    - 35 - Kazmierski (2006) had a ratio of 2 (2 key RFs included in
    - 36 analysis: age and cognitive impairment included in
    - 37 analysis)
  - 38 ○ Age and polypharmacy
    - 39 - Ely (2007) had a ratio of 8 (2 RFs: age included in the
    - 40 analysis and polypharmacy constant because patients in
    - 41 ICU; illness severity also included)

- 1 - Rolfson (2003) had a ratio of 8 (age was constant due to  
2 narrow age range, and polypharmacy constant because  
3 elective cardiac operations in older patients)
- 4 ○ Cognitive impairment and polypharmacy
- 5 - Pisani (2007) had a ratio of 9 (cognitive impairment  
6 included in the analysis and polypharmacy constant  
7 because patients in ICU; illness severity also included)
- 8
- 9 • Probably confounded: not enough of important risk factors taken into  
10 account in the multivariate analysis (1/4), but did have a ratio of events  
11 to covariates of at least 10
- 12 ○ Cognitive impairment
- 13 - Furlaneto (2006) had a ratio of 15 (cognitive impairment  
14 included in the analysis)
- 15 - Pompei (2002) had a ratio of 16 and 21 (cognitive  
16 impairment included in the analysis; comorbidity also  
17 included)
- 18
- 19 • Probably confounded: not enough of the important risk factors taken into  
20 account in the multivariate analysis (1/4), and did not have high enough  
21 ratio of events to covariates
- 22 ○ Age
- 23 - Caeiro (2001) had a ratio of 7 (age included in the  
24 analysis)
- 25 - Levkoff (1988) had a ratio of 6 (age included in the  
26 analysis)
- 27 - Yildizeli (2005) had ratio of less than 1 (age included in  
28 the analysis)
- 29 ○ Cognitive impairment
- 30 - Zakriya (2008) had a ratio of 8 [patients with dementia  
31 were excluded but method of assessment not stated; illness  
32 severity also included (as American Society of  
33 Anesthesiologists, ASA grade)]
- 34
- 35 • Confounded: no important risk factors taken into account in the multivariate  
36 analysis (0/4) and did not have a high enough ratio of events to  
37 covariates
- 38 ○ Edlund (2001) had a ratio 4 for incident delirium
- 39

1 The McCusker (2001) study reporting delirium severity used analyses at various  
 2 times reflecting different states (repeated measures multivariate analyses, using  
 3 the previous most recent severity score as a factor in the multivariate analysis).  
 4 The GDG considered this to be an acceptable method.

5  
 6 Overall, the risk of bias was considered for each cohort study, and ratings were  
 7 given of high, moderate and low quality, and biased/confounded.  
 8

9 • Six studies were judged to be biased and therefore not considered further:

- 10 ○ Edlund (2001): no key risk factors
- 11 ○ Furlaneto (2006): 57% prevalent delirium included
- 12 ○ Levkoff (1988): inadequate method of delirium assessment;  
 13 retrospective
- 14 ○ Pompei 1994 (Yale): 56% prevalent delirium included
- 15 ○ Redelmeier (2008): inadequate method of delirium assessment;  
 16 retrospective
- 17 ○ Yildizeli (2005): not enough patients for multivariate analysis  
 18 (ratio less than 1); retrospective

19 • Twelve studies were given a low overall rating and were treated with  
 20 caution (evaluated in sensitivity analyses) (Andersson 2001; Caeiro  
 21 2004; Inouye 1993; Kazmierski 2006; Korevaar 2005; Leung 2007;  
 22 McCusker 2001; Pompei 1994 (Chicago); Santos 2004; Sheng 2006;  
 23 Veliz-Reissmüller 2007; Zakriya 2008)

24 • Fifteen studies had a moderate rating; (Böhner 2003; Bucerius 2004;  
 25 Goldenberg 2006; Ely 2007; Hofsté 1997; Inouye 2007; Levkoff 1992;  
 26 Lin 2008; Ouimet 2007; Pandharipande 2006; Pisani 2007; Ranhoff  
 27 2006; Rolfson 1999; Rudolph 2007; Schor 1992)

28 • No studies had a high rating  
 29

30 **6.4.3 Risk factors investigated by the cohort studies (multivariate analyses)**

31 The following risk factors have been investigated in the included studies:  
 32

33 **6.4.3.1 Patient characteristics**

- 34 • Age (21 studies)
- 35 • Cognitive impairment and/or dementia (14 studies)
- 36 • Sensory impairment (7 studies)
- 37 • Polypharmacy (2 studies)
- 38 • Dehydration (5 studies)
- 39 • Severity of illness (5 studies)
- 40 • Comorbidity (4 studies)

- 1 • Sex (7 studies)
- 2 • Electrolyte disturbance (2 studies)
- 3 • Depression (6 studies)
- 4 • Infection (5 studies)
- 5 • Fracture on admission (1 study)
- 6 • Mobility (1 study)
- 7 • Continence (1 study)
- 8 • Constipation (no studies)
- 9 • Sleep deprivation (no studies)
- 10
- 11 **6.4.3.2 Environmental**
- 12 • Pre-hospital setting (3 studies)
- 13 • Hospital unit: ICU, surgery, medical, oncology, long-term care, mixed (1
- 14 study)
- 15 • Recent room change (1 study)
- 16 • Room type: private, semi-private, ward (1 study)
- 17 • Stimulation: based on the distance of the room from the nurses station (1
- 18 study)
- 19 • Same room (1 study)
- 20 • Single room (1 study)
- 21 • Surroundings not well lit (1 study)
- 22 • Surroundings sound too noisy/quiet (1 study)
- 23 • Radio/TV on (1 study)
- 24 • Clock/watch (1 study)
- 25 • Calendar (1 study)
- 26 • Personal possessions present (1 study)
- 27 • Wearing glasses (1 study)
- 28 • Using hearing aid (1 study)
- 29 • Family present (1 study)
- 30 • Isolation (because of infection risk) (1 study)
- 31
- 32 **6.4.3.3 Procedural**

- 1           • Type of surgery (5 studies)
- 2           • Iatrogenic interventions (2 studies)
- 3           • Physical restraint (2 studies)
- 4

#### 5   **6.4.4 Outcomes**

6           The studies measured the following outcomes:

- 7           • Incidence of delirium
- 8           • Duration of delirium
- 9           • Severity of delirium

10

### 11   **6.5 Results**

#### 12   **6.5.1 Patient related risk factors**

##### 13   **6.5.1.1 Setting**

###### 14       Pre-hospital setting as a risk factor for the incidence of delirium

15       Two studies included pre-hospital setting in their multivariate analysis (Andersson  
16       2001, low; Schor 1992) and one study (Levkoff 1992) reported results  
17       separately for patients from long-term care and from the community, and also  
18       carried out a multivariate analysis in which pre-hospital long-term care was  
19       included (the other factors were age, sex, pre-existing cognitive impairment and  
20       illness severity; and patients with severe sensory impairment were excluded).

21

22       The Andersson (2001) study (low rating) found no significant effect of sheltered  
23       housing relative to the person's own home, and Schor (1992) (moderate rating)  
24       found no significant effect of pre-hospital long-term care (the other risk factors  
25       were age, prior cognitive impairment, fracture on admission, sex, infection, pain  
26       (poorly controlled), neuroleptic use, and narcotic use). In neither case were data  
27       reported, although the Schor (1992) study reported the odds ratio adjusted for  
28       age and sex only - which is a low evidence rating - OR 2.54 (95%CI 1.38 to  
29       4.67), and was statistically significant. The Levkoff (1992) study (moderate  
30       rating), however, found a statistically significant effect of long-term care on the  
31       incidence of delirium developing in hospital: OR 2.16 (95%CI 1.15 to 4.1).

32

33       The Levkoff (1992) study mostly analysed the data using separate analyses for  
34       the two pre-hospital groups of long-term care and the community, and as will be  
35       seen in subsequent risk factor analyses, there were large differences between  
36       the two groups. The GDG stated that dementia and comorbidity would likely be  
37       higher in people from long-term care settings.

38

###### 39       Setting as a risk factor for increased severity of delirium

40       For severity of delirium, one large study (McCusker 2001: low; n=587 time  
41       dependent states) considered the effect of different hospital units, using a  
42       repeated measures multivariate analysis. At any given time, patients could be in

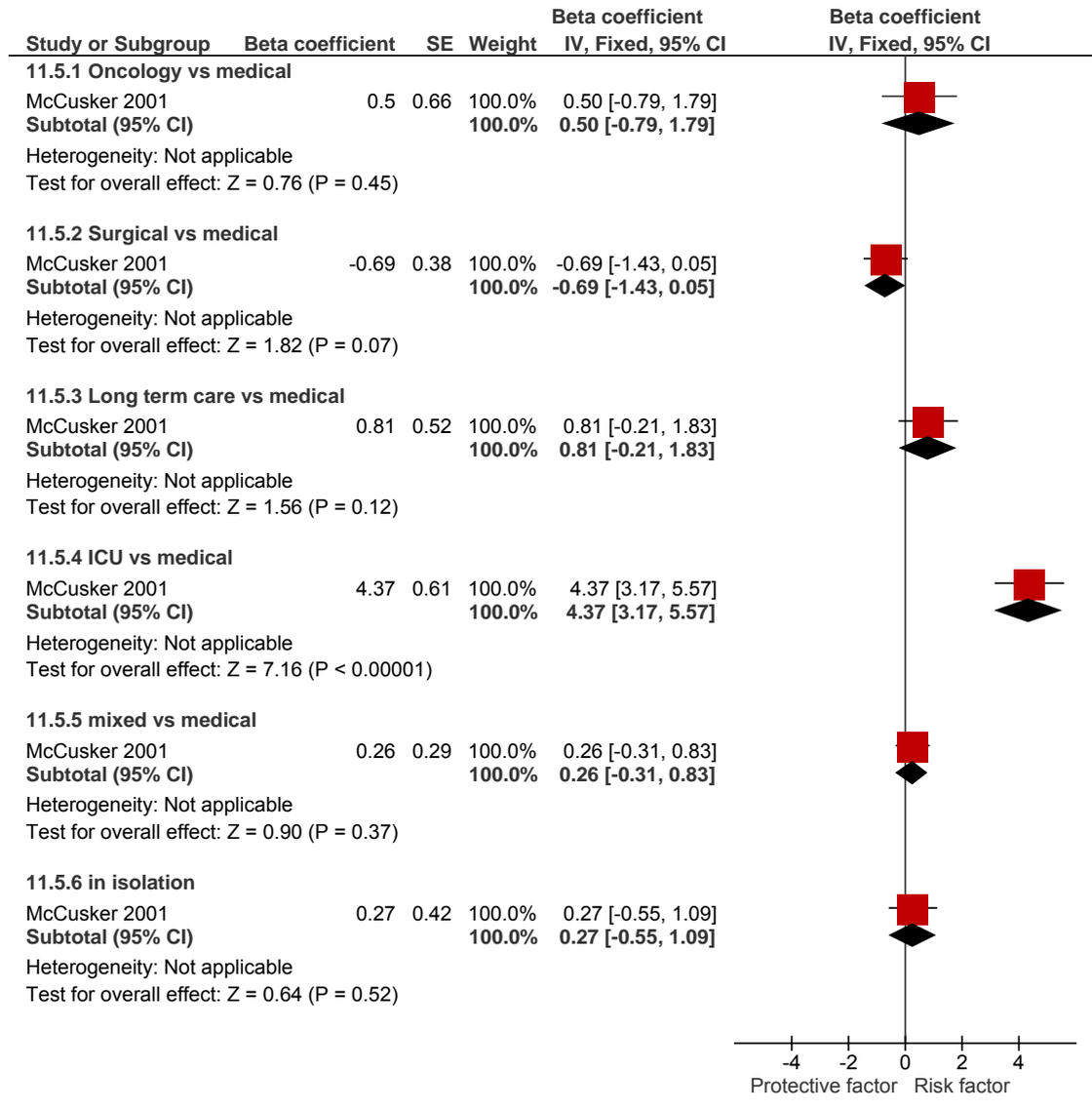
1 long-term care, long-term care /medical, or in hospital wards (subdivided into  
2 general medical, oncology, surgery and ICU). Numbers of patients who had  
3 spent time in each unit were as follows:

- 4 • ICU ( $20/587 = 3\%$ )
- 5 • Surgery ( $81/587 = 14\%$ )
- 6 • General medical ( $281/587 = 48\%$ )
- 7 • Oncology ( $20/587 = 3\%$ )
- 8 • Long-term care ( $34/587 = 6\%$ )
- 9 • Mixed long-term care/medical ( $151/587 = 26\%$ )

10  
11 Thus, we would expect some uncertainty around the results for ICU (3%),  
12 oncology (3%) and long-term care (6%). Results from the multivariate analysis  
13 (with medical ward as the reference) are reported in figure 6.1 and show  
14 significant differences only for patients in ICU. However, this is likely to be of  
15 limited reliability because only a small proportion was in ICU. Furthermore, the  
16 GDG considered it likely that the ICU status was a proxy measure for  
17 polypharmacy and/or severity of illness, neither of which were included in the  
18 multivariate analyses.

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Figure 6.1: hospital unit as a risk factor



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**Summary of setting as a risk factor for delirium**

The evidence regarding the risk factor, long-term care setting prior to hospitalisation, is inconsistent for the incidence of delirium. The evidence is inconclusive for the effect of setting on the severity of delirium, although patients in ICU may be at higher risk than patients in medical wards.

**6.5.1.2 Age**

Seventeen studies presented data on age in their multivariate analyses, see table 6.5 (Andersson 2001 (low rating); Böhner 2003; Bucerius 2004; Caeiro 2004 (low); Ely 2007; Goldenberg 2006; Hofsté 1997; Kazmierski 2006 (low); Leung 2007 (low); Levkoff 1992; McCusker 2001 (low); Pandharipande 2006; Ranhoff 2006; Rudolph 2007; Santos 2004 (low); Schor 1992; Sheng 2006 (low)) (figures 6.2 and 6.3). Four other studies also included age as a risk factor

1 in their multivariate analyses, but did not report any data (Korevaar 2005 (low  
2 rating); Inouye 1993 (low), Ouimet 2007 (moderate), Veliz-Reissmüller  
3 2007(low). It was stated that age was not a significant risk factor in the studies  
4 by Ouimet 2007 and Inouye 1993.

5  
6 One study carried out a 'Markov regression', which was a regression analysis  
7 that included the patient's cognitive state 24 hours previously. The study  
8 reported transitions to delirium and plotted graphically the probability of  
9 developing delirium versus age (Pandharipande 2006).

10  
11 One of the studies investigated the duration of delirium (Ely 2007) (figure 6.4)  
12 and one investigated the severity of delirium (McCusker 2001; low) (figure 6.5);  
13 the rest evaluated incidence of delirium.

14  
15 The standard error for the Böhner (2003) study was calculated from its p-value:  
16 confidence intervals were not reported for the odds ratio (but were for the beta  
17 coefficient).

18  
19 Table 6.5: patient ages in 17 studies that conducted multivariate analyses

Study	Age range	Study	Age range
Bucerius	54-75 (+/- SD)	Caeiro	24-86
Rudolph	63-75	Schor	73-88 (+/-)
Santos	66-78	McCusker	76-90 (+/-)
Leung	66-78 (+/-)	Ranhoff	60-94
Ely	31-79	Sheng	65-95
Kazmierski	25-81	Levkoff inst	80-95 (+/-)
Hofste	29-83	Andersson	65-96
Bohner	NS	Goldenberg	66-98
Levkoff com	71-85 (+/-)	Pandharipande 2006	25-90 (graph)

20 +/- indicates that the range was calculated from the mean +/- one standard  
21 deviation

22  
23 We note that, of these studies, nine were in patients admitted for surgery  
24 (Andersson 2001; Böhner 2003; Bucerius 2004; Goldenberg 2006; Hofsté  
25 1997; Kazmierski 2006; Leung 2007; Rudolph 2007; Santos 2004), three were  
26 in patients admitted to ICUs (Ely 2007; Pandharipande 2006; Ranhoff 2006),  
27 three were conducted in patients from medical wards (Caeiro 2004; McCusker  
28 2001; Sheng 2006), and the remaining two studies were in patients from both  
29 medical and surgical wards (Levkoff 1992; Schor 1992).  
30

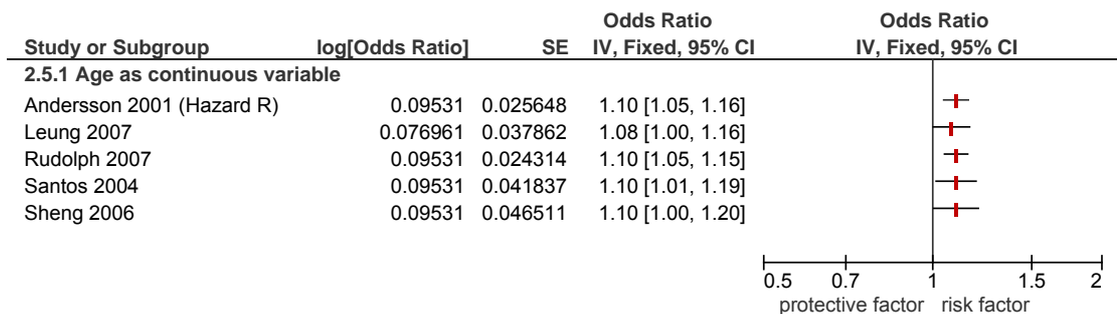
### 31 **Age as a risk factor for the incidence of delirium**

32 Fifteen studies investigated age as a risk factor for the incidence of delirium.

- 1 • Five studies evaluated age as a continuous variable (Andersson 2001, low;  
2 Leung 2007, low; Rudolph 2007; Santos 2004, low; Sheng 2006, low);  
3 the age range across all these studies was 63 to 96 years
- 4 • One study reported the probability of developing delirium as a function of  
5 age, between the ages of 25 and 90 years. Although the study reported  
6 the odds ratio for age as a continuous variable, this was not included in  
7 the analysis because of the non-linearity over the age range  
8 (Pandharipande 2006)
- 9 • Three studies evaluated age over 65 years versus age below 65 years  
10 (Böhner 2003; Caeiro 2004, low; Kazmierski 2006, low)
- 11 • One study evaluated age 70 years and over versus age below 60 years  
12 (Hofsté 1997)
  - 13 ○ We note that the Hofsté (1997) study did not report the category  
14 60 to 69 years in the multivariate analysis (and for the separate  
15 cognitive disorders analysis there are other categorical variables  
16 not reported). Therefore this study should be treated with caution  
17 for age as a risk factor.
- 18 • Four studies evaluated age over 80 versus age below 80 years  
19 (Goldenberg 2006 (age over 81); Levkoff 1992 community and  
20 institution; Ranhoff 2006; Schor 1992)
- 21 • The study by Bucerius (2004) evaluated three comparisons of categorical  
22 age variables (which we have inverted to allow for comparison with the  
23 other studies): over 70 versus under 50, over 70 versus 50-59 years, and  
24 over 70 versus under 60

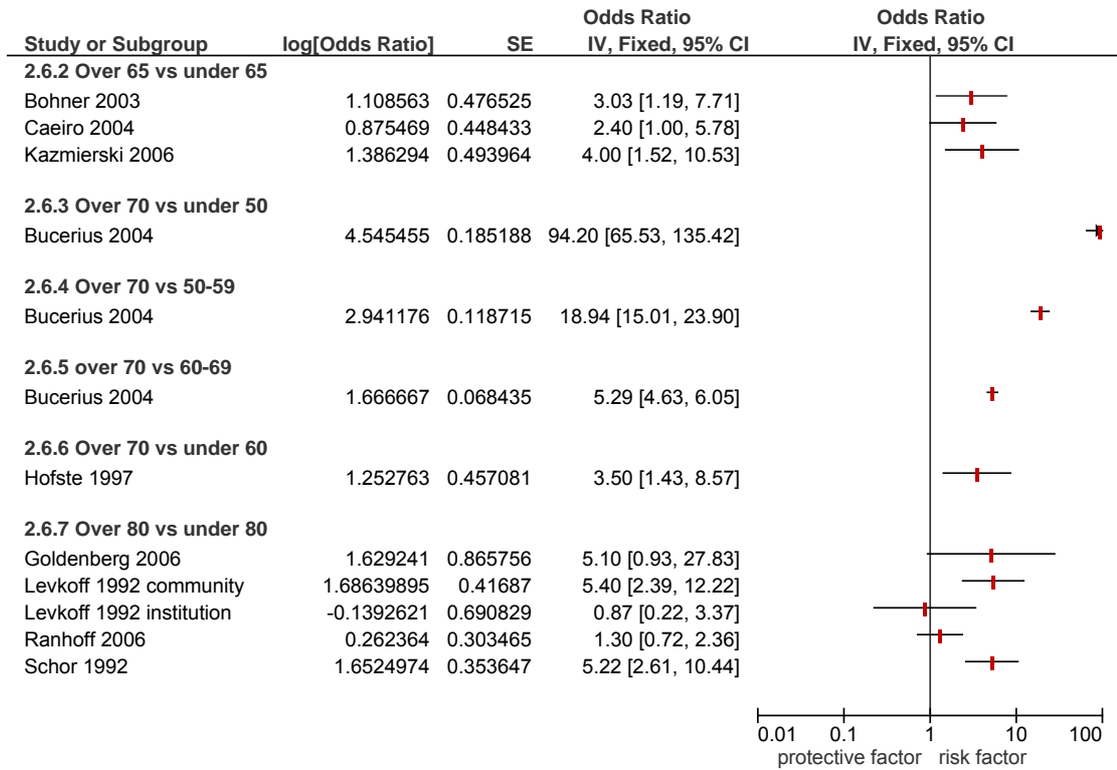
25 The results are reported in Figures 6.2 and 6.3a, with a sensitivity analysis  
26 (excluding low quality studies) shown in figure 6.3b.

27  
28 Figure 6.2: age as a risk factor: incidence of delirium



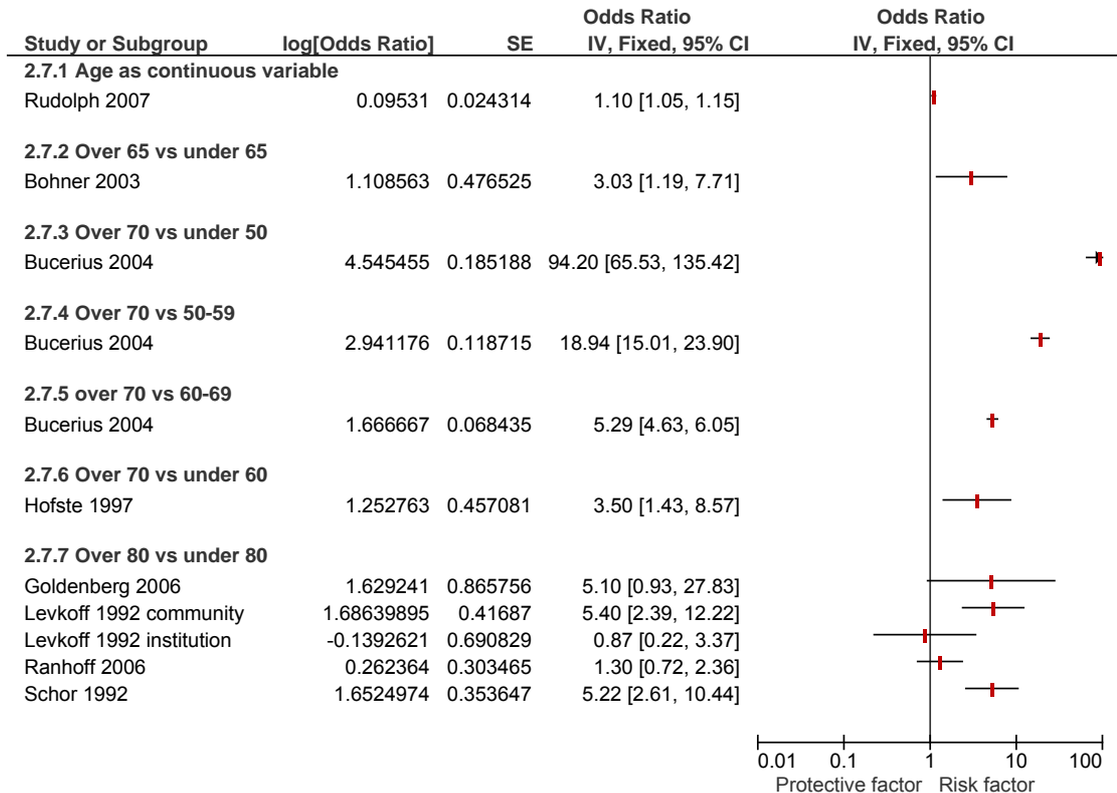
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1 Figure 6.3a: age as a risk factor: incidence of delirium



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3

4 Figure 6.3b: age: incidence of delirium excluding studies with a low rating



5

The sensitivity analysis in figure 6.3b showed no important differences compared with figures 6.2 and 6.3a, and so it was decided to use all the data.

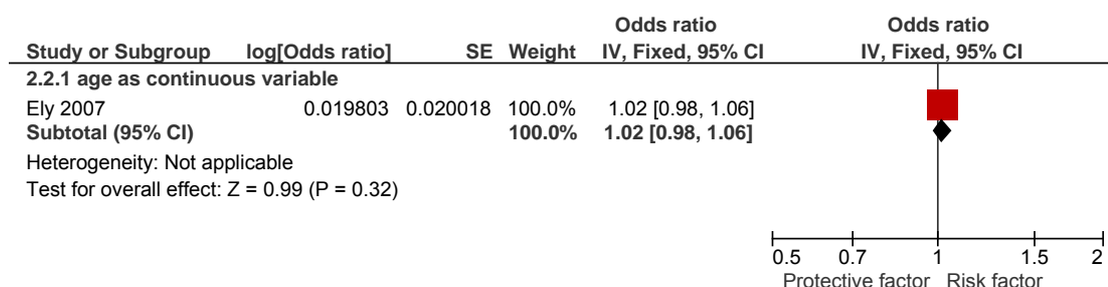
For the age cut-off of 80 years, there was heterogeneity. However, the GDG noted that the mean age in the Levkoff (1992) institution group was 87.4 years and only 11.4% patients were younger than 80 years; suggesting that the age range may not have been large enough to allow conclusions to be derived. The Ranhoff (2006) study was the only one investigating the effect of age (on the incidence of delirium) that was conducted in an ICU setting; the GDG suggested that the effects of illness would be likely to overshadow the effects of age in this setting – the study had not included illness severity in the multivariate analysis, although it had taken account of polypharmacy. Following discussion, the GDG agreed that the effect of age over 80 years was best described by the other three studies.

The GDG wished to define a cut-off point for age as a risk factor and noted that the studies reported different age thresholds. Further information was provided by one moderate quality study (Pandharipande 2006), which reported the probability of developing delirium as a function of age. This probability showed a non-linear pattern across the age range 25 to 90 years. Between the ages of 25 and about 48 years there was a steady increase in the probability, then between 48 and 65 years the graph showed a plateau (same probability independent of age). Finally, above 65 years the probability increased rapidly. This study is the only one to demonstrate the importance of age 65 years as a cut off for age as a risk factor.

Age as a risk factor: increased duration of delirium

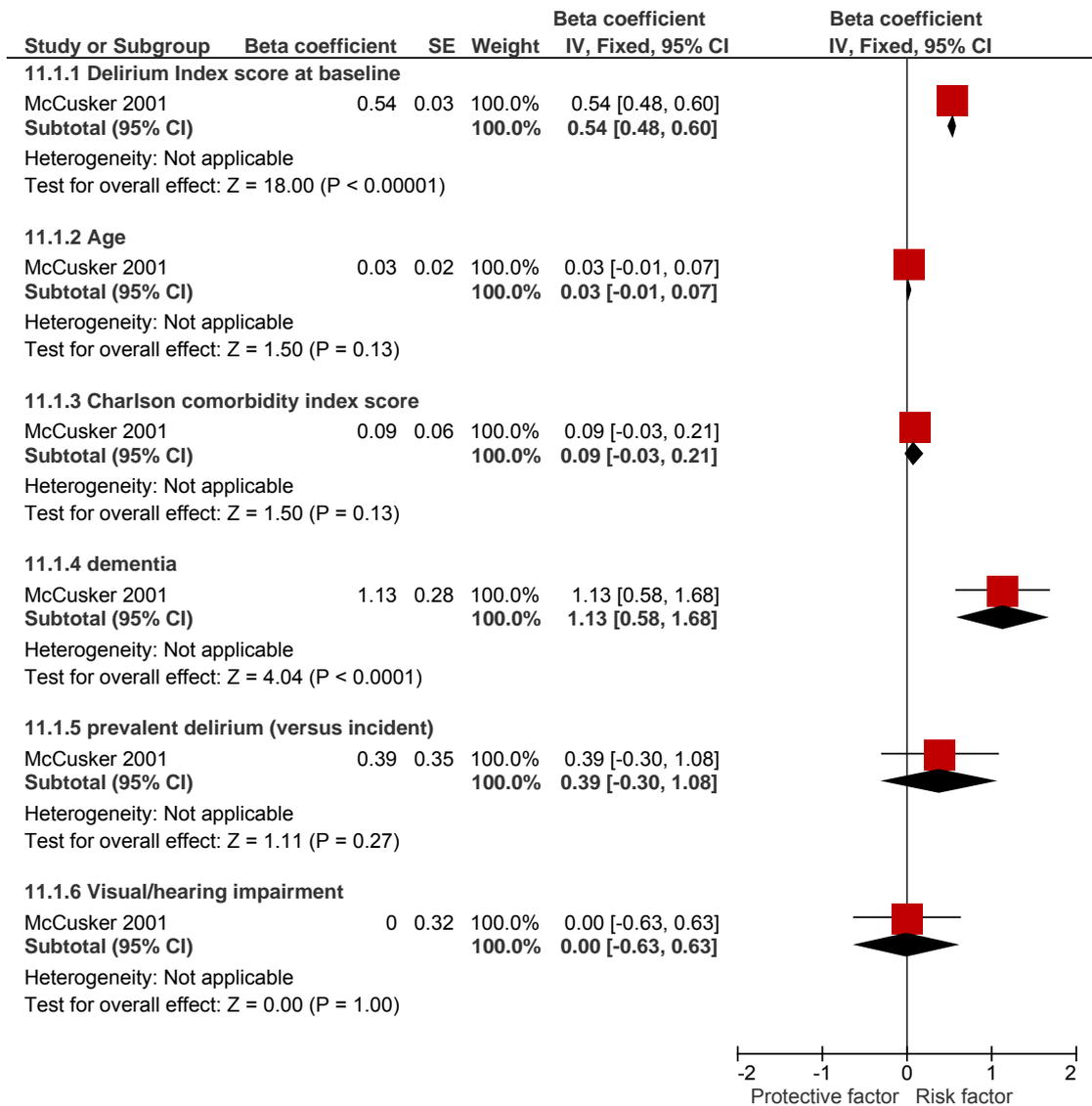
One small study (Ely 2007; n=47) investigated the effect of age as a continuous variable on the duration of delirium, for patients aged 31 to 79 years. We note that this study (with a moderate rating) was conducted in ICU in mechanically ventilated patients. There was no significant effect of age as a continuous variable on the duration of delirium (figure 6.4); OR 1.02 (95%CI 0.98 to 1.06).

Figure 6.4: age as a risk factor: duration of delirium



1 Age as a risk factor: increased severity of delirium  
 2 One large study (McCusker 2001; low, n=444) investigated the effect of age as  
 3 a continuous variable on the severity of delirium, for patients of mean age 83.3  
 4 years (SD 7.0). The effects of different risk factors are shown in figure 6.5,  
 5 reporting the beta coefficient representing the estimated difference in Delirium  
 6 Index scores between the independent variable and the reference category. For  
 7 age, as a continuous variable, there was no significant effect: beta coefficient  
 8 0.03 (95% CI -0.01 to 0.07).  
 9

10 Figure 6.5: patient characteristics as risk factors: severity of delirium



11

1

2 **Summary for age as a risk factor**

3 Thus, the following summary can be given:

- 4 • For age as a continuous variable, the odds ratio for incidence of delirium  
5 ranged from 1.08 to 1.10. This means that for every year increase in  
6 age the odds of having delirium increases by a factor of 1.08 to 1.10.  
7 Taking the 1.10 value, for a 10 year increase in age, the odds increases  
8 by  $(1.10)^{10}$ , which is 2.59. We note that the results are consistent over a  
9 range of studies, and are likely to be valid. The age range covered by  
10 the studies was 63 to 96 years.
- 11 • The odds ratio for delirium incidence for a cut-off point of age 65 years  
12 was 3.03 (95%CI 1.19 to 7.71) for the only study (Böhner 2003) that  
13 was not of low quality (this value was derived from the quoted beta  
14 coefficient of 1.11 (SE 0.468).
- 15 • Age was a significant risk factor for incidence of delirium for most (3/5) of  
16 the studies when a cut-off point of age 80 years was taken, with the OR  
17 ranging from 0.87 (95%CI 0.22 to 3.3) to 5.40 (95%CI 2.4 to 12.3).  
18 There appeared to be significant heterogeneity amongst these studies,  
19 with two studies not showing a significant effect of age (Ranhoff 2006  
20 and Levkoff 1992 institution (in patients who had come from a long-term  
21 care setting)), and three studies showing a similar significant odds ratio  
22 around 5.
- 23 ○ The GDG noted that the mean age in the Levkoff (1992)  
24 institution group was 87.4 years and only 11.4% patients were  
25 younger than 80 years; suggesting that the age range was not  
26 large enough to allow conclusions to be derived.
  - 27 ○ The Ranhoff (2006) study was conducted in an ICU setting; the  
28 GDG suggested that the effects of illness would be likely to  
29 overshadow the effects of age in this setting, and noted that  
30 illness severity was not included in the multivariate analysis for  
31 this study, even though polypharmacy was.
- 32 • One moderate quality study (Pandharipande 2006) examined the  
33 variation across the age range 25 to 90 years, of the probability of  
34 developing delirium, which showed age 65 years to be a point above  
35 which the probability increased rapidly, and this was taken as the age  
36 cut-off.
- 37 • There was no significant effect of age as a continuous variable on the  
38 duration of delirium, over the range 31 to 79 years, in one small study  
39 (n=47) in mechanically ventilated patients in ICU; OR 1.02 (95%CI 0.98  
40 to 1.06)
- 41 • There was no significant effect of age as a continuous variable on the  
42 severity of delirium, for patients of mean age 83.3 years (SD 7.0), in one  
43 large low quality study (n=444); beta coefficient 0.03 (95% CI -0.01 to  
44 0.07).

1

2 **6.5.1.3 Cognitive impairment and/or dementia**

3 Fourteen studies evaluated cognitive impairment and/or dementia in their  
4 multivariate analyses (Böhner 2003; Goldenberg 2006; Inouye 1993, low;  
5 Inouye 2007; Kazmierski 2006, low; Korevaar 2005, low; Levkoff 1992;  
6 McCusker 2001, low; Pisani 2007; Pompei 1994, low; Ranhoff 2006; Schor  
7 1992; Sheng 2006, low; Veliz-Reissmüller 2007, low) (figure 6.7). In the  
8 study by Pompei (1994), data from only one trial (the Chicago hospital)  
9 were reported because the Yale-New Haven hospital data was judged to be  
10 biased.

- 11 • Eight studies used an MMSE score:
  - 12 ○ below 18 cut off for patients at discharge (Ranhoff 2006)
  - 13 ○ below 21-24 cut off depending on education (Pompei 1994)
  - 14 ○ below 24 (Goldenberg 2006; Inouye 1993; Inouye 2007;  
15 Kazmierski 2006)
  - 16 ○ below 25 (Böhner 2003)
  - 17 ○ below 28 (Veliz-Reissmüller 2007)
- 18 • Three studies used IQCODE (Pisani 2007: above 3.3; McCusker 2001:  
19 above 3.5; Korevaar 2005: above 3.9) IQCODE
- 20 • Two studies did not state the assessment method (Schor 1992; Sheng 2006)
- 21 • One study (Levkoff 1992) stated that no systematic standardised method  
22 was used to detect cognitive impairment, with reliance on medical chart  
23 review, which would have led to underreporting

24

25 Of these studies, the GDG did not consider the definition of cognitive impairment  
26 to be reliable in the Veliz-Reissmüller (2007) and Levkoff (1992) studies, so  
27 these were not included in the analysis. Due to the low percentage (8%) of  
28 patients with dementia in the study by Sheng (2006) (table 6.6), the results from  
29 this study were also omitted from the analysis. The Ranhoff (2006) study was  
30 considered in sensitivity analyses because cognitive impairment was assessed at  
31 discharge, in association with activities of daily life measurements.

32

33 We note that of the remaining studies, three were in patients admitted for  
34 surgery (Böhner 2003; Goldenberg 2006; Kazmierski 2006), two were in  
35 patients admitted to ICUs (Pisani 2007; Ranhoff 2006), and the other studies  
36 were in patients from both medical/surgical wards (Inouye 2007; Korevaar  
37 2005; McCusker 2001; Pompei 1994; Schor 1992).

38

39 One of the studies investigated persistent delirium (Inouye 2007) (figure 6.7)  
40 and one investigated the severity of delirium (McCusker 2001) (figure 6.5); the  
41 rest evaluated incidence of delirium.

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43 We note that the Inouye (1993) study excluded people with severe underlying  
44 dementia.

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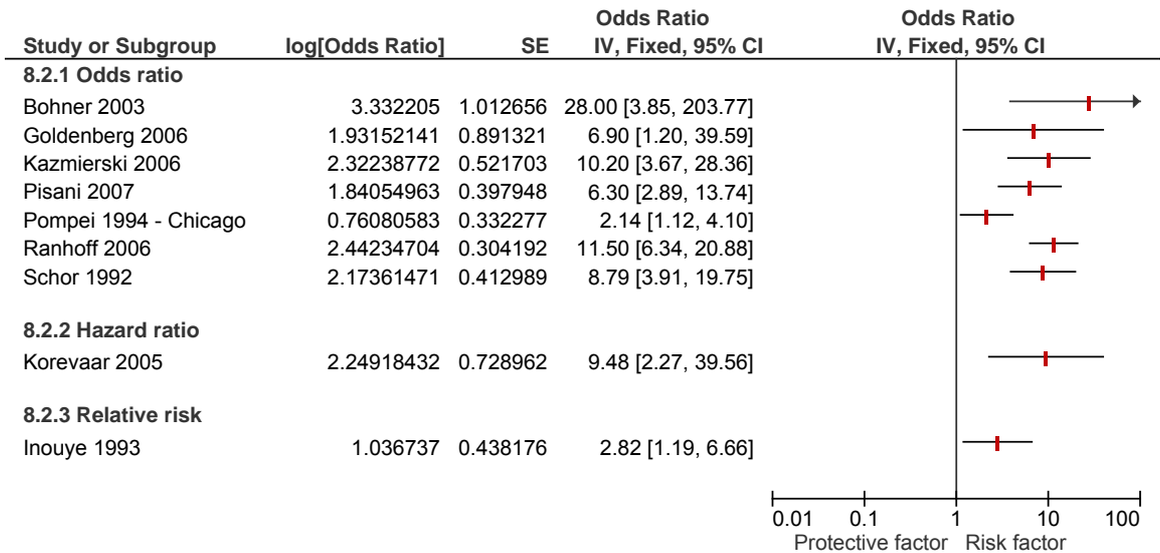
The standard error for the Böhner (2003) study was calculated from its p-value: confidence intervals were not reported.

Table 6.6: cognitive impairment and/or dementia in 11 studies that conducted multivariate analyses

Study	Cognitive impairment / dementia	Study	Cognitive impairment / dementia
<b>Goldenberg</b>	<b>69%</b>	<b>Pisani</b>	<b>31%</b>
<b>Inouye 1993</b>	<b>36%</b>	<b>Pompei-Chicago</b>	<b>37%</b>
<b>Inouye 2007</b>	<b>39%</b>	<b>Ranhoff</b>	<b>30%</b>
<b>Kazmierski</b>	<b>53% &amp; 16%</b>	<b>Schor</b>	<b>19%</b>
<b>Korevaar</b>	<b>43%</b>	<b>Sheng</b>	<b>8%</b>
<b>McCusker 2001</b>	<b>60%</b>	<b>Bohner</b>	<b>Not reported</b>

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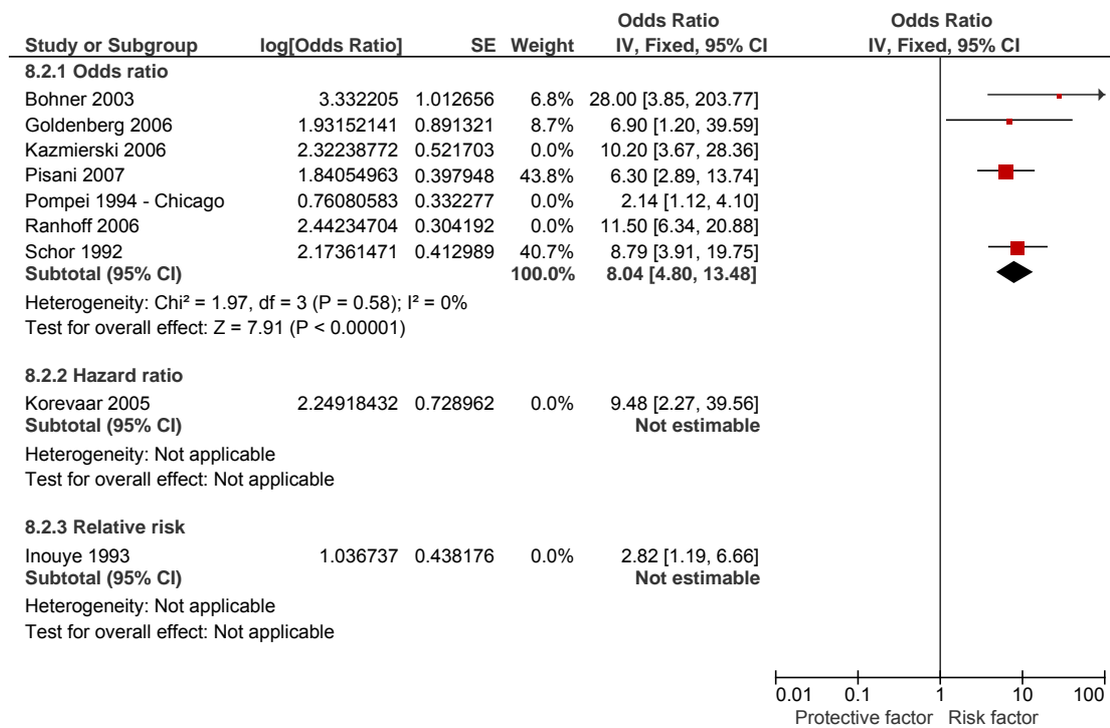
Figure 6.6a: cognitive impairment and/or dementia as a risk factor: incidence of delirium



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Figure 6.6b: cognitive impairment and/or dementia: incidence of delirium excluding studies with a low rating, and also Ranhoff (2006)

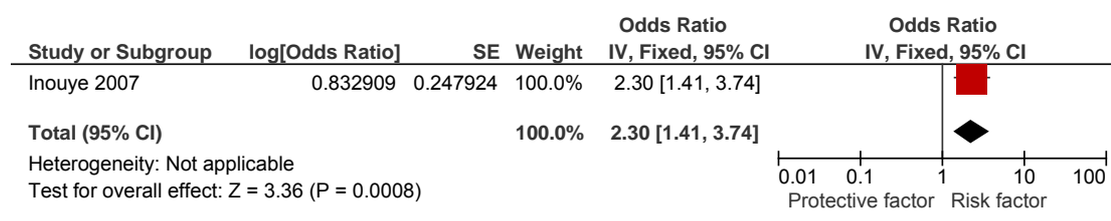
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There was some heterogeneity in figure 6.6a which was removed when only the higher quality studies were analysed (figure 6.6b), so the sensitivity analysis was considered more reliable. There was a large significant effect of cognitive impairment on the risk of delirium. The odds ratio ranged from 6.3 (95%CI 2.9 to 13.8) to 11.5 (95%CI 6.1 to 20.1) with an apparent outlier (Böhner 2003) at OR 28.0 (p value 0.001; beta coefficient 3.33 (SE 0.927)).

#### Cognitive impairment and/or dementia as a risk factor for the incidence of persistent delirium

One moderate quality study investigated the effect of cognitive impairment on the incidence of persistent delirium (Inouye 2007) in 491 patients. We note that these results are from a subpopulation of patients with delirium (n=443). Cognitive impairment was a significant risk factor for persistent delirium (figure 6.7); OR 2.3 (95%CI 1.4 to 3.7).

Figure 6.7: cognitive impairment and/or dementia as a risk factor: persistent delirium



### Cognitive impairment and/or dementia as a risk factor for increased severity of delirium

One large low quality study (McCusker 2001; n=444) investigated the effect of dementia (IQCODE score at least 3.5). Figure 6.5 shows a significant effect; the beta coefficient for the mean difference in delirium severity score is 1.13 (95% CI 0.58 to 1.68).

#### Summary for cognitive impairment/dementia as a risk factor

- Restricting the analysis to the studies that were of higher quality, there was a large significant effect of cognitive impairment on the risk of delirium. The odds ratio ranged from 6.3 (95%CI 2.9 to 13.8) to 11.5 (95%CI 6.1 to 20.1) with an apparent outlier (Böhner 2003) at OR 28.0 (p value 0.001; beta coefficient 3.33 (SE 0.927)).
- For persistent delirium, the odds ratio was 2.30 (95% CI 1.41 to 3.74). We note that these results are from a subpopulation of patients with delirium.
- There was a statistically significant effect of cognitive impairment on the severity of delirium; the beta coefficient for the mean difference in severity of delirium was 1.13 (95% CI 0.58 to 1.68) in one large low quality study.

#### **6.5.1.4 Sensory impairment**

Seven studies included sensory impairment in their multivariate analyses (Andersson 2001, low; Inouye 1993, low; Inouye 2007; McCusker 2001, low; Ranhoff 2006; Sheng 2006, low; Schor 1992).

##### Sensory impairment as a risk factor for incidence of delirium

Two studies presented data on vision impairment in their multivariate analyses (Andersson 2001 – low; Inouye 1993 - low). One other study also evaluated impaired vision as a risk factor in multivariate analysis, but did not report the non-significant results (Sheng 2006, low), and another study (Schor 1992) carried out an analysis adjusted for age and sex for each of vision and hearing loss. Since this Schor (1992) analysis included only age as a key risk factor, we gave it a low quality rating. Results for this study were included in Figure 8 for

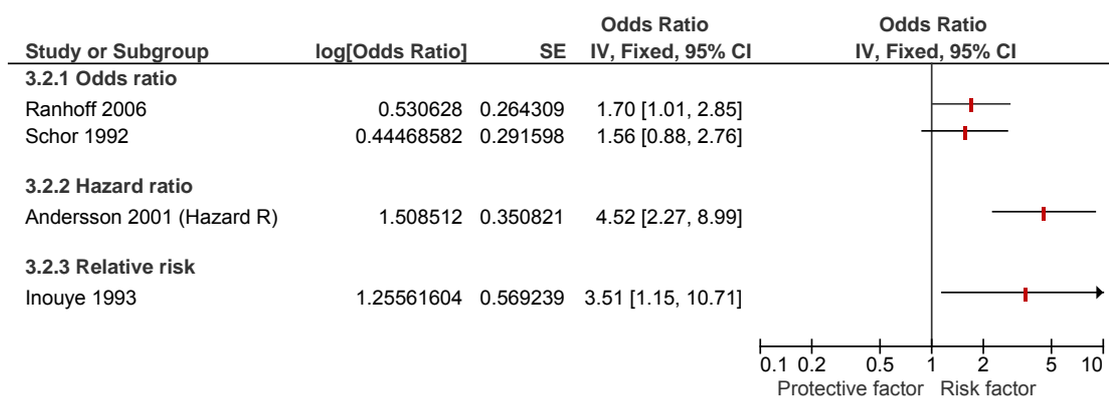
1 vision impairment; hearing impairment had a non significant adjusted odds ratio  
2 of 1.62 (95%CI 0.85 to 3.06).

- 3 • In Andersson (2001) (low rating; n=457 patients), 31% of the surgical  
4 patients had vision impairment and 39% had hearing impairment.
- 5 • In Inouye (1993) (low rating; n=107), 6% of patients in the medical wards  
6 had vision impairment and 54% hearing impairment.
- 7 • In Ranhoff (2006) (moderate rating; n=401), 29% of the ICU patients had  
8 vision impairment (hearing impairment was not reported).
- 9 • In Schor (1992) (low rating for this risk factor; n=291), 33% of patients (in  
10 medical and surgical wards) had vision impairment and 21% hearing  
11 impairment
- 12 • In Sheng (2006) (low rating; n=156), 18% of the patients (in medical  
13 wards) had vision impairment (hearing impairment was not reported)

14  
15 The proportion of only 6% in the Inouye (1993) study is considered likely to lead  
16 to inaccuracy. In both the Andersson (2001) and Inouye (1993) studies, the  
17 authors reported results for impaired vision only; hearing impairment was  
18 included in their multivariate analyses, but the non-significant results were not  
19 reported.

20  
21 Figure 6.8 shows a significant effect of vision impairment on the incidence of  
22 delirium. In the absence of the low quality studies, the remaining large study  
23 (Ranhoff 2006; n= 401) showed a small effect for patients in ICU: OR 1.70  
24 (1.01 to 2.85). We note that this study did not define what was meant by vision  
25 impairment.

26  
27 Figure 6.8: impaired vision as a risk factor: incidence of delirium

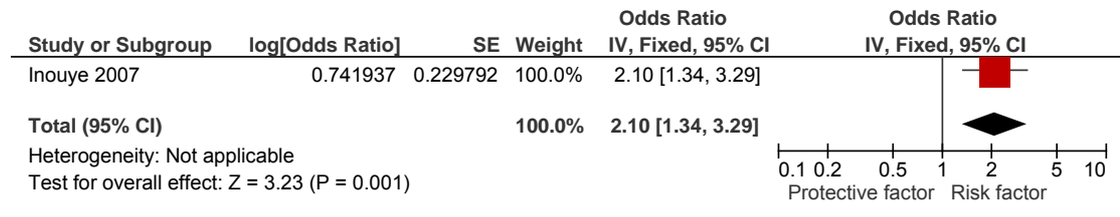


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31  
32 Sensory impairment as a risk factor for incidence of persistent delirium

33 One large, moderate rated study included vision impairment as a risk factor  
34 (Inouye 2007) in 443 patients; 38% of patients in the medical wards had vision

1 impairment (hearing impairment was not reported). There was a significant  
2 effect (figure 6.9), OR 2.1 (95%CI 1.3 to 3.2).

3  
4  
5 Figure 6.9: impaired vision as a risk factor: persistent delirium



6  
7  
8  
9 Sensory impairment as a risk factor for increased severity of delirium

10 One large low quality study (McCusker 2001; n=444) investigated the effect of  
11 sensory impairment; 20% of the patients in the medical wards were reported to  
12 have vision/hearing impairment. Figure 6.5 shows there was no significant effect;  
13 the beta coefficient for the mean difference in delirium severity score is 0 (95%  
14 CI -0.63 to 0.63).

15  
16 Summary for sensory impairment as a risk factor

- 17
- 18 • Restricting the analysis for delirium incidence to the study that was of higher  
19 quality (Ranhoff 2006), this large ICU study showed a small effect of  
20 vision impairment: OR 1.70 (1.01 to 2.85). We note that this study did  
not define what was meant by vision impairment.
  - 21 • For persistent delirium, there was a significant effect in a study that defined  
22 vision impairment carefully; OR 2.1 (95% CI 1.3 to 3.3). We note that  
23 these results are from a subpopulation of patients with delirium.
  - 24 • The beta coefficient for the mean difference in severity of delirium for  
25 vision impairment was not significant in one large low quality study: 0.0  
26 (95% CI -0.63 to 0.63)
  - 27 • There was very limited evidence that hearing impairment was not an  
28 important risk factor for delirium incidence from low quality studies
- 29  
30

31 **6.5.1.5 Polypharmacy**

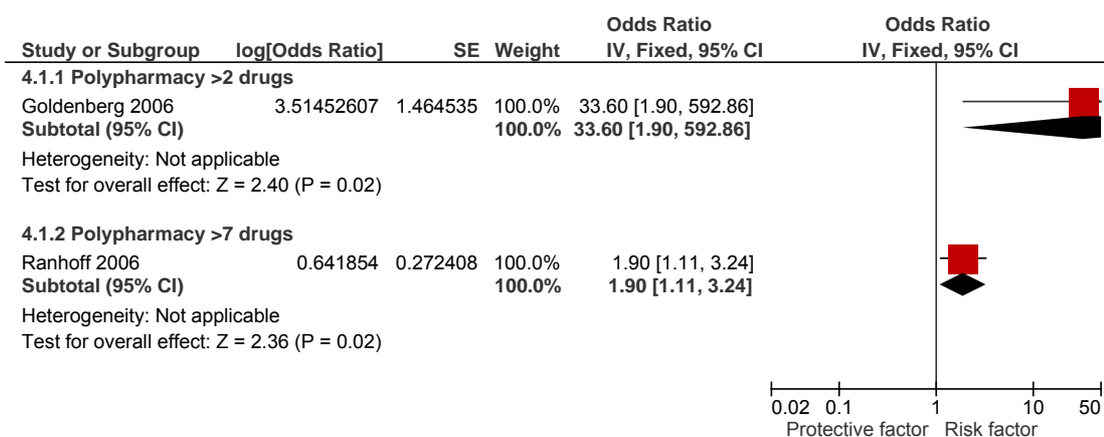
32 Polypharmacy as a risk factor for incidence of delirium

33 Two studies presented data on the number of drugs as a risk factor for the  
34 incidence of delirium in their multivariate analyses (Goldenberg 2006; Ranhoff  
35 2006). In neither case was illness severity or comorbidity included in the  
36 multivariate analyses.

37  
38 In the study by Goldenberg (2006), the use of more than three medications  
39 (other than vitamins) was defined to represent multiple medication use, with 87%  
40 polypharmacy use in this sample. In the study by Ranhoff (2006), the authors  
41 evaluated the maximum concurrent number of drugs (including laxatives) as the

1 following dichotomous variable: 7 or more drugs versus fewer than 7. The mean  
 2 number of drugs used was 8.5 (SD 3.4) in patients with prevalent delirium, 8.0  
 3 (SD3.2) in patients with incident delirium, and 7.3 (SD 3.1) in patients without  
 4 delirium. These studies both had moderate ratings. We note that the small study  
 5 (n=77) by Goldenberg (2006) was in patients admitted for surgery, whereas  
 6 the large study (n=401) by Ranhoff (2006) was conducted in ICU patients, a  
 7 setting in which patients are likely to receive multiple medications. Figure 6.10  
 8 shows a significant effect of polypharmacy on the incidence of delirium for both  
 9 studies, but the confidence interval is very wide for the study with a cut-off point  
 10 of 3 drugs.

11 Figure 6.10: polypharmacy: incidence of delirium  
 12



13 Summary for polypharmacy as a risk factor  
 14  
 15

- 16 • There was little evidence on polypharmacy as a risk factor.
- 17 • The odds ratio was 33.60 (95% CI 1.9 to 591.6) in the study by  
 18 Goldenberg (2006), and 1.9 (95% CI 1.1 to 3.2) in the study by Ranhoff  
 19 (2006).
- 20 • We note that 87% of the patients in the study by Goldenberg (2006) had  
 21 taken more than 3 medications.
- 22 • The GDG stated that more than 7 drugs in an ICU setting was not a useful  
 23 clinical risk factor to assess.

24  
 25 **6.5.1.6 Dehydration**

26 Dehydration as a risk factor for incidence of delirium

27 A widely accepted laboratory measure of dehydration is the disproportionate  
 28 rise in blood urea nitrogen (BUN) to creatinine. This was measured in two studies  
 29 (Inouye 1993, low; Pisani 2007, moderate). Three other studies (Kazmierski  
 30 2006, low; Korevaar 2005, low; Santos 2004, low) recorded the blood urea  
 31 content only; this measure is not considered to have high specificity for  
 32 dehydration.  
 33

1 Three studies presented data on dehydration as a risk factor for the incidence of  
2 delirium in their multivariate analyses (figure 6.11). All of these studies had low  
3 quality ratings. We note that the study by Santos (2004) was in patients  
4 admitted for surgery, and the studies by Inouye (1993) and Korevaar (2005)  
5 were in medical wards.

6 • In the study by Inouye (1993), a baseline blood urea nitrogen/creatinine  
7 ratio of 18 or more was used as an index of dehydration; 67% in the  
8 group with delirium were dehydrated compared with 39% in the group  
9 without delirium (data calculated)

10 • In the study by Korevaar (2005), the mean baseline urea nitrogen (mmol/l)  
11 concentration was 15.9 mmol/l (SD 13.6) in patients with delirium after  
12 acute admission compared with 10.6 mmol/l (SD 6.2) in patients without  
13 delirium

14 • In the study by Santos (2004), the pre-operative blood urea level ranged  
15 from 15-127 mg/dl; it was on average, 50.63 mg/dl (SD 23.26) in  
16 patients with delirium, and 41.85 (SD 14.39) in patients without delirium

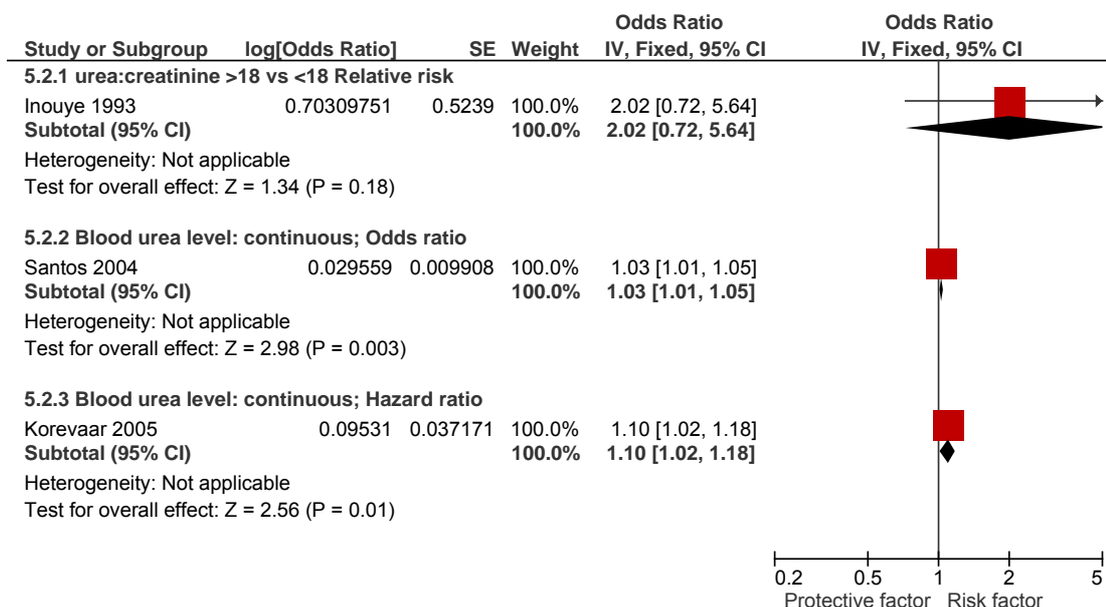
17  
18 In addition, two studies included dehydration as a risk factor in their multivariate  
19 analyses, but did not report the non-significant results (Kazmierski 2006, low;  
20 Pisani 2007, moderate).

21 • In the study by Kazmierski (2006), 5/30 (17%) of delirious patients had a  
22 pre-operative serum urea concentration greater than 50 mg/dl  
23 compared to 6/230 (7%) in patients without delirium; 8% overall

24 • In the study by Pisani (2007), 148/214 (69%) patients with delirium, and  
25 54/90 (60%) patients without delirium, had a ratio of serum urea  
26 nitrogen to creatinine greater than 18 (measured in the first 48 hrs of ICU  
27 admission).

28  
29  
30

1 Figure 6.11: dehydration as a risk factor: incidence of delirium

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34 Summary of dehydration as a risk factor

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- 7
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- 10
- The GDG stated that a urea/creatinine ratio of 18 is difficult to interpret and depends on the units used (e.g. mmol/l), and a high urea level is not specific for dehydration
  - One low quality study recorded the outcome representative of dehydration (urea/creatinine ratio) and the confidence interval was too wide to determine if dehydration was a risk factor for delirium.

11

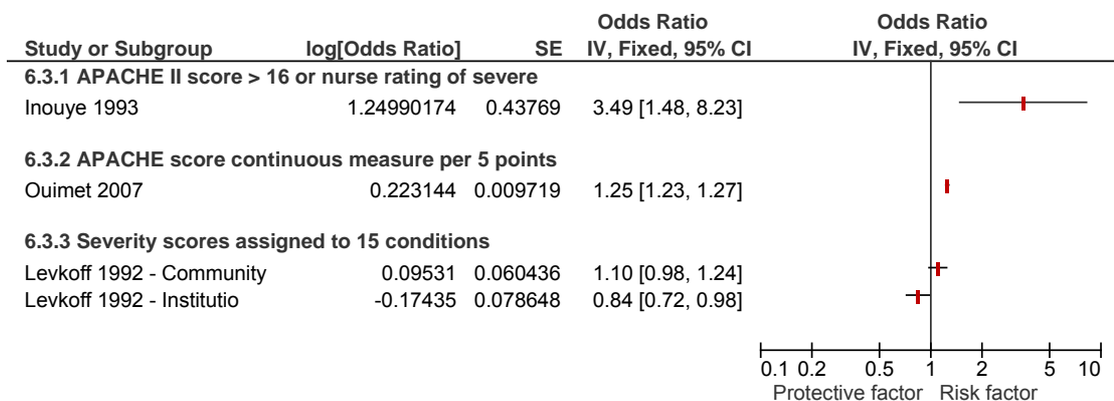
12 **6.5.1.7 Severity of illness**13 Illness severity as a risk factor for incidence of delirium

14 Three studies presented data on illness severity as a risk factor for the incidence of delirium in their multivariate analyses: Inouye (1993) (low), Levkoff (1992) (moderate) and Ouimet (2007) (moderate) (figure 6.12); the Ouimet (2007) study was conducted in ICU. A further ICU study included illness severity as a risk factor in their multivariate analysis, but the non-significant results were not reported (Pisani 2007, moderate). In none of the studies were polypharmacy or comorbidity included in the multivariate analyses.

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- In the study by Inouye (1993), a composite score defined by a nurse rating of 'severe' or an Acute Physiology and Chronic Health Evaluation (APACHE) II score of more than 16 was considered to represent severe illness. In this study, 44% in group with delirium and 10% in group without delirium had 'severe illness' (data calculated). This study was conducted in a medical ward.

- 1           • The Ouimet (2007) study in ICU also used the APACHE II score (0 to 71
- 2           maximum possible) as a continuous variable; the mean score at baseline
- 3           was 16.5 (SD 8.2), range 0 to 59
- 4           • The APACHE II score was also used in the Pisani (2007) study; the mean
- 5           score was 24.7 (SD 6.1) in patients with delirium compared to 20.0 (SD
- 6           5.6) in patients without delirium
- 7           • In the study by Levkoff (1992), an illness severity score was calculated by
- 8           summing the severity scores assigned to 15 medical conditions; they
- 9           ranged from 1 for conditions that were not likely to have an impact on
- 10          the process of care, to 4 for conditions that were imminently life
- 11          threatening (baseline data were not reported). This study was conducted
- 12          in both medical and surgical wards. The GDG noted that this was an
- 13          unvalidated scale, and treated these results with caution.

Figure 6.12: illness severity as a risk factor: incidence of delirium



16

17

18          For the two studies using validated scales (Inouye 1993, low and Ouimet 2007),

19          there was a significant effect of illness severity on the incidence of delirium. The

20          results from the Levkoff 1992 study were considered to be paradoxical by the

21          GDG, and they noted that this study used an unvalidated scale, The GDG

22          decided to remove this study and the low quality one (Inouye 1993) in a

23          sensitivity analysis (not shown). The remaining very large study (n=764), Ouimet

24          2007, showed a significant effect of illness severity as a continuous variable: OR

25          1.25 (95%CI 1.23 to 1.27) per 5 point increase in APACHE II score, or 1.049

26          (95%CI 1.028 to 1.070) per point increase, which is a fairly large effect. The

27          former means that for every 5 points on the APACHE II scale, the odds of

28          delirium increases by 1.25. We note that this remaining study was conducted in

29          ICU patients.

30

31          Illness severity as a risk factor for increased duration of delirium

32          One small, moderate quality study conducted in mechanically ventilated patients

33          in ICU (Ely 2007; n=53) examined the effect of illness severity on the duration of

34          delirium. Illness severity was determined using the APACHE II score, and this had

35          mean scores of 26.8 (SD 8.0) to 27.8 (SD 5.3).

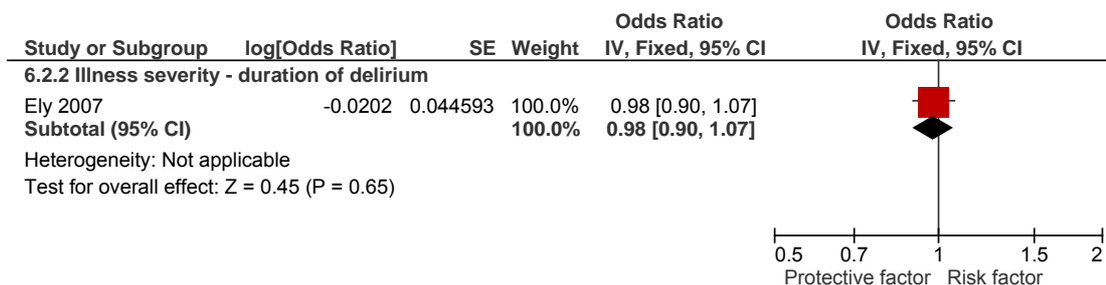
36

37          Results are shown in figure 6.13, and there is no significant effect of illness

38          severity as a continuous factor on the duration of delirium.

39

1 Figure 6.13: illness severity as a risk factor: duration of delirium

2  
34 Summary of illness severity as a risk factor

5 The following summary can be given:

- 6
- 7 • Illness severity had a significant effect on the incidence of delirium in one
  - 8 large study conducted in ICU; for APACHE II scores as a continuous
  - 9 variable, the odds ratio was 1.25 (95% CI 1.23 to 1.27) per 5 point
  - 10 increase, or 1.049 (95%CI 1.028 to 1.070) per point increase.
  - 11 • One low quality non-ICU study showed severity of illness to be a risk factor
  - 12 for the incidence of delirium; patients were assessed to have severe
  - 13 illness if they had an APACHE II score of more than 16
  - 14 • Illness severity did not show a significant effect on the duration of delirium
  - 15 in one small study in mechanically ventilated patients in ICU

16 **6.5.1.8 Comorbidity**17 Comorbidity as a risk factor for incidence of delirium18 Two studies presented data on comorbidity (Andersson 2001; Pompei 1994);  
19 both had a low quality rating.

- 20
- 21 • In Andersson (2001), 10% of patients with 'acute confusional state' (ACS)
  - 22 had four or more diseases compared to 1% of patients without ACS.
  - 23 • In Pompei (1994), we considered the number of Major Diagnostic
  - 24 Categories (MDCs) to be indicative of comorbidity. MDCs related to a
  - 25 major body system (e.g. circulatory or respiratory system), or conditions
  - 26 that impact on more than one body system (e.g. sepsis or major trauma)
  - 27 (a patient with hypertension, ischaemic heart disease, and aortic vascular
  - 28 sclerosis would have three diagnoses but only one MDC). The mean
  - 29 number of MDCs in patients with delirium was 4.2 (SD 1.6) and 2.9 (SD
  - 30 1.5) in patients without delirium.

31 We note that the study by Andersson (2001) was in patients admitted for

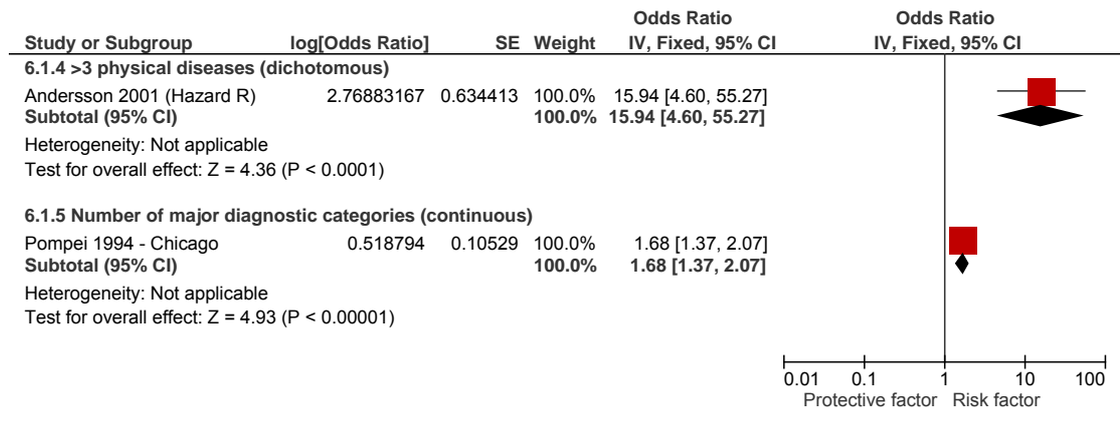
32 surgery, and the study by Pompei (1994) was conducted in patients from both

33 surgical and medical wards. In neither study was polypharmacy or illness

34 severity taken into consideration in the analysis.

35  
36 There was a significant effect of comorbidity on delirium incidence (figure 6.14).  
37

1 Figure 6.14: comorbidity as a risk factor: incidence of delirium



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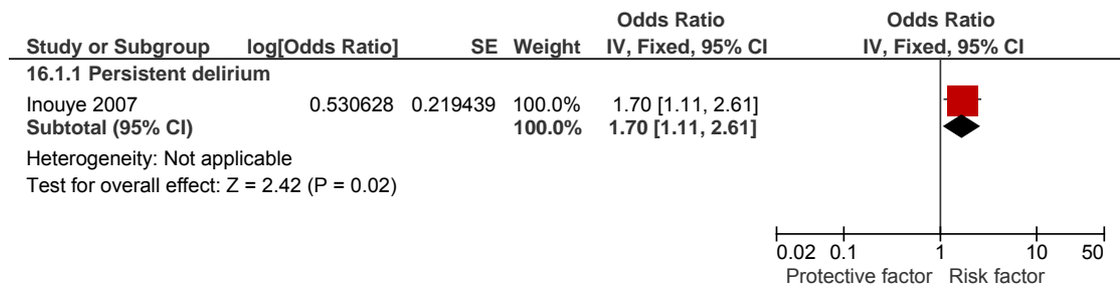
5

6 Comorbidity as a risk factor for incidence of persistent delirium

7 One large, moderate quality study analysed comorbidity as a risk factor (Inouye  
8 2007) in 443 patients. The study was conducted in patients in medical wards, of  
9 whom 29% had a Charlson Comorbidity score of 4 or more, with a mean  
10 baseline score of 2.7 (SD 2.1); the study did not include illness severity or  
11 polypharmacy in the analysis. There was a significant effect of comorbidity on  
12 the incidence of persistent delirium (figure 6.15): OR 1.7 (95%CI 1.1 to 2.6).  
13

14

15 Figure 6.15: comorbidity as a risk factor: persistent delirium



16

17

18

19

20 Comorbidity as a risk factor for increased severity of delirium

21 One large, low quality study (McCusker 2001; n=444) investigated the effect of  
22 comorbidity on the severity of delirium; the study did not include illness severity  
23 or polypharmacy in the analysis. The study was conducted in patients in medical  
24 wards, for whom the mean baseline Charlson Comorbidity score was 2.7 (SD  
25 2.0).

26 Figure 6.5 shows no significant effect; the beta coefficient for the mean  
27 difference in delirium severity score is 0.09 (95% CI -0.03 to 0.21).  
28

28

### 1 Summary of comorbidity as a risk factor

- 2 • Both studies that evaluated incidence of delirium had a low rating, and  
3 their results should be treated with caution, but both showed a significant  
4 effect of comorbidity on delirium incidence
- 5 • For persistent delirium, there was a significant effect of comorbidity (as  
6 measured by the Charlson comorbidity index) in a large moderate  
7 quality study; OR 1.7 (95% CI 1.1 to 2.6). We note that these results are  
8 from a subpopulation of patients with delirium
- 9 • In one large, low quality study, the beta coefficient for the mean difference  
10 in severity of delirium for comorbidity (as measured by the Charlson  
11 comorbidity index) was not significant: 0.09 (95% CI -0.03 to 0.21)

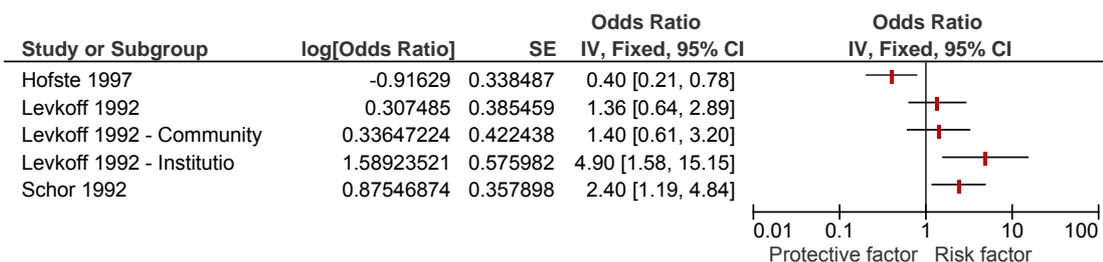
### 13 **6.5.1.9 Sex (gender)**

#### 14 Sex as a risk factor for incidence of delirium

15 Three studies presented data on sex as a risk factor for the incidence of delirium  
16 in their multivariate analyses (Hofsté 1997; Levkoff 1992; Schor 1992) (figure  
17 6.16a). Proportion of male patients in each study is shown in figure 6.16b. All  
18 studies had a moderate quality rating (Hofsté 1997). In addition, four studies  
19 included sex as a risk factor in multivariate analyses, but the non-significant  
20 results were not reported (Andersson 2001 (low); Inouye 1993 (low); Kazmierski  
21 2006 (low); Rudolph 2007 (moderate)).

22  
23 The studies were conducted in surgical patients (Andersson 2001; Kazmierski  
24 2006; Hofsté 1997; Rudolph 2007), and medical/surgical patients (Inouye  
25 1993; Levkoff 1992; Schor 1992).

29 Figure 6.16a: sex (male) as a risk factor: incidence of delirium



32 Table 6.16b: percentage of males in studies that conducted multivariate  
33 analyses

Study	Male	Study	Male
Schor	33%	Inouye	46%
Andersson	34%	Rudolph	53%
Levkoff -	29%	Hofsté	73%

comm

Levkoff- inst 35%

Kazmierski

76%

Summary of sex as a risk factor

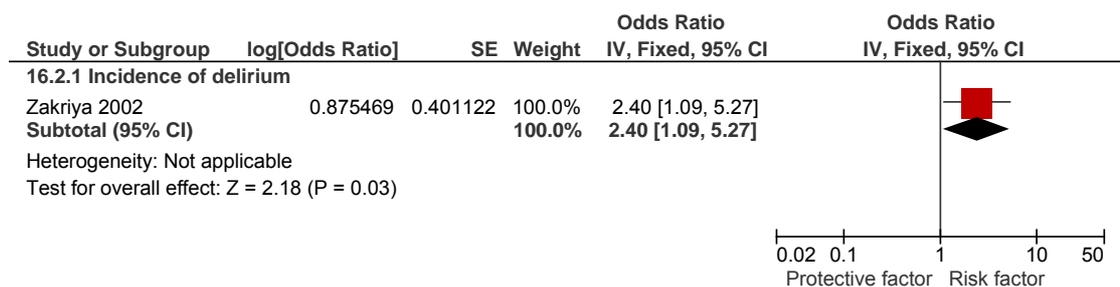
- The odds ratio for male sex ranged from 0.4 (95% CI 0.2 to 0.8) to 4.9 (95%CI 1.6 to 15.3).
- There was heterogeneity amongst these studies with one study showing a significant effect of the risk factor, male sex, one study showing a protective effect of male sex and one study showing a non-significant effect (Levkoff 1992) (community and institutional settings combined).
- The evidence was unable to show if sex is a clinically important risk factor.

**6.5.1.10 Electrolyte disturbance**

One low quality study presented data on electrolyte disturbance as a risk factor for the incidence of delirium in surgical patients in their multivariate analysis (Zakriya 2008) (figure 6.17). In addition, one study included electrolyte disturbance as a risk factor in multivariate analysis, but the non-significant results were not reported (Korevaar 2005). Both studies had a low quality rating.

The study by Zakriya (2008) considered abnormal serum sodium (Na+) (below 135 or above 148 mEq/l) to be indicative of electrolyte disturbance. Overall, 22% of the patients had abnormal serum sodium (data not reported for patients with and without delirium).

Figure 6.17: electrolyte disturbance as a risk factor: incidence of delirium



Due to the low rating of this study, the results should be treated with caution.

Summary

There was low quality evidence to suggest that electrolyte disturbance is a risk factor for delirium, but the absence of other important risk factors in the analysis made this uncertain.

### 1 6.5.1.11 Depression

#### 2 Depression as a risk factor for incidence of delirium

3 Four studies presented data on depression as a risk factor for the incidence of  
4 delirium in their multivariate analyses (Böhner 2003; Inouye 1993; Kazmierski  
5 2006; Pompei 1994) (figure 6.18). The study by Böhner (2003) had a moderate  
6 rating; the three other studies had low ratings. Two further studies included  
7 depression as a risk factor in multivariate analyses, but the non-significant results  
8 were not reported (Leung 2007 (low); Pisani 2007 (moderate)).  
9

10 We note that these studies were conducted in all settings: surgical patients  
11 (Böhner 2003; Kazmierski 2006; Leung 2007) medical/surgical wards (Inouye  
12 1993; Pompei 1994) and ICU patients (Pisani 2007).  
13

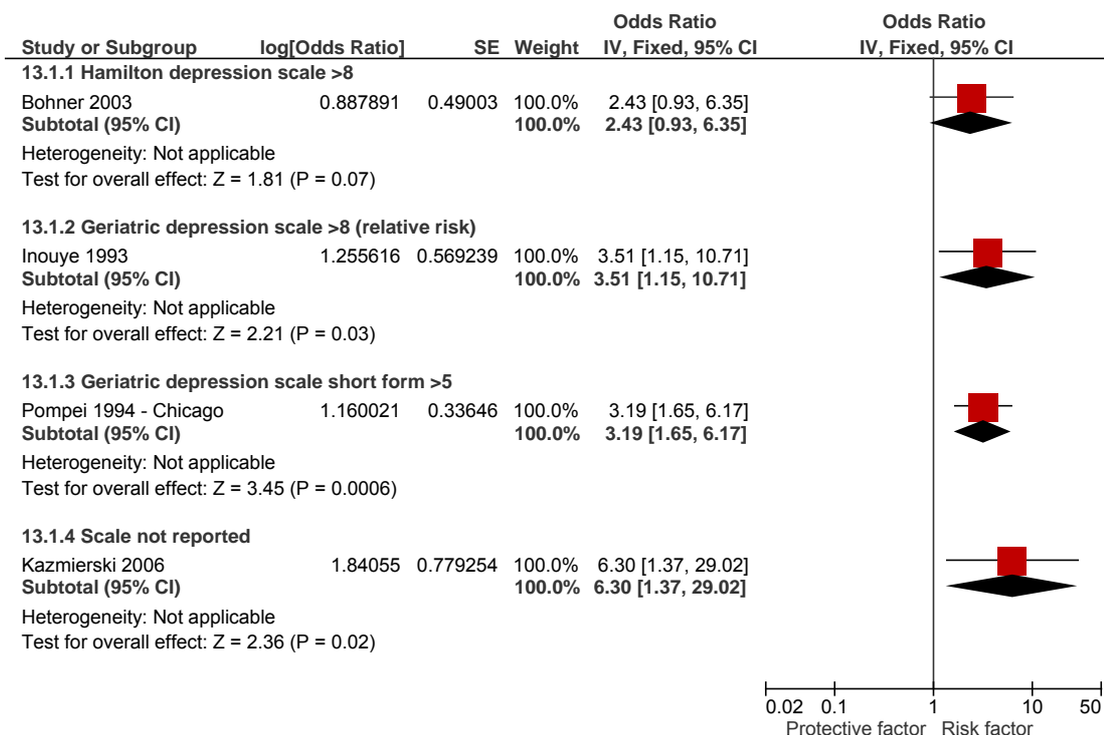
- 14 • In the study by Böhner (2003), a score of more than 8 using the Hamilton  
15 Depression Scale was indicative of depression; patients with delirium had  
16 a mean score of 8.16 (5.50) and patients without delirium had a mean  
17 score of 5.32 (5.52)
- 18 • In the study by Inouye (1993), depressive symptoms were considered  
19 present if the Geriatric Depression Score was 8 or more; 63% in the  
20 group with delirium and 44% in the group without delirium were  
21 depressed at baseline (data calculated).
- 22 • The method of defining depression was not reported in the study by  
23 Kazmierski (2006); 13% in the group with delirium, and 5% in the group  
24 without delirium had major depression.
- 25 • In the study by Pompei (1994), a score of 5 or more using the short form of  
26 the Yesavage Geriatric Depression scale was considered indicative of  
27 depression; of the Chicago sample, 41% with delirium and 17% without  
28 delirium were depressed
- 29 • In the study by Leung (2007), the authors evaluated depression using the  
30 Geriatric Depression Score: 12% had a score of 6 or higher
- 31 • The study by Pisani (2007) reported that 33% of the patients with delirium  
32 had a history of depression compared with 16% of patients without  
33 delirium (the scale used to measure depression was not reported).

34  
35 The standard error for the Böhner (2003) study was calculated from its p-value:  
36 confidence intervals were not reported for the odds ratio.  
37

38 The GDG noted that the scales used to measure depression were not diagnostic  
39 tools for that condition, and the cut-off points were not necessarily appropriate.  
40 The GDG also noted that in these studies, only Inouye (1993) also included illness  
41 severity in the multivariate analysis, and there was likely to be some confounding  
42 by physical illness. Thus, although there appeared to be a significant effect of  
43 depression as a risk factor for delirium, the GDG was not confident in this result.  
44 Considering only the higher quality study (Böhner 2003), the effect was just non-  
45 significant; OR 2.43 (95%CI 0.93 to 6.35) or beta coefficient 0.89 (SE 0.483;  
46 p=0.066).

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Figure 6.18: depression as a risk factor: incidence of delirium



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**Summary of depression as a risk factor**

Although there appeared to be a significant effect of depression on the incidence of delirium, the majority of the studies were low quality, and there was likely to be some confounding. Restricting the analysis for delirium incidence to the study that was of higher quality (Bohner 2003), this moderate sized study showed an almost significant effect of depression OR 2.43 (95%CI 0.93 to 6.35) or beta coefficient 0.89 (SE 0.483). The GDG considered that even this result could be confounded by physical illness and was not confident in its validity.

**6.5.1.12 Infection**

**Infection as a risk factor for incidence of delirium**

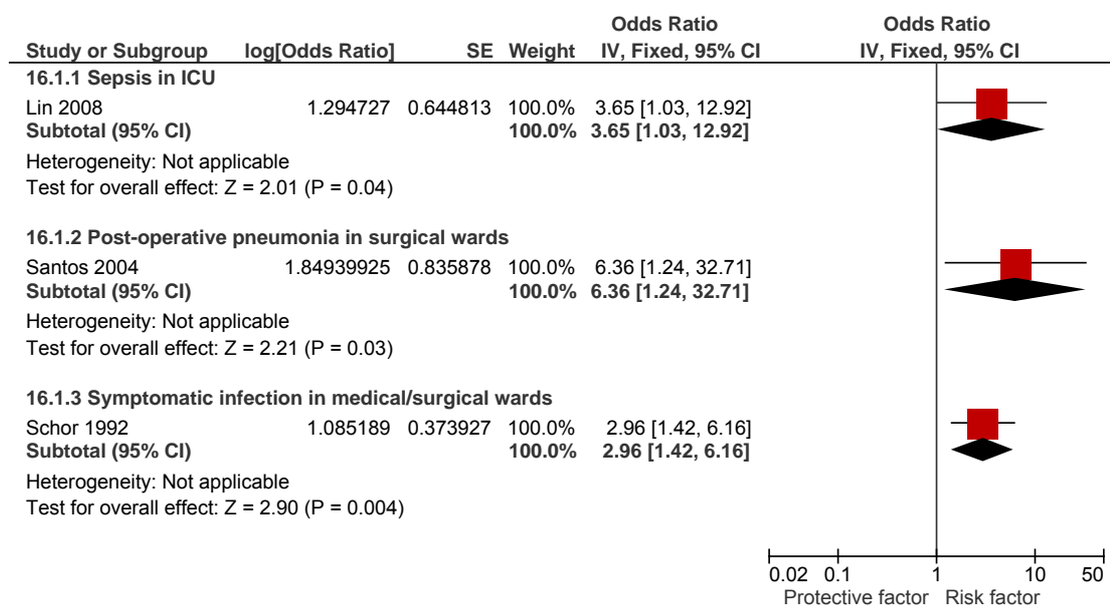
Three studies presented data on infection as a risk factor for the incidence of delirium in their multivariate analyses (Lin 2008; Santos 2004; Schor 1992). Two studies had a moderate rating (Lin 2008; Schor 1992), and one had a low rating (Santos 2004). One other study included infection as a risk factor in the multivariate analysis, but the non-significant results were not reported (Sheng 2006 (low)).

We note that these studies were conducted in all settings: surgical patients (Santos 2004), medical/surgical wards (Schor 1992; Sheng 2006) and ICU patients (Lin 2008).

1 The study by Lin (2008) reported that 80% of patients with delirium had sepsis  
 2 (defined by the American College of Chest Physicians and the Society of Critical  
 3 Care Medicine) and 57% without delirium had sepsis. The study by Santos  
 4 (2004) reported that 19% patients with delirium and 3% of patients without  
 5 delirium had post-operative pneumonia. The study by Schor (1992) reported  
 6 that 37% with delirium and 17% without delirium had symptomatic infection. The  
 7 study by Sheng reported that 15% of the patients with delirium had urinary tract  
 8 infection compared to 4% of patients without delirium.

9  
 10 Figure 6.19 shows that infection is a significant risk factor for delirium, although  
 11 the confidence intervals are wide. A sensitivity analysis without the low quality  
 12 study (Santos 2004) makes little difference.

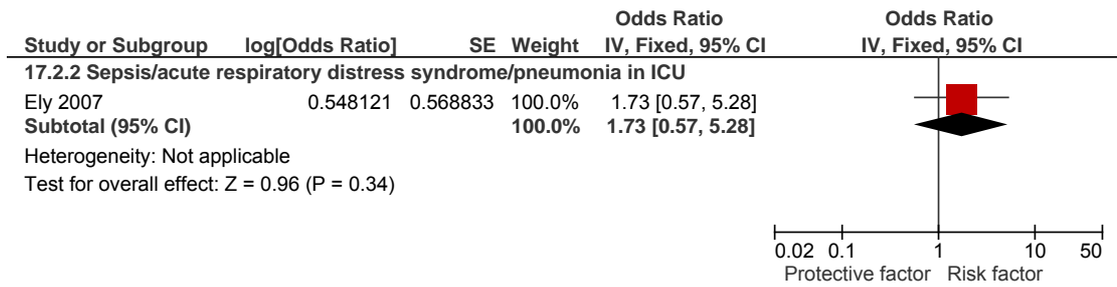
13  
 14  
 15 Figure 6.19: infection as a risk factor: incidence of delirium



16  
 17  
 18  
 19 Infection as a risk factor for increased duration of delirium

20 One small, moderate quality study in mechanically ventilated patients in ICU  
 21 patients evaluated infection as a risk factor for the duration of delirium (Ely  
 22 2007). The study reported that, overall, 15% had sepsis and 23% had  
 23 pneumonia. Figure 6.20 shows no significant effect of infection on the duration of  
 24 delirium, although the CI is wide in this small study.

1 Figure 6.20: infection as a risk factor: duration of delirium



2  
3

Summary of infection as a risk factor

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- Three moderate quality and one low quality studies showed a similar trend, indicating that infection is a risk factor for delirium, despite the different types of infection evaluated; the odds ratio ranged from 2.96 (95%CI 1.42 to 6.16) to 6.36 (95%CI 1.24 to 32.71).

8  
9  
10

- Evidence from one small study mechanically ventilated patients in ICU showed no significant relationship between infection and duration of delirium.

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12

13 **6.5.1.13 Fracture on admission**

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21

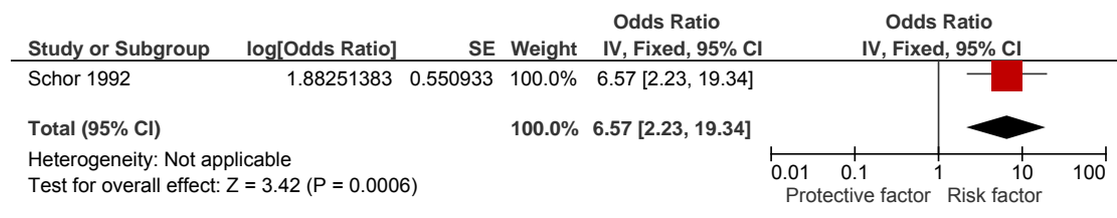
One moderate quality study in 291 patients (Schor 1992) included fracture on admission as a risk factor for delirium. The study did not report what type of fractures were found, but there were 8.3% of patients with a fracture (8.3% of patients were also admitted to orthopaedic surgery). This is a relatively small percentage so there is likely to be some inaccuracy in the results. There was a significant effect of fractures on admission on the incidence of delirium (figure 6.21); OR 6.57 (95%CI 2.23 to 19.33).

22  
23  
24  
25  
26  
27

This conclusion was supported by a second study (Andersson 2001, low quality), which showed that emergency hip fracture surgery was a significant risk factor for delirium incidence, compared with elective surgery for knee arthritis or hip arthritis (see procedural risk factors, section 6.5.3); OR 4.74 (95%CI 1.76 to 12.80).

28  
29

Figure 6.21: fracture on admission as a risk factor



30  
31

Summary of fracture as a risk factor

32  
33  
34

In summary, there was a significant effect of fractures on admission on the incidence of delirium in a single study, but there is some uncertainty associated

1 with the effect; OR 6.57 (95%CI 2.23 to 19.33). The conclusion was supported  
2 by evidence from a low quality study comparing emergency hip fracture surgery  
3 with elective surgery for knee or hip arthritis.  
4

#### 5 **6.5.1.14 Immobility**

6 One low quality study included immobility (ability to walk without aid before  
7 admission) as a risk factor for the incidence of delirium in multivariate analysis,  
8 but the non-significant results were not reported (Andersson 2001). This study  
9 had a low rating. The study reported that 29% of patients with delirium were  
10 able to walk without an aid before admission compared to 46% of patients  
11 without delirium.  
12

##### 13 Summary

14 There is a lack of evidence on immobility as a risk factor for the incidence of  
15 delirium.  
16

#### 17 **6.5.1.15 Incontinence**

18 One low quality study included urinary and faecal incontinence as risk factors for  
19 the incidence of delirium in multivariate analysis, but the non-significant results  
20 were not reported (Sheng 2006 (low)). In this study 31% of patients with  
21 delirium and 13% of patients without delirium had urinary incontinence, and  
22 23% with delirium and 8% without delirium had faecal incontinence.  
23

##### 24 Summary

25 There is a lack of evidence on continence as a risk factor for the incidence of  
26 delirium.  
27

#### 28 **6.5.2 Environmental risk factors**

29 One low quality study presented various environmental factors in their  
30 multivariate analysis of delirium severity (McCusker 2001). This study reporting  
31 delirium severity used analyses at various times reflecting different states  
32 (repeated measures multivariate analyses, using the previous most recent  
33 severity score as a factor in the multivariate analysis). The proportions of each of  
34 these states as a function of the number of different states for that variable are  
35 given below.  
36

37 Some of the measures are subjective: for example, the research assistant  
38 decided whether the patient's surroundings were too noisy or whether the room  
39 was well lit. Other risk factors were more objective: e.g. whether or not various  
40 orientation aids were present and whether physical restraints were used. The  
41 study reported that the inter-rater reliability was assessed for these  
42 environmental observations in 29 patients and 75-100% agreement was found.

- 43 • Recent room change (173/617 = 28%)

- 1 • Stimulation: based on the distance of the room from the nurses station: high  
2 (105/573 = 18%), moderate (243/573 = 42%), low (225/573 =  
3 39%)
- 4 • In same room (403/590 = 68%)
- 5 • Single room (124/509 = 24%)
- 6 • Surroundings' not well lit (61/504 = 12%)
- 7 • Surroundings' too noisy/quiet versus normal (159/421 = 38%)
- 8 • Radio/TV on (72/513 = 14%)
- 9 • Clock/watch absent versus present (294/585 = 50%)
- 10 • Calendar absent versus present (430/498 = 86%)
- 11 • Personal possessions absent versus present (421/538 = 78%)
- 12 • Not wearing glasses (375/587 = 64%)
- 13 • Not using hearing aid (433/470 = 92%)
- 14 • Family absent when carrying out assessment versus present (426/558 =  
15 76%)
- 16 • In isolation because of screening for infection control (52/490 = 11%)

17  
18 The results of the multivariate analyses are reported in figures 6.22 to 6.24.  
19 Most environmental risk factors showed no significant effect on the severity of  
20 delirium, but there was reported to be a significant effect for the following:

- 21 • Greater number of room changes
- 22 • Absence of a clock or watch
- 23 • Not wearing reading glasses

24  
25 The GDG noted that in the UK, however, the number of moves is often influenced  
26 by management, rather than clinical reasons, and commented that it was unclear  
27 why the patients had been moved in this study.

28  
29 The study also carried out exploratory analyses and noted two statistically  
30 significant interactions:

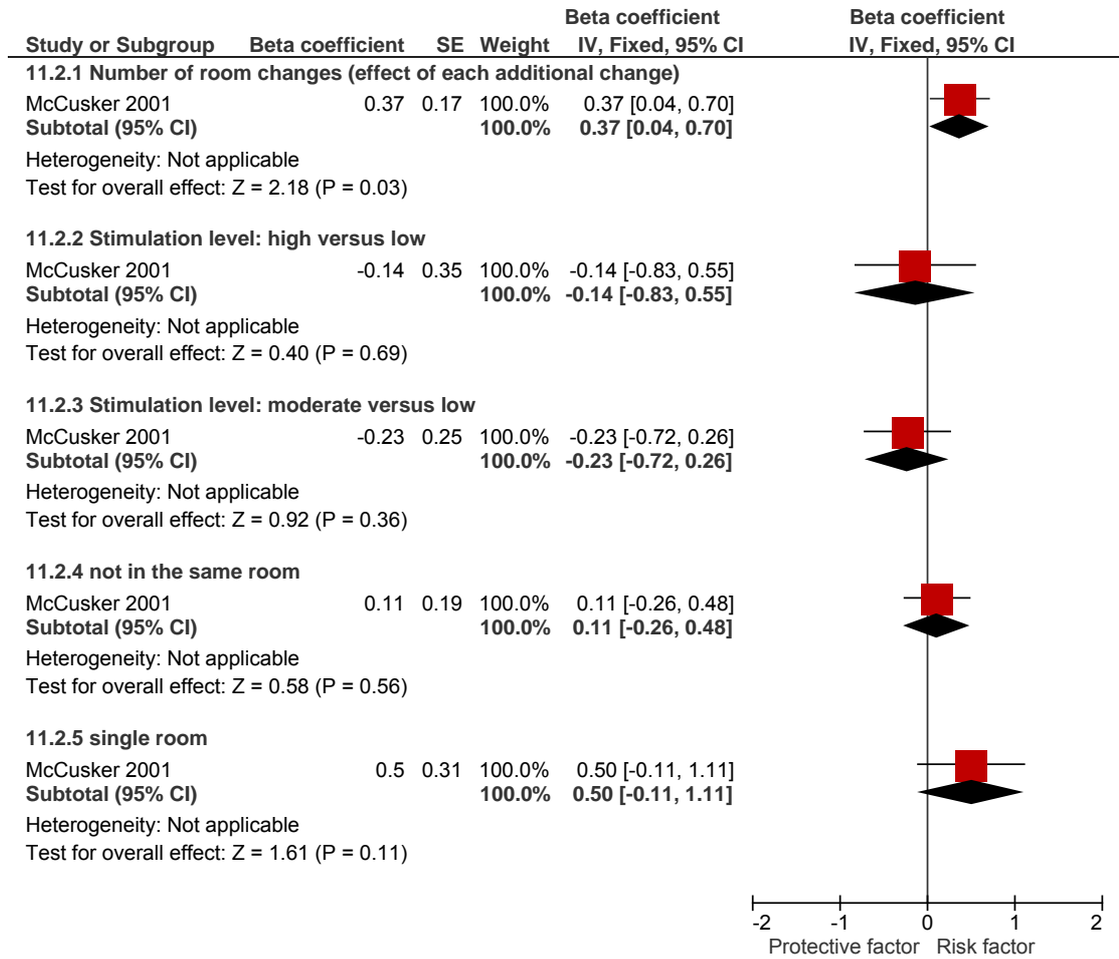
- 31 • The number of room changes was affected by the level of stimulation: a  
32 higher number of room changes had a strong impact on the severity of  
33 delirium only if the patient was in a room with high stimulation
- 34 • Moderate stimulation had a greater impact on patients in a unit with mixed  
35 medical and long-term care patients than in a medical ward

36

37 However, the authors stated that a large number of interactions were tested so  
38 that these results should be interpreted with caution.

39

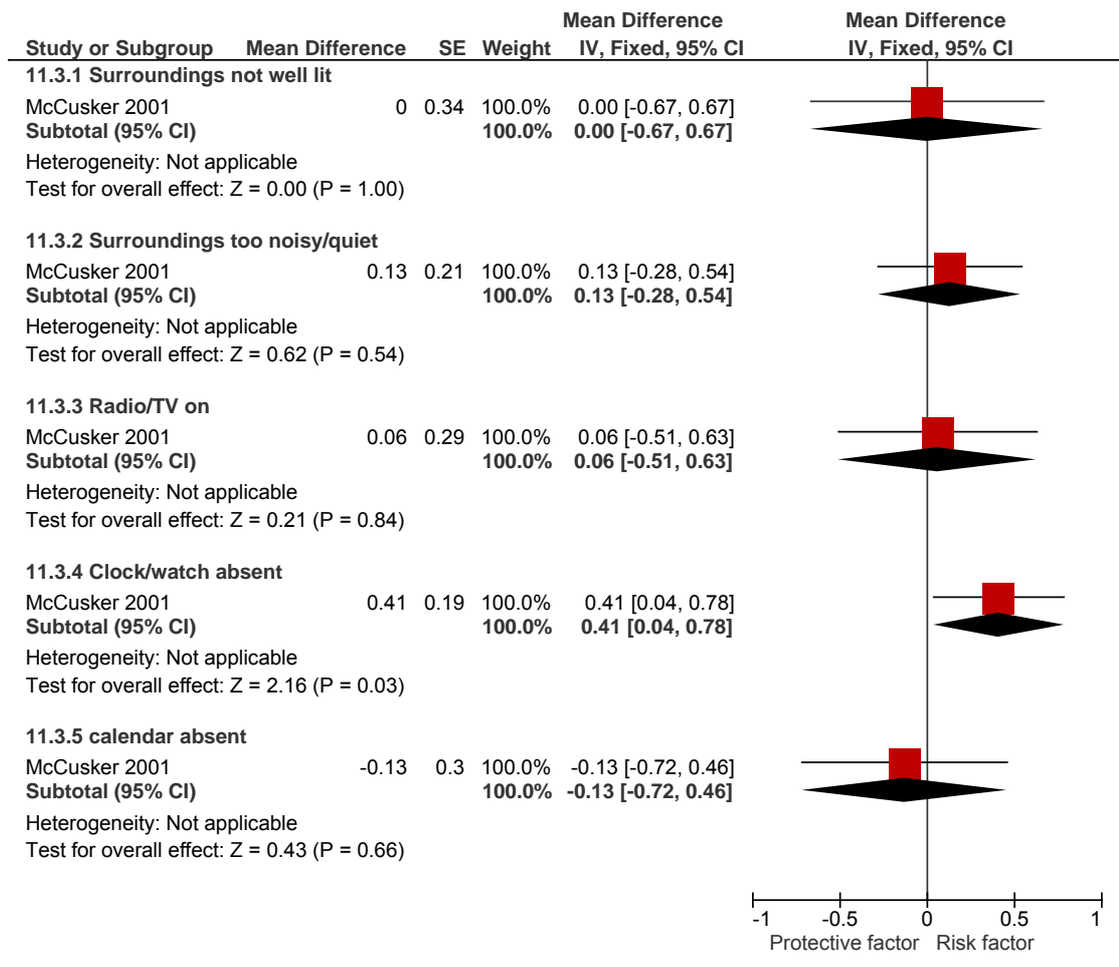
1 Figure 6.22: environmental risk factors: severity of delirium



2

3

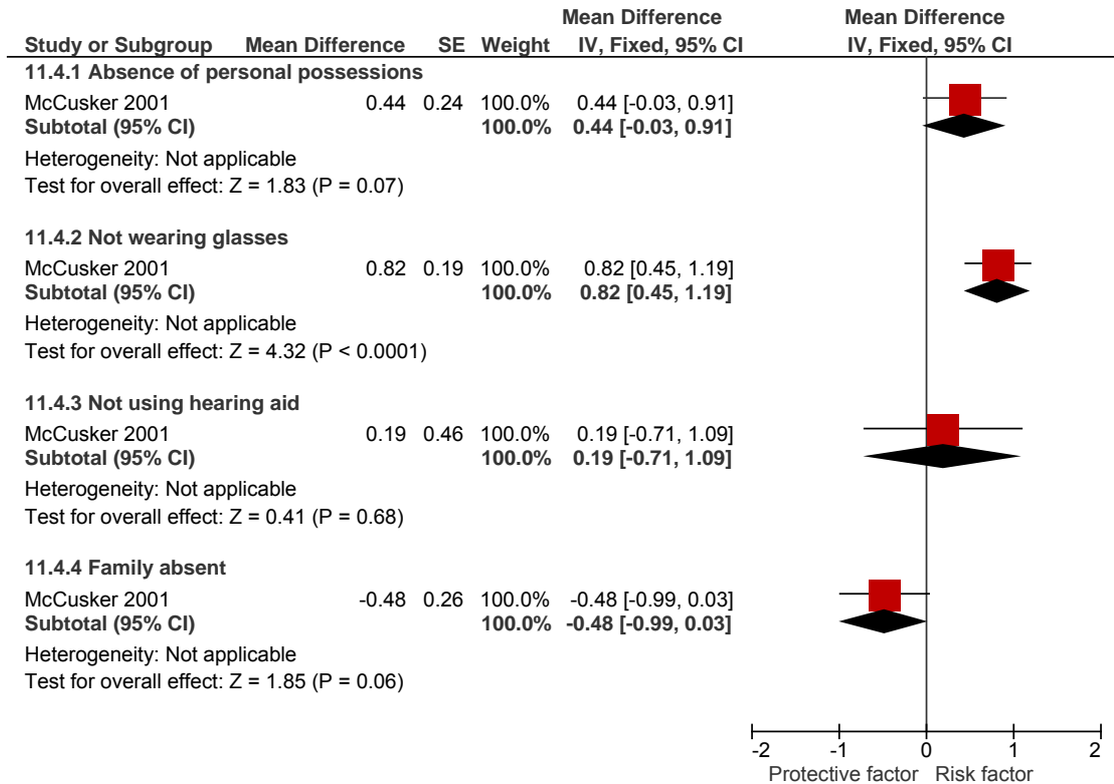
1 Figure 6.23: environmental risk factors: severity of delirium (NB scale -1 to +1)



2

3

1 Figure 6.24: environmental risk factors: severity of delirium



2  
3  
4

5 Summary of environmental risk factors for the severity of delirium

- 6 • In one large, low quality study, the beta coefficient for the mean difference  
7 in severity of delirium was significant for the following factors:
- 8 ○ The number of room changes: beta coefficient 0.37 (95% CI 0.04  
9 to 0.70)
  - 10 ○ The absence of a clock or watch: beta coefficient 0.41 (95% CI  
11 0.04 to 0.78)
  - 12 ○ Not wearing reading glasses: beta coefficient 0.82 (95% CI 0.45  
13 to 1.19)
- 14 • In one large, low quality study, the beta coefficient for the mean difference  
15 in severity of delirium did not appear to be significant for the following  
16 factors: level of stimulation, single room, surroundings not well lit,  
17 surroundings too noisy or quiet, radio/TV on, calendar absent, absence  
18 of personal possessions, not using a hearing aid, family member present.
- 19 • We note that this study also controlled for age, dementia, baseline delirium  
20 severity; age, dementia, comorbidity, and visual or hearing impairment.

1

2 **6.5.3 Procedural risk factors**

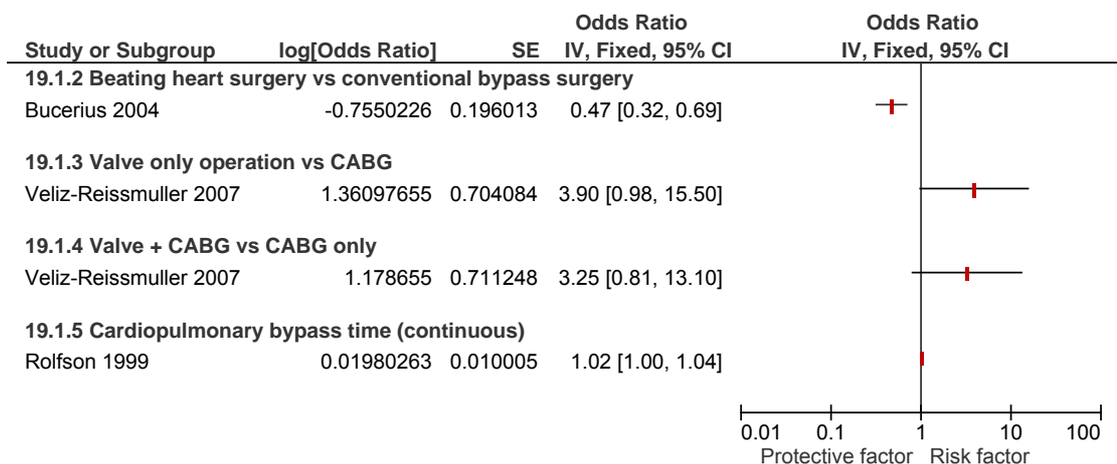
3

4 **6.5.3.1 Type of surgery**

5 Five studies evaluated surgery as a risk factor for the incidence of delirium in  
 6 their multivariate analyses (Andersson 2001; Bucerius 2004; Rolfson 1999;  
 7 Rudolph 2007; Veliz-Reissmüller 2007) (figure 6.25). Two studies had a low  
 8 rating (Andersson 2001; Veliz-Reissmüller 2007); the remaining studies had a  
 9 moderate rating. Three of these studies evaluated cardiac surgery. None of the  
 10 studies included illness severity in their multivariate analyses, although the  
 11 Andersson (2001) study included comorbidity.  
 12

- 13 • The study by Bucerius (2004) compared patients who underwent beating  
 14 heart surgery (no cardiopulmonary bypass) with those who underwent  
 15 bypass (conventional) surgery.
- 16 • The study by Veliz-Reissmüller (2007) compared patients who underwent  
 17 valve operation plus coronary bypass grafting (CABG) with CABG only.
- 18 • The study by Rolfson (1999) evaluated the duration of cardiopulmonary  
 19 bypass (minutes).
- 20 • The GDG suggested that differences in the type of operation may be a  
 21 proxy for illness severity  
 22

23 Figure 6.25: cardiac surgery risk factors: incidence of delirium



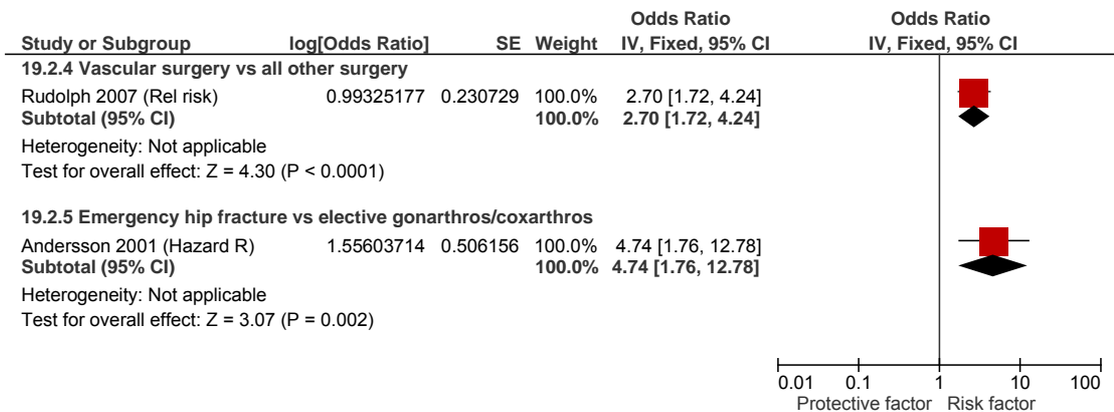
24  
25

26 Figure 6.26 presents the results for three studies: one low quality study  
 27 evaluated the risk of delirium in emergency hip fracture surgery patients versus  
 28 patients admitted for elective surgery for knee arthritis or hip arthritis (Andersson  
 29 2001). The GDG concluded that this risk factor was connected with the  
 30 underlying condition (i.e. hip fracture), rather than the type of surgery.  
 31

1 One moderate quality study compared vascular surgery with all other surgery  
 2 (abdominal, orthopaedic, genitourinary, thoracic and other) (Rudolph 2007), and  
 3 showed that vascular surgery puts the patient at greater risk of delirium than  
 4 other forms of surgery.  
 5

6 The GDG stated that vascular surgery may be a proxy for other factors, such as  
 7 undiagnosed vascular dementia or cerebral damage.  
 8  
 9  
 10  
 11  
 12

13 Figure 6.26: type of surgery a risk factor: incidence of delirium



14  
15  
16 Summary of surgical procedural factors as risk factors for delirium incidence

- 17 • One moderate quality study showed a significant protective effect on the  
18 incidence of delirium for beating heart surgery compared with  
19 conventional bypass surgery.
- 20 • One moderate quality study showed that vascular surgery was a significant  
21 risk factor for delirium incidence, compared with other types of (non-  
22 cardiac) surgery.
- 23 • One moderate quality study showed a borderline significant effect of  
24 cardiopulmonary bypass time as a risk factor
- 25 • None of the studies included illness severity in their multivariate analyses  
26 and the GDG concluded that the effects were likely to be a proxy for  
27 illness severity

1 **6.5.3.2 Iatrogenic interventions and medical restraint**

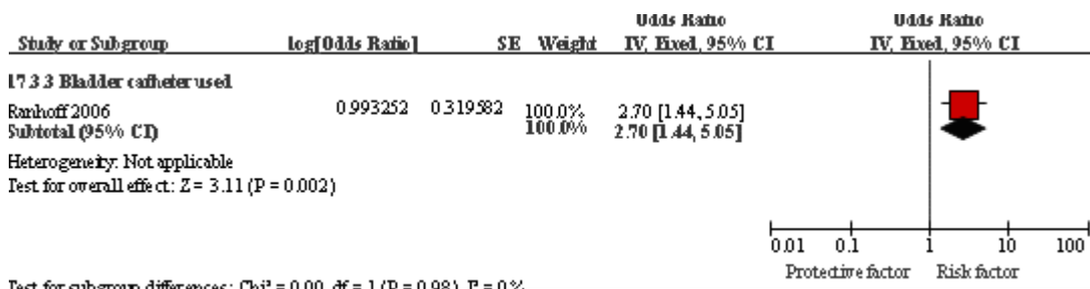
2 Iatrogenic interventions

3 Two studies evaluated iatrogenic interventions as risk factors for the incidence of  
4 delirium in their multivariate analysis (Andersson 2001, low; Ranhoff 2006)  
5 (figure 6.27).  
6

7 Both studies evaluated if a fitted bladder catheter was a risk factor. In the study  
8 by Ranhoff (2006), 81% of patients started to have prevalent delirium, and  
9 80% of patients with incident delirium, used a bladder catheter (data were not  
10 reported for Andersson 2001). The study by Andersson (2001) did not report  
11 the non-significant results for the use of bladder catheter for emergency surgery  
12 patients in their multivariate analysis.  
13

14 The study by Andersson (2001) was conducted in surgical patients and had a  
15 low rating while the study by Ranhoff (2006) was conducted in ICU patients and  
16 had a moderate rating.  
17  
18  
19  
20  
21  
22  
23

Figure 6.27: iatrogenic intervention as a risk factor: incidence of delirium



- 24
- 25 • Due to the low rating of the Andersson (2001) study, the results for this
  - 26 study should be treated with caution.
  - 27 • The GDG noted that the risk factor examined in the Ranhoff (2006) study
  - 28 was in-situ bladder catheter in ICU, rather than a bladder catheter being
  - 29 introduced, but they found the clinical interpretation of this study difficult.
- 30

31 Medical restraint

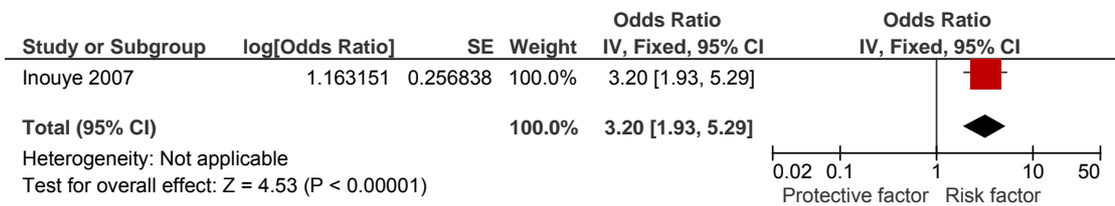
32 One low quality study presented data on medical restraint in their multivariate  
33 analysis for the severity of delirium (McCusker 2001; figure 6.29).  
34 Medical restraint was stated to include intravenous and oxygen tubing, and  
35 occurred in 320/658 (49%) patient states. This was a significant risk factor;  
36 beta coefficient 0.41 (95% CI 0.04 to 0.78).  
37

1 **6.5.3.3 Physical restraint**

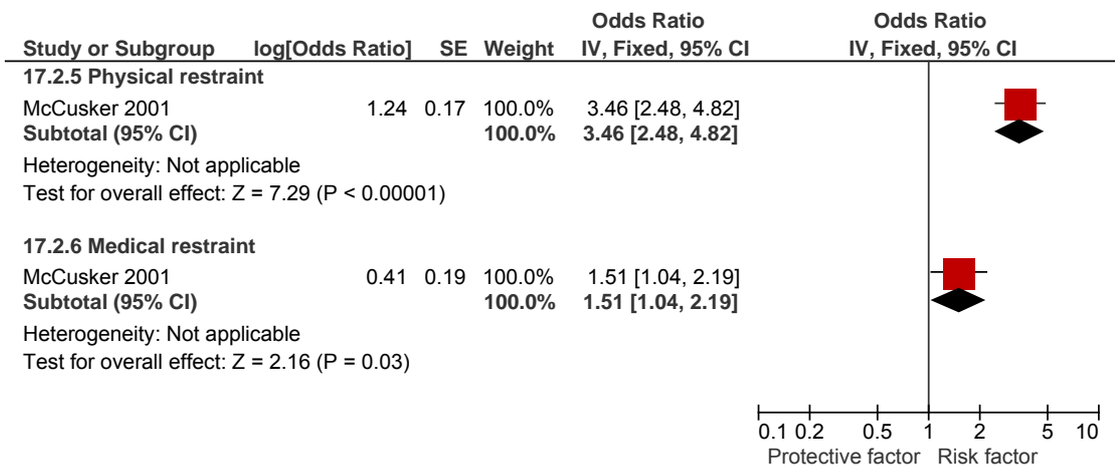
2 Two studies presented data on physical restraint in their multivariate analyses  
 3 (Inouye 2007; McCusker 2001) (figures 6.28 and 6.29). The Inouye (2007) study  
 4 was of moderate rating, but the McCusker (2001) study was considered to be of  
 5 low quality; both were conducted in medical wards. In the Inouye (2007) study,  
 6 restraint use during delirium occurred in 15% of the patients. In the McCusker  
 7 (2001) study, physical restraint was examined as a risk factor for delirium  
 8 severity and occurred in 303/658 (44%) patient states; more detailed  
 9 information was not reported.

10 Both studies reported a significant effect of physical restraint on delirium  
 11 persistence (OR 3.20 (95%CI 1.93 to 5.29) and the severity of delirium (beta  
 12 coefficient 1.24 (95% CI 0.91 to 1.57)).  
 13  
 14

15 Figure 6.28: physical restraint during delirium: persistent delirium



16  
 17  
 18 Figure 6.29: physical and medical restraint as a risk factor for the severity of  
 19 delirium



- 20  
 21  
 22 • For persistent delirium, the odds ratio was 3.2 (95% CI 1.9 to 5.2). We  
 23 note that these results are from a subpopulation of patients with delirium.  
 24 • The beta coefficient for the mean difference in severity of delirium was  
 25 0.21 (95% CI 0.08 to 1.54).  
 26

## 1 Summary

- 2 • There was moderate quality evidence that a bladder catheter used in ICU  
3 patients was a risk factor for the incidence of delirium, but the GDG was  
4 uncertain how to interpret this information
- 5 • There was low quality evidence that medical restraint was a risk factor for  
6 the severity of delirium
- 7 • There was low quality evidence that physical restraint was a risk factor for  
8 the severity of delirium and moderate evidence that it was a risk factor  
9 for persistent delirium

### 11 **6.5.4 Overall summary**

12 Of the many risk factors examined for the incidence of delirium, the GDG  
13 concluded that they had some confidence in the results for the following risk  
14 factors:

- 15 • Age as a continuous variable
- 16 • Age over 65 years
- 17 • Age over 80 years
- 18 • Cognitive impairment
- 19 • Vision impairment
- 20 • Illness severity
- 21 • Fracture on admission
- 22 • Infection
- 23 • Physical restraint

24  
25 The GDG had less confidence in the results for the following risk factors:

- 26 • Comorbidity
- 27 • Vascular surgery

28  
29 The GDG noted that the following risk factors had inconsistent or uncertain  
30 results:

- 31 • Depression
- 32 • Hearing impairment
- 33 • Polypharmacy
- 34 • Dehydration
- 35 • Sex
- 36 • Electrolyte disturbance
- 37 • Immobility

- Incontinence
- Bladder catheter

The dichotomous results for the risk factors for delirium incidence are summarised on a forest plot, ordered by size of effect (figure 6.30). This is intended to give a visual summary and the values are represented by the highest quality study or the midpoint. The corresponding values for persistent delirium are shown on figure 6.31.

Figure 6.30: risk factors for incidence of delirium

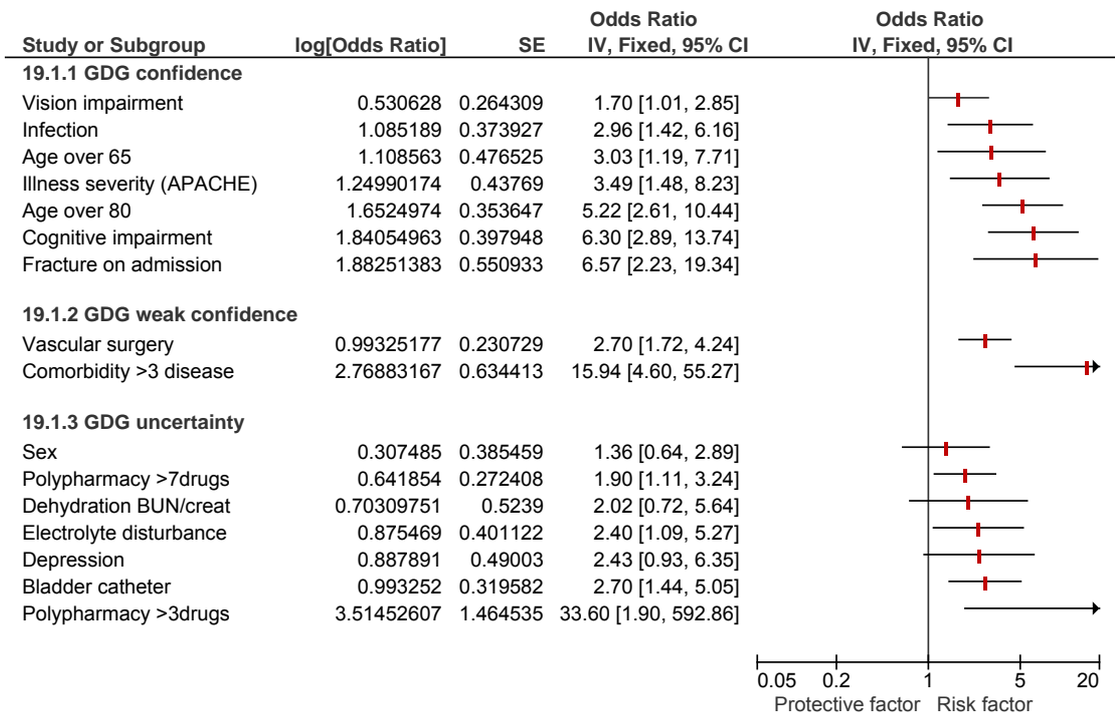
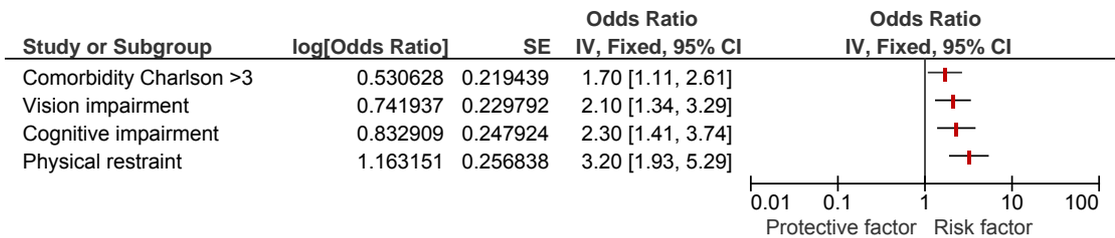


Figure 6.31: risk factors for persistent delirium



1

# 1 **7 Risk factors for delirium: pharmacological**

## 2 **agents**

### 3 **7.1 Clinical introduction**

4 Delirium often occurs in individuals who are already on medications either for  
5 longstanding conditions or acute illness. Some medications seem to be associated  
6 with higher incidence of delirium. It appears that many classes of drugs are  
7 implicated in the development of delirium. By identifying those drugs  
8 responsible, clinicians would not necessarily avoid their use altogether but  
9 potentially consider alternatives or be more judicious in their use. Also by  
10 identifying pharmacological risk factors, staff or carers looking after the  
11 individual would be more vigilant for the signs of the development of delirium. It  
12 is not known whether it is the individual's drugs that pose a risk, or the  
13 combinations of the different types of drugs.

14 The knowledge of the propensity of different drugs or groups of drugs to  
15 contribute to the development of delirium will help clinicians to reduce the  
16 individual's risk at many stages in the patient's journey e.g. admission to a new  
17 in-hospital care setting, on admission to long-term care or on routine review by  
18 the patients General Practitioner.

### 20 **7.2 Selection criteria**

21 Selection criteria were as outlined in the general methods section apart from the  
22 types of risk factor.

#### 24 **7.2.1 Types of study design**

25 The study designs for pharmacological agents as risk factors were to be RCTs  
26 (because they are interventions) or cohort studies. If neither of these designs  
27 were available for a particular risk factor, case control studies could also be  
28 included.

#### 30 **7.2.2 Types of pharmacological risk factor**

31 Any pharmacological agent used that was reported to be a risk factor for  
32 delirium was to be considered.

33

### 1 7.2.3 Types of comparison

2 The following comparisons were to be included:

- 3 • Intervention versus placebo / no intervention
- 4 • Intervention 1 + intervention 2 versus intervention 2 alone
- 5 • Drug A versus drug B (both drugs in same class)
- 6 • Drug class A versus drug class B
- 7 • Dose 1 versus dose 2

8  
9 It was decided to combine the two types of comparison: (i) intervention versus  
10 placebo / no intervention and (ii) intervention 1 + intervention 2 versus  
11 intervention 2 alone, and examine this assumption using sensitivity analyses.

12

### 13 7.2.4 Type of outcome measure

14 The types of outcome measure were to be:

- 15 • Incidence of delirium [also recording when incidence was measured]
- 16 • Severity of delirium
- 17 • Duration of delirium

18

### 19 7.2.5 Stratification and subgroup analyses

20 We planned to stratify the studies by class of drug.

21

22 The following subgroups were to be considered:

- 23 • Type of pharmacological agent
- 24 • Dose

25

## 26 7.3 Description of studies

27 Twenty-eight papers were evaluated for inclusion. Six studies were excluded  
28 and are listed in Appendix G with reasons for exclusion.

29

30 We included 22 reports of 21 studies (Agostini 2001; Beaussier 2006; Christe  
31 2000; Centorrino 2003; Dubois 2001; Foy 1995; Han 2001; Herrick 1996;  
32 Holroyd 1994; Kim 1996; Leung 2006; Marcantonio 1994; Morrison 2003;  
33 Nitschke 1996; Pandharipande 2006; Pandharipande 2008; Papaioannou  
34 2005; Pisani 2007; Pisani 2009; Scott 2001; Shulman 2005; Williams-Russo  
35 1992), for which full data extraction was carried out. One study (Pisani 2007)

1 had more than one report (Pisani 2007; Pisani 2009); hereafter, these studies  
2 are referred to by the first named report, but separately in the methodological  
3 quality assessment and results section. Three further studies (Oh 2008; Shiba  
4 2009; Van Rompaey 2009) were identified in the update searches; these studies  
5 were considered to be low quality and did not add to the body of evidence so  
6 these were not analysed in depth.

7

### 8 **7.3.1 Study Design**

9 The 22 reports had different study designs: nine were RCTs (Beaussier 2006;  
10 Christe 2000; Herrick 1996; Kim 1996; Leung 2006; Nitschke 1996;  
11 Papaioannou 2005; Scott 2001; Williams-Russo 1992), nine reports of eight  
12 studies were prospective cohort studies (Agostini 2001; Dubois 2001; Foy 1995;  
13 Han 2001; Morrison 2003; Pandharipande 2006; Pandharipande 2008; Pisani  
14 2007; Pisani 2009); three were retrospective cohort studies (Centorrino 2003;  
15 Holroyd 1994; Shulman 2005) and one was a case control study (Marcantonio  
16 1994). The Leung (2006) study also carried out a multivariate analysis on the  
17 study population for risk factors other than those randomised, and is treated as  
18 a prospective cohort study for the other risk factors. The Han (2001) study  
19 reported that patients were those diagnosed with delirium enrolled in what the  
20 authors reported as 'an RCT of a delirium geriatric service or in an observational  
21 cohort study of outcomes of delirium' [references not provided for either study in  
22 the text].

23

24 One study was conducted in the UK (Scott 2001). Twelve were conducted in the  
25 USA (Agostini 2001; Centorrino 2003; Holroyd 1994; Kim 1996; Leung 2006;  
26 Marcantonio 1994; Morrison 2003; Nitschke 1996; Pandharipande 2006;  
27 Pandharipande 2008; Pisani 2007; Williams-Russo 1992); four in Canada  
28 (Dubois 2001; Han 2001; Herrick 1996; Shulman 2005); one was in France and  
29 Switzerland (Beaussier 2006); one in Switzerland (Christe 2000); one in Greece  
30 (Papaioannou 2005); and one in Australia (Foy 1995).

31

32 Two studies received funding from a pharmaceutical company (Christe 2000;  
33 Kim 1996 [also non pharmaceutical funding]) and eleven studies had non-  
34 pharmaceutical based funding (Herrick 1996; Leung 2006; Marcantonio 1994;  
35 Morrison 2003; Pandharipande 2006; Pandharipande 2008; Nitschke 1996;  
36 Papaioannou 2005; Pisani 2007; Shulman 2005; Williams-Russo 1992). The  
37 remaining studies did not state how they were funded.

38

39 Five studies had fewer than 100 to 200 patients (Beaussier 2006: n=59; Christe  
40 2000: n=65; Nitschke 1996: n=92; Papaioannou 2005: n=50; Williams-Russo  
41 1992: n=60); five studies had 100 or more patients (Centorrino 2003: n=139;  
42 Holroyd 1994: n=114; Kim 1996: n=127; Herrick 1996: n=136;  
43 Pandharipande 2008: n=100); five studies had more than 200 patients (Dubois  
44 2001: n=216; Han 2001: n=278; Leung 2006: n=228; Marcantonio 1994:  
45 n=245; Pandharipande 2006: n=275), and six studies were large studies

(Agostini 2001: n=426; Scott 2001: n=420; Foy 1995: n=418; Morrison 2003: n=541; Pisani 2007: n=304; Shulman 2005: n=10230).

### 7.3.2 Population

The mean age (table 7.1) where reported, ranged from 40.8 (Centorrino 2003) to 83 years (Han 2001). The age ranges varied, and are shown in table 1.

Table 7.1: patient ages. Unless otherwise specified, all data are presented as mean (range);  $\pm$  indicates that the range was estimated from the mean  $\pm$  1 standard deviation. IQR = interquartile range.

Study	Age (range) years	Study	Age (range) years
Agostini (2001)	80 (73.2 to 86) $\pm$	Morrison (2003)	range not reported
Beaussier (2006)	77.5 (72 to 83) $\pm$	Marcantonio (1994)	73 (65 to 81)
Centorrino (2003)	40.8 (26.7 to 54.9) $\pm$	Nitschke (1996)	66.6 (65 to 69)
Christe (2000)	Median 84 (63 to 98)	Pandharipande (2006)	55.5 (38.5 to 72.5) $\pm$
Dubois (2001)	64.8 (49.3 to 79.7) $\pm$	Pandharipande (2008)	median: 48 (IQR 36 to 60)
Han (2001)	83.4 (76.1 to 90.7) $\pm$	Papaoiannou (2005)	median : 68
Foy (1995)	70.2 (59 to 88)	Pisani (2007)	74.6 (67 to 81) $\pm$
Herrick (1996)	72 (65 to 85)	Pisani (2009)	75 (67 to 83) $\pm$
Holroyd (1994)	74.1 (65 to 92)	Scott (2001)	60.8 (49.6 to 68.1) $\pm$
Kim (1996)	66 (24 to 86)	Shulman (2005)	74.7 (67.8 to 81.5)
Leung (2006)	74 ( 65 to 95)	Williams-Russo (1992)	68 (48 to 84)

One study (Morrison 2003) did not report the mean age, but stated that 9% of the patients had a mean age less than 70 years, 26% were between the ages of 70 to 79 years and 65% were 80 years or older.

The studies varied in the proportions of patients reported to have **cognitive impairment** at baseline. In addition, the GDG decided that, when this was not clearly stated, it was unlikely that patients undergoing elective cardiac surgery would have cognitive impairment at baseline. This gave the following subgroups:

- Three studies reported patients with cognitive impairment/dementia were excluded

- 1                   ○ one study (Christie 2000) reported that patients with moderate to  
2                   severe cognitive impairment were excluded at baseline;
- 3                   ○ one study (Pandharipande 2006) reported patients with severe  
4                   dementia and psychosis were excluded;
- 5                   ○ one study (Shulman 2005) reported that patients with a past  
6                   diagnosis of dementia were excluded *a priori*.
- 7                   ● Fourteen studies reported that some patients had cognitive impairment at  
8                   baseline (Agostini 2001; Beaussier 2006; Christie 2000; Foy 1995; Han  
9                   2001; Herrick 1996; Holroyd 1994; Kim 1996; Leung 2006;  
10                  Marcantonio 1994; Morrison 2003; Nitschke 1996; Papaioannou 2005;  
11                  Pisani 2007).
- 12
- 13                  Information on cognitive impairment status was not reported in the remaining  
14                  studies (Centorrino 2003; Dubois 2001; Pandharipande 2008; Scott 2001). The  
15                  Scott (2001) study included patients undergoing CABG and the GDG advised  
16                  that these patients were unlikely to have cognitive impairment at baseline.
- 17
- 18                  Cognitive impairment/dementia was assessed using different scales:
- 19                  ● Nine studies assessed cognitive impairment based on the MMSE score  
20                  (Agostini 2001; Beaussier 2006; Christie 2000; Foy 1995; Herrick 1996;  
21                  Kim 1996; Holroyd 1994; Nitschke 1996; Papaioannou 2005);
- 22                  ○ Two studies reported excluding patients with a preoperative  
23                  MMSE score of 23 or below (Foy 1995; Papaioannou 2005).
- 24                  ● Two studies (Herrick 1996; Nitschke 1996) reported the cognitive  
25                  impairment change scores.
- 26                  ● Two studies (Leung 2006; Marcantonio 1994) used the Telephone Interview  
27                  For Cognitive Status (TICS)
- 28                  ● One study (Williams-Russo 1992) used the Mattis Dementia Rating Scale
- 29                  ● One study (Pandharipande 2006) used the Blessed Dementia Rating Scale
- 30                  ● Two studies used the IQCODE (Han 200; Pisani 2007: short version).
- 31                  ● One study (Morrison 2003) based its assessment on the diagnosis or history  
32                  of memory impairment or a dementing illness or if one or more errors  
33                  were made in answering a four item screening test (assessing orientation  
34                  [place and time]; circumstances of the fracture [place, time, circumstance];  
35                  immediate recall of the nature and purpose of the research study; recall  
36                  of the name or position of the person administering informed consent)
- 37                  ● One study did not state what scale was used to assess cognitive impairment  
38                  (Shulman 2005).
- 39

1 Six studies reported the mean MMSE score (range 0 to 30) and cognitive  
2 impairment status was deduced from the scores. In one study the mean MMSE  
3 score indicated that some patients had no cognitive impairment (Beaussier 2006)  
4 and in five studies some patients had some cognitive impairment (Agostini 2001;  
5 Christie 2000; Kim 1996; Holroyd 1994; Papaioannou 2005).

- 6 • The mean Blessed Dementia Rating Scale (range: 0 to 17, with 17  
7 indicating worst; score of 4 or higher representing threshold for  
8 dementia) reported in one study (Pandharipande 2006) indicated  
9 patients had low presence of dementia
- 10 • In two studies (Leung 2006; Marcantonio 1994) the mean TICS score was  
11 reported (range 0 to 41; cutoff score not reported in either study)  
12 indicating that some of the patients may be cognitively impaired.
- 13 • One study (Williams-Russo 1992) reported the mean Delirium rating scale  
14 (DRS) score (range: 36 item; 5 subscales; score less than 123 points is the  
15 cut off for dementia) and range and reported two patients would be  
16 classified as mildly demented pre-operatively.
- 17 • One study (Pisani 2007) reported the 31% [ 94/304] of the patients  
18 scored above 3.3 in the IQCODE (range: 1 to 5; with 1 indicating much  
19 improved compared to 10 years ago and 5 indicating much worse  
20 compared to 10 years ago).

21  
22 Sensory impairment at baseline was reported in four studies (Han 2001;  
23 Pandharipande 2006; Pisani 2007; Shulman 2005) and not reported in the  
24 remaining studies. Levels of sensory impairment are given in table 7.2. The  
25 studies did not generally give much information on how sensory impairment was  
26 assessed:

- 27 • sensory impairment was patient reported (Pisani 2007)
- 28 • assessed clinically at enrolment for presence or absence (Han 2001)
- 29 • not reported (Pandharipande 2006; Shulman 2005)

30  
31 One study (Papaioannou 2005) reported excluding patients with severe  
32 auditory or visual disturbances.

33  
34  
35 Table 7.2: levels of sensory impairment

Study	Visual impairment	Hearing impairment
<b>Han 2001</b>	<b>19.8%</b>	
<b>Pandharipande 2006</b>	<b>58%</b>	<b>16%</b>
<b>Pisani 2007</b>	<b>10.5%</b>	<b>17%</b>
<b>Shulman 2005</b>	<b>1.6%</b>	<b>10.6%</b>

36  
37  
38 Fourteen reports of 13 studies reported medications taken; some patients were  
39 taking several drugs; table 7.3.

1  
2

Table 7.3: mean number and/or types of medications

Study	Mean number of medications/ Types of medications
Agostini (2001)	5.4 (SD 3.1) and 5.6 (SD 3.2) medications for the diphenhydramine-exposed and non-exposed groups, respectively. Type of medications not stated
Centorrino (2003)	At least one centrally active drug: benzodiazepine, antipsychotic, antidepressants, anticonvulsant, lithium or a combination (97%)
Christie (2000)	Benzodiazepines (49%), antidepressants (15%), neuroleptics (11%), opioids (11%);
Dubois (2001)	Benzodiazepines, lorazepam, propofol, opioids (fentanyl, meperidine), steroids, antipsychotics (haloperidol or other), corticosteroids
Han 2001	Atypical antipsychotics, anticholinergics, benzodiazepine (not all types of medications listed)
Holroyd (1994)	Treatment with psychotropic medication (various tricyclics (58.8%), antipsychotics (27.2%) serotonin reuptake inhibitors (13.2%), anticholinergic medication (8.8%), methylphenidate (8.8%), bupropion (8.8%), carbamazepine (8.8%), MAOIs (5.1%), thyroid augmentation (3.5%), valproate (3.5%), verapamil (1.8%)
Morrison (2003)	Benzodiazepines or other sedatives and hypnotics, opioids (including meperidine)
Pandharipande (2006)	Opioids (morphine or fentanyl), sedatives (lorazepam, propofol or midazolam), antipsychotics (haloperidol or olanzapine), anticholinergics (atropine, diphenhydramine, bupropion hydrochloride, metoclopramide, prochlorperazine, promethazine)
Pandharipande (2008)	Sedatives, opioids, anticholinergics, antipsychotics, general anaesthesia, histamine blockers, antiarrhythmics, NSAIDs, steroids, antidepressants
Pisani (2007)	History of benzodiazepines or narcotics as an outpatient (25%); and narcotics before ICU admission (20%)
Pisani (2009)	Benzodiazepine or opioids use on admission (25%); during study: benzodiazepine or opioid use (81%), medium to high potency anticholinergic medication use (32%), haloperidol use at any point during the ICU stay (32%), steroid use at any point during ICU stay (52%)
Scott (2001)	All patients received 250 ml of 20% mannitol and 8 mmol of magnesium sulphate
Shulman (2005)	13.66 (SD 8.04) ; number of drugs taken in year prior to first treatment for drug of interest
Williams-Russo (1992)	Medications for psychiatric illness (4%)

3  
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10  
11

One study (Kim 1996) examining the role of H2 antagonists on delirium reported patients taking an H2 antagonist preoperatively were excluded. In two studies (Foy 1995; Pisani 2007) evaluating the use of benzodiazepines, use of benzodiazepines within the month prior to admission was confirmed in 26% of the patients in one study (Foy 1995) and use of benzodiazepines or narcotics was confirmed in 25% of the patients in another study (Pisani 2007).

1 The studies were conducted in the following settings:

- 2 • Four studies in medical wards (Agostini 2001; Centorrino 2003; Foy 1995;  
3 Han 2001);
- 4 • Four studies in the ICU (Dubois 2001; Pandharipande 2006;  
5 Pandharipande 2008; Pisani 2007);
- 6 • Eleven studies were in a surgical setting (Beaussier 2005; Christie 2000;  
7 Herrick 1996; Kim 1996; Leung 2006; Marcantonio 1994; Morrison  
8 2003; Nitschke 1996; Papaioannou 2005; Scott 2001; Williams-Russo  
9 1992);
- 10 • One study (Holroyd 1994) evaluated outpatients;
- 11 • One study (Shulman 2005) did not clearly describe the setting.

12  
13 Type of surgery ranged from cardiac surgery (Kim 1996; Scott 2001); colon  
14 resection surgery (Beaussier 2006; Nitschke 1996), gastrointestinal endoscopy  
15 (Christie 2000) orthopaedic surgery (Herrick 1996; Morrison 2003) general or  
16 orthopaedic surgery (Marcantonio 1994) and mixed types of surgery (Leung  
17 2006:spine/orthopaedic, gynaecological and others; Papaioannou 2005;  
18 gynaecological, orthopaedic, urological, and vascular).

19  
20 Eight studies reported some patients were admitted with multiple diagnoses:

- 21 • cardiopulmonary diseases (Agostini 2001; Christie 2000)
- 22 • hypertension, chronic obstructive pulmonary disease, (Dubois 2001)
- 23 • Central nervous system (CNS) and mental disorders, circulatory, respiratory  
24 (Foy 1995)
- 25 • respiratory, gastrointestinal haemorrhage, sepsis, neurologic, diabetes  
26 mellitus, metabolic abnormalities, acute renal failure and cardiac causes  
27 (Pisani 2007)
- 28 • diabetes mellitus, cardiovascular or respiratory diseases (Papaioannou  
29 2005)
- 30 • sepsis/acute respiratory distress syndrome, pneumonia, myocardial  
31 infarction/congestive failure, chronic obstructive pulmonary disease  
32 (COPD), GI bleeding, drug overdose, hepatic or renal failure,  
33 malignancy, other (Pandharipande 2006)
- 34 • haemorrhage, airway or facial trauma, chest trauma, colonic or gastric  
35 trauma, gastric surgery, neurosurgical trauma, hepatobiliary-pancreatic  
36 surgery, orthopaedic surgery, septic shock or acute respiratory distress  
37 syndrome (ARDS), other (Pandharipande 2008)

38

39 Comorbidities were not reported in the remaining studies.

40

### 1 7.3.3 Pharmacological risk factors

2 The following pharmacological risk factors have been investigated in the  
3 included studies, either in RCTs or in multivariate analyses in prospective cohort  
4 studies; other designs/methods of analysis were included only if there were no  
5 other data. Where reported, the indication for the drug is given if it was  
6 possible that the drug was given to treat delirium.

7

#### 8 7.3.3.1 Benzodiazepines

9

- Midazolam

- one RCT (Christe 2000) used midazolam as a sedative for endoscopy
- two cohort studies (Pandharipande 2006; Pandharipande 2008); both used midazolam as a sedative to reduce anxiety in mechanically ventilated patients

15 • Lorazepam: two cohort studies (Pandharipande 2006; Pandharipande  
16 2008) used lorazepam as a sedative to reduce anxiety in mechanically  
17 ventilated patients

18 • Benzodiazepines (short acting: oxazepam, lorazepam, triazolam,  
19 midazolam, and temazepam) given postoperatively (reason not stated):  
20 one case control study (Marcantonio 1994)

21 • Benzodiazepines (long acting: chlordiazepoxide, diazepam, flurazepam)  
22 given postoperatively (reason not stated): one case control study  
23 (Marcantonio 1994)

24 • Benzodiazepines (not specified): three prospective cohort studies (Foy  
25 1995, prescribed pre-hospital usually for insomnia; Leung 2006, given  
26 postoperatively (reason not stated); Pisani 2007, given before ICU  
27 admission (reason not stated))

28

29 The Pandharipande (2008) study reported that patients may have received  
30 sedative medications as consequence of delirium. The GDG considered this  
31 study likely to be confounded and this study is not considered further.

32

#### 33 7.3.3.2 Antipsychotics:

34

- Clozapine: one retrospective cohort study (Centorrino 2003)

35

- Haloperidol: one cohort study (Pisani 2009), haloperidol indication unclear,  
36 but 70% of patients had agitation on the first day they received  
37 haloperidol

38

### 1 7.3.3.3 Anticholinergics

- 2 • Antihistamines with anticholinergic activity:
  - 3 ○ Diphenhydramine given 24h postoperatively: one prospective
  - 4 cohort study (Agostini 2001) and one case control study
  - 5 (Marcantonio 1994)
  - 6 ○ Benztropine: one retrospective cohort study (Shulman 2005)
- 7 • All medications with anticholinergic activity:
  - 8 ○ All drugs with anticholinergic activity given 24h postoperatively
  - 9 (antihistamines, tricyclic antidepressants, antiemetics, some
  - 10 neuroleptics): one case control study (Marcantonio 1994)
  - 11 ○ Anticholinergics (including antipsychotics and benzodiazepines),
  - 12 purpose not stated, but 43% haloperidol: one cohort study (Han
  - 13 2001)
  - 14 ○ The GDG judged this classification of 'all anticholinergics' to be
  - 15 too vague, so this risk factor was not considered further.
  - 16

### 17 7.3.3.4 H2-receptor antagonists

- 18 • Cimetidine (high dose intravenous) versus ranitidine: one RCT (Kim 1996)
  - 19 ○ The GDG noted that the IV form of cimetidine is rarely used in the
  - 20 UK any more, although low dose oral cimetidine can be bought
  - 21 over the counter. However, this study using a high dose
  - 22 intravenous route did not approximate to the over the counter
  - 23 medicine. Therefore this study was not considered further.
- 24 • H2 blockers (type and dose not specified): one cohort study
- 25 (Pandharipande 2008)
- 26

### 27 7.3.3.5 Mood stabilising drugs

- 28 • Lithium: two retrospective cohort studies (Holroyd 1994; Shulman 2005)
  - 29 ○ Lithium (dose not reported) for mean duration of 7.5 years (SD
  - 30 2.1) (Holroyd 1994) and mean follow up duration of 8.2 months
  - 31 (new users) (Shulman 2005)
  - 32 ○ Valproate: one study; mean follow up duration of 7.5 months
  - 33 (new users) (Shulman 2005)
  - 34

### 35 7.3.3.6 Non Steroidal Anti-inflammatory Drugs (NSAIDs)

- 36 • Ketorolac tromethamine: one RCT (Nitschke 1996)

1

2 **7.3.3.7 Opioids**

- 3 • Morphine: one RCT (Preoperative intrathecal morphine in addition to  
4 postoperative patient controlled analgesia (PCA) morphine (Beaussier  
5 2006)
- 6 • Morphine: two cohort studies (Pandharipande 2006; Pandharipande 2008)
- 7 • Opioids via PCA: two RCTs (Herrick 1996; Nitschke 1996) and one  
8 prospective cohort study (Leung 2006)
- 9 • Opioids general: two cohort studies (Dubois 2003: morphine, fentanyl or  
10 other; Morrison 2003)
- 11 • Meperidine via epidural and via PCA: one case control study (Marcantonio  
12 1994)
- 13 • Meperidine : one cohort study (Morrison 2003)
- 14 • Fentanyl: one case control study (Marcantonio 1994)
- 15 • Fentanyl: one cohort study (Pandharipande 2008)
- 16 • Oxycodone: one case control study (Marcantonio 1994)
- 17 The Pandharipande (2008) study reported that patients may have received  
18 sedative medications as consequence of delirium. The GDG considered this  
19 study likely to be confounded and this study is not considered further.

20

21 **7.3.3.8 Anaesthesia/Analgesia**

- 22 • Thoracic epidural anaesthesia versus opioid analgesia: one RCT (Scott  
23 2001)
  - 24 ○ Bupivacaine plus clonidine perioperatively versus patient  
25 controlled analgesia morphine pump postoperatively; all patients  
26 had general anaesthesia
- 27 • Continuous epidural bupivacaine plus fentanyl (Williams Russo 1992)
- 28 • Nitrous oxide with oxygen versus oxygen: one RCT (Leung 2006)
- 29 • General anaesthesia versus regional anaesthesia: one RCT (Papaioannou  
30 2005)
- 31 • Anaesthetics (unspecified): one cohort study (Pandharipande 2008)

32

33 **7.3.3.9 More than one drug class**

- 34 • Benzodiazepine or opioids : one cohort study (Pisani 2009)

1

2 **7.3.4 Comparisons**3 For the cohort studies the reference for most of these drugs was the absence of  
4 the drug, apart from the following:

- 5
- Leung (2006): PCA opioids relative to oral opioids
6   - Shulman (2005): benzotropine and valproate relative to lithium
7   - Morrison (2003): low dose (below 10 mg) and moderate dose (10 to 30
8 mg) relative to high dose opioid (above 30 mg/day morphine9 equivalent)

10

11 For the RCTs, the following comparisons were carried out:

12

13 **7.3.4.1 Benzodiazepine comparisons**

14 Benzodiazepines versus placebo/no treatment

- 15
- Midazolam (30 µg/kg IV) versus placebo (saline 0.9% IV) (Christe 2000).

16 **7.3.4.2 Opioid comparisons**

- 17
- Opioid versus placebo
    - Intrathecal morphine injected via the 4-5 interspace versus
18 placebo (subcutaneous saline 3 ml injected at the L4-L5 level);
19 both groups also had PCA morphine(300 µg of preservative-free20 morphine [100 µg /ml] (Beaussier 2006)
- 21
- Opioid 1 versus opioid 2
  - PCA fentanyl (10 µg/dose) versus PCA morphine (1 mg/dose)
22 (Herrick 1996)
- 23
- Opioid route of administration 1 versus route 2
  - PCA morphine versus IM morphine (Nitschke 1996)
24   - The doses, intervals and lockout levels for PCA morphine were
25 determined individually based on patients' weight, age and
- 26 serum creatinine level. Dosing interval: every 4 hours for IM27 morphine28
- The doses, intervals and lockout levels for PCA morphine were
- 29 determined individually based on patients' weight, age and30 serum creatinine level. Dosing interval: every 4 hours for IM31 morphine

31

32 **7.3.4.3 Analgesia comparisons**

- 33
- Type of analgesia 1 versus type 2
    - Thoracic epidural anaesthesia perioperatively versus PCA
34 morphine postoperatively (Scott 2001)
35

34

35



1 An adequate method of randomisation was reported in five studies (computer  
2 generated: Beaussier 2006; Leung 2006; Papaioannou 2005; table of random  
3 numbers: Christe 2000; drawing lots: Scott 2001). The remaining three studies  
4 (Herrick 1996; Nitschke 1996; Williams-Russo 1992) did not state the method of  
5 randomisation.

6 An adequate method of allocation concealment was reported in three studies in  
7 which an independent member of staff performed the randomisation (Beaussier  
8 2006; Scott 2001) or this was carried out in the hospital pharmacy (Christe  
9 2000). A partially adequate method of allocation concealment was reported in  
10 two studies (sealed envelope: Leung 2006; Nitschke 1996) and was not  
11 reported or unclear in the remaining studies.

12 Two studies (Leung 2006; Nitschke 1996) reported that the outcome assessors  
13 were blinded to the interventions, one study (Scott 2001) reported blinding was  
14 not maintained and blinding was not clearly stated in the remaining studies.

15 Five studies (Beaussier 2006; Christe 2000; Kim 1996; Leung 2006; Scott 2001)  
16 described an *a-priori* power calculation. In one study (Leung 2006) the sample  
17 size was calculated for the primary outcome, the incidence of delirium. In order  
18 to detect a 50% reduction in delirium for the patients not receiving N<sub>2</sub>O, 114  
19 patients were needed at 80% power,  $p=0.05$ .

20 The remaining studies reported sample size calculations for other outcomes.  
21 Further details are in Appendix E.

22 One study (Christe 2000) reported delirium as an adverse event following  
23 sedation with midazolam or placebo (saline) for an upper gastrointestinal  
24 endoscopy.

25 Six studies reported loss to follow up of less than 20% (Beaussier 2006; Christe  
26 2000; Kim 1996; Nitschke 1996; Papaioannou 2005; Scott 2001)

27 Two studies (Leung 2006; Papaioannou 2006) reported an intention to treat  
28 analysis, two studies (Beaussier 2006; Scott 2001) carried out an available case  
29 analysis and analysis details were not reported or unclear in the remaining  
30 studies.

31 The Papaioannou (2006) study reported conducting both an intention to treat  
32 analysis and a per protocol analysis to examine the effect of type of  
33 anaesthesia on the MMSE score. It was unclear whether an intention to treat or  
34 per protocol analysis was conducted for analysing the incidence of delirium.

35 All studies included in the review demonstrated baseline comparability of the  
36 groups on characteristics such as age, gender, duration of surgery, weight, and  
37 type of surgery.

38  
39 The method of assessment of delirium was:

40 • **adequate** in three studies (CAM: Beaussier 2006; Leung 2006; DSMIII:  
41 Papaioannou 2005);

- 1           • **inadequate** in four studies (Christe 2000: a 3 point decline in MMSE scores  
2           and medical chart review; Herrick 1996: medical chart review; Nitschke  
3           1996: MMSE; Scott 2001: the GDG agreed that 'confusion' was an  
4           inadequate definition of delirium.

5  
6           The overall risk of bias was assessed for the RCTs. Five studies were considered  
7           to have potential for bias and were not considered further: four used an  
8           inadequate method of assessment of delirium (Christe 2000; Herrick 1996;  
9           Nitschke 1996; Williams-Russo 1992) and one (Scott 2001) reported an  
10          inadequate *definition* of delirium. The remaining study (Papaioannou 2005) did  
11          not describe allocation concealment blinding of outcome assessors was not  
12          stated. This study was therefore considered at increased risk of bias.

13

#### 14   **7.4.2 Cohort studies**

15          There were seven reports of six prospective cohort studies (Agostini 2001;  
16          Dubois 2001; Foy 1995; Morrison 2003; Pandharipande 2006; Pisani 2007;  
17          Pisani 2009); three were retrospective cohort studies (Centorrino 2003; Holroyd  
18          1994; Shulman 2005) and one was an RCT that was analysed as a cohort study  
19          for the benzodiazepine risk factor (Leung 2006). In the Centorrino (2003)  
20          study, in patients with more than one admission within the study period, one entry  
21          was randomly selected for analysis without knowledge of delirium.

22  
23          None of the cohort studies were considered to be truly representative of the  
24          population (i.e. adults in surgical and/or medical wards in hospital or long-term  
25          care).

26  
27          In all studies, the non-exposed cohorts were drawn from the same community as  
28          the exposed cohort.

29  
30          Levels of missing data were as follows:

- 31           • Three studies (Dubois 2001; Pisani 2007; Shulman 2005) reported less than  
32           20% missing data, that is, acceptable levels of missing data;  
33           • The remaining studies did not report on missing data.

34  
35          One study (Shulman 2005) reported patients with inconsistent data (0.1%  
36          [11/10230]) were excluded; the Pisani (2007) study reported imputing missing  
37          values (missing: 0.3% for visual impairment to 26% bilirubin)

38  
39          One study (Foy 1995), reported an *a priori* sample size calculation and  
40          calculated that 400 patients would give a power of 98% to detect a relative  
41          risk of 4 for the development of cognitive impairment in the benzodiazepine

1 group. Of the 964 patients screened, 568 patients met the eligibility criteria and  
 2 418 patients were available for analysis. The study reported separate results  
 3 for the development of cognitive impairment and delirium.

4  
 5 The studies varied in the number of patients with prevalent delirium (delirium at  
 6 baseline): further details are given in Appendix D.

7 • Four reported that none of the patients had delirium at baseline (Agostini  
 8 2001; Foy 1995; Morrison 2003 (patients with delirium not enrolled);  
 9 Shulman 2005)

10 • Two studies reported that some of the patients had delirium at baseline  
 11 (Dubois 2001: 4% [9/216]; Pandharipande 2006: at least 33% with  
 12 delirium [66 +/198] )

13 • One study reported these patients were excluded (Dubois 2001);

14 • Three reports of two studies reported the number of patients who  
 15 developed delirium following admission (Morrison 2003: 16% [87/541];  
 16 Pisani 2007: 70.4% [214/304] within first 48h of ICU admission; Pisani  
 17 2009: 79% [239/304] during the ICU stay)

18  
 19 One study (Pandharipande 2006) reported the number of patients who  
 20 experienced delirium during ICU admission who were administered antipsychotics  
 21 [88%: 66/75] and anticholinergic drugs [83%: 52/63]. Information on delirium  
 22 status is missing for 30% (60/198) of the patients.

23  
 24 The method of delirium assessment used was:

25 • **Adequate** in four studies:

- 26 ○ Assessed with CAM-ICU and the Richmond Agitation Sedation  
 27 Scale (Pandharipande 2006)
- 28 ○ Assessed with CAM-ICU on weekdays and medical chart review  
 29 at weekends (Pisani 2007)
- 30 ○ Assessed with CAM on weekdays and medical chart reviewed at  
 31 weekends (for key words: for example, 'delirious/delirium'  
 32 'agitated/agitation' to supplement the CAM observations);  
 33 delirium was diagnosed if either the CAM or the medical record  
 34 chart criteria were met (Morrison 2003)
- 35 ○ MMSE scores and nurse assessed checklists to assess orientation,  
 36 overall cognitive function, level of alertness and personal care  
 37 and staff description of nocturnal events to assess criteria  
 38 according to DSM IIR criteria (Foy 1995);

39  
 40 • **Partially inadequate** in two studies:

- 41 ○ Assessed by intensivist and confirmed by a formal psychiatric  
 42 assessment (Dubois 2001)

- 1                   ○ Multivariate analysis only for 'cognitive decline', which consisted  
2 of commonly accepted delirium symptoms in addition to  
3 standardised, validated instruments including CAM for delirium  
4 and MMSE (Agostini 2001)

5

6                   • **Inadequate** in two studies:

- 7                   ○ Assessed from medical charts, and from a 3 point severity scale  
8 [mild, moderate, severe]. (Centorrino 2003)
- 9                   ○ Informa0-+
- 10                  ○ tion on delirium (classified as a side effect) was extracted by the  
11 author in a chart using a structured instrument (no further  
12 information on the instrument) (Holroyd 1994).

13

14                  The method of assessment was not reported in one study (Shulman 2005).

15

16                  Confounders taken into account

17                  We considered whether the cohort studies took account of particular  
18 confounders, either in the study design or the multivariate analysis. The GDG had  
19 identified, by consensus, three risk factors to be important: age, sensory  
20 impairment, and cognitive impairment.

21                  Studies were summarised according to the number of key risk factors included in  
22 the multivariate analysis and the ratio of events to covariates (the GDG  
23 considered a ratio of 1 or less to be flawed and a ratio of 2 or 3 to be possibly  
24 confounded). We assumed that the key risk factors were the same for severity of  
25 delirium and duration of delirium.

26                  Eight reports of nine studies conducted multivariate analyses (Agostini 2001;  
27 Dubois 2001; Foy 1995; Morrison 2003; Pandharipande 2006; Pisani 2007;  
28 Pisani 2009; Shulman 2005). Two studies conducted only univariate analyses  
29 (Centorrino 2003; Holroyd 1994) and these are not considered further. Further  
30 details of the factors included in the multivariate analysis are given in Appendix  
31 F.

- 32
- 33                  • One study had all/most (3 or 2) of the important risk factors taken into  
34 account in the multivariate analysis or they were held constant and had a  
35 ratio of events to variables of 10 or more:

- 36                  ○ Shulman (2005): valproate vs lithium: ratio: 12 [72/6];  
37 benztropine vs lithium: 16 [93/5]; key factors were taken into  
38 account: age, hearing and visual impairment; patients with  
39 dementia were excluded so treated as a constant

- 1           • Two studies had all/most (3 or 2) of the important risk factors taken into  
2           account in the multivariate analysis or they were held constant but had  
3           insufficient ratio of events to variables:
- 4           ○ Morrison (2003): ratio: 5 [87/16]; key risk factors taken into  
5           account: age, cognitive impairment.
- 6           ○ Pandharipande (2006): ratio ranging from: 4 [66/17] to  
7           7[118/17]; key risk factors taken into account: age, visual and  
8           hearing deficits, dementia
- 9           ○ The study reported the number of patients who experienced  
10          delirium for two subgroups: those who received antipsychotics  
11          (66/75) and those who received anticholinergics (52/63); it is  
12          unclear whether any of the patients were prescribed both drugs.  
13          We estimated the incidence of delirium, with incidence ranging  
14          from 33% (66/198: the minimum number who had delirium) to  
15          60% (118/198; assuming that patients received either  
16          antipsychotics or anticholinergics).
- 17          • Six reports of seven studies were possibly confounded: not enough of the  
18          important risk factors were taken into account in the multivariate analysis:
- 19          ○ Agostini 2001) ratio: 31 [122/4] had one key risk factor (age) in  
20          the analysis and patients with profound dementia were excluded.
- 21          ○ Foy (1995) ratio: 2[21/12]; one key risk factor was taken into  
22          account: age
- 23          ○ Leung (2006) ratio:18 [90/5] had one key risk factor taken into  
24          account: age
- 25          ○ Pisani (2007) ratio: 9 [214/23] had one key factor taken into  
26          account: dementia (IQCODE score greater than 3.3)
- 27          ○ Pisani (2009) ratio: 30 [304/10]; key risk factor taken into  
28          account: dementia (IQCODE score greater than 3.3)
- 29          ○ Dubois (2001 ratio: 5 [38/7] had no risk factors taken into  
30          account

31

#### 32 **7.4.2.1 Overall quality for the cohort studies**

- 33          • Two cohort studies were considered to be biased and were not considered  
34          further:
- 35          ○ Retrospective study and the method of assessment for delirium  
36          was not reported (Shulman 2005);
- 37          ○ None of the key risk factors were taken into account (Dubois  
38          2001)
- 39          • Five reports of four cohort studies were given a low overall quality and  
40          treated with caution (evaluated in sensitivity analysis):
- 41          ○ Only one key risk factor was taken into account (Agostini 2001;  
42          Foy 1995; Leung 2006; Pisani 2007; Pisani 2009); and Foy  
43          (1995) also had a ratio of 2.

- 1 • Two studies (Morrison 2003; Pandharipande 2006) were given a  
2 moderate quality rating.  
3

#### 4 7.4.3 Case control studies

5 The case control study (Marcantonio 1994) was not considered to be truly  
6 representative of the population (i.e. adults in surgical and/or medical wards in  
7 hospital or long-term care). The Marcantonio (1994) study was in a surgical  
8 setting and the non-exposed cohort was drawn from the same community as the  
9 exposed cohort.

10  
11 The study did not report on missing data or on an *a priori* sample size  
12 calculation. The study reported 9% (117/1341) of the patients had delirium at  
13 baseline (Marcantonio 1994).

14  
15 The method of delirium assessment was adequate (CAM).

#### 16 Confounders taken into account

17  
18 We considered whether the case control study took account of particular  
19 confounders, either in the study design or the multivariate analysis. Cases and  
20 controls were matched for: age; poor cognitive function; poor physical function;  
21 self reported alcohol abuse; markedly abnormal preoperative serum sodium,  
22 potassium or glucose levels; aortic aneurism surgery; and noncardiac thoracic  
23 surgery. Thus matching was carried out on two of the key risk factors (age and  
24 cognitive impairment). A matched analysis was carried out with drugs being  
25 analysed by a logistic regression method so that the effect of each was obtained  
26 independently.

27  
28 Overall, the case control study was both considered to be of low quality because  
29 of its design and was considered only if there were no other data.

### 30 31 7.5 Results

32 We consider below the effects of different risk factors on the incidence, duration  
33 and severity of delirium. Results from RCTs and prospective cohort studies are  
34 reported mainly and case control studies where there is no other evidence.

#### 35 36 7.5.1 Benzodiazepines as a risk factor for the incidence of delirium

37 Two low quality prospective cohort studies (Leung 2006; Pisani 2007), one  
38 moderate quality prospective cohort study (Pandharipande 2006) and one case

1 control study (Marcantonio 1994) reported the effect of benzodiazepines on the  
2 incidence of delirium.

3

#### 4 **7.5.1.1 Benzodiazepine dose as a continuous variable**

##### 5 Midazolam

6 One moderate quality cohort study (Pandharipande 2006) evaluated the use of  
7 midazolam (sedative for mechanically ventilated patients to reduce anxiety) as a  
8 risk factor for delirium. The analysis considered the transition from normal,  
9 delirious or comatose states during the previous 24h to either normal or delirious  
10 states in the following 24h. the Pandharipande (2006) study reported that there  
11 were small numbers of patients receiving midazolam.

12

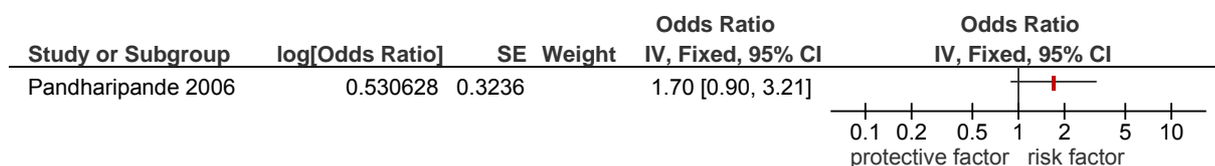
13 The Pandharipande (2006) study reported the effect of dose (in mg) of  
14 midazolam in the previous 24 hours, as a continuous variable, on the incidence of  
15 delirium [OR 1.70 (95% CI 0.90 to 3.21); figure 7.1].

16

17 There was no significant effect of midazolam on the incidence of delirium.

18

19 Figure 7.1: Midazolam as a risk factor for development of delirium



20

##### 21 Lorazepam

22 One moderate quality cohort study (Pandharipande 2006 ) evaluated the use of  
23 lorazepam (as a sedative for mechanically ventilated patients to reduce anxiety)  
24 as a risk factor for delirium. The multivariate analysis considered the transition  
25 from normal, delirious or comatose during the previous 24h to either normal or  
26 delirious status in the following 24h. The number of patients who received  
27 lorazepam was not reported.

28

29 The Pandharipande (2006) study reported the effect of dose (in mg) of  
30 lorazepam in the previous 24 hours, as a continuous variable, on the incidence of  
delirium (figure 7.2).

31

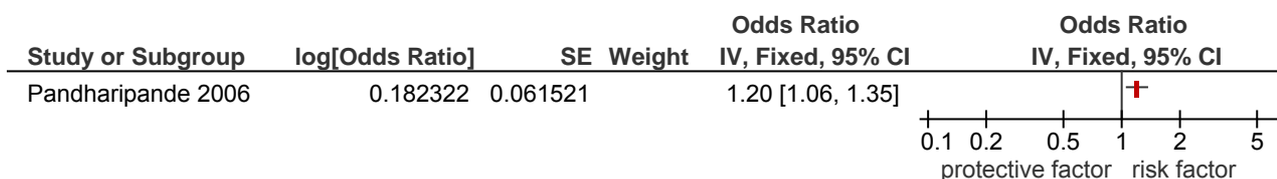
32 The study reported that administration of lorazepam in the previous 24h resulted  
33 in a 20% increased risk in transition to delirium in the range 0 to 40 mg [OR 1.2  
34 (95% CI 1.06 to 1.35)]. The study also reported that the incremental risk was  
35 large at low doses and the risk of delirium versus dose reached a plateau at 20  
mg. It is unclear how this affected the multivariate analysis.

36

1

2

Figure 7.2: lorazepam as a risk factor for development of delirium



3

#### 4 7.5.1.2 Benzodiazepines as dichotomous variables

5 Three low quality cohort studies (Foy 1995; Leung 2006; Pisani 2007) and one  
6 case control study (Marcantonio 1994) evaluated the use of benzodiazepines as  
7 a dichotomous risk factor for delirium. The Foy (1995) study evaluated as a risk  
8 factor the use of benzodiazepines within 5 days of admission, the Marcantonio  
9 (1994) study and the Leung (2006) study evaluated postoperative use on day 1  
10 and days 1 or 2 respectively and Pisani (2007) evaluated use before admission  
11 to the ICU.

12

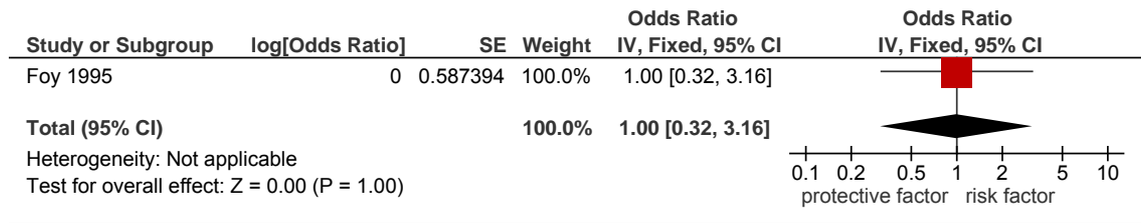
13 The Marcantonio (1994) study reported exposure to long-acting agents,  
14 including chlordiazepoxide, diazepam and flurazepam, compared with short-  
15 acting agents, including oxazepam, lorezapam, triazolam, midazolam and  
16 temazepam. Type of benzodiazepines in the Foy (1995) study were diazepam,  
17 oxazepam, temazepam, nitrazepam, bromazepam, flunitrazepam, and  
18 clorazepate, usually these were prescribed for insomnia. Type of  
19 benzodiazepine was not specified in two studies (Leung 2006; Pisani 2007).  
20 Indications for benzodiazepine use were not reported. The GDG decided that  
21 the studies in which benzodiazepines were given postoperatively were likely to  
22 be confounded: it was anticipated that a new prescription of a benzodiazepine  
23 would be given for agitation. Therefore, these studies were not considered  
24 further.

25 In the remaining study (Foy 1995), the incidence of delirium was 5% (21/418)  
26 and exposure to benzodiazepines was indicated by self-report in 23%  
27 (96/418) of the patients.

28 The odds ratio was 1.0 (95% CI 0.3 to 3.0) indicating use of benzodiazepines 5  
29 days before admission was not a significant risk factor for delirium (figure 7.3).

30

1 Figure 7.3: benzodiazepines as a risk factor for delirium



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3  
4

5 **7.5.2 Antipsychotics**

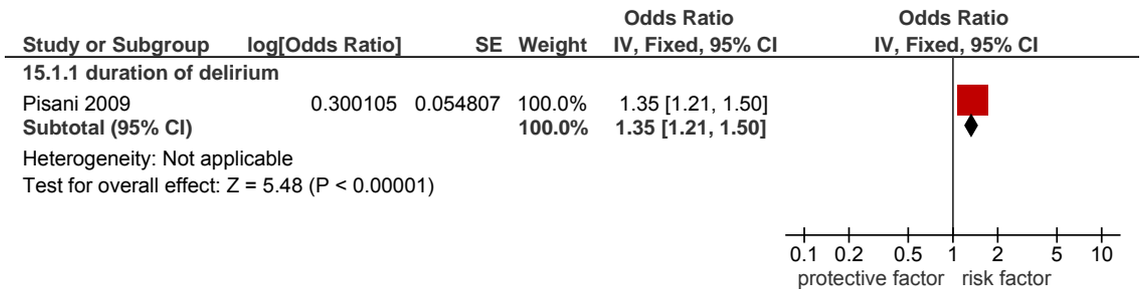
6

7 **7.5.2.1 Haloperidol as a risk factor for increased duration of delirium**

8 One low quality cohort study (Pisani 2009) evaluated use of haloperidol as a  
9 risk factor for increased duration of delirium in ICU. The study reported that  
10 haloperidol was a significant risk factor for the increased duration of delirium  
11 (OR 1.35 (95% 1.21 to 1.50) (figure 7.4). The study stated that the haloperidol  
12 indication was unclear, but 70% of patients had agitation on the first day they  
13 received haloperidol. The GDG considered this study likely to be confounded.

14  
15

Figure 7.4: Haloperidol as a risk factor for duration of delirium



16  
17

NB: Scale 0.1 to 10

18

19 **7.5.3 Anticholinergics**

20 Two studies examined specific drugs with anticholinergic activity as a risk factor  
21 for delirium: one prospective cohort study (Agostini 2001) and one case control  
22 study (Marcantonio 1994) evaluated diphenhydramine. The GDG advised that  
23 diphenhydramine should be classified as an antihistamine with anticholinergic  
24 activity.

25  
26  
27  
28  
29

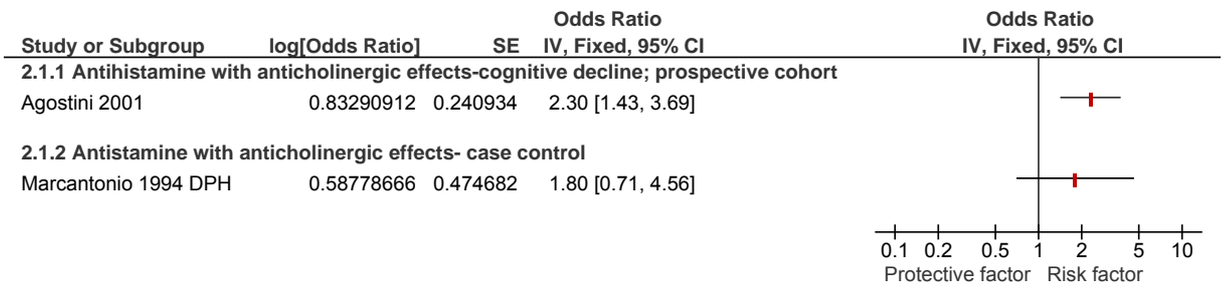
One low quality prospective cohort study (Agostini 2001) with 426 patients reported a multivariate analysis (controlling for age, gender and baseline delirium risk) for the risk of cognitive decline in diphenhydramine-exposed group. Cognitive decline was assessment was based on CAM rating for delirium,

MMSE scores and presence of delirium symptoms. The number of patients meeting the CAM delirium criteria and decline in MMSE score ( $\geq 3$  points) was 13% (9/71) in patients receiving the 25mg dose, 17% (7/43) in patients receiving 50mg dose, and 8% (25/312) in patients who did not receive diphenhydramine. 67% of the patients (59/114) were administered the drug for one day and 1 patient received the drug for six consecutive days. Mean number of doses per patient was 2.1 (SD 1.6), and the maximum cumulative daily dose given was 100 mg. Indications for use of diphenhydramine included sleep (68%) and agitation (0.4%).

The Marcantonio 1994 (study) reported results for diphenhydramine administered to 7.3% of the patients (18/245). Of the 22 patients receiving all anticholinergics, 68% (15/22) received a low-dose (defined as one therapeutic dose or less; for example, 25mg for diphenhydramine). The remaining patients (7/22) were administered a higher dose, given in either single or multiple doses. Indications for the use of diphenhydramine were not reported.

The odds ratio ranged from 1.80 (95% CI 0.71 to 4.56) to 2.30 (95% CI 1.43 to 3.69) for antihistamines (with anticholinergic activity); figure 7.5. We note that both studies had a potential for bias.

Figure 7.5: antihistamines with anticholinergic activity

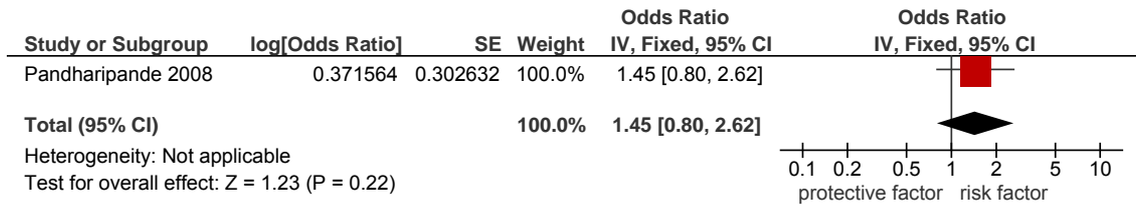


#### 7.5.4 H2 receptor antagonists (H2 blockers)

One cohort study (Pandharipande 2006) evaluated whether exposure to histamine blockers (type not specified) in the previous 24 hours was a risk factor for delirium. The number of patients who received H2 blockers was not reported. There was no significant effect of H2 blockers as a risk factor for delirium [OR 1.45 (95% CI 0.80 to 2.62); figure 7.6].

1  
2

Figure 7.6: exposure to H2 blockers on the incidence of delirium

3  
4

## 5 7.5.5 Opiate analgesics

6 Six studies evaluated opioid analgesics as a risk factor for delirium: four  
7 evaluated the effects of individual opioids (cohort studies: Morrison 2003;  
8 Pandharipande 2006; Pandharipande 2008; case control: Marcantonio 1994);  
9 one considered the class of opioids (cohort study: Morrison 1994); one RCT  
10 examined the added effect of morphine (Beaussier 2006); one cohort study  
11 (Leung 2006) compared PCA postoperative opioid analgesia versus oral  
12 administration. The case control study (Marcantonio 1994) examined the effect  
13 of different types of opioid (meperidine, morphine, fentanyl and oxycodone);  
14 because there are higher quality studies reporting the effects of meperidine,  
15 morphine and fentanyl, only the results for oxycodone are presented.

16

### 17 7.5.5.1 Effect of individual opioids

18 Two prospective cohort studies (Morrison 2003; Pandharipande 2006) and one  
19 case control study (Marcantonio 1994) evaluated the effect of exposure to  
20 individual opioids on the incidence of delirium. The Pandharipande (2006) study  
21 reported the effect of dose of the individual opioid in the previous 24 hours, as a  
22 continuous variable, on the incidence of delirium. The Pandharipande (2006)  
23 study accounted for the delirium status for only 69% of the patients. The study  
24 reported the number of patients who experienced delirium for two subgroups:  
25 those who received antipsychotics (66/75) and those who received  
26 anticholinergics (52/63); it is unclear whether any of the patients were  
27 prescribed both drugs. We estimated the incidence of delirium, with incidence  
28 ranging from 33% (66/198: the minimum number who had delirium) to 60%  
29 (118/198; assuming that patients received either antipsychotics or  
30 anticholinergics).

#### 31 Opioids as continuous variables

##### 32 *Fentanyl*

33 One moderate quality cohort study (Pandharipande 2006) evaluated the effects  
34 of administration of fentanyl (every unit dose in mcg) in the previous 24h on  
35 delirium status. Details on doses and number of patients who were administered  
36 the drugs were not reported.

37 The study showed no significant effect of fentanyl as a risk factor for the  
38 incidence of delirium. The confidence interval is wide (figure 7.7a).

1  
2*Morphine*

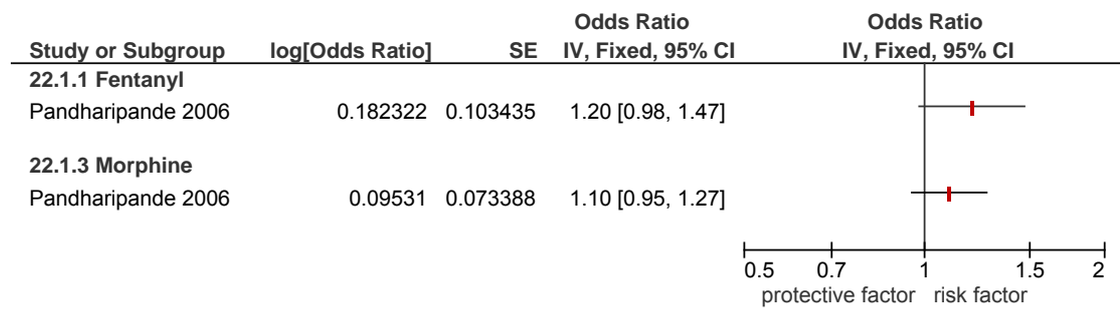
3 One moderate quality cohort study (Pandharipande 2006) evaluated the effect  
4 of morphine on the incidence of delirium. Details on doses and number of  
5 patients who were administered the drugs were not reported. Exposure of  
6 morphine (every unit dose in mg) in the previous 24h on delirium status was  
7 reported (OR 1.10 (95% CI 0.95 to 1.27). The confidence interval is wide.

8 Although this is not a significant effect (OR 1.10), This means that for every  
9 increment of a unit dose (in mg) of morphine, the odds of having delirium could  
10 increase by a factor of 1.10. Therefore for a 10 mg dose increase, the odds  
11 increase by  $(1.10)^{10}$ , which is 3.00, with the odds ratio ranging from  $(0.95)^{10}$  to  
12  $(1.27)^{10}$ , which is 2.59 to 3.56.

13 The Pandharipande (2006) study showed no significant effect of morphine on the  
14 incidence of delirium (figure 7.7a).

15

16 Figure 7.7a: Effect of individual opioids on delirium

17  
18

19

20 NB: Scale 0.5 to 2

21

22 Opioids as dichotomous variable

23

*Meperidine*

24 One moderate quality study (Morrison 2003) evaluated meperidine use as a risk  
25 factor for the development of delirium following admission for hip fracture. 21%  
26 of the delirious patients (27/129) received meperidine following admission.  
27 Meperidine is a significant risk factor: RR 2.4 (95% CI 1.3 to 4.5); figure 7.77.

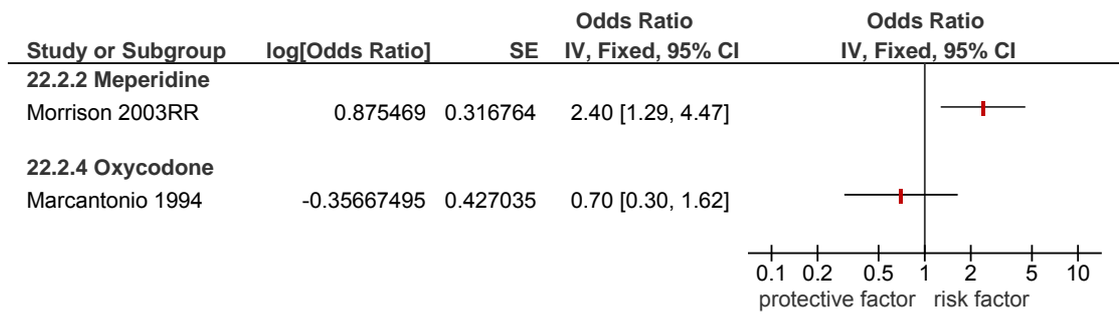
28

1 Oxycodone

2 One case control study (Marcantonio 1994) examined the effect of oxycodone  
 3 administered during a 24 hour period on the incidence of delirium; 10% of the  
 4 patients with delirium (9/91) received oxycodone. Details on dose were not  
 5 reported, nor were indications for the use of oxycodone. There was no significant  
 6 effect on the incidence of delirium of oxycodone: RR 0.70 (95% CI 0.30 to  
 7 1.62); figure 7.7b.

8

9 Figure 7.7b Effect of individual opioids on delirium



10

11

12 **7.5.5.2 Effect of all opioids: dose effect**

13 The Morrison (2003) study evaluated the effect on delirium incidence of three  
 14 different dose ranges (less than 10 mg; 10 mg to 30 mg; above 30 mg)  
 15 different total daily doses of parenteral morphine sulphate equivalents; doses of  
 16 all opioids, including continuous infusions and PCA were converted to equivalent  
 17 dosage. The total daily opioid dose for delirious patients was calculated for the  
 18 24 hours preceding the delirious episode and the highest 24h cumulative opioid  
 19 dose for the first 3 postoperative days for non-delirious patients. The total  
 20 number of patients who received opioid at the following dose ranges were as  
 21 follows: below 10 mg: 38% (204/541); 10 to 30 mg: 36% (192/541); above  
 22 30 mg 23% (145/541). The study reported the pattern of opioid use in  
 23 cognitively intact patients (44%: 242/541).

24

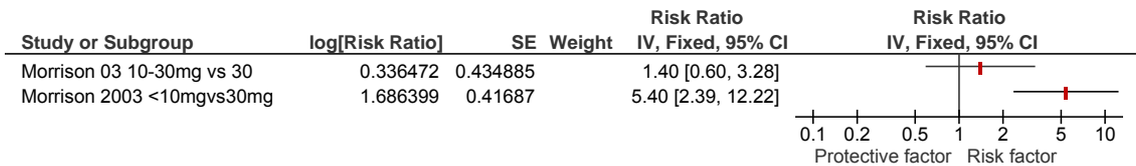
25 There was a significant effect of parenteral morphine sulphate equivalents on the  
 26 incidence of delirium observed in patients receiving low doses (below 10 mg  
 27 compared with the reference above 30mg): RR 5.40 (95% CI 2.39 to 12.22).  
 28 There was no significant effect of the medium dose (10 to 30 mg) parenteral  
 29 morphine sulphate equivalents on the incidence of delirium: RR 1.40 (95% CI  
 30 0.60 to 3.28); figure 7.8.

31

32 The authors suggested that it is the untreated pain, as opposed to a low dose of  
 33 opioid, that is the risk factor for developing delirium; the GDG concurred.

34

1 Figure 7.8: Effect of opioids on the incidence of delirium

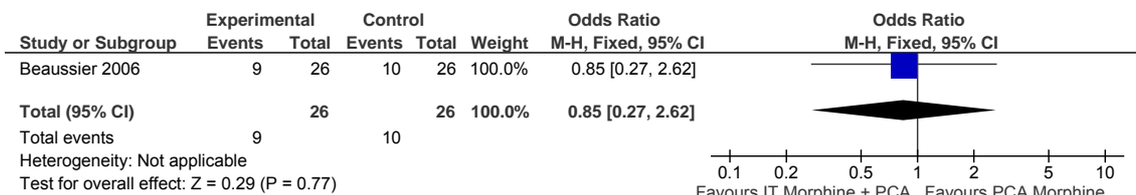


3  
4

### 5 7.5.5.3 Preoperative morphine in addition to postoperative patient controlled analgesia

6 One RCT (Beaussier 2006) compared the additional effect of preoperative  
7 intrathecal morphine on the incidence of delirium in 52 older people recovering  
8 from major colorectal surgery. The study compared intrathecal (IT) morphine 0.3  
9 mg (preoperatively) followed by patient controlled analgesia (PCA) morphine  
10 (postoperatively), versus preoperative subcutaneous saline plus PCA morphine  
11 postoperatively in the control group. The incidence of delirium was 35% (9/26)  
12 and 38% (10/36) in the IT morphine plus PCA morphine group and the placebo  
13 plus PCA morphine group, respectively. The CI is wide, indicating a low level of  
14 precision. The result is imprecise (figure 7.9).

15  
16 Figure 7.9: effect of intrathecal morphine + PCA morphine versus placebo +  
17 PCA morphine



19  
20

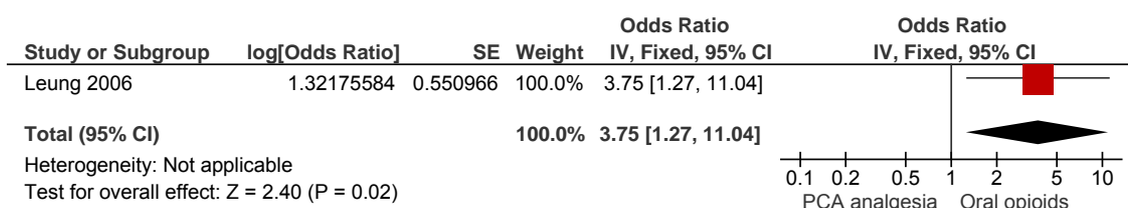
### 21 7.5.5.4 Comparison of different routes of administration of opioids postoperatively

22 One low quality prospective cohort study (Leung 2006) compared the effects of  
23 different routes of delivery of postoperative opioids (PCA opioids versus oral  
24 opioids) on the incidence of delirium during recovery.

25  
26 The multivariate analysis (adjusted for age, anaesthesia type, dependence on  
27 performing at least one ADL, postoperative analgesia, use of benzodiazepines)  
28 showed a higher risk of delirium in patients who received PCA, compared with  
29 oral opioids (figure 7.10). PCA administration of opioids was a significant risk  
30 factor for delirium compared with oral opioids; OR 3.75 (95% CI 1.27, 11.04);  
31 the CI is wide, indicating some uncertainty in the magnitude of the effect (figure  
32 7.10). No details were given regarding the oral opioids, and the doses were not  
33 reported for either route.

34

1 Figure 7.10: Effect of PCA opioid analgesics versus oral opioids



2  
3

4 **7.5.6 Anaesthesia**

5 Three studies (Leung 2006; Papaioannou 2005; Pandharipande 2008)  
6 investigated the effects of anaesthesia on delirium: one RCT at higher risk of bias  
7 (Papaioannou 2005) compared general with regional anaesthesia (epidural or  
8 spinal), one RCT (Leung 2006) compared nitrous oxide and oxygen versus  
9 oxygen alone and one cohort study (Pandharipande 2008) evaluated the effect  
10 of anaesthetics on the incidence of delirium.

11

12 **7.5.6.1 General anaesthesia versus regional anaesthesia**

13 One RCT (Papaioannou 2005) compared the incidence of delirium in patients  
14 receiving general anaesthesia (n=25) versus those receiving regional  
15 anaesthesia (epidural or spinal) (n=25) for orthopaedic, urological, vascular or  
16 gynaecological surgery. Details on type of anaesthetic agents and dose were  
17 not stated. Duration of anaesthesia was over 120 min in over half the cases.  
18 Benzodiazepines were not administered for premedication or intraoperative  
19 sedation.

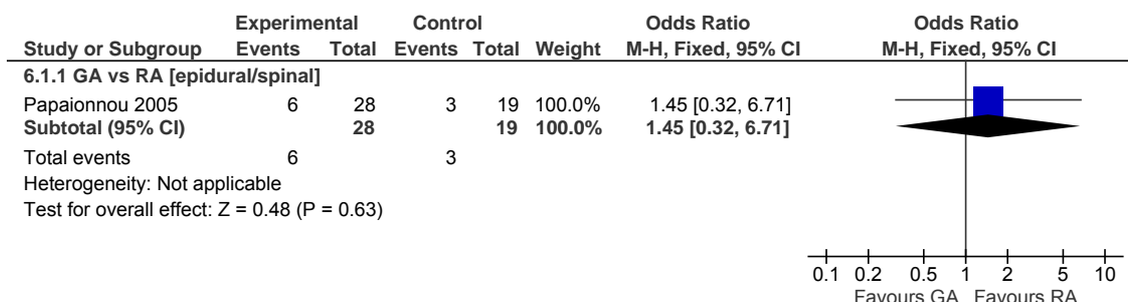
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21 The incidence of delirium was 21% (6/28) and 16% (3/19) in the general and  
22 regional groups, respectively in the Papaioannou (2005) study. There was no  
23 significant effect of type of anaesthesia on delirium, although the results are  
24 very imprecise. (figure 7.11).

25

26 Figure 7.11: Effect of general anaesthesia versus regional anaesthesia on  
27 delirium

28

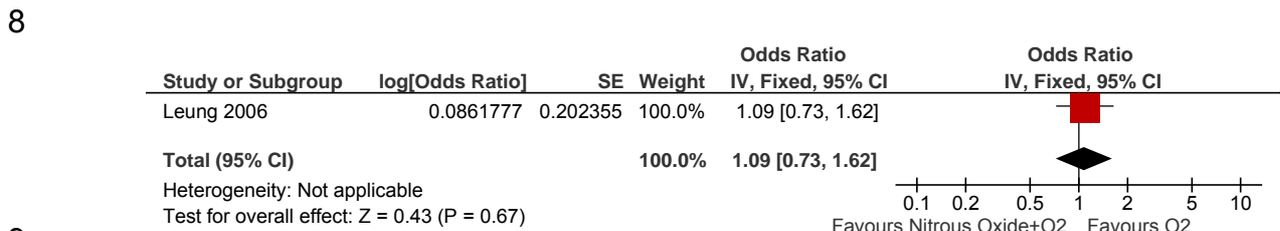


29  
30

### 1 7.5.6.2 N<sub>2</sub>O plus oxygen versus oxygen

2 In one RCT (Leung 2006) 228 patients were randomised to receive nitrous oxide  
3 plus oxygen or oxygen alone to evaluate if there was a difference in the  
4 incidence of delirium during recovery from general anaesthesia. There was no  
5 significant difference (figure 7.12), although the results are imprecise.

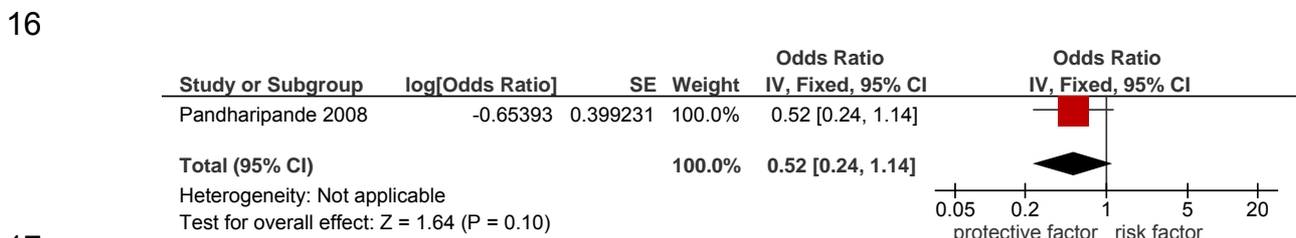
6  
7 Figure 7.12: Effect of N<sub>2</sub>O plus O<sub>2</sub> versus O<sub>2</sub> on delirium



### 10 7.5.6.3 Anaesthesia

11 One study (Pandharipande 2008) reporting the effect of exposure to  
12 anaesthetics (type not reported) on the incidence of delirium showed no  
13 significant effect; OR 0.52 (95% CI 0.23 to 1.16); figure 7.13.

14  
15 Figure 7.13: Effect of anaesthetics on delirium



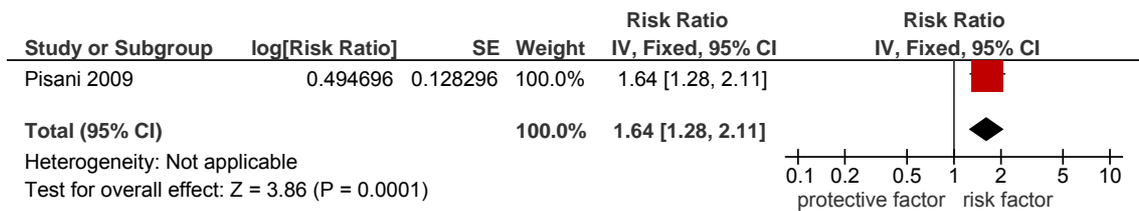
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### 20 7.5.7 Effect of benzodiazepines or opioids on the duration of delirium

21 One study (Pisani 2009) evaluated the use of benzodiazepines or opioids as a  
22 risk factor for the duration of delirium; 81% (247/304) of the patients were  
23 administered benzodiazepines or opioids. There was a significant effect of use  
24 of these drugs on the duration of delirium in ICU, but results were not reported  
25 separately for the two classes of drugs; RR 1.64 (95% CI 1.27 to 2.10); figure  
26 7.14. The GDG considered the results from this study set in the ICU had limited  
27 applicability when compared to other hospital populations. The GDG noted that  
28 in the ICU patient group, the methods of administration, dose, indication and  
29 intention of drug use is often very different to other hospital populations.

30

Figure 7.14: Effect of benzodiazepines or opioids on the duration of delirium



## 7.6 Evidence summary / statements

- There is moderate quality evidence to show no significant effect of midazolam on the incidence of delirium.
- There is moderate quality evidence to show there is a significant effect of lorazepam as a risk factor for the incidence of delirium.
- There is low quality evidence indicating that the use of benzodiazepines 5 days before admission was not a significant risk factor for the incidence of delirium.
- There is low quality evidence to show that diphenhydramine (an antihistamine with anticholinergic activity) is a significant risk factor for the incidence delirium; there is some uncertainty with this result.
- There is very low quality evidence to show diphenhydramine (an antihistamine with anticholinergic activity) is not a significant risk factor for the incidence delirium.
- There is moderate quality evidence to show no significant effect of H2 blockers on the incidence of delirium.
- There is inconsistent evidence for the effect of individual opioids on delirium.
  - There is moderate quality evidence to show no significant effect of fentanyl on the incidence of delirium.
  - There is moderate quality evidence to show meperidine is an important risk factor for the incidence of delirium.
  - There is moderate quality evidence to show no significant effect of morphine on the incidence of delirium.
  - There is very low quality evidence to show no significant effect of oxycodone on the incidence of delirium.
- There is moderate quality evidence to show untreated pain is a significant risk factor for the incidence of delirium.
- There is moderate quality evidence from one RCT to show preoperative morphine in addition to patient controlled analgesia in the postoperative period is not a significant risk factor for delirium. There is some uncertainty with this result.

- 1 • There is low quality evidence showing patient controlled administration of  
2 opioids was a significant risk factor for delirium compared with oral  
3 opioids. There is some uncertainty with this result.
- 4 • There is moderate quality evidence from one RCT to show there was no  
5 significant effect of type of anaesthesia (general compared with regional  
6 anaesthesia) on delirium. There is much uncertainty with this result.
- 7 • There is moderate quality evidence from one RCT to show no significant  
8 difference in the incidence of delirium in patients receiving nitrous oxide  
9 plus oxygen or oxygen alone.
- 10 • There is low quality evidence to show anaesthesia is not an important risk  
11 factor for the incidence of delirium.
- 12 • There is low quality evidence to show use of benzodiazepines or opioids  
13 is a significant risk factor for the duration of delirium in ICU.

## 1 8 Consequences of delirium

### 2 8.1 Clinical introduction

3 Delirium has the potential to have an effect on a wide range of outcomes for the  
4 delirious person themselves, their family or carers, and health and social care  
5 organisations. Some of these may be a direct result of damage caused by the  
6 inflammatory response to delirium, whereas others may be a consequence of  
7 delirium affecting motor control and behaviour. In addition, many outcomes may  
8 also be affected by the index condition that is causing the delirium. Establishing  
9 the effect delirium has on outcomes can be challenging, with many potential  
10 confounding variables to be considered. This review examines the evidence for  
11 an independent effect of delirium on outcomes affecting individuals (such as  
12 mortality, the development of dementia, falls) and organisations (length of  
13 hospital stay, institutionalisation) which will help to demonstrate the impact of  
14 delirium and identify areas for improvement.

### 15 16 8.2 Description of studies

17 Thirty six papers were evaluated for inclusion and 24 reports of 19 studies were  
18 included (Andrew 2005; Balas 2009; Bourdel-Marchasson 2004; Dolan 2000;  
19 Drame 2008; Ely 2004; Francis 1990; Francis 1992; Holmes 2000; Nightingale  
20 2001; Inouye 1998; Leslie 2005; Levkoff 1992; Lin 2004; Lin 2008;  
21 Marcantonio 2000; Givens 2008; Marcantonio 2002; McAvay 2006; O’Keeffe  
22 1997; Pitkala 2005; Rockwood 1999; Rudolph 2008; Thomason 2005). Twelve  
23 studies were excluded and reasons for exclusion are listed in Appendix G. One  
24 study (Bickel 2008) was subsequently identified. The study has not been  
25 reported in depth as it was of low quality and would have been excluded in the  
26 sensitivity analysis.

27 Three studies had more than one report, which differed in the outcomes reported  
28 (Francis 1990 and Francis 1992; Holmes 2000 and Nightingale 2001;  
29 Marcantonio 2000, Marcantonio 2002 and Givens 2008). Hereafter, these  
30 studies are referred to by the first named reports, but are reported separately  
31 where appropriate and reported separately in the results section. One report  
32 (Lin 2008) included some of the same patients included in the Lin (2004) study  
33 but reported different outcomes and are reported separately. Two studies  
34 (Leslie 2005; McAvay 2006) included some of the same patients but reported  
35 different outcomes and are reported individually.

36  
37 This review examines the evidence for the consequences associated with  
38 presence of prevalent or incidence delirium, increased delirium duration and  
39 increased delirium severity. The following are reported:

- 40 • Dementia/cognitive impairment/cognitive dysfunction:
  - 41 ○ Cognitive impairment at discharge (Ely 2004);
  - 42 ○ Cognitive dysfunction at 7 days (Rudolph 2008);

- 1                   ○ Cognitive dysfunction at 3 months (Rudolph 2008);
- 2                   ○ Dementia at 3 years (Rockwood 1999).
- 3                   • New admission to institution
- 4                   ○ At discharge (Balas 2009; Bourdel-Marchasson 2004; Inouye
- 5                   1999; Levkoff 1992);
- 6                   ○ 3 months (Inouye 1999);
- 7                   ○ 6 months (O’Keeffe 1997);
- 8                   ○ 2 years (Pitkala 2005).
- 9                   • Mortality
- 10                  ○ In hospital (Inouye 1998; O’Keeffe 1997);
- 11                  ○ In ICU ( Lin 2004);
- 12                  ○ In ICU and hospital (Lin 2008; Thomason 2005);
- 13                   - 1 month (Marcantonio 2000);
- 14                   - 6 weeks (Drame 2008);
- 15                   - 3 months (Inouye 1998);
- 16                   - 6 months (Ely 2004 [incidence and duration of delirium];
- 17                   Francis 1990; Holmes 2000; Levkoff 1992; Marcantonio
- 18                   2000; O’Keeffe 1997);
- 19                   - 1 year (Leslie 2005 [incidence and severity of delirium];
- 20                   Pitkala 2005);
- 21                   - 2 years (Dolan 2000; Francis 1992; Nightingale 2001;
- 22                   Pitkala 2005);
- 23                   - 3 years (Rockwood 1999).
- 24                  • Length of stay
- 25                  ○ Hospital (Ely 2004 [incidence and duration of delirium]); Francis
- 26                  1990; Holmes 2000; Levkoff 1992; Thomason 2005; O’Keeffe
- 27                  1997);
- 28                   - The Holmes (2000) study reported the risk of being
- 29                   discharged sooner, which corresponds to decreased risk of
- 30                   remaining in hospital. This outcome will be grouped with
- 31                   studies reporting length of stay and the key confounding
- 32                   factors identified for length of stay would be applicable
- 33                   for this outcome.
- 34                  ○ ICU (Thomason 2005);
- 35                  ○ Post ICU (Ely 2004 [incidence and duration of delirium]).
- 36                   - The Ely (2004) study defined post ICU stay as length of
- 37                   stay after first ICU discharge.
- 38

1 The GDG agreed, *post-hoc*, that the following outcomes, including composite  
2 ones, identified during the course of the review, should also be included:

- 3 • Hospital acquired complications (O’Keeffe 1997);
- 4 • Cognitive dysfunction (Rudolph 2008 [incidence and duration of delirium]);
  - 5 ○ The GDG agreed that for incidence of delirium, cognitive
  - 6 dysfunction can be grouped with studies reporting dementia and
  - 7 cognitive impairment and that the key confounding factors
  - 8 identified for dementia would be applicable for this outcome.
- 9
- 10 • Mortality or new admission to institution
  - 11 ○ At discharge (Inouye 1998);
  - 12 ○ At 1 month (Givens 2008; Marcantonio 2000; Marcantonio 2002
  - 13 [severity of delirium]);
  - 14 ○ At 3 months (Inouye 1998)
  - 15 ○ At 6 months (Givens 2008; Marcantonio 2000; Marcantonio
  - 16 2002 [severity of delirium])
  - 17 ○ At 1 year (McAvay 2006);
  - 18 ○ At 2 years (Pitkala 2005)
- 19
- 20 • Mortality or functional decline at discharge and at 6 months (Andrew 2005
- 21 [duration of delirium])
- 22

23 One additional study (Francis 1992) reported the outcome ‘loss of independent  
24 living’ defined as ‘patients institutionalised or needing assistance on 1 of 4 ADL’.  
25 The GDG thought that for this outcome, patients needing assistance on 1 of 4  
26 ADL may be confounded by stroke (10% of patients with cerebrovascular  
27 diseases) and advised that this outcome should not be included in the review.

28 The Rudolph (2008) study also reported a subgroup analysis for two different  
29 durations of delirium, not allowing for duration of delirium in the multivariate  
30 analysis. This outcome will not be considered in this review.

31 The general characteristics of the studies including methodological quality are  
32 discussed for all studies first. These are reported separately for each outcome,  
33 where appropriate, and the results are reported separately for each  
34 consequence.

35

### 36 **8.3 Characteristics of included studies**

#### 37 **8.3.1 Study Design**

38 All the studies were prospective cohort studies and funding, where reported, was  
39 non industry.

1 Three studies reported patients were part of either the intervention and/or  
2 control group in a trial (Leslie 2005: intervention and control groups enrolled in a  
3 delirium prevention intervention (Inouye 1999); McAvay 2006: control group of  
4 Delirium Prevention Trial (Inouye 1999); Marcantonio 2000: intervention and  
5 control arms of a trial described as a randomised trial on prevention of delirium  
6 [proactive geriatric consultation]).

7 Two studies were conducted in the UK (Holmes 2000; O’Keeffe 1997), ten  
8 studies in the USA (Balas 2009; Dolan 2000; Ely 2004; Francis 1990; Inouye  
9 1998; Leslie 2005; Levkoff 1992; Marcantonio 2000; McAvay 2006; Thomason  
10 2005), two in Canada (Andrew 2005; Rockwood 1999), two in France (Bourdel-  
11 Marchasson 2004; Drame 2008), one in Finland (Pitkala 2005) and two in  
12 Taiwan (Lin 2004; Lin 2008). One study (Rudolph 2008) was multinational and  
13 recruited patients from eight countries: UK, Denmark, France, Germany, Greece,  
14 the Netherlands, Spain and USA.

15 Six reports of five studies had fewer than 200 patients (Andrew 2005: n=77;  
16 Balas 2009: n=117; Lin 2004: n=131; Lin 2008: n=143; Marcantonio 2000  
17 n=126; Marcantonio 2002: n=122); nine studies had between 200 and 500  
18 patients (Bourdel-Marchasson 2004: n=427; Ely 2004: n=275; Francis 1990:  
19 n=229; Levkoff 1992: n=325; McAvay 2006: n=433; O’Keeffe 1997: n=225;  
20 Pitkala 2005: n=425; Rockwood 1999: n=203; Thomason 2005: n=261); three  
21 studies had between 500 patients and 1000 patients (Dolan 2000: n=682;  
22 Holmes 2000: n=731; Inouye 1998: n=727) and two studies recruited more than  
23 1000 patients (Drame 2008: n=1036; Rudolph 2008: n=1218).

24 One study was conducted in both hospital and long-term care; the latter was the  
25 setting for 53% of the patients (Pitkala 2005). All the remaining studies were  
26 conducted in hospitals. Patients were in different types of wards:

- 27 • medical (Bourdel-Marchasson 2004; Dolan 2000; Drame 2008; Francis  
28 1992; Leslie 2005; McAvay 2006; O’Keeffe 1997; Rockwood 1999).  
29 Where reported, the principal diagnoses of patients admitted to medical  
30 wards were:
  - 31 ○ hip fracture (Dolan 2000);
  - 32 ○ cancer, coronary artery disease, congestive heart failure, chronic  
33 lung disease, cerebrovascular disease, diabetes, hypertension  
34 (Francis 1992);
  - 35 ○ pneumonia, chronic lung disease, congestive heart failure, ischemic  
36 heart disease, gastrointestinal disease, diabetes mellitus or  
37 metabolic disorder, cancer, cerebrovascular disease, renal failure,  
38 anaemia, and other conditions (Leslie 2005).
- 39
- 40 • surgical (Marcantonio 2000; Rudolph 2008). For these patients, the surgery  
41 was:
  - 42 ○ hip fracture repair (Marcantonio 2000) ;
  - 43 ○ non cardiac surgery (Rudolph 2008).

- 1 • ICU (Balas 2009; Ely 2004; Lin 2004; Thomason 2005). Patients were in  
2 ICU for the following reasons:
- 3 ○ mechanically ventilated patients (Ely 2004; Lin 2004);
- 4 - Principal admission diagnoses of sepsis and/or acute  
5 respiratory distress syndrome (46%), pneumonia,  
6 myocardial infarction/congestive heart failure, hepatic or  
7 renal failure, chronic obstructive pulmonary disease,  
8 gastrointestinal bleeding, malignancy, drug overdose, and  
9 other diagnoses not stated (Ely 2004);
- 10 - Principal admission diagnoses of pneumonia (34%), chronic  
11 lung disease, cerebrovascular disease, cancer, congestive  
12 heart failure, ischemic heart disease, gastrointestinal  
13 disease, diabetes mellitus or metabolic disorder, drug  
14 intoxication and other diagnoses not stated (Lin 2004);
- 15 ○ non-ventilated [non invasive] patients. (Thomason 2005);
- 16 - Diagnostic admission for pulmonary (27%),  
17 gastrointestinal, metabolic, cardiac,  
18 haematology/oncology, neurological, renal, and other  
19 reasons not stated.
- 20 ○ surgical ICU (Balas 2009)
- 21 - 42.1% received mechanical ventilation at sometime during  
22 Surgical Intensive Care Unit (SICU) admission
- 23 - Type of surgery included general (colorectal, surgical  
24 oncology and gastrointestinal surgery), vascular, and  
25 trauma/emergency surgery.
- 26 • mixture of medical and surgical wards (Inouye 1998; Levkoff 1992).
- 27 ○ reasons for admission included:
- 28 - cancer, coronary artery disease, cardiac arrhythmias,  
29 congestive heart failure, chronic lung disease, pneumonia,  
30 gastrointestinal, cerebrovascular disease diabetes, renal  
31 disease and other conditions not reported (40%); number  
32 of surgical patients and type of surgery was not reported  
33 (Inouye 1998);
- 34 - circulatory system disease (29.2%), digestive system  
35 disease, respiratory system disease, fracture, cancer,  
36 genitourinary system disease, endocrine, nutritional and  
37 metabolic diseases, diseases of skin or other reasons not  
38 stated. Type of surgery was not reported (Levkoff 1992).
- 39 • mixture of medical (32%), surgical (19%) and geriatric wards (48%)  
40 (Andrew 2005).
- 41
- 42 Eight studies reported the settings from which patients were admitted:
- 43 ○ community (Dolan 2000; Francis 1990);
- 44 ○ emergency units (Drame 2008);

- 1           ○ community (65%) and the remaining patients from long-term care
- 2           (Levkoff 1992);
- 3           ○ community (41%), nursing homes(4%) and the remaining
- 4           admission were unclear (Inouye 1998);
- 5           ○ 6.1% from nursing home (Leslie 2005);
- 6           ○ community (93%) and the remainder from nursing homes
- 7           (Marcantonio 2000);
- 8           ○ community (81%) and remaining patients from long-term care or
- 9           residential home care (O'Keeffe 1997).

### 11 8.3.2 Population

12           The mean age, where reported, ranged from 55 years (Ely 2004) to 82.1 years

13           (Holmes 2000). The age range was reported in four studies (Andrew 2005;

14           Drame 2008; Holmes 2000; McAvay 2006) and the range was estimated from

15           the mean  $\pm$  1 standard deviation in the remaining studies (table 8.1).

16

17

18           Table 8.1: patient ages

Study	Mean age and range (years)	Study	Mean age and range (years)
Andrew 2005	78.5 (64 to 93)	Leslie 2005	80 (73.5 to 86.5) <sup>±</sup>
Balas 2009	75.4 (69.1 to 81.7) <sup>±</sup>	Levkoff 1992	81.4 (73.7 to 89.1) <sup>±</sup>
Bourdel-Marchasson 2004	85 (78.4 to 92.4) <sup>±</sup>	Lin 2004	73.6 (70.5 to 77.4) <sup>±</sup>
Dolan 2000	82 (72.6 to 90.1) <sup>±</sup>	Lin 2008	76 (64 to 85.5)
Drame 2008	85 (75 to 103)	McAvay 2006	80 (70 to 99)
Ely 2004	55 (37 to 73) <sup>±</sup>	Marcantonio 2000	79 (71 to 87) <sup>±</sup>
Francis 1992	78 (72.1 to 85.0) <sup>±</sup>	O'Keeffe 1997	82 (76 to 88) <sup>±</sup>
Holmes 2000	82.1 (65 to 99)	Rudolph 2008	69 (62.9 to 76.3) <sup>±</sup>
Inouye 1998	78.9 (72 to 85.8) <sup>±</sup>	Thomason 2005	52.5 (32 to 74) <sup>±</sup>

19           (±) indicates that range was calculated from the mean  $\pm$  1 standard deviation

20

21           The age range was not stated and could not be calculated in two studies (Pitkala

22           2005; Rockwood 1999). The Pitkala (2005) study, however, reported that

23           patients younger than 70 years were excluded and that 59% were over 85

24           years. In the Rockwood (1999) study patients over 65 years were enrolled and

1 the mean age of 79 years was reported. In the Francis (1990) study patients  
2 over 70 years were enrolled and had a mean age of 78 years.

3 Where reported, all studies included both males and females. Two studies  
4 (Holmes 2000; Pitkala 2005) had less than 20% male patients, twelve studies  
5 had less than 50% (Andrew 2005; Dolan 1997; Drame 2008; Francis 1990;  
6 Inouye 1998; Leslie 2005; Levkoff 1992; Marcantonio 2000; McAvay 2006;  
7 O’Keeffe 1997; Rockwood 1999; Thomason 2005) and five studies had 50% or  
8 more male patients (Balas 2009; Ely 2004; Lin 2004; Lin 2008; Rudolph 2008).  
9 The Bourdel-Marchasson (2004) study did not report the number of male and  
10 female patients enrolled.

11 Fifteen studies reported including patients with cognitive impairment (Andrew  
12 2005; Balas 2009; Bourdel-Marchasson 2004; Drame 2008; Francis 1990;  
13 Holmes 2000; Inouye 1998; Leslie 2005; Levkoff 1992; Lin 2008; McAvay  
14 2006; Marcantonio 2000; O’Keeffe 1997; Pitkala 2005; Rockwood 1999), one  
15 study (Dolan 2000) reported patients with cognitive impairment were excluded,  
16 three studies (Lin 2004; Lin 2008; Rudolph 2008) reported that patients with  
17 dementia were excluded, and cognitive impairment was not reported in one  
18 study (Thomason 2005). Cognitive impairment ranged from 24% (Levkoff 1992)  
19 to 75% (Bourdel-Marchasson 2004). Assessment of cognitive impairment was  
20 based on the following scales:

- 21 • MMSE (range 0 to 30) (Holmes 2000; Inouye 1998; McAvay 2006; Pitkala  
22 2005; Rudolph 2008);
  - 23 ○ one study (Inouye 1998) used a cut off score of 20 or below to  
24 define dementia; a cut off score of below 24 were used in two  
25 studies (Ely 2004; McAvay 2006); patients with score of 24 or  
26 below were excluded in one study (Rudolph 2008) and the cut-  
27 off point was not reported in one study (Holmes 2000);
    - 28 - The Inouye (1998) multicentre study used a 21 point scale  
29 MMSE at one of the three sites, and scores on the 21 point  
30 scale were adjusted to a denominator of 30 points;
  - 31 ○ the Pitkala (2005) study used a score below 20 to define  
32 moderate cognitive impairment;
- 33 • Blessed’s Dementia Rating Scale (Francis 1990; Leslie 2005; Lin 2008;  
34 Marcantonio 2000; O’Keeffe 1997);
  - 35 ○ The cut-off point was 4 or more in three studies (Francis 1990;  
36 Marcantonio 2000; O’Keeffe 1997); 2 or more in one study  
37 (Leslie 2005; modified version of Blessed scale); 3 or higher (Lin  
38 2008)
- 39 • DSM III-R criteria (Andrew 2005);
- 40 • cognitive status (MMSE, Blessed dementia rating scale) and functional  
41 assessment (Barthel Index, Physical Self-Maintenance Scale) to screen for  
42 cognitive impairment and assessment of dementia by geriatrician  
43 (Rockwood 1999);
- 44 • based on family interviews and physicians and checked if existed with  
45 respect to DSM-IV criteria (Bourdel-Marchasson 2004);

- 1           • IQCODE (Balas 2009);
- 2           • medical chart review or assessment of a senior practitioner (Drame 2008);
- 3           • medical chart review (Levkoff 1992).

4  
5 Further details are reported in Appendix D.

6  
7 Ten studies reported comorbidity scores, using the Charlson Comorbidity Index:  
8 (Bourdel-Marchasson 2004; Dolan 2008; Drame 2008; Ely 2004; Leslie 2005;  
9 McAvay 2006; Marcantonio 2000; O’Keeffe 1997; Pitkala 2005; Thomason  
10 2005). Further details are reported in Appendix D.

11 Eight studies reported severity of illness assessed with an established scale  
12 (APACHE II: Balas 2009; Ely 2004; McAvay 2006; Leslie 2005; Inouye 1998;  
13 Thomason 2005; APACHE III: Lin 2004; Lin 2008). Two studies used a clinician  
14 based rating (Francis 1992; Levkoff 1992), severity of illness based on a rating  
15 scale (range 1 to 9, with 1 = not ill and 9 = moribund) (Francis 1992) and a sum  
16 of severity scores, calculated based on severity scores assigned to 15 medical  
17 conditions: one study (Levkoff 1992).

18 One study (Holmes 2000) reported using a researcher-rated scale, the modified  
19 Burvill scale to record concurrent physical illness (range: 0 to 6, with 0  
20 representing no physical illness and 6 representing severe chronic physical  
21 illness).

22 Further details are reported in Appendix D.

23

### 24 **8.3.3 Incidence of delirium and its method of assessment**

25 Overall rates of delirium ranged from 8% (Bourdel-Marchasson 2004; Rudolph  
26 2008) to 48% (Thomason 2005).

27 All of the patients in one study (Andrew 2005: n=77) had delirium; this study  
28 was looking at the effects of increased duration of delirium.

29 The studies varied in whether they investigated the effects of prevalent delirium  
30 (occurring on admission to hospital) or incident delirium (appearing during the  
31 course of the hospital stay) or both.

32           • Nine studies included only prevalent delirium as a risk factor (Andrew  
33 2005; Dolan 2005; Drame 2008; Holmes 2000; Inouye 1998; Lin 2004  
34 (ICU study using delirium developed in first 5 days); Lin 2008 (ICU study  
35 using delirium developed in first 5 days); Pitkala 2005 (only recorded  
36 prevalent delirium; Rockwood 1999 (only recorded prevalent delirium))

37           • Four studies (Balas 2009; Leslie 2005 (patients with prevalent delirium  
38 were excluded); McAvay 2006 (patients with prevalent delirium were

1 excluded); Marcantonio 2000 (reported to be incident delirium)) included  
2 only incident delirium rates

3 • One study (Bourdel-Marchasson 2004) included both prevalent and  
4 incident delirium. and analysed them separately

5 • Four studies (Ely 2004; Francis 1990; Rudolph 2008; Thomason 2008)  
6 reported both incident and prevalent delirium, but combined them as  
7 'delirium' in the analysis

8 • Two studies (Levkoff 1992; O'Keeffe 1997) reported both prevalent and  
9 incident delirium and combined these in some analyses (Levkoff 1992:  
10 mortality, length of stay; O'Keeffe 1997: mortality; length of stay;  
11 hospital acquired complications) but both reported only incident delirium  
12 for discharge to an institution.

13

14 Rates of delirium ranged from 8% (Rockwood 1999:16/203) to 82% (Ely 2004:  
15 183/224).

16 The Bourdel-Marchasson (2004) study reported four categories of delirium: for  
17 patients classified as having prevalent delirium [8%:34/427] if the diagnosis of  
18 delirium was within the first 4 days of stay, subsequent delirium was classified as  
19 incident [3.5%:15/427], prevalent subsyndromal delirium [20.6%:88/427] and  
20 incident subsyndromal delirium [14%:60/427]. Patients having one or more  
21 CAM symptoms but not fulfilling the CAM algorithm were termed 'subsyndromal  
22 delirium'. Results for patients with only prevalent and incident delirium will be  
23 reported in this review.

24 In addition to examining the consequences of either prevalent and/or incident  
25 delirium, the GDG wanted to investigate the effect of persistent delirium on  
26 adverse outcomes. Persistent delirium was classified in accordance with the  
27 definition provided in the McAvay (2006) study. These authors defined persistent  
28 delirium as 'patients who met full criteria for delirium at the discharge interview,  
29 or had full delirium during the hospitalisation and partial symptoms at  
30 discharge'.

31 Four studies reported information on persistent delirium (Levkoff 1992;  
32 Marcantonio 2000; McAvay 2006; O'Keeffe 1997).

33 Persistent delirium rates were reported for the following time periods:

34 • discharge: ranged from 17% (Levkoff 1992: 54/325) to 32% (O'Keeffe  
35 1997 [24%: 8/33 of those with prevalent delirium; 37%: 17/46 of those  
36 with incident delirium]);

37 • 1 month: 29% (Marcantonio 2000: 15/52);

38 • 3 months: 16.2% (Levkoff 1992);

39 • 6 months: ranged from 6% (Marcantonio 2000: 3/52) to 13.3% (Levkoff  
40 1992);

41 • 1 year: 43% (McAvay 2006: 24/55).

42

1 In the Levkoff (1992) study only the percentages of patients with resolved  
2 delirium were reported from which the persistent delirium rates were calculated.

3 The method of assessment of persistent delirium differed from baseline  
4 assessment in one study (Levkoff 1992). At 3 and 6 months follow-up, relatives  
5 or carers were interviewed to determine if symptoms persisted. This was deemed  
6 an inadequate method of assessment.

7 In one study (Rockwood 1999), the study population was also separated into  
8 patients with delirium and dementia at baseline (11%: 22/203), prevalent  
9 dementia only (8%:17/203) and patients with neither delirium nor dementia  
10 (73%:148/203). For the outcome, dementia as a consequence of delirium, results  
11 were only presented for the combined groups, patients with delirium and  
12 patients with neither delirium nor dementia.

13 In one study (Ely 2004), 67% (123/183) of patients who had delirium for a  
14 median of 2 days (IQR 1 to 3) were in a coma for a median of 2 days (IQR 1 to  
15 4).

16 The method of assessment of delirium varied between the studies. The GDG  
17 considered that 19 studies had an adequate method of assessment; two had a  
18 partially adequate method; one had a partially inadequate method and one  
19 was inadequate:

## 20 **Adequate**

- 21 • Ten studies used either the Confusion Assessment Method (CAM) (Bourdel-  
22 Marchasson 2004; Inouye 1998; Leslie 2005; Marcantonio 2000;  
23 McAvay 2006) or a variation (CAM-ICU: Balas 2009 ; Ely 2004;  
24 Thomason 2005; Chinese version of CAM ICU: Lin 2004; Lin 2008).
- 25 • One study (Balas 2009) reported patients were considered delirious if  
26 patient scored positive on the CAM-ICU and the RASS (score  $\geq$  -3)
- 27 • Three studies (Drame 2008; Pitkala 2005; Rockwood 1999) reported that  
28 delirium was classified based on DSM-IV criteria
- 29 • Two studies (Andrew 2005; Francis 1990) reported that delirium was  
30 classified based on DSM III.
- 31 • One study (Rockwood 1999) study used the Delirium Rating Scale
- 32 • One study (Holmes 2000) used the MMSE to identify patients with cognitive  
33 impairment and the Delirium Rating Scale was used to differentiate  
34 between delirium and dementia
- 35 • One study (Levkoff 1992) used the Delirium Symptom Interview (DSI) which  
36 assesses the domains of delirium specified in DSM III
- 37 • One study (O’Keeffe 1997) used the Delirium Assessment Scale (DAS),  
38 based on the DSM-III criteria for delirium

39

### Partially inadequate

- One study (Rudolph 2008) reported that delirium was classified based on DSM III.
  - The method of delirium assessment was not consistent: patients were assessed with MMSE and medical records until postoperative day 3 and from day 4 until discharge, evaluation was based on the medical and nurse chart
  - Criterion 5 of the DSM-III was not a requirement [‘evidence, from the history, physical examination, or laboratory tests of a specific organic factor judged to be etiologically related to the disturbance’]. Primary caregiver or other informant was interviewed to identify symptoms that were new or had worsened within the week before hospital admission.

### Inadequate

- One study (Dolan 2000) had a review of medical notes and/or proxy interview using CAM [proxies were family members or friends who could report on the patient’s health]

The GDG considered the Dolan (2000) study to be biased because the method of assessment was based on review of medical notes and/or interview with proxy. The GDG agreed that the three studies (Levkoff 1992; O’Keefe 1997; Rudolph 2008) which used the DSM III (or methods based on DSM III) for assessment were acceptable if the method of assessment remained consistent throughout the duration of the study. However, in comparing with other studies, these studies should be treated with caution.

#### 8.3.3.1 Assessment of severity

One study (Marcantonio 2002) used the Memorial Delirium Assessment Scale (MDAS) (range 0 to 30, with 30 indicating high severity) to assess severity of delirium and used 12.44 [the median of the average MDAS score for all patients with delirium] as the cut-off point between mild and severe delirium. Results were presented by severity of delirium.

#### 8.3.4 Methodological quality of included studies

One study (Pitkala 2005) was considered to be truly representative of the population (i.e. adults in long-term and hospital settings) and the remaining studies were considered to be somewhat representative of the population.

The non-exposed cohort was drawn from the same community as the exposed cohort.

### 1 8.3.4.1 Missing data by outcome

#### 2 • Dementia

- 3 ○ One study (Rockwood 1999) reported less than 20% missing  
4 data (i.e. acceptable levels) for the outcome dementia;
- 5 ○ One study (Rudolph 2008) reported less than 20% missing data  
6 (i.e. acceptable levels) for the outcome postoperative cognitive  
7 dysfunction at 7 days
- 8 ○ One study (Rudolph 2008) reported less than 20% missing data  
9 for the outcome postoperative cognitive dysfunction at 3 months,  
10 and here the authors showed that the 19% of missing data was  
11 not missing at random because those with delirium were twice as  
12 likely not to complete the testing, which indicates potential for  
13 bias;
- 14 ○ One study (Ely 2004) was considered to have too high levels of  
15 missing data for the outcome cognitive impairment (28%) – these  
16 patients were not tested because of their inability to complete  
17 testing or because of rapid discharge. This also indicates  
18 potential for bias.

#### 19 • New admission to institution

- 20 ○ Five studies (Balas 2009; Bourdel-Marchasson 2004; Inouye  
21 1998: at discharge; O’Keeffe 1997; Pitkala 2005) reported less  
22 than 20% missing data (i.e. acceptable levels). In one study (Balas  
23 2009) the missing data were due to patients remaining in hospital  
24 at the time of study closure and voluntary withdrawal from the  
25 study. In the remaining studies, the missing data were due to  
26 deaths
- 27 ○ One study (Inouye 1998) had about 20% missing data at 3  
28 months follow up, but most of these were due to death or being  
29 lost to follow up: the missing group reportedly did not differ  
30 significantly from the completing group;
- 31 ○ The level of missing data was not reported in one study (Levkoff  
32 1992).

#### 33 • Mortality

- 34 ○ Seven reports of 6 studies had no missing data (Holmes 2000  
35 [Nightingale 2001]; Inouye 1998- discharge; Levkoff 1992;  
36 Marcantonio 2000: 1 month; O’Keeffe 1997; Rockwood 1999;);
- 37 ○ Eleven studies stated there was less than 20% missing data (i.e.  
38 acceptable levels) (Dolan 2000; Ely 2004; Drame 2008; Francis  
39 1990; Inouye 1998: 3 months; Leslie 2005; Lin 2004; Lin 2008;  
40 Marcantonio 2000: 6 months; Pitkala 2005; Thomason 2005).

- 1           • Length of stay
- 2           ○ Three studies (Ely 2004: hospital; O’Keeffe 1997: hospital;
- 3           Thomason 2008: hospital and ICU) reported less than 20%
- 4           missing data (i.e. acceptable levels);
- 5           ○ One study (Ely 2004: post ICU) had 29% missing data because
- 6           of deaths in ICU and patients in a persistent coma. The former
- 7           (10%) may have biased the outcome, but was at a low level;
- 8           ○ Holmes (2001) reported no missing data.
- 9           • Hospital acquired complications
- 10          ○ One study (O’Keeffe 1997) had no missing data.
- 11          • Mortality or new admission to institution
- 12          ○ Three reports of two studies (Givens 2008 at 1 month and 6
- 13          months; Marcantonio 2000: 1 month; McAvay 2006: 1 year) had
- 14          no missing data;
- 15          ○ Two studies (Inouye 1998- discharge; 3 months; Marcantonio
- 16          2000: 6 months) reported less than 20% missing data.
- 17          • Mortality or functional decline
- 18          ○ One study (Andrew 2005) reported no loss to follow up for the
- 19          outcome at discharge and less than 20% loss to follow up at 6
- 20          months.

#### 21    **8.3.4.2 Assessment of delirium**

22           As discussed above, the GDG considered that 19 studies had an adequate

23           method of assessment; one had a partially inadequate method (Rudolph 2008)

24           and one was inadequate (Dolan 2000).

25

#### 26    **8.3.4.3 Outcome of interest at baseline**

- 27           • Dementia
- 28           ○ One study (Rockwood 1999) excluded patients with dementia
- 29           from the analysis.
- 30           ○ One study (Ely 2004) assessing cognitive impairment reported the
- 31           baseline modified Blessed Dementia rating score [range: 0 to 17]
- 32           (mean (SD): 0.23(SD0.8): 0.14 (SD 0.6) for the delirious and non-
- 33           delirious groups, respectively, indicating none of the patients
- 34           were likely to have dementia.
- 35           ○ One study (Rudolph 2008) assessing postoperative cognitive
- 36           dysfunction reported that patients with a score of less than 23 on
- 37           the MMSE were excluded but did not provide baseline scores for
- 38           the neuropsychological tests used to assess postoperative
- 39           cognitive dysfunction.

- 1           • New admission to institution
- 2           ○ Five studies (Bourdel-Marchasson 2004; Inouye 1998; Levkoff
- 3           1992; O’Keeffe 1997; Pitkala 2005) reported patients in long-
- 4           term care settings at admission were excluded from the analysis
- 5           for this outcome.
- 6           ○ In one study (Balas 2009) patients in long-term care setting at
- 7           admission [3.5%: 4/114] were included in the analysis
- 8           • Hospital acquired complications (falls, pressure sores, urinary incontinence
- 9           and any other complication)
- 10          ○ One study(O’Keeffe 1997) reported patients with a pressure
- 11          sore corresponding to Grade 2 of Shea’s classification (Shea
- 12          1975) on admission were excluded; patients with frequent
- 13          incontinence or with a catheter on admission were excluded from
- 14          the analysis; history of falls was not reported;
- 15          • Mortality or new admission to institution
- 16          ○ Mortality: not applicable;
- 17          ○ New admission to institution:
- 18               - One study (McAvay 2006) excluded patients admitted to
- 19               hospital from a nursing home
- 20               - Three reports of two studies (Inouye 1998; Marcantonio
- 21               2000; Marcantonio 2002) reported new admission to
- 22               institutions for patients who had not been previously
- 23               institutionalised at time of admission
- 24          • Mortality or functional decline
- 25          ○ mortality: not applicable;
- 26          ○ functional decline: the mean baseline Barthel index score was
- 27          86.6 (range 42 to 100) indicating some patients had less
- 28          likelihood of living independently prior to hospitalisation (Andrew
- 29          2005)

30  
31

#### 32 **8.3.4.4 Confounders taken into account:**

33           The overall quality rating of the study was made taking into account the number

34           of key risk factors, the method of delirium assessment, missing data in addition to

35           the ratio of events to covariates.

36           All the included studies conducted multivariate analyses. The Marcantonio (2000)

37           and Givens (2008) studies reported the same outcomes but adjusted for

38           different variables in the multivariate analysis.

39           In relation to the events to covariate ratio, the GDG provided the following

40           guidance:

Delirium: full guideline DRAFT (November 2009)

- 1           • ratio of 1 or less: biased;
- 2           • ratio of 2 or 3: possibly confounded and rated as low quality;
- 3           • ratio of 4 to 7: moderate quality feature;
- 4           • ratio of 8 to 10: high quality feature.

5  
6           The rest of this section examines the ratio of events to covariates and the number  
7           of key risk factors for each outcome.

8  
9           A. Risk factor: presence of prevalent or incident delirium

10  
11           1. *Dementia/cognitive impairment/cognitive dysfunction*

12  
13           The GDG identified age, depression, and cognitive impairment as the key  
14           confounding factors. None of the studies included depression in the analyses, and  
15           studies were not downgraded if this risk factor was missing.

- 17           • One study had 2/3 of the important risk factors taken into account in the  
18           multivariate analysis, or held constant and the ratio of events to variables  
19           was at least 10. Patients with an MMSE score of 23 or less were  
20           excluded from the study.

- 21           ○ Rudolph (2008) ratio: 66 [265/4]; [7 days postoperative  
22           dysfunction]; 24 [94/4]; [3 months postoperative dysfunction];  
23           key factor was age, and cognitive impairment was constant

- 25           • Two studies had 2/3 of the important risk factors taken into account in the  
26           multivariate analysis but had an insufficient ratio of events to variables.

- 27           ○ Ely (2004) ratio: 5 [63/12]; key risk factors were: age, cognitive  
28           impairment (dementia);
- 29           ○ Rockwood (1999) ratio: 8 [32/4]; key factor was: age ; patients  
30           with dementia excluded from analysis

31  
32  
33           2. *Progression of dementia*

34  
35           The GDG identified age and gender as the key confounding factors. There were  
36           no studies identified reporting this outcome.

37  
38           3. *New admission to an institution*

39  
40           The GDG identified ADL, cognitive impairment, and depression as the key  
41           confounding factors. None of the studies included depression in the analyses.

- 43           • Three studies had 2/3 of the important risk factors taken into account in the  
44           multivariate analysis and had a ratio of number of events to variables of  
45           at least 10.

- 1                   ○ Bourdel-Marchasson (2004) ratio: 10 [117/12]; key factors  
2                   were: ADL, cognitive impairment [prevalent and incident delirium]  
3                   ○ Inouye (1998) ratio:11 [77/7]; [3 month follow up]; key factors  
4                   were: ADL, cognitive impairment  
5                   ○ Pitkala (2005) ratio: 10 [72/7]; key factors were: ADL, cognitive  
6                   impairment [dementia]  
7  
8                   ● Three studies had 2/3 of the important risk factors taken into account in the  
9                   multivariate analysis but had insufficient ratio of events to variables.  
10                  ○ Inouye (1998) ratio: 9 [60/7]; [at discharge]; key factors were:  
11                  ADL, cognitive impairment  
12                  ○ O’Keeffe (1997) ratio:5 [35/7]; key factors were: ADL, cognitive  
13                  impairment  
14                  ○ Balas (2009) ratio: 3 [35/13] ; key factors were: ADL, dementia  
15  
16                  ● One study had only one of the important risk factors taken into account in  
17                  the multivariate analysis and had an insufficient ratio of events to  
18                  variables.  
19                  ○ Levkoff (1992) ratio: 6 [30/5]; key factor was: cognitive  
20                  impairment

21  
22

#### 4. Falls

23                  The GDG identified age, gender, polypharmacy, and cognitive impairment as  
24                  the key confounding factors. There were no studies identified reporting this  
25                  outcome. Falls are, however, included as part of the hospital acquired  
26                  complications outcome.  
27

28

#### 5. Hospital admission (for those who were in long-term care)

29                  The GDG identified age, gender, cognitive impairment, severity of illness and/or  
30                  comorbidity as the key confounding factors. There were no studies identified  
31                  reporting this outcome.

32  
33

#### 6. Post discharge care

34                  The GDG identified ADL, living alone and cognitive impairment as the key  
35                  confounding factors. There were no studies identified reporting this outcome.  
36

37

#### 7. Post traumatic stress disorder

38                  There were no studies identified reporting this outcome.  
39  
40

1           8. *Pressure Ulcers*

2           The GDG identified age, gender, and immobility as the key confounding factors.  
3           There were no studies identified reporting this outcome. Pressure ulcers are part  
4           of the hospital acquired complications outcome.  
5

6           9. *Mortality*

7           The GDG identified age, cognitive impairment, and severity of illness as the most  
8           important confounding factors.  
9

- 10           • Three studies had all 3 important risk factors taken into account in the  
11            multivariate analysis and had a ratio of events to variables of at least  
12            10
- 13            ○ Inouye (1998): ratio: 14 [98/7] [3 months]; key risk factors were:  
14            age, severity of illness, cognitive impairment [dementia]
  - 15            ○ Levkoff (1992): ratio:12 [59/5]; key factors were: age, cognitive  
16            impairment, and severity of illness
  - 17            ○ Nightingale (2001): ratio: 38 [ 347/10] [2 years]; key risk  
18            factors: age, dementia, physical illness [report of Holmes 2000]
- 19
- 20           • Four studies had 2/3 of the important risk factors taken into account in the  
21            multivariate analysis and had a ratio of events to variables of at least  
22            10
- 23            ○ Dolan (2000): ratio: 62 [369/6]; key factors were: age, cognitive  
24            impairment [cognitive impairment held constant as patients  
25            with cognitive impairment excluded]
  - 26            ○ Drame (2008): ratio: 11 [135/12]; key factors were: age,  
27            cognitive impairment [dementia]
  - 28            ○ Pitkala (2005): ratio: 15 [106/7][ 1 year]; ratio:28 [198/7] [2  
29            years]; key factors were: age, cognitive impairment [dementia]
  - 30            ○ Rockwood (1999): ratio: 11 [101/9]; key factors were: age,  
31            cognitive impairment
- 32
- 33           • Four studies had all of the important risk factors taken into account in the  
34            multivariate analysis but had an insufficient ratio of events to variables.
- 35            ○ Holmes (2000): ratio: 9 [195/ 22] [6 months]; key factors were:  
36            age, dementia, physical illness
  - 37            ○ Ely (2004): ratio:6 [69/12]; key factors were: age, severity of  
38            illness, dementia
  - 39            ○ Inouye (1998): ratio 5 [35/7][discharge]; key risk factors were:  
40            age, severity of illness, cognitive impairment [dementia]
  - 41            ○ O’Keeffe (1997): ratio: 3 [22/7] [in hospital]; 7 [49/7] [for 6  
42            months]; key factors were: age, severity of illness, cognitive  
43            impairment [dementia]

1

2

3

- Three studies had 2/3 of the important risk factors taken into account in the multivariate analysis but had an insufficient ratio of events to variables.

4

5

- Thomason (2005): ratio: 5 [32/7]; key factors were: age, severity of illness

6

7

8

9

- Francis (1990): ratio: 4 [24/6]; key factors were: cognitive impairment, severity of illness [Unclear which factors were adjusted for in the multivariate analysis therefore used the factors reported for length of stay analysis]

10

11

- Marcantonio (2000): ratio:1 [3/5] [1 month]; ratio: 3 [15/5] [6 months]; key factors were: age, cognitive impairment

12

13

14

- Two studies had only one of the important risk factors taken into account in the analysis and had a ratio of events to variables of at least 10

15

16

- Francis (1992): ratio: 14 [55/4]; key factor was: cognitive impairment

17

18

- Leslie (2005): ratio: 35 [208/6]; key factor was: age

19

20

- Two studies had only one of the important risk factors taken into account in the analysis and had an insufficient ratio of events to variables

21

22

23

- Lin (2004): ratio:6 [40/7]; key factor was: severity of illness, although patients with a history of chronic dementia were excluded from the study

24

25

- Lin (2008): ratio: 6 [59/10]; key factor was: age

26

### 10. Impact on carers

27

28

The GDG identified cognitive impairment and disability as the important confounding factors.

29

There were no studies identified reporting this outcome.

30

31

32

### 11. Length of stay

33

34

The GDG identified age, comorbidity and/or severity of illness as the important confounding factors

35

36

- Five studies had all of the important risk factors taken into account in the multivariate analysis and had ratio of at least 10

37

38

- Ely (2004): ratio: 19 [224/12] [length of stay-hospital]; key factors were: age, comorbidity and severity of illness

39

40

- Ely (2004): ratio: 16 [196/12] [Post-ICU stay]; key factors were: age, comorbidity and severity of illness

- 1                   ○ Levkoff (1992): ratio: 42 [211/5] [community]; 23 [114/5]  
2                   [institution]; key factors were: age, severity of illness
- 3                   ○ Holmes (2000): ratio: 33 [731/22] [risk of discharge sooner, i.e.  
4                   decreased risk of remaining in hospital]; key factors were: age,  
5                   physical illness
- 6                   ○ O’Keeffe 1997 ratio: 32 [225/7]; key factors were: age,  
7                   severity of illness, comorbidity
- 8                   ○ Thomason (2005): ratio: 37 [260/7]; [length of stay-hospital and  
9                   length of stay-ICU]; key factors were: age, comorbidity and  
10                  severity of illness

11

- 12                  • One study had one of the important risk factors taken into account in the  
13                  multivariate analysis and had ratio of at least 10

- 14                  ○ Francis (1990): ratio: 38 [229/6]; key factor was: severity of  
15                  illness

16

17

### 12. Quality of life

18                  The GDG identified cognitive impairment and disability as the important  
19                  confounding factors. There were no studies identified reporting this outcome.

20

21

22                  13. Hospital acquired complication [urinary incontinence, falls, pressure sores or  
23                  any other complications]

24                  The GDG identified age, gender, polypharmacy, cognitive impairment [factors  
25                  previously identified for falls] and/or age, gender, immobility [factors previously  
26                  identified for pressure sores] as the important confounding factors

27

- 28                  • One study had 2/5 of the confounding factors taken into account in the  
29                  multivariate analysis but had a ratio of at least 10

- 30                  ○ O’Keeffe (1997): ratio: 32 [225/7]; key factors were: age,  
31                  cognitive impairment

32

### 33                  14. Mortality or new admission to institution

34                  The GDG identified ADL, age, cognitive impairment, comorbidity, severity of  
35                  illness as the important confounding factors

- 36                  • Three studies had all/most (4 or 5) of the important risk factors taken into  
37                  account in the multivariate analysis and had ratio of at least 10

- 38                  ○ Inouye (1998): ratio: 14 [95/7] at discharge; ratio: 24 [165/7]  
39                  at 3 months; key factors were: ADL, age, cognitive impairment  
40                  [dementia], severity of illness

- 41                  ○ McAvay (2006) ratio: 22 [198/9] key factors were: ADL, age,  
42                  comorbidity, dementia, severity of illness

- 1                   ○ Pitkala (2005): ratio: 48 [336/7] ; key factors were: age, ADL,  
2                    dementia, comorbidity [outcome: mortality or *residing in institution*  
3                    *at 2 years*]  
4  
5                   • One study had all/most (4 or 5) of the important risk factors taken into  
6                    account in the multivariate analysis but had insufficient ratio of events to  
7                    variables.  
8                   ○ Marcantonio (2000): ratio: 7 [33/5] [mortality or admission to  
9                    nursing home at 1 month]; ratio:6 [28/5] [mortality or admission  
10                  to nursing home at 6 months]; key factors were: age, cognitive  
11                  impairment, ADL, comorbidity  
12  
13                  • One report of the Marcantonio (2000) study had 3/5 of the important risk  
14                  factors taken into account in the multivariate analysis but had insufficient  
15                  ratio of events to variables.  
16                  ○ Givens (2008): ratio 5 [33/7] [mortality or admission to nursing  
17                  home at 1 month]; key factors were: age, ADL, comorbidity  
18                  ○ Givens (2008): ratio: 4 [ 28/7] [mortality or admission to nursing  
19                  home at 6 months]; key factors were: age, ADL, comorbidity  
20

## 21 **B. Risk Factor: Increased duration of delirium**

22  
23 For this risk factor it was assumed that the other key risk factors for the various  
24 outcomes were the same as for the incidence of delirium

### 25 26 *1. Mortality*

- 27                  • One study had all of the important risk factors taken into account in the  
28                  multivariate analysis but had insufficient ratio of events to variables.  
29                  ○ Ely (2004) ratio:6 [69/12]; key factors were: age, severity of  
30                  illness, dementia

### 31 32 *2. Length of stay*

- 33                  • One study had all of the important risk factors taken into account in the  
34                  multivariate analysis and had ratio of at least 10  
35                  ○ Ely (2004): ratio: 19 [224/12] [Length of stay: hospital]; key  
36                  factors were: age, comorbidity and severity of illness  
37                  ○ Ely (2004): ratio: 16 [196/12] [Length of stay: Post-ICU stay];  
38                  key factors were: age, comorbidity and severity of illness

1

2 *3. Mortality or Functional decline*3 The GDG identified age, cognitive impairment and severity of illness as the key  
4 confounding factors for the composite outcome mortality or functional decline.

- 5 • One study had not enough risk factors (1/3) taken into account in the
- 
- 6 multivariate analysis but the ratio of events to covariate was at least 10
- 
- 7 ○ Andrew (2005): ratio: 12 [48/4] [6 months]; key factor was: age
- 
- 8

- 9 • One study had not enough risk factors (1/3) taken into account in the
- 
- 10 multivariate analysis and the ratio of events to covariate was insufficient
- 
- 11 ○ Andrew (2005): ratio: 8 [32/4] [discharge]; key factor was: age
- 
- 12

13 C. Risk Factor: Severity of delirium

14

15 For this risk factor it was assumed that the same key risk factors applied as for  
16 the incidence of delirium17 *1. Mortality*

- 18 • One study had 1/3 confounding factors for mortality but the ratio of events
- 
- 19 to covariates was at least 10
- 
- 20 ○ Leslie 2005 ratio: 30 [208/7]; key factor was: age
- 
- 21

22 *2. Mortality or new admission to institution (for people who were in hospital)*

- 23 • One report of the Marcantonio (2000) study had 2 of the 5 confounding
- 
- 24 factors for mortality or nursing home placement but had an insufficient
- 
- 25 ratio of events to variables.
- 
- 26 ○ Marcantonio (2002): ratio: 7 [22/3] [1 month]; ratio: 6 [17/3] [6
- 
- 27 months]; key factors were: ADL and cognitive impairment
- 
- 28

29 **8.3.4.5 Overall quality assessment**30 Overall, the risk of bias was considered for each cohort study for each outcome,  
31 and a rating was given of high, moderate, low quality, and biased/confounded.

32

33 Four studies were judged to be biased for the following outcomes and therefore  
34 not considered further:

- 35 • Mortality (Dolan 2000: 2 years; Marcantonio 2000: 1 month)
- 
- 36 • Dementia (Cognitive impairment: Ely 2004 at discharge; Cognitive
- 
- 37 dysfunction: Rudolph 2008)
- 
- 38

1 The Marcantonio (2000) study was considered biased because there were more  
2 variables than events for the mortality outcome (at 1 month); the Dolan (2000)  
3 study was considered biased for the outcome mortality (at 2 years) because the  
4 method of delirium assessment was judged to be inadequate; the Rudolph  
5 (2008) study for the outcome cognitive dysfunction because of partially  
6 inadequate method of assessment of delirium and for the outcome cognitive  
7 dysfunction at 3 months, the study had missing data that was influenced by the  
8 presence of the prognostic factor; the Ely (2004) study had 29% missing data,  
9 which was attributed to an unexpected discharge or an inability to complete  
10 testing; inability to complete testing may have been related to the presence of  
11 delirium.

12  
13 Thirteen reports of ten studies were given a low overall rating for the following  
14 outcomes and were treated with caution:

- 15 • Hospital acquired complications (O’Keeffe 1997)
- 16 • New admission to institution (Balas 2009; Levkoff 1992)
- 17 • Mortality (Francis 1990 - 6 months [Francis 1992- 2 years]; Leslie 2005  
18 [incidence and severity of delirium]; Lin 2004; Lin 2008; Marcantonio  
19 2000: 6 months; O’Keeffe 1997: in hospital; Thomason 2005)
- 20 • Mortality or new admission to institution (Givens 2008: 1 month and 6  
21 months)
- 22 • Mortality or new admission to institution (Marcantonio 2002; severity of  
23 delirium)
- 24 • Mortality or functional decline (Andrew 2005; duration of delirium)
- 25 • Length of stay (Francis 1990)

26  
27 Ten studies were given a moderate rating for the following outcomes:

- 28 • Dementia (Rockwood 1999)
- 29 • New admission to institution (Bourdel-Marchasson 2004; Inouye 1998:  
30 discharge and 3 months; O’Keeffe 1997)
- 31 • Mortality (Drame 2008: 6 week; Ely 2004 [incidence and duration of  
32 delirium]; Holmes 2000 - 6 months; Inouye 1998: discharge; 3 months;  
33 Levkoff 1992; O’Keeffe 1997: 6 months; Pitkala 2005: 1 year and 2  
34 years; Rockwood 1998)
- 35 • Length of stay (Ely 2004:post ICU [incidence and duration of delirium];  
36 Levkoff 1992)
- 37 • Mortality or new admission to institution (Inouye 1998: 3 months;  
38 Marcantonio 2000- 1 month and 6 months; Pitkala 2005- 2 years )

39

1 Eight reports of 7 studies were given a high rating for the following outcomes:

- 2 • New admission to institution (Pitkala 2005)
- 3 • Mortality (Nightingale 2001- 2 years)
- 4 • Length of stay (Ely 2004: hospital [incidence and duration]; Holmes 2000
- 5 [discharged from hospital earlier]; O’Keeffe 1997; Thomason 2005:
- 6 hospital and ICU)
- 7 • Mortality or new admission to institution (Inouye 1998: discharge; McAvay
- 8 2006 - 1 year; Pitkala 2005: mortality or residing in long-term care at 2
- 9 years)

## 12 8.4 RESULTS

14 Two studies (Andrew 2005; Ely 2004) reported the dependence of adverse  
 15 consequences on the duration of delirium; two studies (Leslie 2005; Marcantonio  
 16 2002) reported the effects of increased severity of delirium and the remaining  
 17 studies examined incidence of delirium as a prognostic factor.

18 Factors included in the multivariate analyses are given in Appendix F.

19 The following outcomes have been investigated:

- 20 • Risk Factor: Presence of prevalent and incident delirium
  - 21 ○ Dementia (1 study)
  - 22 ○ Progression of dementia (no studies)
  - 23 ○ New admission to an institution (6 studies)
  - 24 ○ Hospital admission (for those who were in long-term care) (no
  - 25 studies)
  - 26 ○ Post discharge care (no studies)
  - 27 ○ Pressure Ulcers (no studies) but see hospital acquired
  - 28 complications
  - 29 ○ Falls ( no studies) but see hospital acquired complications
  - 30 ○ Mortality (16 reports of 14 studies)
  - 31 ○ Impact on carers (no studies)
  - 32 ○ Length of stay (6 studies)
  - 33 ○ Quality of life (no studies)
  - 34 ○ Hospital acquired complications (1 study)
  - 35 ○ Mortality or new admission to an institution (5 reports of 4 studies)
- 36
- 37 • Risk factor: Increased duration of delirium
  - 38 ○ Mortality (1 study)

- 1                   ○ Length of stay (1 study)
- 2                   ○ Mortality or functional decline (1 study)
- 3
- 4                   ● Risk factor: Severity of delirium
  - 5                   ○ Mortality (1 study)
  - 6                   ○ Mortality or new admission to an institution (1 study)
- 7

#### 8 8.4.1 Risk factor: presence of prevalent or incident of delirium

9

##### 10 8.4.1.1 Dementia

11

12 One moderate quality study (Rockwood 1999) reported dementia as a  
13 consequence of delirium at 3 year follow-up.

14 The Rockwood (1999) study reported 21% (32/154) of the patients developed  
15 dementia; the median follow-up period in the Rockwood (1999) study was 32.5  
16 months.

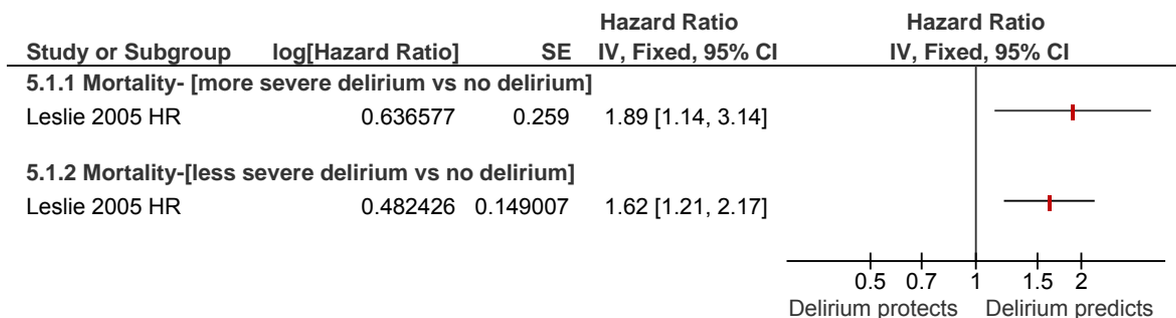
17 Cognitive impairment was evaluated with MMSE (range 0 to 30), the Blessed  
18 dementia rating scale (range 0 to 17; higher score indicative of greater degree  
19 of dementia) and dementia was determined by a geriatrician. Information on  
20 patients who had died by follow-up was obtained through the IQCODE  
21 interviews from proxy informants. The study did not clarify who the proxies were.

22 This study in 203 patients showed that dementia was a significant consequence  
23 of delirium at 3 years follow up [OR 5.97 (95% CI 1.83 to 19.54)]; the  
24 confidence interval is wide (figure 8.1)

25

26 Figure 8.1: dementia as a consequence of delirium

27



28

29

NB: Scale 0.05 to 20

30

1 **8.4.1.2 New admission to institution**

2  
3 Six studies (Balas 2009; Bourdel-Marchasson 2004; Inouye 1998; Levkoff 1992;  
4 O’Keeffe 1997[incident delirium only]; Pitkala 2005) reported new admissions  
5 to an institution following discharge. Two studies (Balas 2009; Levkoff 1992)  
6 were low quality, three were moderate quality (Bourdel-Marchasson 2004;  
7 Inouye 1998; O’Keeffe 1997 [incident and prevalent delirium]) and one study  
8 was high quality (Pitkala 2005).

9 The studies reported new admission to an institution following discharge from  
10 hospital (Inouye 1998; Levkoff 1992), at 3 months (Inouye 1998), 6 months  
11 (O’Keeffe 1997) and during 2 years (Pitkala 2005).

12 The number of patients (with delirium) admitted to an institution ranged from 3%  
13 (20/692) at discharge (Inouye 1998) to 36% (Pitkala 2005: 72/200) at 2  
14 years.

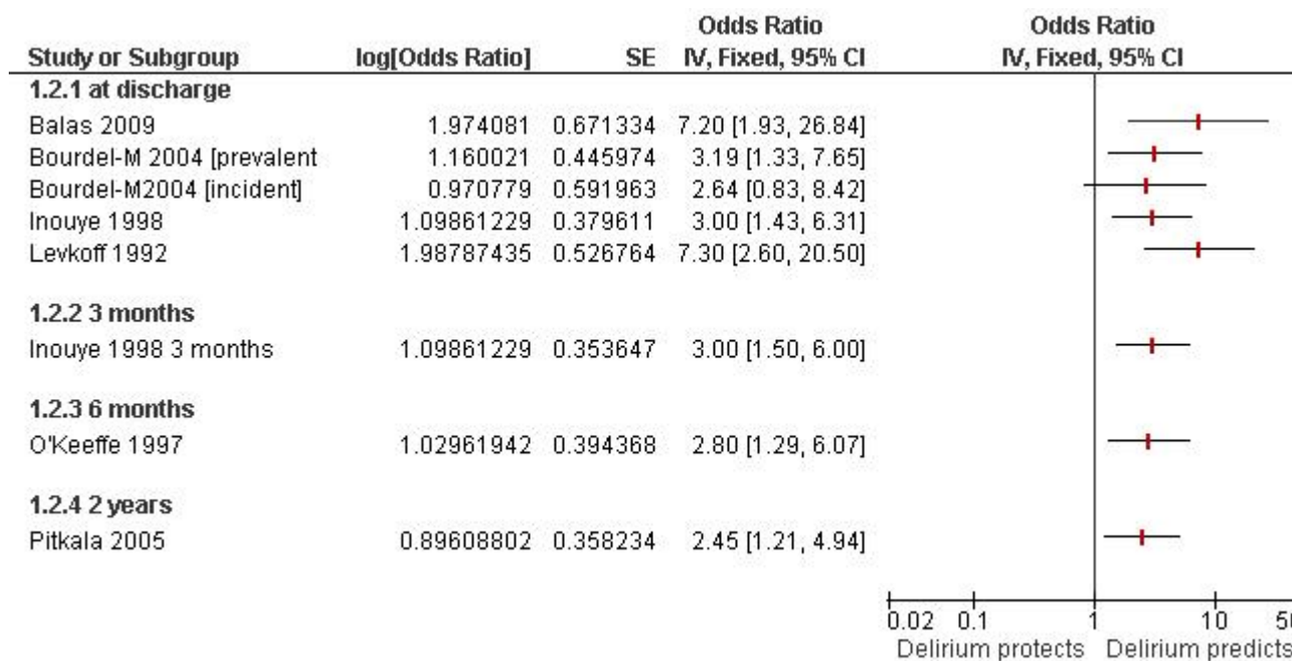
15 The studies varied in their consideration of the key risk factors (ADL, cognitive  
16 impairment). Further information on these factors is reported in Appendix F.  
17 None of the studies reported including depression as a factor in the multivariate  
18 analysis.

19 Two studies (Inouye 19998; O’Keeffe 1997) reported excluding deaths for this  
20 outcome; one study (Balas 2009) reported patients who died within 24 hours of  
21 SICU admission were not considered for enrollment and one study (Bourdel-  
22 Marchasson 2004) reported the number of patients discharged either back to  
23 community or institution taking into account the number of deaths.

24 The odds ratio was generally around 2.8 and appeared to be fairly  
25 independent of when this was measured. The results suggest that new admission  
26 to an institution is a significant consequence of delirium (figure 8.2a).

27  
28

1 Figure 8.2a: new admission to institution as a consequence of delirium

2  
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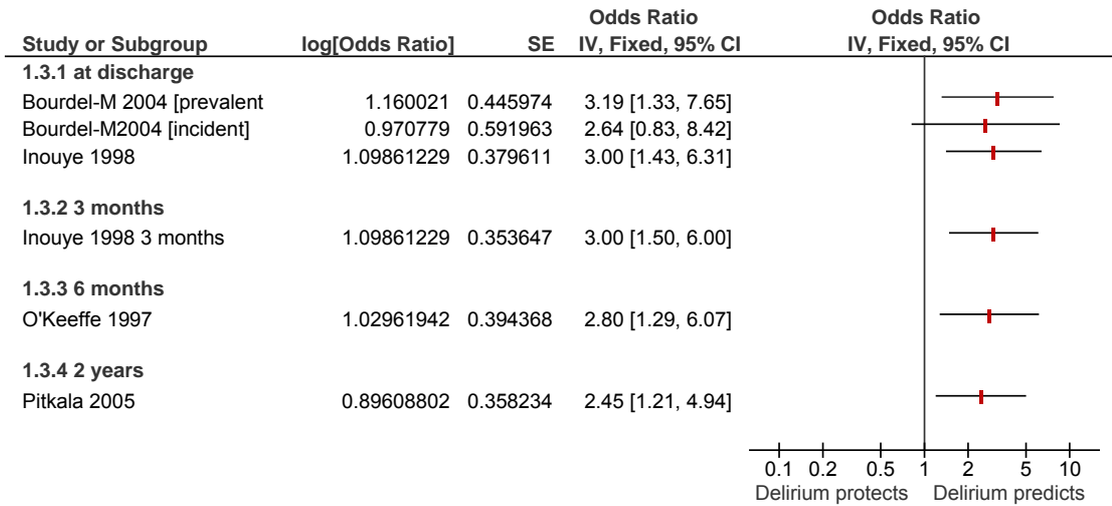
NB: Scale 0.05 to 20

4

5 A sensitivity analysis was undertaken (figure 8.2b) excluding the low quality  
6 studies. Three moderate quality study studies (Bourdel-Marchasson 2004  
7 (n=427); Inouye 1998 (n=727); O'Keeffe 1997 (n=225)) and one high quality  
8 study (Pitkala 2005 (n=425)) were included. At discharge, the odds ratio  
9 ranged from 2.64 (95% 0.83 to 8.45) (Bourdel-Marchasson 2004: incident  
10 delirium) to 3.19 (95% CI 1.33 to 7.64) (Bourdel-Marchasson 2004: prevalent  
11 delirium). One study (Pitkala 2005) showed a significant effect of delirium on  
12 new institutionalisation at 2 years following discharge [adjusted OR 2.45 (95%  
13 CI 1.2 to 4.9)].

1  
2

Figure 8.2b: new admission to institution [moderate quality studies]



3  
4

NB: Scale 0.1 to 10

5

6 **8.4.1.3 Mortality**

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Sixteen reports of 14 studies (Drame 2008; Ely 2004; Francis 1990 [Francis 1992:2 years]; Holmes 2000 [Nightingale 2001: 2 years]; Inouye 1998; Leslie 2005; Levkoff 1992; Lin 2004; Lin 2008; Marcantonio 2000; O'Keeffe 1997; Pitkala 2005; Rockwood 1999; Thomason 2005) reported mortality following delirium. Most studies did not state the cause of death, with the exception of two studies (Lin 2004; Drame 2008) which reported death from all causes.

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Eight reports of seven studies were of low quality (Francis 1990: 6 months [Francis 1992: 2 years]; Leslie 2005; Lin 2004; Lin 2008; Marcantonio 2000: 6 months; O'Keeffe 1997: in hospital; Thomason 2005) and treated with caution; there were 8 studies of moderate quality (Drame 2008; Ely 2004; Holmes 2000: 6 months; Inouye 1998: hospital and 3 months; Levkoff 1992; O'Keeffe 1997: 6 months; Pitkala 2005; Rockwood 1998) and one report of the Holmes (2000) study was rated as high quality (Nightingale 2001: 2 years).

21

22  
23

Information on the key factors (age, cognitive impairment, severity of illness) adjusted for in the multivariate analysis are presented in Appendix F.

24  
25  
26  
27  
28

Three studies reported death in hospital (O'Keeffe 1997; Inouye 1998; Thomason 2005). Of these, only the results from the O'Keeffe (1997) study will be considered as the GDG stated that only UK results are applicable for this outcome at discharge, however, the other studies are also shown on the forest plot for information.

29  
30  
31

Of the studies reporting mortality following discharge from hospital or ICU, eight reports of seven studies included hospital deaths (Drame 2008; Ely 2001; Francis 1990; Inouye 1998; Marcantonio 2000; Holmes 2000; Nightingale

1 2001; O’Keeffe 1997), three studies excluded death in hospital (Francis 1992  
2 2.6% [6/229]; Leslie 2005: 1.5% [14/919]; Rockwood 1999 12.9% [32/247  
3 enrolled]) and was unclear in two studies (Levkoff 1992; Pitkala 2005)

4 The number of patients who were in long-term care when they died was  
5 considered for the following time points:

6 • 6 weeks

- 7 ○ In one study (Drame 2008), 17% of the patients [218/1306]  
8 were admitted from long-term care. It is unclear how many  
9 patients were discharged back into long-term care or if there  
10 were any new admissions and how many people died in long-  
11 term care.

12 • 3 months

- 13 ○ In one study (Inouye 1998), of the 4% [29/77] patients admitted  
14 from long-term care it was unclear how many patients were  
15 discharged back into long-term care. Of those newly admitted to  
16 long-term care at discharge 8.7% [60/692], it is unclear how  
17 many people died there in the follow up period of 3 months. At 3  
18 month follow-up, all deaths in hospital and at 3 months were  
19 excluded for the outcome new admission to long-term care.

20 • 6 months

- 21 ○ In one study (Ely 2004) it was unclear if any patients were  
22 admitted to long-term care following discharge from ICU.
- 23 ○ One study (Francis 1990) reported 7% (16/226: 16% vs 3.4%  
24 for the delirious and non delirious groups, respectively) of the  
25 patients were discharged to nursing homes, personal-care homes  
26 and rehabilitation facilities. The study also reported the  
27 percentages at 6 month follow-up [12% and 5% for the delirious  
28 and non delirious groups, respectively]. It is unclear how many  
29 patients in long-term care died.
- 30 ○ In one study (Holmes 2000), of the patients who were diagnosed  
31 with delirium and living in non-residential setting at admission  
32 [76%: 82/108], 23% [19/63] were discharged to a residential  
33 or nursing home. It is unclear how many of these patients in long-  
34 term care died during the 6 month follow up period.
- 35 ○ The Levkoff (1992) study reported 15% [30/203] of the  
36 community-dwelling patients with incident delirium were  
37 discharged to institution. It is unclear how many patients died in  
38 long-term care.
- 39 ○ The Marcantonio (2000) study reported the composite outcome  
40 mortality or new nursing home placement. The proportion of  
41 patients who either died or were placed into nursing home [new  
42 admissions] was 22% [28/126] at 6 months. The proportion of  
43 patients who died was 12% [15/126] at 6 months.

- 1           • 1 year
- 2           ○ In the Leslie (2005) study, of the 222 patients who died during
- 3           the study period, 9.5% (21/222) were nursing home residents at
- 4           admission. It is unclear whether all patients were discharged back
- 5           into long-term care and subsequently how many died there.
- 6           ○ In the Pitkala (2005) study, of the 53% [224/425] patients
- 7           assessed in long-term care, it is unclear how many of these
- 8           patients died in the first year during the course of the study.
- 9           • 2 years
- 10          ○ In Francis (1992) it is unclear how many of the patients
- 11          discharged to long-term care (as reported in Francis 1990) were
- 12          followed up or how many died in the long-term care setting.
- 13          ○ Pitkala 2005- Of the 53% [224/425] patients assessed in long-
- 14          term care or the 36% of the patients [72/200] newly admitted to
- 15          long-term care during the course of the 2 years, it is unclear how
- 16          many of these patients died in long-term care. The study
- 17          reported that 79% of the patients [336/425] were residing in
- 18          institutional care or died during 2 years.
- 19          • 3 years
- 20          ○ One study (Rockwood 1999) reported that, of the patients
- 21          [101/203] who died during the 3 year follow-up, 79% (30/38)
- 22          had delirium. Of the patients with delirium who died, the study
- 23          reported 70% of the patients (21/30) were in institutional care.

24

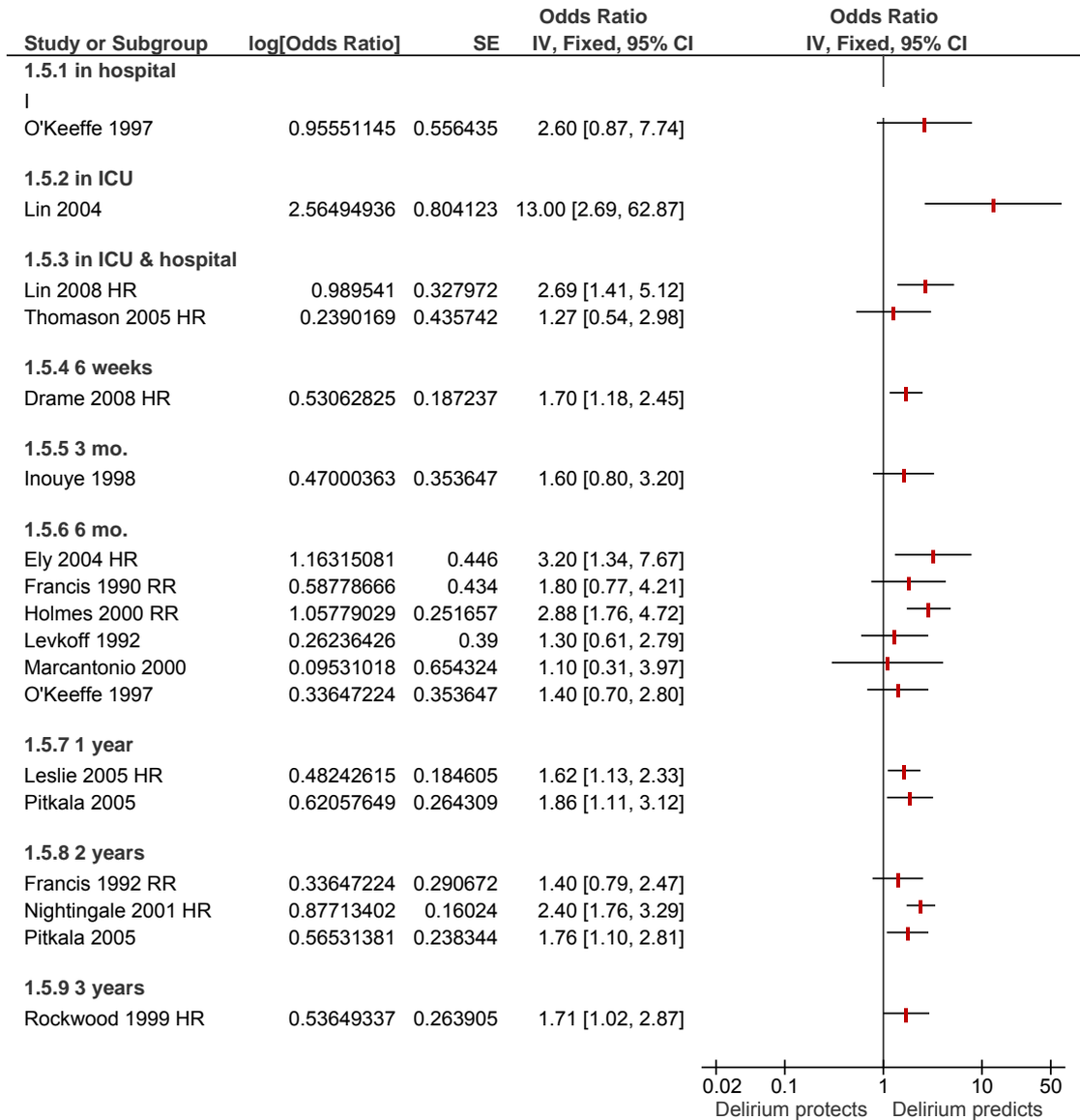
25          The risk of mortality as a consequence of delirium varied with time as shown in

26          the forest plot (figure 8.3a).

27

28

1 Figure 8.3a: mortality as a consequence of delirium

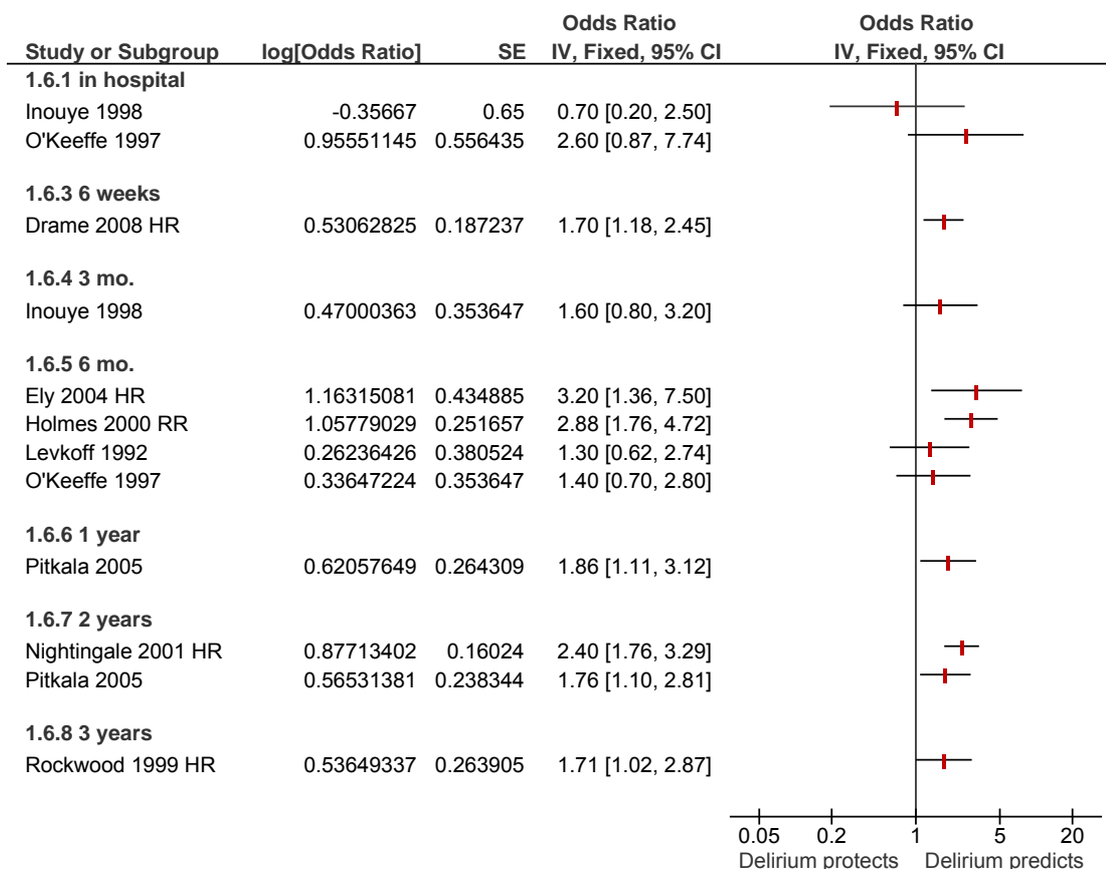


2 NB: Scale 0.02 to 50  
3

4  
5 Excluding the low quality studies, the following results were found (figure 8.3b).

6  
7

1 Figure 8.3b: mortality as a consequence of delirium; high and moderate quality  
 2 studies and restricting to the UK hospital study



3  
 4 NB: Scale 0.05 to 20

5  
 6 There is a significant effect of delirium incidence on mortality, which appears to  
 7 be independent of time.

8  
 9

10 **8.4.1.4 Length of stay**

11  
 12 Two high quality studies (Holmes 2000; O'Keefe 1997), one moderate quality  
 13 study (Levkoff 1992) and one low quality study (Francis 1990) reported length  
 14 of stay in hospital. Two high quality studies (Ely 2004; Thomason 2005) reported  
 15 length of stay in hospital (including the period in ICU), one high quality study  
 16 (Thomason 2005) reported length of stay in the ICU and one (Ely 2004)  
 17 reported length of stay post ICU (moderate quality for this outcome). The Ely  
 18 (2004) study defined post ICU length of stay as the time after first ICU  
 19 discharge.

20 The Holmes (2000) study, reported the relative risk of being discharged earlier,  
 21 which corresponds to a decreased length of stay.

1  
2 Three studies (Francis 1990; Levkoff 1992; O’Keeffe 1997) reported length of  
3 stay, adjusted for confounding factors in a multivariate analysis and gave p-  
4 values. The Levkoff (1992) study reported that delirium contributed to a longer  
5 length of stay both for patients admitted from the community ( $t=4.03$ ;  
6  $p=0.0001$ ; 30.9 days and 7.4 days for the delirious and non delirious groups,  
7 respectively) and from long-term care ( $t=4.48$ ;  $p=0.0001$ ; 10.6 days and 6.9  
8 days for the delirious and non delirious groups, respectively). The Francis (1990)  
9 study reported that delirious patients stayed in the hospital longer than the non  
10 delirious group (12.1 days versus 7.2 days, for the delirious and non delirious  
11 groups, respectively;  $p<.001$ ). The O’Keeffe (1997) study reported that  
12 delirium was the only significant predictor of duration of hospital stay in a  
13 multivariate analysis (accounting for 6.7% of the variance; adjusted  $t=3.8$ ,  
14  $p<.001$ ). The mean length of stay was 21 days and 11 days, for the delirious  
15 and non delirious groups, respectively ( $p<.001$ ).

16 The median length of stay in hospital and interquartile range (IQR) were  
17 reported in the Ely (2004) study [21 days (IQR 19 to 25): 11 days (IQR 7 to 14)  
18 for the delirious and non delirious groups, respectively] and the Thomason (2005)  
19 study [median 5 days (IQR 2 to 8) and 3 days (IQR 2 to 6) for the delirious and  
20 non delirious groups, respectively]. In the Ely (2004) study, length of stay was  
21 measured from admission for prevalent delirium patients and from time of  
22 diagnosis for incident delirium patients.

23 The median length of stay in ICU and interquartile range (IQR) was reported in  
24 the Thomason (2005) study [median 4 days (IQR 3 to 5) and 3 days (IQR 2 to 4)  
25 for the delirious and non delirious groups, respectively].

26 The median length of post ICU stay and interquartile range (IQR) was reported  
27 in the Ely (2004) study [median 7 days (IQR 4 to 15.5) and 5 days (IQR 2 to 7)  
28 for the delirious and non delirious groups, respectively].

29 One study (Holmes 2000) reporting discharge from hospital, showed the  
30 likelihood of discharge was decreased in the presence of delirium, leading to an  
31 increased length of stay [RR 0.53 (95% CI 0.41 to 0.68); figure 8.4a].

32 The adjusted hazard ratio ranged from 1.41 (95% CI 1.05 to 1.89) to 2.0 (95%  
33 CI 1.4 to 3.0) showing increased length of stay in hospital to be a significant  
34 consequence of delirium for patients who had been in ICU (figure 8.4b).

35 There was no significant effect on length of stay in ICU [HR 1.29 (95% CI 0.98 to  
36 1.69)] but there was an effect of delirium on post-ICU stay [HR 1.6 (95% CI 1.1  
37 to 2.3); figure 8.4b].

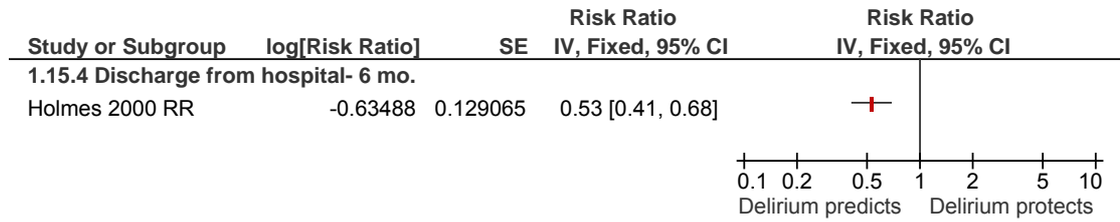
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1 Figure 8.4a: length of stay (discharge from hospital) as a consequence of  
2 delirium

3

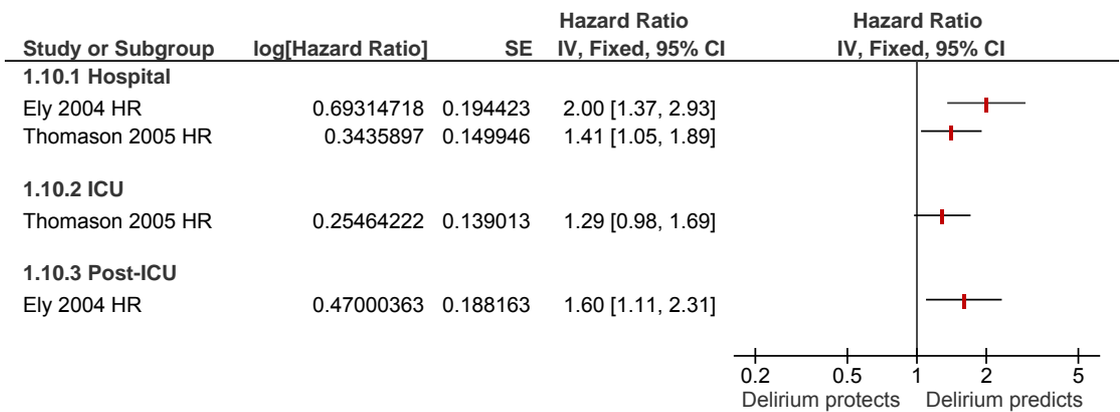


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5 NB: Scale 0.1 to 10

6

7 Figure 8.4b: length of stay as a consequence of delirium



8

9 NB: Scale 0.2 to 5

10

11 **8.4.1.5 Hospital acquired complication [urinary incontinence, falls, pressure sores or any**  
12 **other complication]**

13

14 One low quality study (O’Keeffe 1997) reported results for hospital acquired  
15 complications. The percentages of patients with complications were as follows:  
16 urinary incontinence: 46% (86/206); falls: 12.4% (28/225); pressure sores: 4%  
17 (8/202) or any other complications: 44% (100/225). The multivariate analysis  
18 adjusted for age, chronic cognitive impairment, severity of illness, comorbidity,  
19 disability score and length of stay.

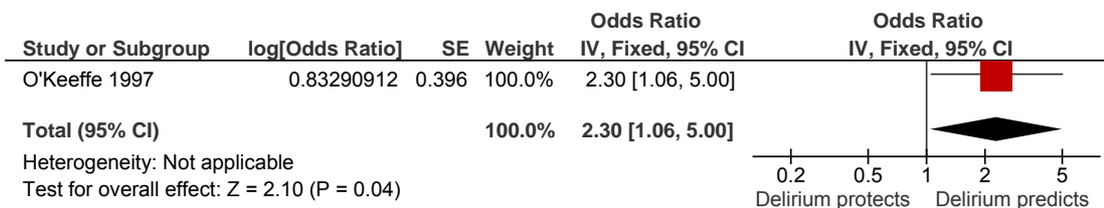
20

21 The study reported that falls, pressure sores (corresponding to grade 2 Shea  
22 classification) and urinary incontinence (new onset or worsening after admission  
23 to hospital) were identified based on interviews with nursing staff. The authors  
24 defined a fall as ‘unintentionally coming to rest on ground ... not as a result of  
25 an obvious major intrinsic event (such as stroke or syncope) or overwhelming  
hazard.’

26

The result showed that hospital acquired complications is a significant consequence of delirium [OR 2.3 (95% CI 1.7 to 5.0); figure 8.5].

Figure 8.5: hospital acquired complications as a consequence of delirium



NB: Scale 0.2 to 5

#### 8.4.1.6 Mortality or new admission to institutions

Five reports of four studies (Inouye 1998; McAvay 2006; Marcantonio 2000 [Givens 2008]; Pitkala 2005) reported a composite outcome of mortality or new admission to institution. The Givens (2008) report of the Marcantonio (2000) study and the Marcantonio (2000) study reported results for the same cohort but the multivariate analyses were adjusted for different factors. The Givens (2008) report only gave the adjusted odds ratio and p values. The standard error was calculated, on a trial and error basis, based on the reported p values.

Three studies were high quality (Inouye 1998 at hospital discharge; McAvay 2006; Pitkala 2005), two were of moderate quality (Inouye 1998 at 3 months; Marcantonio 2000), and the Givens (2008) report of the Marcantonio (2000) study was low quality. The Pitkala (2005) study reported mortality or residing in institution at 2 years.

Rates of the composite outcomes (mortality and new admission to institution) and the rates for each outcome, where reported, were as follows:

In hospital: 13% (Inouye 1998:95/727; mortality: 5% [35/727]; new admission: 9% [60/692])

- 1 month: 26% (Marcantonio 2000: 33/126; mortality: 2% [ 3/126] )
- 3 months: 25% (Inouye 1998: 165/663; mortality: 14% [98/680]; new admission: 13% [77/600] )
- 6 months: 23% (Marcantonio 2000: 28/123; mortality: 12% [15/123]);
- 1 year: (McAvay 2006)
  - delirium at discharge: 83% [ 20/24]; ( mortality: 38% [9/24]; new admission: 79% [19/24]);
  - delirium resolved: 68% [21/31]; (mortality: 26% [8/31]; new admission: 45% [14/31]);

- 1                                   ○ never delirious: 42% [157/378]; (mortality: 20% [75/378]; new  
2                                   admission: 29% [111/378]).

3

4                   At discharge from hospital, one multicentre study set in the US (Inouye 1998 -  
5                   high quality) showed there was a significant effect of delirium on the composite  
6                   outcome, mortality or new admission to institution [OR 2.1 (95% CI 1.1 to 4.0)]  
7                   however, the confidence interval is fairly wide.

8                   At three months, one moderate quality study (Inouye 1998) showed a significant  
9                   effect of delirium [OR 2.6 (95% CI 1.4 to 4.5)]; however, the confidence  
10                   interval is fairly wide.

11                   One moderate quality study (Marcantonio 2000) and one low quality study  
12                   (Givens 2008 showed a significant effect at one month with adjusted odds ratio  
13                   ranging from 3.0 (95% CI 1.1 to 8.4)] to 4.26 (95% CI 1.49 to 12.16), however,  
14                   the confidence interval was wide.

15                   There was no significant effect shown at 6 months.

16                   The McAvay (2006) study reported the results at 1 year for those with delirium  
17                   at discharge, resolved delirium and never delirious. There was a significant  
18                   effect at 1 year [patients with delirium at discharge compared with those never  
19                   delirious] [HR 2.64 (95% CI 1.60 to 4.35)] but the confidence interval is wide. In  
20                   patients with delirium resolved compared with those never delirious and in  
21                   patients with delirium at discharge compared with delirium resolved there was  
22                   no significant effect at 1 year (figure 8.6).

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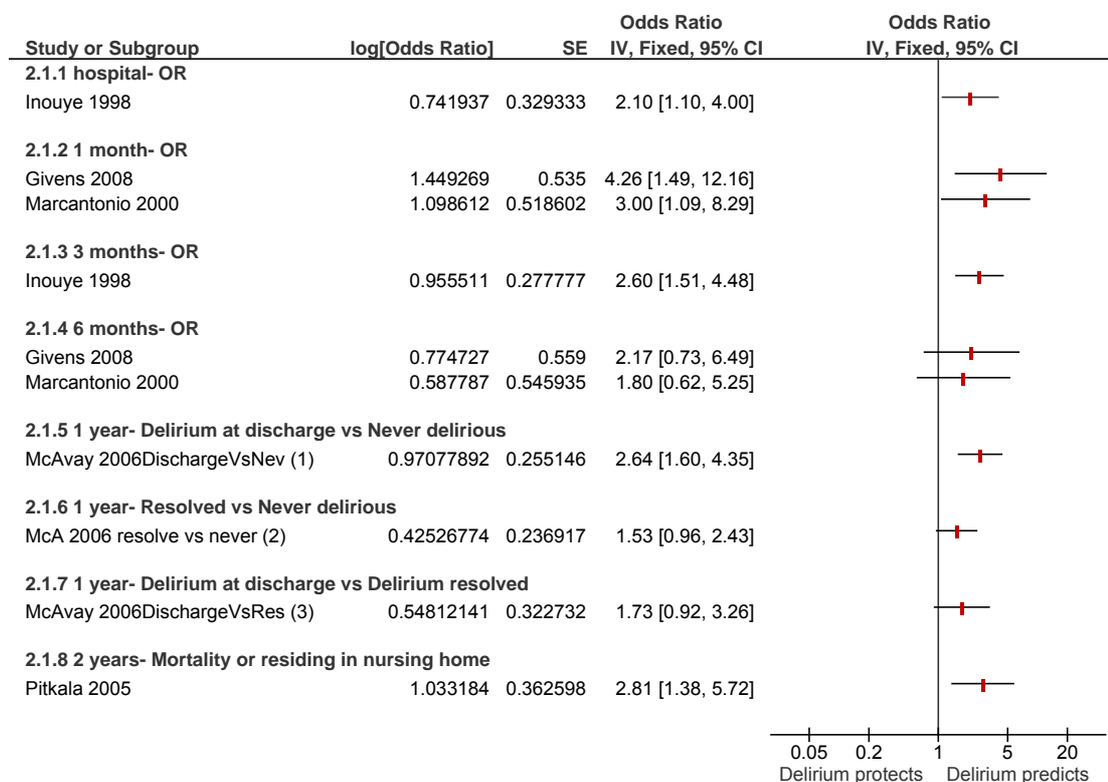
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Figure 8.6: mortality or new admission to institution as a consequence of delirium

(1) HR  
(2) HR  
(3) HR5  
6  
7

NB: Scale 0.05 to 20

8

## 8.4.2 Risk Factor: Increased duration of delirium as a continuous variable

10

### 8.4.2.1 Mortality

12

One moderate quality study (Ely 2004) reported mortality at 6 months as a consequence of duration of delirium. The study used duration of delirium as a continuous risk factor in the multivariate analysis. The results relate to each additional day of delirium for ICU patients.

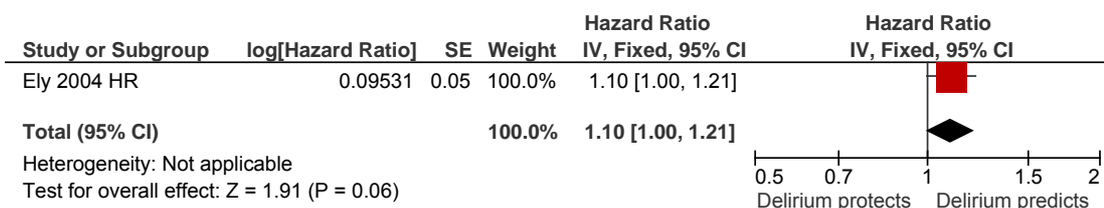
17

There was a borderline significant effect of duration of delirium on mortality [HR 1.1 (95% CI 1.0 to 1.3); figure 8.7]. For each extra day with delirium, the hazard ratio increases by 1.10, so that if there were 3 extra days it would become  $(1.10)^3$  (i.e. 1.33).

20

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3

Figure 8.7: mortality as a consequence of increased duration of delirium



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5  
6

NB: Scale 0.5 to 2

7

8

9 **8.4.2.2 Length of stay**

10

11 One study (Ely 2004) reported length of stay (hospital [high quality] and post-  
12 ICU stay[moderate quality]) as a consequence of increased duration of delirium.

13

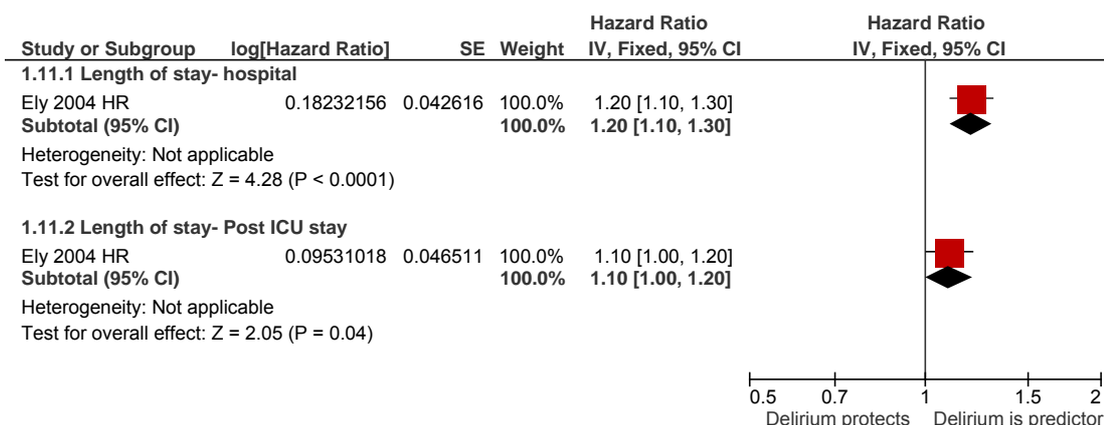
14 The study used duration of delirium as a continuous risk factor in the multivariate  
analysis. The results relate to each additional day of delirium for ICU patients.

15

16 The length of ICU plus hospital stay was significantly greater for patients who  
17 had longer periods of delirium [HR 1.20 (95% CI 1.1 to 1.3)] and the post-ICU  
stay was of borderline significance [HR 1.10 (95% CI 1.0 to 1.2); figure 8.8].

18

19 Figure 8.8: length of stay as a consequence of increased duration of delirium



20  
21

NB: Scale 0.5 to 2

22

23

### 8.4.2.3 Mortality or functional decline

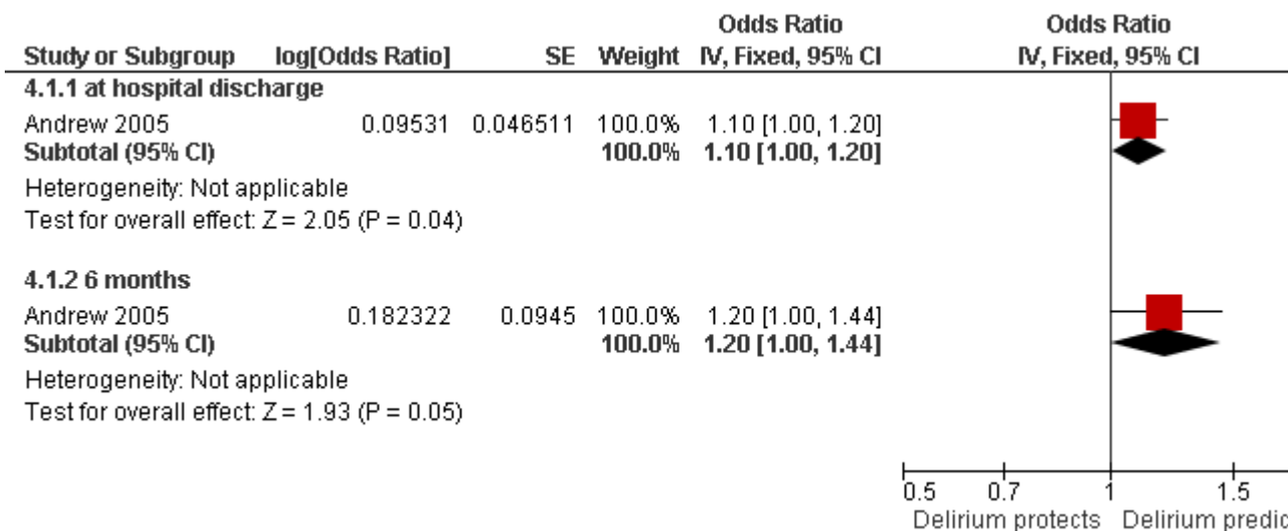
One low quality study (Andrew 2005) reported a composite outcome of incomplete functional recovery or death following an episode of delirium. Functional decline was defined as a decrease by at least 10 points on the Barthel Index (BI) compared with the baseline BI score.

The results were presented for duration of delirium, adjusted for age, gender, and frailty. Frailty was assessed on the geriatric severity score (ranging from healthy and independent to terminally ill). Further information on these factors are presented in Appendix F. Mean duration of delirium was 6.3 days (range 1 to 35). The mean pre morbid (baseline) Barthel Index score was 86.6 (range 42 to 100), with an 8.9 point decrease at discharge and 12.7 decline in score at 6 months.

The study reported that at discharge the mortality rate was 8% (6/77) and functional decline was reported in 37% (26/71) of the patients. At 6 months, 68% of the patients (48/71) had an outcome of death or functional decline.

Mortality or functional decline was a borderline significant consequence of increased duration of delirium at hospital discharge [OR 1.1 (95% CI 1.0 to 1.2)] and at 6 months [OR 1.2 (95% CI 1.0 to 1.4); figure 8.9].

Figure 8.9: mortality or functional decline as a consequence of increased duration of delirium



NB: Scale 0.5 to 2

1 **8.4.3 Risk factor: severity of delirium as a categorical outcome**

2 **8.4.3.1 Mortality**

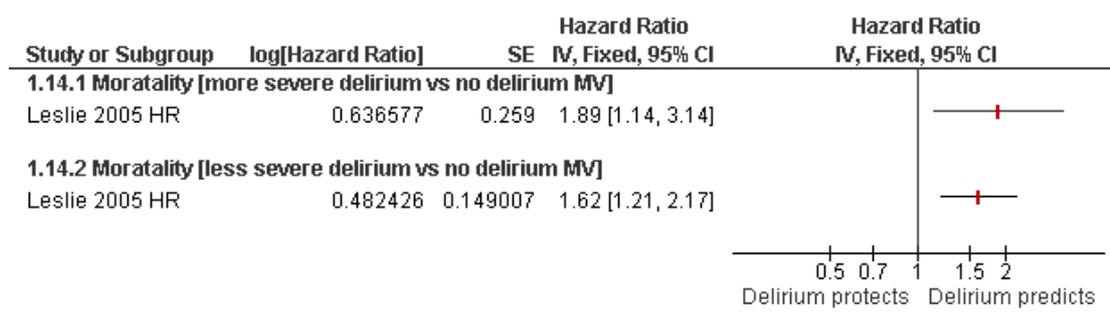
3  
4 One low quality study (Leslie 2005) reported the effect of severity of delirium,  
5 assessed during hospitalisation, on mortality at 1 year.

6  
7 The mortality rate of patients with more severe delirium was 40% (16/40),  
8 30.3% (80/264) for those with less severe delirium and 18.5% (110/596) for  
9 those who were never delirious.

10  
11 At 1 year, increased severity (assessed during hospitalisation) had a significant  
12 effect on mortality compared with no delirium [HR 1.89 (95% CI 1.13 to 3.14)].  
13 Less severe delirium (assessed during hospitalisation) also had a significant effect  
14 [HR 1.62 (95% CI 1.21 to 2.17; figure 8.10).  
15

16

17 Figure 8.10: mortality (at 1 year) as a consequence of delirium (severity)



18

19 NB: Scale 0.5 to 2

20

21 2. Mortality or New admission to institution

22

23 One low quality study (Marcantonio 2002) reported mortality or discharge to a  
24 care home at 1 month and 6 months. The study examined the effect of severity  
25 of delirium in patients with CAM defined delirium and those with non-delirious  
26 symptoms (some had subsyndromal delirium). The results for the former group  
27 (n= 49) are reported here.

28

29 Mortality or new admission to institution at 1 month was 33% (8/24) and 56%  
30 (14/25) for the mild and severe delirium groups, respectively. At 6 months  
31 mortality or new admission to institution was 17% (4/24) and 52% (13/25) for  
32 the mild and severe delirium groups, respectively

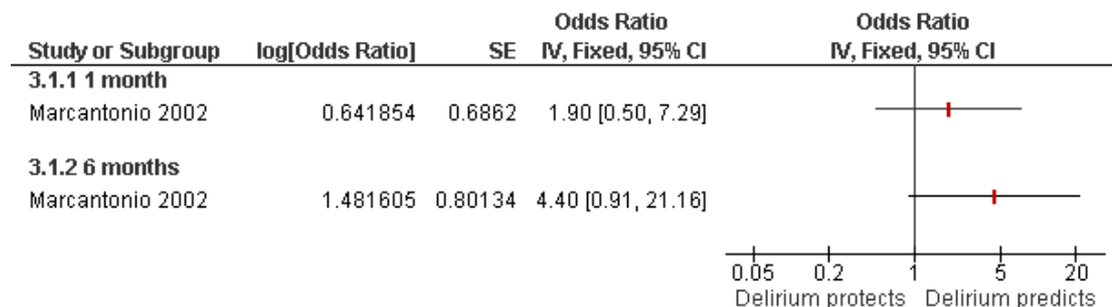
33

34 At 1 month, severe delirium compared with delirium had no significant effect on  
35 mortality or nursing home placement [OR 1.90 (95% CI 0.50 to 8.0)]. At 6  
36 months, the confidence interval is very wide [OR 4.4 (95%CI 0.9 to 21.1; figure  
37 8.11], and there is too much uncertainty to draw conclusions.  
38

38

39

Figure 8.11: mortality or new admission to institution (at 1 month and 6 months) as a consequence of delirium severity



NB: Scale 0.05 to 20

## 8.5 Clinical evidence statements

- There is high quality evidence to show that:
  - the likelihood of discharge was decreased in the presence of delirium, leading to an increased length of stay in hospital.
  - an increased length of stay in hospital is a significant consequence of delirium for patients who had been in ICU.
  - there is no significant effect of delirium on the length of stay in ICU.
  - post-ICU stay is a significant consequence of delirium.
  - there is a significant effect of delirium on the composite outcome, mortality or new admission to institution at discharge from hospital; there is some uncertainty around this result.
  - there is a significant effect of persistent delirium on the composite outcome, mortality or new admission to institution, at 1 year; there is some uncertainty around this result.
  - there is significant effect of delirium on the composite outcome, mortality or new admission to institution, at 2 years; there is some uncertainty around this result.
- There is moderate quality evidence to show that:
  - dementia is a significant consequence of delirium at 3 year follow-up.
  - new admission to institution is a significant consequence of delirium, which appears to be independent of time.

- 1                   ○ mortality is a significant consequence of delirium, which appears  
2                   to be independent of time.
- 3                   ○ there is a significant effect of delirium on the composite outcome,  
4                   mortality or new admission to institution at 3 months following  
5                   discharge from hospital; there is some uncertainty around this  
6                   result.
- 7                   ○ show there is a borderline significant effect of duration of  
8                   delirium on mortality.
- 9                   ○ there is a significantly increased length of ICU plus hospital stay  
10                  for patients who had longer periods of delirium.
- 11                  ○ there is a borderline significant effect on increased length of  
12                  post-ICU for patients who had longer periods of delirium.
- 13
- 14                  • There is low to moderate quality evidence to show that:
- 15                  ○ there is a significant effect of delirium on the composite outcome,  
16                  mortality or new admission to institution, at one month following  
17                  discharge from hospital; there is some uncertainty around this  
18                  result.
- 19                  ○ there is no significant effect of delirium on the composite outcome,  
20                  mortality or new admission to institution, at 6 months following  
21                  discharge from hospital; there is some uncertainty around this  
22                  result.
- 23
- 24                  • There is low quality evidence to show that:
- 25                  ○ hospital acquired complications [pressure sores, falls, urinary  
26                  incontinence or any other complication] are a significant  
27                  consequence of delirium.
- 28                  ○ mortality or functional decline was a borderline significant  
29                  consequence of increased duration of delirium at discharge from  
30                  hospital and at 6 months following discharge.
- 31                  ○ mortality was a significant consequence of increased severity of  
32                  delirium (assessed during hospitalisation).
- 33                  ○ an increased severity of delirium had no significant effect on the  
34                  composite outcome, mortality or new admission to institution, at 1  
35                  month following discharge from hospital.
- 36                  ○ an increased severity of delirium had no significant effect on the  
37                  composite outcome, mortality or new admission to institution, at 6  
38                  months following discharge from hospital; there is too much  
39                  uncertainty around this result.

## 9 Non-pharmacological prevention

### Clinical introduction

Prevention of any harmful condition is clearly desirable, and delirium is no exception. Unfortunately, the introduction of delirium prevention protocols into routine care has been slow, partly because the existing research evidence base is fragmented and not well known to clinicians. Delirium prevention is similar in many respects to the issue of pressure sore prevention in the 1980s when the NHS was content to spend considerable amounts on the treatment of pressure sores and largely ignore prevention strategies. The prevention of pressure sores required specific and well-supported clinical policies to foster a new culture of prevention with the adoption of new procedures and skills in routine care.

A useful practical approach to the understanding of delirium has been to consider patient vulnerability (risk factors) in relation to stressor events (delirium precipitants). Thus, the precipitants do not alone cause an episode of delirium; they interact with the underlying risk factors. This clinical model suggests that interventions designed to reduce the impact of selected delirium risk factors might be associated with a reduction in delirium incidence. This section reviews the evidence for this approach – for single risk factors (single component interventions), and for multiple risk factors (multi-component interventions).

## 9A) Single component prevention: hydration and music

### 9A. 1 HYDRATION FOR THE PREVENTION OF DELIRIUM (LONG-TERM CARE SETTING)

#### 9.1 Description of studies

##### 9.1.1 Study Design

Two papers were evaluated for inclusion and both were included: one (Mentes 2003) described a cluster randomised trial: four nursing homes were randomised to intervention or control groups; and the other (Robinson 2002) was a before-and-after study, in which the patients were monitored 2 weeks pre-intervention, then received 5 weeks of the intervention, followed by 2 weeks post-intervention study.

1  
2 Both studies were conducted in the USA and both received funding from non-  
3 industry sources. There were 49 patients in the Menten (2003) study and 51 in  
4 the Robinson (2002) study.

### 5 6 **9.1.2 Population**

7  
8 Both studies took place in a long-term care setting. In the Menten (2003) study,  
9 patients with acute confusion at baseline were excluded. Nine of 24 people in  
10 the intervention group and two of 25 in the control group had a diagnosis of  
11 cognitive impairment, although it was not specified how this was diagnosed or  
12 defined. In the Robinson (2002) study, it was unclear how many participants had  
13 cognitive impairment. Sensory impairment was not reported in either study.

14  
15 In the Menten (2003) study, the mean number of drugs daily was 6.4 in the  
16 intervention group compared with 7.1 among controls (not significantly different)  
17 and in the Robinson (2002) study 80% (41/51) had more than four drugs  
18 prescribed. It was not stated whether all eligible patients were included in either  
19 study.

20  
21 The mean age in the Menten (2003) study was around 82 years and it was 83.5  
22 years in the Robinson (2002) study. The Menten (2003) study included 22 men  
23 and 27 women, and the Robinson (2002) study had 8 men and 43 women.  
24 Ethnicity was reported in the Menten (2003) study: all participants were  
25 Caucasian except for one who was African American. The Robinson (2002) study  
26 did not report ethnicity.  
27

### 28 **9.1.3 Interventions**

29  
30 In the Menten (2003) study, the intervention was an 8-week hydration  
31 management intervention. This was based on calculating a daily individual fluid  
32 goal for each participant adjusted for his or her weight. For the intervention  
33 group, methods for ensuring that a participant met their goals included a  
34 standardised 180 ml fluid intake with each medication administration, fluid  
35 rounds morning and evening and 'happy hours' or 'tea time' twice a week in the  
36 late afternoon. The control group patients' fluid goals were also assessed and  
37 they received 'usual care', described as 'standard nursing care'.

38  
39 The Robinson (2002) study gave the participants a hydration programme which  
40 consisted of the following components: a caregiver knowledgeable in techniques  
41 of fluid administration; an individualised plan of care incorporating the most  
42 effective techniques to administer fluids; a colourful beverage cart with colourful  
43 pitchers and glasses to enhance residents' interest in drinking; and a choice from  
44 2 beverages at each encounter. Residents had a goal of 8 oz twice per day, but  
45 47% did not achieve this goal every time.  
46  
47

#### 1 9.1.4 Comparison

2  
3 Hydration intervention versus usual care; outcomes recorded after 8 weeks  
4 (Mentes 2003). Concurrent medications were not reported in the Mentes (2003)  
5 study.  
6  
7  
8

### 9 9.2 Methodological quality

10

#### 11 9.2.1 RCTs

12

13 In the RCT (Mentes 2003), the method of randomisation to intervention or control  
14 was at the level of the nursing home and was by coin toss. Allocation  
15 concealment was unclear. No account was taken in the analysis of the fact that  
16 this was a cluster randomised trial, and there are likely to be unit of analysis  
17 errors.  
18

19

20 It was assumed that patients were not blinded to treatment allocation. Blinding of  
21 outcome assessors was unclear. In the intervention group, the assessments  
22 appeared to be carried out by the research nurses involved in delivery of the  
23 intervention (i.e. not blinded), but in the control group, the assessment was  
24 carried out by the research nurses blinded to the patient's fluid goals; whether  
25 they were aware of the research question is not clear.

26

27 The study did not report an *a priori* sample size calculation and its small size and  
28 short duration suggest that it may have been underpowered.

29

30 The authors demonstrated baseline comparability of the groups on some  
31 measures (age, gender, number of diagnoses, mean number of daily  
32 medications, depression), but significant differences between the groups on  
33 several measures although there were confounders would be likely to negate  
34 differences between interventions. The intervention group scores on the  
35 NEECHAM Confusion Scale indicated that they were more at risk for delirium  
36 than the control group (mean 26.4 versus 28.4,  $p=0.005$ ). This scale ranges from  
37 0 to 30, where a score of less than 25 indicates confusion, and 26 to 27  
38 indicates at risk of confusion. The treatment group had more patients with a  
39 diagnosis of cognitive impairment (9 versus 2,  $p=0.02$ ) and the treatment group  
40 were more physically frail than the control group (mean scores 79.4 versus  
41 112.2,  $p<0.001$ ) on the Functional Independence Measure (FIM) instrument;  
42 (scale score ranges from 0 to 126; not specified for long-term care but higher  
43 values indicate better function). In addition, the mean length of stay for the  
44 intervention group in long-term care was 22.9 months compared with 94.9  
45 months for control group patients.

46

47 It is noted that, cognitive impairment, a risk factor for delirium, was greater at  
baseline for the intervention group than the control group. The risk factors review

1 had inconsistent evidence regarding whether long-term care was a risk factor for  
2 delirium, and functional status was not investigated as a risk factor for delirium.

3  
4 All patients were followed up for the 8 weeks of the trial and all patients' data  
5 were analysed.

6  
7  
8 The primary outcome measure for the study was 'hydration-linked events',  
9 defined as acute confusion, urinary tract infection, upper respiratory infection,  
10 pneumonia or influenza, preceded by a urine specific gravity of 1.020 or above  
11 and decreased fluid intake as measured by intake records.

12  
13  
14 Delirium assessment was triggered if a participant exhibited a sudden change in  
15 mental status, or a cognitive or behavioural change. A participant was  
16 considered acutely confused if he or she scored lower than baseline on the  
17 MMSE and lower than 25 on the NEECHAM Confusion Scale. The GDG  
18 considered the MMSE to be an inadequate method of assessment of delirium.

19  
20 The differences in baseline comparability between the groups, the randomisation  
21 by nursing home with only four nursing homes involved and the delirium  
22 assessment method mean that this study is at higher risk of bias.  
23

## 24 **9.2.2 Non-randomised study**

25  
26 The Robinson (2002) study was a before-and-after, prospective study. It was  
27 unclear if all eligible participants were included. In addition, the method of  
28 assessing delirium was not reported and, indeed, results for this outcome were  
29 not given. Overall, the nature of the design meant this was poor quality  
30 evidence.  
31  
32  
33

## 34 **9.3 Results**

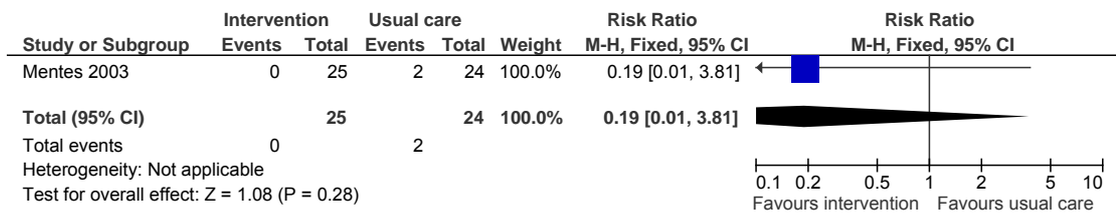
### 36 **9.3.1 Hydration intervention versus usual care**

37

#### 38 **9.3.1.1 Incidence of delirium**

39  
40 The Mentes (2003) study reported no delirium in the treatment group during the  
41 8 weeks of treatment compared with 2 people in the control group (figure 9.1).  
42 The confidence interval is very wide and is consistent with both significant benefit  
43 and significant harm due to the small number of events and so there is  
44 uncertainty about the effect of the intervention on this outcome.  
45

1 Figure 9.1: Acute confusion.



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### 9.3.1.2 Other outcomes

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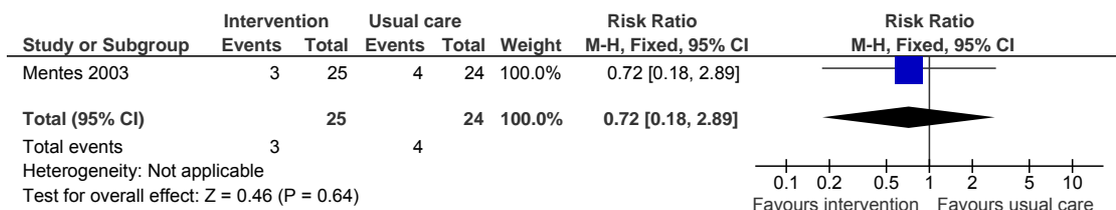
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12

13

The primary outcome measure of the Mentés (2003) study was 'all hydration-linked events', and these were urinary tract infections (1 in the control group), upper respiratory infections (2 in the control group), pneumonia (1 each in the intervention and control groups) and influenza (2 in the intervention group) (figure 9.2). The results are again very imprecise.

13 Figure 9.2: Hydration-linked events.



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## 9.4 Clinical evidence statements

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There is very low quality evidence showing that a hydration intervention had no significant effect on the incidence of delirium, and did not have a significant effect on hydration linked events (urinary tract infection, upper respiratory, pneumonia, influenza); however, there is a lot of uncertainty around these results.

## 1 9.5 Health economic evidence

### 2 9.5.1 Single component non-pharmacological intervention for the prevention of 3 delirium in a long-term care setting

4

5 One economic evaluation study was included as evidence (Robinson 2002). This  
6 was a before-and-after study of 51 older adults in the USA. The aim of the  
7 study was to determine the effect of a specific program on the level of  
8 hydration, and on the prevention of conditions associated with dehydration,  
9 namely, delirium, urinary tract infections, respiratory infections, falls, skin  
10 breakdown, and constipation. Patients in the intervention group were enrolled in  
11 a hydration programme to improve hydration. The programme included a  
12 hydration assistant to administer fluid, an individualised plan of care  
13 incorporating the most effective techniques to administer fluid, a colourful  
14 beverage cart with colourful pitchers and glasses to enhance residents' interest in  
15 drinking, and a choice from 4 beverages at each encounter. The goal was for  
16 each resident to consume an additional 8-ounce beverage mid-morning and mid-  
17 afternoon, which would increase fluid intake to 1.5L daily.

18 Patients in the control group received usual gray coloured institutional carts,  
19 white foam cups and limited variety of beverages. The cost of colourful cups and  
20 assorted beverages was \$154 per week, and \$3 per resident per week. The  
21 average cost of employee time per week per resident was \$8. The intervention  
22 resulted in a cost savings of \$103 over the 5 week period as a result of fewer  
23 negative outcomes for patients. There was no report on the delirium incidence or  
24 severity, mortality or HRQoL. This study did not adequately report the main  
25 outcomes of interest. The results of this study are not directly applicable.

26

27

## 28 9A. 2. HYDRATION FOR THE PREVENTION OF DELIRIUM (HOSPITAL 29 SETTING)

30

### 31 9.6 Description of studies

32

33 One paper was included (O'Keeffe 1996).

34

35

#### 36 9.6.1 Study Design

37

38 This study was an RCT conducted in the UK. The study did not report on funding,  
39 and 60 patients were included.

40

41 The study compared the effectiveness and tolerability of two methods of  
42 delivering fluids; it was not concerned with preventing delirium. The study is  
43 therefore included as indirect evidence, which may inform GDG discussion.

44

### 9.6.2 Population

The study took place in an acute geriatric unit. Patients suffering from mild dehydration or poor oral intake, requiring parenteral fluids for at least 48 hours and who had cognitive impairment were included. Cognitive impairment was defined as disorientation for time and place or an MMSE score of 20 or less. Patients were excluded if there was clinical evidence of poor tissue perfusion or if the amount of fluid administered would be critical (e.g. in those with renal or heart failure).

The mean age was 82.5 years and 38% were male. Ethnicity was not reported.

### 9.6.3 Interventions

In the O’Keeffe (1996) study the patients were randomised to receive either subcutaneous or intravenous fluids. Up to 2 litres of fluid were permitted in a 24 hour period.

### 9.6.4 Comparison

Subcutaneous fluids versus Intravenous fluids; outcomes recorded at 48 hours. Concurrent medications were not reported.

### 9.6.5 Outcome measures

The review’s primary outcome measure was incidence of delirium. However, this included study did not give this outcome, but reported on agitation, serum urea and serum creatinine levels at 48 hours and the incidence of local oedema.

## 9.7 Methodological quality

The O’Keeffe (1996) study reported an adequate method of randomisation (table of random numbers) and a partially adequate method of allocation concealment (sealed envelope).

Blinding of patients would not have occurred due to the method of intervention. Blinding of outcome assessors was unclear.

The study reported an *a priori* sample size calculation. In order to detect a difference in serum urea of 1.5mmol/l between the two groups, at 80% power and 5% significance level, it was estimated that a sample size of 56 patients would be required; the study included 60 patients.

Baseline comparability was reported on age, gender, serum urea, serum creatinine levels, and baseline agitation levels. Agitation levels were assessed by a doctor using the modified Cohen-Mansfield Agitation Inventory based on personal observations and discussion with nurses or carers regarding the behaviour of the patient during the previous 48 hours.

1 There was less than 20% missing data, one patient in the subcutaneous group  
 2 died and one patient in the intravenous group was switched to the subcutaneous  
 3 route after 24 hours because of difficulties with venous access. These patients  
 4 were not included in the analysis.

5  
 6 Overall, the study was considered not to be at higher risk of bias, although it  
 7 only reported indirect outcomes.  
 8  
 9

10 **9.8 Results**

11  
 12 **9.8.1 Subcutaneous versus intravenous hydration**

13  
 14 **9.8.1.1 Agitation**

15  
 16 There was a large significant effect of the method of hydration in relation to  
 17 agitated behaviour, with significantly fewer patients experiencing agitation  
 18 related to the subcutaneous method of hydration; RR 0.46 (95% 0.28 to 0.76)  
 19 (figure 9.31). There was some imprecision in the result.  
 20  
 21  
 22

23 Figure 9.3: agitation



25  
 26 (NB: Scale 0.2 to 5)

27  
 28 **9.8.1.2 Serum urea and creatinine levels**

29  
 30 The study reported the serum urea and serum creatinine levels for both groups at  
 31 48 hours. For serum urea, there was no significant difference between  
 32 interventions; mean difference (MD) -0.27 mmol/l (95% CI - 0.78 to 0.24)].  
 33 There was also no significant difference between the serum creatinine levels at  
 34 48 hours; MD 0.31 µmol/l (95% CI -0.20 to 0.82); figure 9.4.  
 35  
 36  
 37

Figure 9.4: serum levels

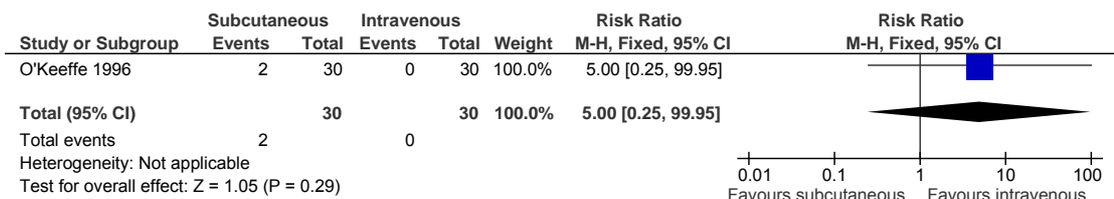


NB: Scale -1 to 1

### 9.8.1.3 Local Oedema

The O'Keeffe (1996) study reported that local oedema was noted in two patients receiving fluids subcutaneously. The confidence interval is very wide due to the small number of events and there is insufficient evidence to draw conclusions about the effect of different hydration strategies on this outcome (figure 9.5).

Figure 9.5: local oedema



NB: Scale 0.01 to 100

## 9.9 Clinical evidence statements

There is no evidence on the effect of subcutaneous versus intravenous fluids on the incidence, duration or severity of delirium. There is moderate quality evidence comparing subcutaneous and intravenous methods of hydration to show significantly lower levels of agitation in patients receiving fluids subcutaneously compared with intravenously, and to show no significant difference in levels of serum urea or serum creatinine levels.

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## **9.10 GDG discussion**

The GDG considered the evidence from this study and decided agitation was not a surrogate outcome for delirium. Although the study was not examining delirium as an outcome, the GDG felt that this study which included patients with cognitive impairment (hence patients at high risk of delirium) would provide an example of strategies for hydration that work.

## **9A. 3. MUSIC THERAPY FOR THE PREVENTION OF DELIRIUM IN A HOSPITAL SETTING**

### **9.11 Description of studies**

Four papers were evaluated for inclusion. Two studies were excluded. Reasons for exclusions are reported in Appendix G .Two papers were included in this review (McCaffrey 2004; McCaffrey 2006).

#### **9.11.1 Study Design**

No studies were conducted in the UK; both were conducted in the USA. The study by McCaffrey (2004) used a non-probability convenience sample of 66 patients from a large tertiary care centre in south-east Florida. McCaffrey (2006) had a sample size of 124 patients from a hospital in Florida, but no further details were given. The McCaffrey (2004) study did not report the number of patients in the intervention or control groups.

#### **9.11.2 Population**

Both studies took place in a university hospital setting in the postoperative orthopaedic unit. Postoperative patients included were those undergoing elective hip or knee surgery, who were alert and oriented to provide consent, able to complete preoperative paperwork independently, and able to hear music.

Proportions of patients with low, intermediate and high risks of delirium at baseline were not reported in either of the studies. Neither delirium nor dementia status at baseline was reported.

The mean age of the patients was 75.7 years (SD 6; range 59 to 82 years) in the McCaffrey (2006) study and 73 years (SD 5) in the McCaffrey (2004) study.

1 In the McCaffrey (2006) study, there was a higher proportion of women (64.5%,  
2 80/124) than men (35.5%, 44/124) and 67% of all patients had knee surgery  
3 (the rest had hip surgery). These details were not reported in the earlier study  
4 (McCaffrey 2004). Ethnicity was not reported in either of the studies.

5

### 6 9.11.3 Interventions

7 The interventions evaluated were:

- 8 • Music therapy: patients in individual rooms were given a bedside compact  
9 disc (CD) player that would automatically play music for a minimum of 1  
10 hour, 3 times/day (McCaffrey 2004) or for a minimum of 1 hour, 4  
11 times/day (McCaffrey 2006). The music started while the patient was  
12 awakening from anaesthesia and continued during the recovery period.
  - 13 ○ The McCaffrey (2004) study stated that the number of times that  
14 the CD could automatically be turned on was three times a day at  
15 the *most*, but that the *minimum* time was 1 hour, three times daily.  
16 In the study by McCaffrey (2006) the CD player would  
17 automatically play CDs for a minimum of 1 hour, 4 times daily.
  - 18 ○ In addition, nurses and family members were asked to turn on the  
19 music when they walked into the orthopaedic unit room.
  - 20 ○ Once awake and oriented, patients received the same instructions  
21 so they could play music when they desired.
  - 22 ○ The first CD placed in the player was chosen by the researcher.  
23 Other musical selections were available to the patients based on  
24 their musical preference.
  - 25 ○ Patients were visited by research assistants to ensure the CD  
26 players were working and that the times for automatic starting of  
27 the CD coincided with the patients' preference, and that the music  
28 playing was what the patient preferred.

29

30 Intervention and control groups in both studies had full access to in-room  
31 televisions, and both groups received standard postoperative care. Patients  
32 were not permitted to bring any electronic music devices into their hospital  
33 rooms.

34

### 35 9.11.4 Comparisons

36 The following comparison was carried out in both studies:

- 37 • Music therapy versus no treatment
  - 38 ○ Both groups received standard postoperative care

- 1                   ○ The total length of postoperative care was 3 days in both the  
2                   intervention and control groups in one study (McCaffrey 2006),  
3                   but was unclear in the other study (McCaffrey 2004).  
4

## 5   **9.12 Methodological quality**

6                   The method of sequence generation was not reported in either study; patients  
7                   were randomly assigned to rooms that had been designated intervention or  
8                   control; this was subject to room availability. Allocation concealment was  
9                   considered to be adequate because the recovery room nurses who assigned  
10                  patients to rooms were said to be unaware of the experimental and control  
11                  group rooms' designation.

12  
13                  Blinding of the outcome assessor was unclear in both studies. It was not possible  
14                  to blind the patients, but the GDG did not consider this to be important. *A priori*  
15                  sample size and power calculations were not reported in either of the studies.

16  
17                  The McCaffrey (2006) study reported limited data on the demographic  
18                  characteristics of the patients. Patients in each group were similar in age,  
19                  proportion of men and women, and proportion of patients with hip and knee  
20                  surgery. This was not reported in McCaffrey (2004).

21  
22                  Only the McCaffrey (2006) study reported on withdrawals. 1.6% (2/126)  
23                  patients were lost to follow-up due to cardiovascular complications during  
24                  surgery, but missing data were not reported for individual groups. The  
25                  McCaffrey (2004) study did not report whether an intention to treat (ITT)  
26                  analysis was carried out, and McCaffrey (2006) used an available case  
27                  analysis.

28  
29                  Both studies evaluated 'acute confusion' as a primary outcome, which was  
30                  identified with delirium: nurses kept computerised notes, recording signs and  
31                  symptoms of delirium. These nurse-identified signs and symptoms of delirium and  
32                  confusion were reviewed retrospectively by researchers with the orthopaedic  
33                  nursing staff to achieve consistency. In the McCaffrey (2004) study, the number  
34                  of episodes of confusion and delirium were entered as a numerical score for that  
35                  patient and the McCaffrey (2006) study recorded the number of patients with at  
36                  least one episode of acute confusion. The GDG did not consider this to be a  
37                  reliable measure of delirium assessment and so these studies were regarded with  
38                  caution.

39                  Overall, these studies were considered to have a higher risk of bias because  
40                  neither had a validated method of assessing delirium incidence.

41

## 1 9.13 Results

### 2 9.13.1 Music therapy plus standard postoperative care versus standard 3 postoperative care

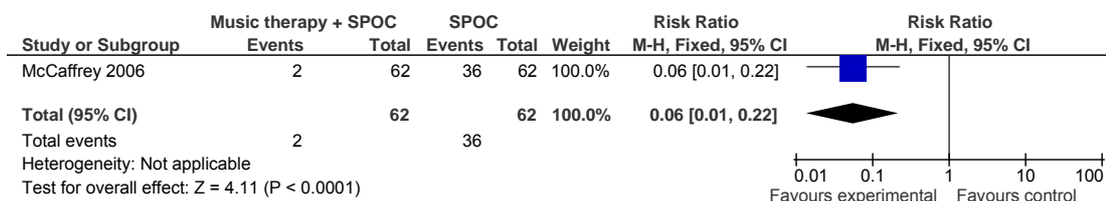
#### 5 9.13.1.1 Incidence of delirium

6 The McCaffrey (2004) study reported that patients receiving music therapy had  
7 significantly fewer periods of confusion or delirium during their hospitalisation  
8 than patients who received no additional therapy, and gave a p-value of 0.001  
9 without detailing the results.

10  
11 The McCaffrey (2006) study in 124 patients demonstrated that significantly  
12 fewer patients experienced acute confusion in the music therapy group. Although  
13 the CI was very wide, the results were not considered to be imprecise as far as  
14 decision making was concerned (figure 9.6); RR 0.06 (95% CI 0.01 to 0.22). This  
15 corresponds to an NNT of 2 (95% CI 2 to 3) for a control group rate of 58%.

16  
17

Figure 9.6: number of patients with delirium



18  
19

NB: forest plot scale 0.01 to 100

20

#### 21 9.13.1.2 Activities of daily life

22 Both studies assessed the patient's 'readiness to ambulate' during the  
23 postoperative period (McCaffrey 2004; McCaffrey 2006). An ambulation  
24 readiness profile was conducted by physiotherapists in both studies using  
25 postoperative scores ranging from 1 (indicating that patients were not ready to  
26 ambulate) to 10 (indicating that patients may be ready to ambulate that day or  
27 the next). The scores were based on: pain level; alertness; stable vital signs;  
28 ability to correctly identify person, place and time; ability to comprehend  
29 instructions; and willingness to participate in their own recovery.

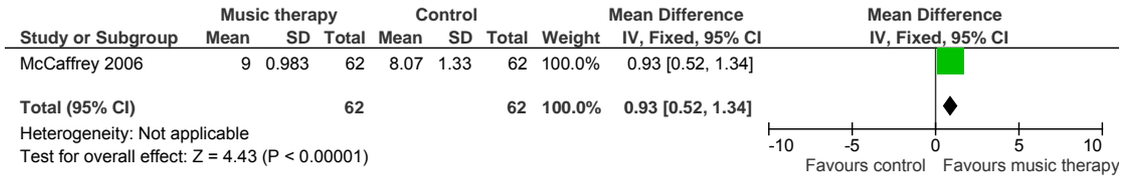
30  
31  
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34

McCaffrey (2004) found that patients receiving music therapy had significantly  
higher scores on the readiness to ambulate scale for the day of surgery than did  
patients who received no additional therapy, and reported a p-value of 0.001.  
No other details were given.

1  
2  
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7

McCaffrey (2006) demonstrated that patients in the music therapy group had significantly higher scores for readiness to ambulate after undergoing surgery than patients in the control group (figure 9.7); MD 0.93 (95%CI 0.52 to 1.34). This is, however, a small effect even though significant.

Figure 9.7: patients readiness to ambulate after undergoing surgery



8  
9

NB: Scale -10 to 10

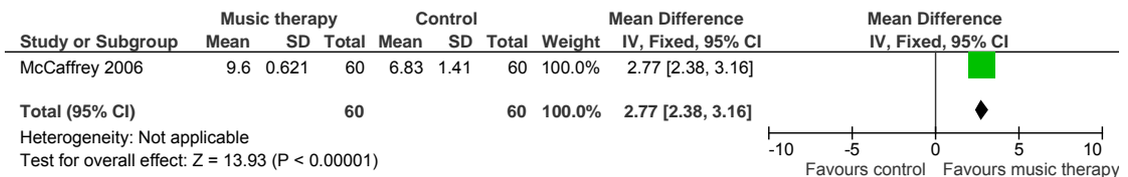
10

11 **9.13.1.3 Patient satisfaction**

12 The McCaffrey (2006) study also measured patient satisfaction: the researcher  
13 phoned each patient 2 weeks after discharge from hospital to determine their  
14 satisfaction with their postoperative experience in the hospital. A scale of 1–10  
15 was used (1 representing the worst experience and 10 the best experience they  
16 could imagine). Analysis showed a significantly higher score for the intervention  
17 group (figure 9.8); MD 2.77 (95%CI 2.38 to 3.16) for a control group score of  
18 6.83.

19  
20

Figure 9.8: patient satisfaction



21  
22

NB: Scale -10 to 10

23

24 **9.14 Evidence statements**

- 25 • There is low quality evidence from one RCT comparing music therapy with  
26 usual care which showed:
- 27 ○ a significantly lower incidence of delirium in the group receiving  
28 music therapy.
  - 29 ○ a higher score for readiness to ambulate after undergoing  
30 surgery in the music therapy group.
  - 31 ○ a higher score in patient satisfaction in the music therapy group.

## 1 **9 B) Multicomponent prevention**

### 2 **9.15 Description of studies**

3 Fourteen papers were evaluated for inclusion. Two studies were excluded  
4 because there were fewer than 20 patients in each arm (Astaneh 2007;  
5 Schindler 1989). Three other studies were excluded and are listed in Appendix  
6 G with reasons for exclusion. Nine reports of studies were included (Bogardus  
7 2003; Gustafson 1991; Harari 2007a; Inouye 1999; Landefeld 1995;  
8 Lundström 2005; Marcantonio 2001; Wanich 1992; Wong 2005), The Bogardus  
9 (2003) study was a six month follow up, post hospital discharge, of a sample of  
10 patients (705/852 (83%)) from the Inouye (1999) study. It appears that these  
11 patients were representative of the original sample; 133/852 (16%) had died.

#### 12 **9.15.1 Study Design**

13 Three studies (Landefeld 1995; Lundström 2005; Marcantonio 2001) were RCTs  
14 and six had a non-randomised design: the latter included two non-randomised  
15 controlled trials (Inouye 1999; Wanich 1992), and three historical controlled  
16 trials (Gustafson 1991; Harari 2007a; Wong 2005).

17 The unit of randomisation in all the RCTs was the patient. One of the non-  
18 randomised controlled studies (Wanich 1992) allocated patients to different  
19 wards (but did not say how this was done), and the Inouye (1999) study  
20 allocated patients by forming matched pairs, matched on age within 5 years,  
21 sex, and base-line risk of delirium (intermediate or high).

22 In the historical controlled trials (Gustafson 1991; Harari 2007a; Wong 2005),  
23 all eligible patients were enrolled at two different time periods. All the studies  
24 compared a group of participants in the period before the intervention was  
25 given with a group who were given the intervention.

26  
27 One study (Harari 2007a) was conducted in the UK. Four studies were carried  
28 out in the USA (Inouye 1999; Landefeld 1995; Marcantonio 2001; Wanich  
29 1992); two were in Sweden (Gustafson 1991; Lundström 2005), and one was  
30 conducted in Australia (Wong 2005). With the exception of Wong (2005), all of  
31 the studies were supported by research grants not associated with industry.  
32 Wong (2005) did not state a funding source.

33 One included study had fewer than 100 patients (Wong 2005: n = 99). Two  
34 studies had more than 100, but fewer than 200 patients (Harari 2007a: n =  
35 108; Marcantonio 2001: n = 126). Five studies enrolled more than 200 patients  
36 (Gustafson 1991: n = 214; Inouye 1999: n = 852; Landefeld 1995: n = 651;  
37 Lundström 2005: n = 400; Wanich 1992: n = 235).

38

## 1 9.15.2 Population

2 All of the studies took place in hospital settings. In four studies, patients were  
3 undergoing surgery, either for hip fracture (Gustafson 1991; Marcantonio 2001;  
4 Wong 2005), or for hip, knee, or other replacements (Harari 2007a). The Harari  
5 (2007a) study intervention was targeted at-risk patients at higher risk of  
6 adverse events/illness (e.g. those with poorly controlled diabetes) and included  
7 those who had been assessed as being too 'medically unfit' to go on the waiting  
8 list; the control group were not selected in this way. The other studies included  
9 older people with acute medical illness (Inouye 1999; Landefeld 1995;  
10 Lundström 2005; Wanich 1992).

11 Comorbidities in patients undergoing surgery were reported in three studies:  
12 Gustafson (1991) reported that some patients also had cerebrovascular  
13 diseases, cardiovascular diseases, hypertension, diabetes, Parkinson's disease,  
14 renal failure, lung disease, on-going infection, urinary incontinence, constipation,  
15 prostatism, depression, and psychosis. Harari (2007a) reported that some of the  
16 surgical patients had rheumatoid arthritis, heart disease, heart failure, atrial  
17 fibrillation, diabetes, renal impairment, hypertension, chronic lung disease,  
18 prostate or bladder problems and cerebrovascular disease. Wong (2005)  
19 reported that some patients had vascular disease, diabetes, chronic lung disease  
20 and/or depression/anxiety at baseline. Comorbidities were not specifically  
21 stated in Marcantonio (2001); 39% in the intervention group and 33% in the  
22 control group were reported to have high medical comorbidity at baseline  
23 (Charlson index of at least 4).

24 Of the studies that examined older people with acute medical illness, reasons for  
25 hospitalisation included cardiac, respiratory, infection, metabolic, neoplasm,  
26 cerebrovascular, or other diagnoses (Inouye 1999; Landefeld 1995; Lundström  
27 2005; Wanich 1992).

28 Medications taken at baseline were reported by Gustafson (1991) and  
29 Lundström (2005). In the Gustafson (1991) study, drugs or groups of drugs taken  
30 by patients included digitalis, diuretics, antihypertensives, nitroglycerin,  
31 analgesics, steroids, antiasthma, sulfonyleurea, insulin, warfarin, laxatives,  
32 antidepressants, neuroleptics, benzodiazepines, other sedatives, antiparkinson  
33 drugs and other drugs; in this study, 16% of patients were not taking drugs.  
34 Lundström (2005) also reported the proportions of patients taking digitalis,  
35 diuretics, beta-blockers, calcium blockers, insulin, analgesics, benzodiazepines  
36 and neuroleptics. None of the other studies reported details on medicine use at  
37 baseline (Harari 1997a; Inouye 1999; Landefeld 1995; Marcantonio 2001;  
38 Wanich 1992; Wong 2005).

39 All of the studies evaluated older patients. The age range across studies was 50  
40 to 102 years, with the mean age, where given, ranging from 75 to 84 years. In  
41 almost all studies the majority of patients were women (Gustafson 1991: 74%;  
42 Harari 2007a: 60%; Inouye 1999: 61%; Landefeld 1995: 67%; Lundström  
43 2005: 56%; Marcantonio 2001: 79%; Wong 2005: 72%). Wanich (1992)  
44 reported that the sex distribution was approximately equal. Ethnicity was  
45 reported in three studies (Inouye 1999; Landefeld 1995; Marcantonio 2001), in  
46 which 59 to 90% of patients were white. Wanich (1992) only reported that  
47 ethnic distributions were approximately equal.

1 The majority of studies (Gustafson 1991; Harari 2007a; Landefeld 1995;  
2 Lundström 2005; Marcantonio 2001; Wanich 1992; Wong 2005) did not  
3 explicitly report the proportions of patients with low, intermediate and high risks  
4 of delirium at baseline, although it may be inferred that many were at high risk.  
5 For example, the Marcantonio (2001) study included hip fracture patients. The  
6 Inouye (1999) study reported that 72% patients had an intermediate risk of  
7 delirium and 28% had a high risk: patients were defined as having intermediate  
8 risk if they had 1 or 2 risk factors and high risk if they had 3 or 4 risk factors  
9 from the following list: visual impairment, severe illness (APACHE II score more  
10 than 16), cognitive impairment (MMSE score below 24), high blood urea nitrogen  
11 to creatinine ratio of at least 18.

12 In the majority of studies, at least some patients were reported to have  
13 dementia: two studies (Inouye 1999; Lundström 2005) reported on cognitive  
14 function using the MMSE instrument (scale 0-30): Inouye (1999) reported a mean  
15 MMSE score of 24 (SD 5) in the treatment group and 23 (SD 5) in the control  
16 group. In Lundström (2005), patients in the treatment and control groups both  
17 had an average score of 25 (SD 6). It is noted that a score of 20-26 indicates  
18 mild dementia or cognitive impairment. Landefeld (1995) reported using the  
19 MMSE scale for the first 21 items (scale of 0-21); they reported scores of 17 in  
20 both groups, and also reported that 11% had dementia at baseline. Inouye  
21 (1999) reported that 11% of the patients had dementia using a modified  
22 Blessed Dementia Rating Scale (>2), and Marcantonio (2001) reported that  
23 40% of patients had dementia at baseline using the Blessed score (>4).  
24 Lundström (2005) reported that 5% of patients had dementia using DSM-IV  
25 criteria, and Gustafson (1991) reported that 22% in intervention group and  
26 15% in the control group had dementia using the DSM-III criteria. Wanich (1992)  
27 and Wong (2005) reported using the MMSE score, but did not present any data.  
28 Harari (1997a) did not report cognitive function scores.

29 Three studies reported sight and hearing impairment at baseline (Gustafson  
30 1991; Inouye 1999; Lundström 2005):

- 31 • In Gustafson (1991), visual and hearing impairment was reported in 23%  
32 and 25% of the patients respectively (methods of assessment not stated).
- 33 • Inouye (1999) reported that visual and hearing impairment occurred in  
34 23% and 26% of the patients respectively (as evaluated using the  
35 standard Jaeger test, and the Whisper test)
- 36 • Lundström (2005) reported that 2% of the intervention group and 4% of  
37 the intervention group had impaired hearing, and 15% to 17% had  
38 impaired vision. In this study, hearing impairment was considered if a  
39 patient could not hear a normal speaking voice within one metre or  
40 without a hearing aid, and impaired vision was considered if a patient  
41 could not read a newspaper without glasses.

42 It is also noted that 59% of patients in the Inouye (1999) study were dehydrated  
43 on admission.

44

### 1 **9.15.3 Interventions**

2 The interventions were largely education and/or management changes with  
3 structured protocols for patient care. Each intervention is described below.  
4 Additionally, in order to understand and compare the interventions more  
5 effectively we have carried out a themed analysis, breaking down the  
6 interventions by risk factors addressed, and whether or not a multidisciplinary  
7 team and educational interventions are described (table 9.1):

8

#### 9 **9.15.3.1 Education programme and reorganisation of nursing and medical care consisting** 10 **of four parts (Lundström 2005):**

- 11 • Two-day course for staff on geriatric medicine which focused on  
12 assessment, prevention and treatment of delirium and underlying causes  
13 (e.g. urinary tract infection); lectures started before the intervention, with  
14 a follow up during the first month of the study
  - 15 ○ training regarding medical interventions included focus on the  
16 prevention of hypoxaemia, hypercortisolism, and avoidance of  
17 drugs with anticholinergic properties
  - 18 ○ training regarding nursing interventions focused on interaction  
19 with patients with reduced attention and orientation in a stressful  
20 situation and optimisation of care for these patients
- 21 • Staff education on caregiver-patient interaction that focused on patients  
22 with dementia and delirium, particularly with respect to comprehension  
23 and orientation of the patients
- 24 • A patient-allocation nursing care system with individualised care (in which  
25 small teams of nurses had full responsibility for a small number of  
26 patients to promote continuity of care)
- 27 • Monthly guidance for nursing staff, focusing on caregiver-patient  
28 interaction
- 29 • The control ward received usual hospital care organised in a task allocated  
30 way.

31

#### 32 **9.15.3.2 'Elder Life Program' (Inouye 1999; Bogardus 2003)**

33 This programme was implemented by a trained interdisciplinary team, consisting  
34 of a geriatric nurse-specialist, two specially trained Elder Life specialists, a  
35 therapeutic-recreation specialist, a physiotherapy consultant, a geriatrician and  
36 trained volunteers.

- 37 • The performance of each staff member was evaluated quarterly, with  
38 completion of checklists to ensure competency and consistent and  
39 complete adherence to protocols.
- 40 • This multidisciplinary team implemented the following interventions, which  
41 were targeted at particular risk factors:

- 1                   ○ Cognitive impairment; outcome: change in orientation score (first  
2                   10 items on MMSE)
- 3                   - an orientation protocol: schedule/name board; reorienting  
4                   communication
- 5                   - therapeutic activities protocol: cognitively stimulating  
6                   activities 3 times daily (e.g. discussion of current events,  
7                   word games, structured reminiscence)
- 8                   ○ Sleep deprivation; outcome: change in use of sedative drugs for  
9                   sleep
- 10                  - non-pharmacological sleep protocol: at bedtime, warm  
11                  drink, relaxation tapes/music, back massage
- 12                  - sleep-enhancement protocol: unit-wide noise-reduction  
13                  strategies (e.g. vibrating beepers, quiet hallways) and  
14                  schedule adjustments to allow sleep (e.g. medications)
- 15                  ○ Immobility; outcome: change in Activities of Daily Living score
- 16                  - Early-mobilisation protocol: ambulation or active range-  
17                  of-motion exercises 3 times daily; minimising use of  
18                  immobilising equipment (e.g. bladder catheters; physical  
19                  restraints)
- 20                  ○ Visual impairment; outcome: early correction of vision up to 48 h  
21                  after admission
- 22                  - vision protocol (for visually impaired people only): visual  
23                  aids (e.g. glasses and magnifying lenses) and adaptive  
24                  equipment (e.g. large illuminated telephone key pads,  
25                  large print books, fluorescent tape on call bell), with  
26                  daily reinforcement of their use
- 27                  ○ Hearing impairment; outcome: change in Whisper Test score
- 28                  - hearing protocol (for hearing impaired people only):  
29                  portable amplifying devices, earwax disimpaction,  
30                  special communication techniques, with daily  
31                  reinforcement of their use
- 32                  ○ Dehydration; outcome: change in ratio of blood urea nitrogen to  
33                  creatinine
- 34                  - dehydration protocol (for those with evidence of  
35                  dehydration, i.e. ratio of blood urea nitrogen to  
36                  creatinine of at least 18): early recognition of  
37                  dehydration and volume repletion (e.g. encouragement  
38                  of oral fluid intake)

39                  Usual care was standard hospital services provided by a multidisciplinary team.

40

1 **9.15.3.3 Education and multicomponent intervention (Wanich 1992), which consisted of:**

- 2           • Nursing staff education in the month before the start of the study and  
3           repeated once during the study on mental and functional status  
4           assessments, nursing management of deficits in sensory-perceptual  
5           function, mobility and environmental modifications
- 6           • Patient assessment and management plans recorded on charts and shared  
7           with staff and families to assist in nursing care and discharge planning
- 8           • Families education and consultation including reassurance and coping skills;  
9           orientation and personalising the environment
- 10          • 2 geriatricians assigned to intervention group
- 11          • Orientation: provision of orientation cues to patients (e.g. day of week,  
12          current events, a discussion of their condition, information about upcoming  
13          diagnostic or therapeutic measures); updated calendars in every room;  
14          favourite TV programmes determined)
- 15          • Communication (families and nurses taught to communicate clearly and  
16          slowly, and to use repetition and orientation clues)
- 17          • Mobilisation (e.g. getting patients out of bed each day, ambulation daily,  
18          physical and occupational therapy as needed)
- 19          • Sensory stimuli (glasses and hearing aids available and nurses encouraged  
20          patients to use them)
- 21          • Environmental modifications (lighting to decrease sensory deprivation; night  
22          lights used)
- 23          • Medical management (to assess medications suspected of contributing to  
24          delirium, e.g. neuroleptics, antidepressants, narcotic analgesics, sedative-  
25          hypnotics, and their unnecessary use discouraged)
- 26          • Discharge planning (with multidisciplinary team: primary nurse, social  
27          worker, discharge planning nurse, physiotherapist, occupational therapist  
28          and dietitian)
- 29          • The control group received usual care, but also received the physical and  
30          occupational therapy components in similar proportion to the intervention  
31          group.

32

33 **9.15.3.4 'Acute Care for Elders' programme (Landefeld 1995)**

34 This was carried out in a special unit and consisted of:

- 35           • Daily assessment by nurses of physical, cognitive and psychosocial function;  
36           daily review of medical care
- 37           • Daily rounds by multidisciplinary team: medical and nursing directors, a  
38           primary nurse, a social worker, a nutritionalist, a physical therapist and a  
39           visiting-nurse liaison

- 1           • Protocols to improve self-care, continence, nutrition, mobility, sleep, skin
- 2            care, mood, cognition (implemented by the primary nurse based on the
- 3            daily assessment)
- 4           • Specially designed environment (carpeting, handrails, uncluttered hallways,
- 5            elevated toilet seats and door levers)
- 6           • Orientation (large clocks and calendars)
- 7           • Patient-centred care
- 8           • Planning for discharge including early involvement of a social worker and
- 9            home healthcare nurse if indicated
- 10          • Protocols to minimise the adverse effects of selected procedures (eg.
- 11          urinary catheterisation) and medications (e.g. sedative-hypnotic agents)
- 12          The comparator was usual care in another general medical unit.

13

#### 14    **9.15.3.5 'Proactive care of older people undergoing surgery (POPS)' (Harari 1997a)**

15          This was a multidisciplinary, preoperative, comprehensive geriatric assessment  
 16          service with postoperative follow-through:

- 17          • Multidisciplinary team consisting of a consultant geriatrician, a nurse
- 18          specialist in older people, an occupational therapist, a physiotherapist
- 19          and a social worker
- 20          • Preoperative assessment: Abbreviated Mental Test Score, Geriatric
- 21          Depression Scale, Barthel Index, Timed Up and Go, 180° turn, body
- 22          mass index, continence screen, orthostatic blood pressure, pain score, and
- 23          peak expiratory flow rates. Then investigation and treatment targeted
- 24          the identified issues and medical comorbidities were optimised according
- 25          to evidence based practice.
- 26          • Management plans and goals were agreed with the patient, and post-
- 27          discharge plans made preoperatively
- 28          • Most patients had preoperative home visits from the occupational therapist
- 29          and the physiotherapist, providing aid and equipment
- 30          • Preoperative education of patients in optimising postoperative recovery
- 31          including home exercises, good nutrition, relaxation techniques and pain
- 32          management; mean number of preoperative clinic visits was 1.79 (range
- 33          1-4)
- 34          • Postoperative staff education on early detection and treatment of medical
- 35          complications, early mobilisation, pain management, bowel-bladder
- 36          function, nutrition and discharge planning
- 37          • Postoperative early detection and treatment of medical complications,
- 38          early mobilisation, pain management, bowel-bladder function, nutrition
- 39          and discharge planning

- 1 • Follow-up therapy home visit in those with functional difficulties, and  
2 outpatient clinic review in those with ongoing medical problems  
3

4 **9.15.3.6 Quality improvement programme (Plan-do-study-act methodology with**  
5 **interventions introduced incrementally) (Wong 2005)**

- 6 • Project team consisting of a consultant and registrar geriatricians, a  
7 consultant anaesthetist, two clinical nurse managers, a member of the  
8 quality improvement unit, and representatives of allied health staff  
9 (pharmacist, dietitian) met approximately fortnightly to supervise the  
10 programme
- 11 • Staff education on definition of delirium, predisposing and precipitating  
12 factors, investigations (including use of CAM) and management of  
13 delirium
- 14 • Geriatric team made recommendations for each person, based on the  
15 following:
- 16 ○ Regulation of bladder and bowel function (remove indwelling  
17 catheters, screening for constipation, retention) [recommended in  
18 24%]
  - 19 ○ Early detection/treatment of major complications (myocardial  
20 ischaemia, infection, pulmonary embolism, etc) [recommended in  
21 22%]
  - 22 ○ Maintenance of fluid and electrolyte imbalance [recommended in  
23 14%]
  - 24 ○ Discontinuation of unnecessary medications (especially  
25 benzodiazepines, antihistamines, drugs with anticholinergic  
26 effects) [recommended in 14%]
  - 27 ○ Maintenance of adequate oxygen delivery (oxygen and blood  
28 transfusion)
  - 29 ○ Pain management
  - 30 ○ Treatment of agitated delirium (including low dose haloperidol or  
31 lorazepam)
  - 32 ○ Use of appropriate environmental stimuli (soft lighting, avoid  
33 putting delirious patients in the same room)
  - 34 ○ Sensory impairment improvement (glasses, hearing aids)
  - 35 ○ Orientation (clock, calendar)
  - 36 ○ Adequate nutritional intake (dentures used properly, adequate  
37 positioning, dietitian review and intervention)
  - 38 ○ Early mobilisation and rehabilitation
- 39  
40

1 **9.15.3.7 Proactive geriatrics consultation (Marcantonio 2001)**

2 This consisted of:

- 3 • A consultation with a geriatrician that began preoperatively, or within 24  
4 hours postoperatively. Geriatrician made daily visits during  
5 hospitalisation at which time target recommendations were made using  
6 the following (it is noted that the recommendations were only made if the  
7 consultants noticed something that was not already being done):
- 8 • Adequate CNS oxygen delivery
- 9 ○ oxygen therapy to keep saturation above 90%, treatment to  
10 raise systolic bp to above 2/3rds that at baseline or above 90  
11 mm Hg; blood transfusion to keep haematocrit above 30%  
12 [applied to 73%]
- 13 • Fluid/electrolyte balance
- 14 ○ Treatment to restore serum sodium, potassium, glucose to normal  
15 limits
- 16 • Treatment of dehydration or fluid overload
- 17 ○ Detected by examination or blood tests [applied to 43%]
- 18 • Treatment of severe pain (regular paracetamol) and treatment of break  
19 through pain
- 20 • Elimination of unnecessary medication
- 21 ○ Discontinuation of benzodiazepines, anticholinergics, histamines  
22 [applied to 56%]
- 23 ○ Elimination of medication redundancies
- 24 • Regulation of bowel/bladder function
- 25 ○ Removal of urinary catheter by postoperative day 2, with  
26 screening for retention or incontinence [applied to 63%]
- 27 • Nutritional intake
- 28 ○ Dentures used properly [applied to 37%]
- 29 ○ Nutritional supplements
- 30 ○ Temporary nasogastric tube
- 31 • Early mobilisation [applied to 47%] and rehabilitation
- 32 • Prevention, detection and treatment of major postoperative complications
- 33 ○ Including myocardial infarction/ischaemia, pneumonia/COPD,  
34 pulmonary embolism [applied to 50%], urinary tract infection
- 35 • Environmental stimuli
- 36 ○ soft lighting and use of radio/tape recorder
- 37 ○ but wasn't implemented for any patient in practice

- 1           • Sensory stimuli (glasses and hearing aid)
- 2           • Orientation (clock and calendar)
- 3           • Treatment of agitated delirium (including haloperidol or lorazepam)

4

5           The usual care group received management by the orthopaedics team, including  
6           internal medicine or geriatrics consultations, but on a reactive rather than  
7           proactive basis.

8

### 9    **9.15.3.8 Geriatric-anaesthesiologic intervention programme (Gustafson 1991)**

10          This involved the following:

- 11           • Surgical policy (patients were operated on as soon as possible after  
12           admission)
- 13           • Preoperative assessment: for all patients, mostly by a specialist in geriatric  
14           and internal medicine
- 15           • Individualised thrombosis prophylaxis: heart failure patients given Heparin,  
16           rest Dextran (c.f. control group all given Dextran)
- 17           • Diuretics: patients with clinical signs of heart failure were treated with extra  
18           doses of diuretics
- 19           • Oxygen therapy: nasal oxygen given soon after admission (1 l/min).  
20           Oxygen enriched air was given throughout the operation and the first  
21           postoperative day, and then continued or not depending on oxygenation  
22           levels
- 23           • Anaesthetic technique: all patients had sc morphine premedication and  
24           spinal anaesthesia; patients who had systolic blood pressure below 90  
25           mm Hg were aggressively treated with phenylephrine
- 26           • Postoperative assessment: all patients were assessed several times by a  
27           geriatrician
- 28           • Treatment of patients developing delirium for complications associated with  
29           acute coronary syndrome (e.g. anaemia, heart failure, urinary retention)  
30           – this is expected to confound measurements on the duration of delirium  
31           and incidence of delirium at 7 days
- 32           • Wards: all patients admitted to the same ward (but not part of the study  
33           protocol)
- 34           • Nursing care in both groups treated according to task allocation system

35

### 36    **9.15.4           Comparisons**

37          The following comparison was carried out in all studies:

#### 1 **9.15.4.1 Multicomponent intervention versus usual hospital care**

2 In Lundström (2005), 'usual hospital care' was task-oriented care (i.e. the same  
3 nurse handling particular tasks for all patients; meaning that several nurses could  
4 care for each patient each day) – for this study, the intervention was patient  
5 oriented care

6

### 7 **9.16 Methodological quality**

#### 8 **9.16.1 Randomised trials**

9 The method of sequence generation was adequate in two RCTs: Landefeld  
10 (1995) employed a computer-generated sequence and Marcantonio (2001)  
11 used a random numbers table. The Lundström (2005) study did not describe  
12 sequence generation.

13 Allocation concealment was partially adequate in Marcantonio (2001), in which  
14 sealed envelopes were used. The method of allocation concealment was not  
15 stated in Landefeld (1995). The study by Lundström (2005) was an RCT in which  
16 patients were randomly allocated to any ward with an accessible bed (i.e. this  
17 may constitute some selection bias), so that intervention patients and controls  
18 were on different wards. The study stated that the staff and assessors knew to  
19 which wards the patients were allocated, i.e. there was inadequate allocation  
20 concealment.

21 Due to the nature of the interventions, none of the RCTs were patient blinded.  
22 Marcantonio (2001) reported that the outcome assessor was blinded to the  
23 intervention status of the patients, and Landefeld (1995) stated that data were  
24 obtained by means of interviews and the interviewers were not blinded to the  
25 patients' group assignments. The Lundström (2005) study stated that the outcome  
26 assessors were blinded for delirium diagnosis, but were not blinded otherwise.

27 Marcantonio (2001) reported an *a priori* sample size calculation to detect the  
28 incidence of delirium; they required a sample size of 125 to detect a 33%  
29 decrease in risk with 80% power (they had sample size of 126). Landefeld  
30 (1995) and Lundström (2005) did not report *a priori* sample size calculations.

31 In the Landefeld (1995) study, 36% (651/1974) of eligible patients were  
32 randomised; 1143 eligible patients were not enrolled because beds were not  
33 available in the intervention or control wards at the time of their admission. In the  
34 Marcantonio (2001) study, 85% of eligible patients were included; of 149  
35 eligible patients, 23 refused to participate. In Lundström (2005), all eligible  
36 patients were randomised.

37 Marcantonio (2001) and Landefeld (1995) demonstrated baseline comparability  
38 of the groups. In Lundström (2005), there were more females in the intervention  
39 ward ( $p = 0.04$ ), a higher mean age in the control ward ( $p = 0.02$ ), a greater  
40 proportion of patients previously diagnosed with diabetes mellitus on the  
41 intervention ward ( $p < 0.001$ ), and a greater proportion of patients diagnosed

1 with myocardial infarction on the intervention ward ( $p = 0.03$ ). The GDG did not  
2 consider these to be important differences.

3 In the Landefeld (1995) study, 7% of patients in both the intervention and  
4 control groups were lost to follow-up. In both these studies, the authors only  
5 analysed data from available patients. Lundström (2005) and Marcantonio  
6 (2001) reported no missing data, and all patients were included in their  
7 analyses.

8 Two studies evaluated delirium as a primary outcome (Marcantonio 2001;  
9 Lundström 2005). The primary outcome in Landefeld (1995) was the change  
10 from admission to discharge in the number of activities of daily living (ADL) that  
11 patients could perform independently.

12 Marcantonio (2001) evaluated delirium using the CAM diagnostic algorithm.  
13 Marcantonio (2001) also assessed individual symptoms of delirium using the DSI  
14 and severity of delirium was evaluated using the MDAS (scored 0-30, 30 best).  
15 In Lundström (2005), delirium was diagnosed using the DSM-IV criteria. Delirium  
16 was also measured using a modified version of the Organic Brain Syndrome  
17 (OBS) scale, which incorporated the MMSE to assess disorientation, and the Katz  
18 ADL index to assess ADL. Landefeld (1995) only reported a mental status score  
19 based on the Mini-Mental State scale (using a score from 0-21, with higher  
20 scores indicating better cognitive function). This was considered to be a partially  
21 adequate method of measuring delirium.

22 Overall, Lundström (2005) was considered to be at higher risk of bias due to  
23 inadequate allocation concealment, and non-blinding of outcome assessors.  
24 Landefeld (1995) was at higher risk of bias because of non-blinding of outcome  
25 assessors, incomplete recruitment and the use of the MMSE for diagnosis of  
26 delirium. With the exception of Landefeld (1995), the RCTs were relatively small  
27 and not highly powered.

28

## 29 **9.16.2 Non-randomised studies**

30 Five non-randomised studies were included in the review (Gustafson 1991;  
31 Harari 2007a; Inouye 1999; Wanich 1992; Wong 2005).

32 Three studies reported that all eligible patients were recruited consecutively to  
33 the study (Gustafson 1991; Harari 2007a; Wong 2005). The Inouye (1999)  
34 study stated that, of the 2434 patients meeting the inclusion criteria, 1265  
35 (52%) were excluded because of inability to participate in interviews: because  
36 of a hospital stay of less than 48 hours (219); prior enrolment in their study  
37 (324), dementia (154), patient not available, etc. The 1265 excluded patients  
38 did not differ significantly from those included in terms of age, sex, risk of  
39 delirium, but a larger proportion were excluded from the control group than the  
40 intervention. The remaining patients had 250/1169 (21%)  
41 patients/family/physician who refused consent and an additional 67 who could  
42 not be matched. These unmatched patients were significantly older, had a higher  
43 risk of delirium at baseline, and were more likely to be admitted to a usual-care  
44 unit.

1 In the Wanich (1992) study, 117/354 (33%) patients/physicians refused  
2 consent.

3 Inouye (1999) was a non-randomised controlled study, and patients were  
4 allocated to groups by matching on age, sex, and baseline risk of delirium.  
5 Wanich (1992) was also a non-randomised controlled study in which patients  
6 from different wards were compared; it was not stated if the patients were  
7 matched.

8 Gustafson (1991) was a historical controlled trial in which a group of patients  
9 given the intervention in December 1986 to January 1988 were compared with  
10 a group of patients in the same hospital from March 1983 to June 1984.

11 Harari (2007a) was a historical controlled trial in which a group of patients  
12 given the intervention in August 2003 to February 2004 were compared with a  
13 group of patients in the same hospital from May to July 2003.

14 Wong (2005) was a historical controlled trial where baseline data were  
15 collected for 28 days on one group of patients, and further data were collected  
16 on another group of patients during the subsequent three months.

17 Inouye (1999) took account of possible confounders, by matching patients on the  
18 basis of age, sex and baseline risk of delirium; patients were included only if  
19 their risk of delirium was intermediate or high, as defined in the Inouye (1993)  
20 study. This Inouye (1993) study used a predictive model to define intermediate  
21 and high risk, based on risk factors of visual impairment, severe illness, cognitive  
22 impairment and a high ratio of blood urea nitrogen to creatinine. In order to  
23 appraise the accuracy of the matching on the basis of delirium risk, we need to  
24 assess the quality of the predictive model. We note that the prognostic factor  
25 review classified the Inouye (1993) study as low quality and that the predictive  
26 model did not include the full set of risk factors for delirium as identified in the  
27 risk factors review (section 6.2.1). Therefore, we can conclude that the possible  
28 confounders have not been completely accounted for in the matching process,  
29 although this may not be an important difference.

30 The method involved prospective individual matching of patients that had  
31 already been assigned to treatment groups; patients were admitted to one of  
32 three units (two control and one intervention) and matching was carried out using  
33 a computerised algorithm, based on logistic regression methods. The authors  
34 stated that randomisation of patients to intervention or usual care units was not  
35 feasible because of the large number of patients in all medical units at the time  
36 of the study; a pilot study found that beds in the intervention group were often  
37 unavailable. This pilot study does not appear to have been reported. The  
38 authors contend that their method of prospective matched pairing was chosen as  
39 an alternative to randomisation, but we note that the matching is only on the  
40 basis of known confounders whereas randomisation theoretically matches on  
41 known and unknown. There were no significant differences at baseline for age,  
42 sex, race, married, residence in a nursing home, education, APACHE II score,  
43 impairment in activities of daily living, MMSE score, patients with dementia,  
44 immobility, visual impairment, dehydration, comorbidities. However, the authors  
45 stated that contamination between groups was evident, because of the low rates

1 of delirium in the control group, and because it was stated that intervention  
2 protocols were carried across to the usual care wards. This contamination would  
3 have underestimated the effect.

4 In the Harari (2007a) study, the patients in the intervention group were selected  
5 to be at-risk: those on the waiting list, aged 65 years and older, were sent a  
6 preoperative questionnaire and those with any risk factor (e.g. significant  
7 medical problems) were invited to the POPS clinic. The control group was not  
8 selected in this way and patients were included regardless of case-mix. At  
9 baseline, there was a significant difference in renal impairment and  
10 hypertension), but the study used linear multiple regression to adjust for any  
11 baseline differences. We note that the percentages of people with hypertension  
12 were 80% and 52% in the intervention and control groups respectively  
13 ( $p=0.01$ ); there were 22% and 4%.respectively with renal impairment  
14 ( $p=0.007$ ). These are highly significant differences.

15 In the Wanich (1992) study, the intervention group had significantly more people  
16 with cardiac disease and cerebrovascular accidents and the control group had  
17 significantly more with neoplasm as the primary diagnosis. Adjustments were not  
18 made for the delirium outcome. The study also reported some contamination  
19 because some intervention techniques (e.g. medication management and  
20 physiotherapy) were also given to control patients.

21 The Wong (2005) study reported no significant differences in the age, sex,  
22 mental scores, Barthel indices, types of surgery or comorbidities between the  
23 baseline group and the post intervention group.

24 The Gustafson (1991) study reported no significant differences between groups  
25 in impaired vision, impaired hearing, dementia, depression, psychosis, many  
26 comorbidities, but significantly more people in the intervention group had  
27 cerebrovascular diseases and significantly more had urinary incontinence; the  
28 intervention group also received significantly fewer antiparkinsonian drugs, but  
29 significantly more of other drugs (e.g. penicillin); the control group also had more  
30 patients walking without walking aids before the fracture. Gustafson (1991) did  
31 not consider potential confounders in their analyses. Although these are important  
32 differences, it is not clear what would be their effects on delirium risk.

33 The historical comparison studies did not have blinded outcome assessors, nor did  
34 the Wanich (1992) study. However, the Inouye (1999) study reported that  
35 outcome assessors were blinded.

36 All the non-randomised studies, with the exception of Harari (2007a) evaluated  
37 delirium as a primary outcome. The primary outcome in Harari (2007a) was  
38 hospital length of stay.

39 Two studies (Inouye 1999; Wong 2005) reported that delirium had been  
40 assessed using the CAM, and two studies (Gustafson 1991; Wanich 1992)  
41 diagnosed delirium using the DSM-III criteria. One study (Harari 1997a)  
42 assessed delirium as 'acute change in mental status postoperatively with  
43 improvements pre-discharge', but did not say how this was done. Therefore, the  
44 GDG down graded this study.

45

1 Five non-randomised studies reported no missing data and all patients were  
2 included in their analyses. In Inouye (1999), 6 (1%) patients in the intervention  
3 group and 7 (2%) patients in the control group died during hospitalisation, but  
4 information on delirium was available for all patients. In the 6 month follow up  
5 study (Bogardus 2003), baseline data were available for 705/852 (83%)  
6 patients, 133 (16%) of whom had died. This study reported some additional  
7 missing data for some outcomes (for example, only 580 (68% of original  
8 sample) reported cognitive impairment).

9 Overall, none of the non-randomised studies were of high quality: the study by  
10 Inouye (1999) had the best study design, but large numbers of patients were not  
11 recruited and the matching of patients had limitations. The Bogardus (2003)  
12 study was considered at higher risk of bias for some outcomes because of  
13 missing data.

14  
15 All of the other studies were considered to have a higher risk of bias:

- 16 • Harari (2007a) appeared to compare different types of patient, as well  
17 as not using a recognised method of assessing delirium and being a  
18 historical comparison.
- 19 • Two other studies had baseline differences (Gustafson 1991; Wanich  
20 1992), but all the confounders in these studies appeared to disfavour the  
21 intervention group.
- 22 • The Wong (2005) study was considered at risk of bias because of its study  
23 design
- 24 • The Wanich (1992) study also reported some contamination
- 25 • In all studies except Inouye (1999), none of the outcome assessors were  
26 blinded.

## 28 9.17 Results

### 30 9.17.1 Multi-component hospital care versus usual treatment

31 In summarising the results we have decided to indicate with one, two or three  
32 asterisks, studies which are considered to be at some, higher or much higher risk  
33 of bias respectively (i.e. moderate, low and very low quality studies,  
34 respectively). High quality studies have no asterisks. Where possible, we have  
35 separated the higher quality studies (zero or one asterisk) in the forest plots, or  
36 have outlined the forest plots in green.

37

### 1 **9.17.1.1 Incidence of delirium**

2 With the exception of the RCT by Landefeld\*\* (1995) all studies evaluated the  
3 incidence of delirium. This outcome was evaluated differently between studies  
4 (e.g. cumulative incidence versus incidence at defined time point):

- 5 • the Gustafson\*\* (1991) study reported ACS in the postoperative period  
6 from 8 hours to 7 days and at 7 or more days
- 7 • the Harari\*\*\* (2007a) study reported outcomes measured during the  
8 hospitalisation period (mean 11.5 to 15.8 days)
- 9 • the Inouye\* (1999) study appeared to report the rate of incidence of  
10 delirium up to 7 days and the number of patients were calculated from  
11 percentages
- 12 • Lundström\*\* (2005) reported the incidence of delirium at 24 hours, 3 days  
13 and 7 days after admission. For the latter two days, the authors reported  
14 the data as the number of delirious patients on day 3 or 7 divided by  
15 the number with delirium on day 1. In our analyses, we have used the  
16 total number of patients in each group as the denominator
- 17 • the Marcantonio (2001) study reported the cumulative incidence during  
18 hospitalisation (mean about 3 days)
- 19 • the Wanich\*\*(1992) study recorded the incidence of delirium at some time  
20 during their hospital stay (about 9 days), 38/48 within 24 h of admission
- 21 • the Wong\*\* (2005) study recorded delirium in hospital (median stay 8-10  
22 days)

23

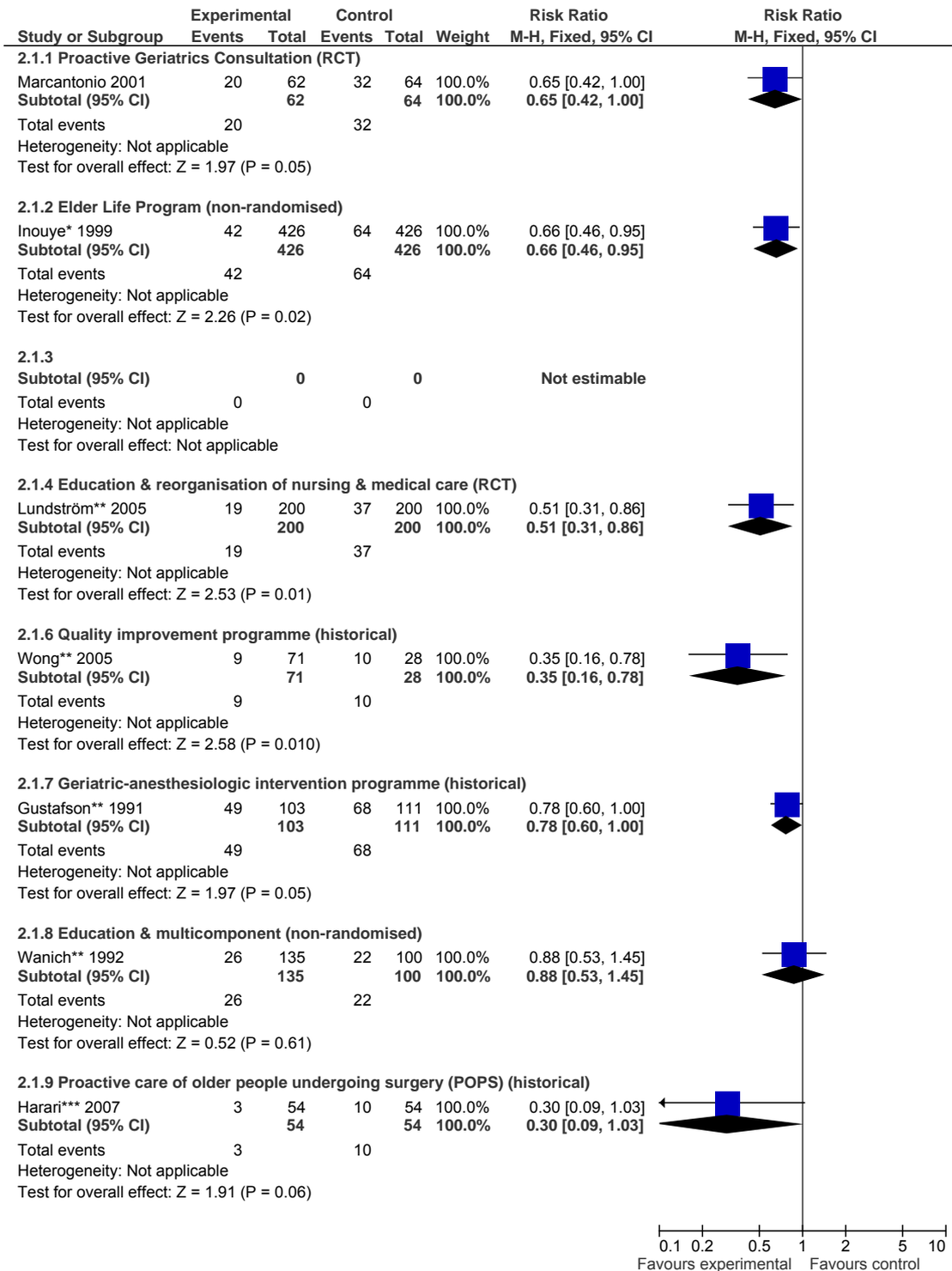
24 Figure 9.9 shows all studies separately for outcomes up to 7 days. Considering  
25 all the studies, we note that, generally, there was a significant effect of  
26 multicomponent interventions on the incidence of delirium. Considering only the  
27 reasonably reliable studies, Marcantonio (2001) and Inouye\* (1999), each had  
28 a relative risk of about 0.66. In general these results were lacking in precision:  
29 the confidence interval was consistent with both a clinically important difference  
30 and no clinically important difference.

31

### 32 **9.17.1.2 Follow up**

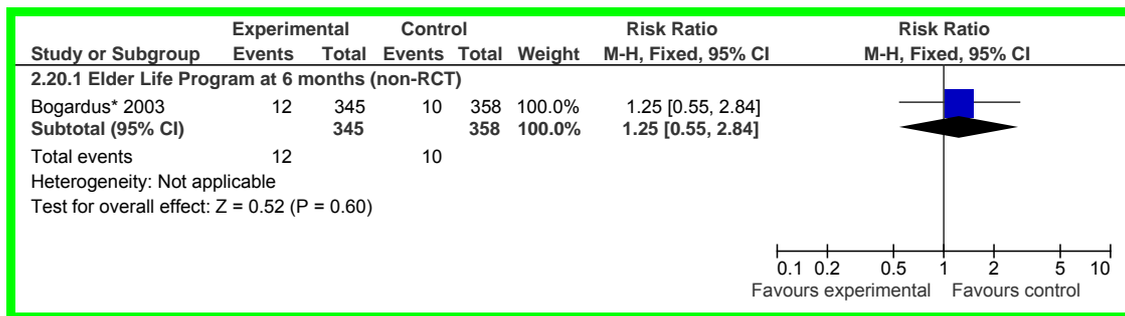
33 The six month follow-up study by Bogardus\* (2003) (following the Inouye\* 1999  
34 study) found no significant difference between the groups (figure 9.10).

1 Figure 9.9: number of patients with delirium in hospital



2  
3  
4

1 Figure 9.10: number of patients with delirium at 6 months follow-up



2

3

4 The confidence limits were consistent with significant harm and significant benefit,  
 5 so the evidence quality was considered to be very low, on the grounds of being  
 6 imprecise.

7

### 8 9.17.1.3 Duration of delirium

9 One RCT reported on the mean number of days with delirium per episode of  
 10 delirium (Marcantonio 2001). The results demonstrate that there was no  
 11 difference in the mean duration of delirium per episode (not per person)  
 12 between the treatment and control group; MD  $-0.20$  days (95%CI  $-0.95, 0.55$ );  
 13 figure 9.11. The results were considered to be precise for this outcome, although  
 14 the study was small.

15

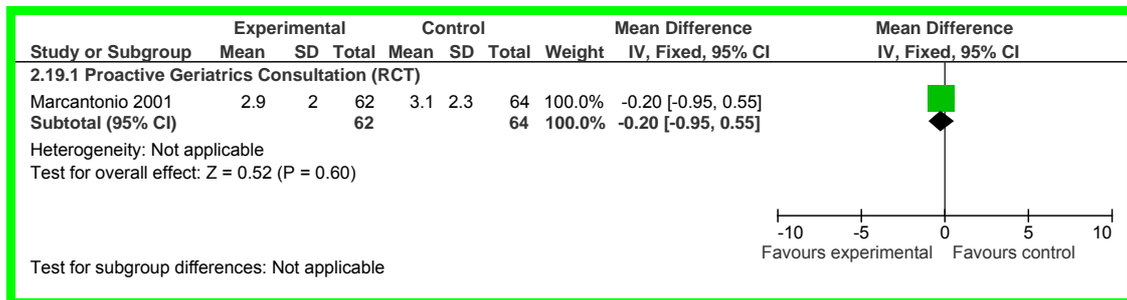
16 One non-randomised study reported on the number of patients with delirium for  
 17 7 days or more (Gustafson\*\* 1991). There was no significant difference  
 18 between groups (figure 9.12).

19

20 The non-randomised study by Inouye\* (1999) reported that the total number of  
 21 days of delirium amongst all patients in each group was significantly lower in the  
 22 intervention group than in the usual-care group (105 versus 161 days,  $p=0.02$ ).

23

1 Figure 9.11: mean duration of delirium

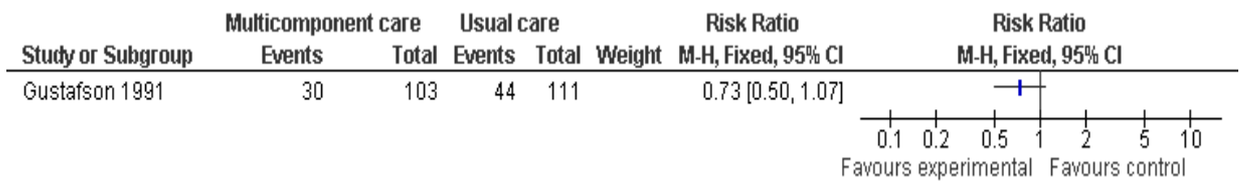


2

3

4

5 Figure 9.12: number of patients with delirium at 7 or more days



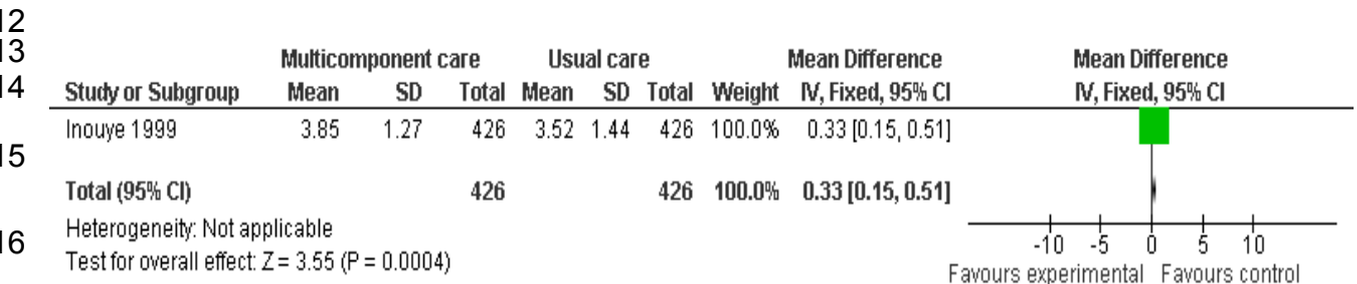
9

1 **9.17.1.4 Severity of delirium**

2 One non-randomised study evaluated severity of delirium (Inouye\* 1999), using  
 3 an additive score for four symptoms (symptom fluctuation, inattention,  
 4 disorganised thinking and an altered level of consciousness), ranging from 0 to 7  
 5 with higher scores indicating increased severity; the GDG were uncertain  
 6 whether this was a validated scale, although it uses individual CAM items.

7 There was no difference in severity of delirium between the intervention and  
 8 control groups (figure 9.13); MD 0.33 (95%CI 0.15 to 0.51); this is a precise  
 9 result.

10  
 11 Figure 9.13: severity scores



17

18 **9.17.1.5 Length of hospital stay**

19 Length of hospital stay was reported in three RCTs (Landefeld\*\* 1995;  
 20 Lundström\*\* 2005; Marcantonio 2001), and five non-randomised studies  
 21 (Gustafson\*\* 1991; Harari\*\*\* 2007a; Inouye\* 1999; Wanich\*\*1992; Wong\*\*  
 22 2005).

23 Three non-randomised trials reported the mean number of hospital days  
 24 (Gustafson\*\* 1991; Harari\*\*\* 2007a; Wanich\*\*1992) (Figure 6).

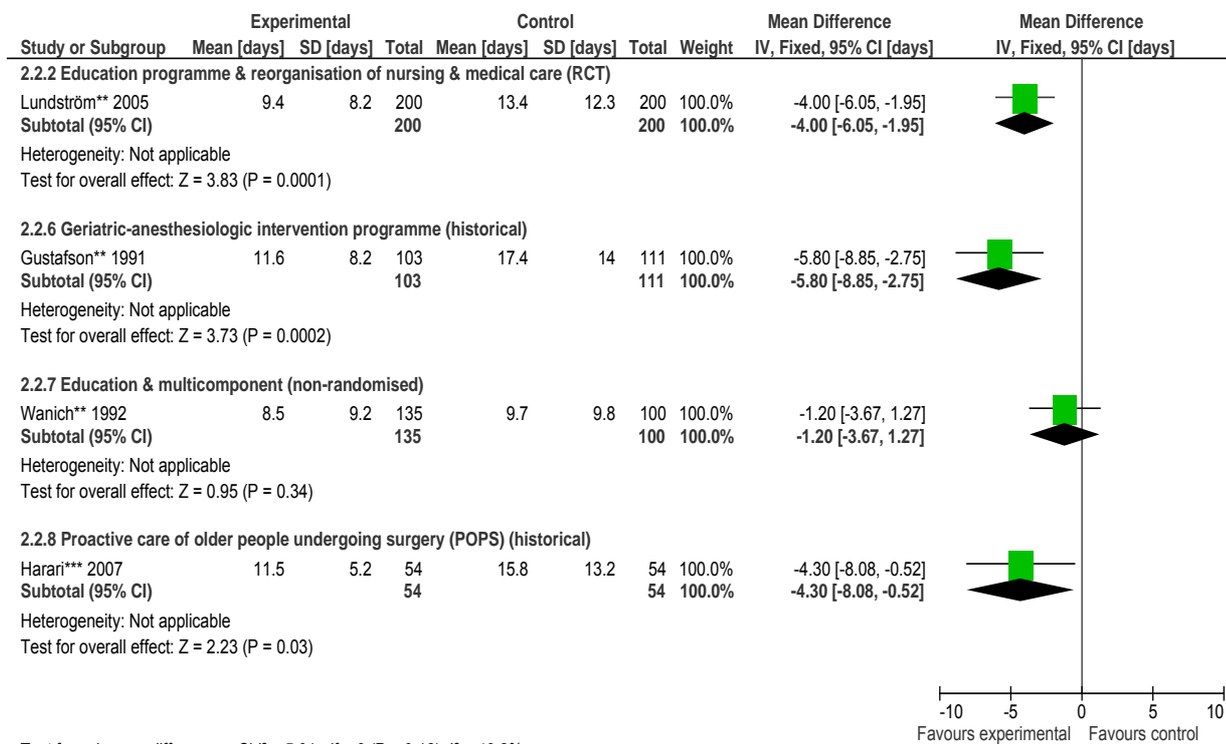
25 Five studies reported the mean length of stay (Gustafson\*\* 1991; Harari\*\*\*  
 26 2007a; Lundström\*\* 2005; Wanich\*\*1992), but in each case, at least one of the  
 27 groups had a skewed distribution.

28 The RCT by Landefeld\*\* (1995) reported mean lengths of hospital stay of 7.3  
 29 and 8.3 days respectively for the intervention and control groups respectively,  
 30 but standard deviations were not reported; the authors also reported that the  
 31 median length of stay (6 days) was the same for both groups. We note that the  
 32 Landefeld\*\* (1995) study did not report the incidence of delirium.

33 The Lundström\*\* (2005) study reported that patients in the treatment ward  
 34 stayed in hospital for significantly fewer days than those in the control group;  
 35 MD -4.05 (95% CI, -6.05, -1.95); figure 9.14. Due to a higher risk of bias,  
 36 however, this result should be interpreted with caution.

37

1 Figure 9.14: length of hospital stay

2  
3

4 With the exception of the Wanich\*\* (1992) study, patients in the intervention  
5 group stayed in hospital for significantly fewer days than patients in the control  
6 group. In the Wanich\*\* (1992) study there was no significant difference in  
7 hospital stay.

8  
9

Four studies reported median length of stay:

- 10 • The Marcantonio (2001) RCT found no significant difference in length of  
11 hospital stay; both groups had a median stay of 5 days (with an  
12 interquartile range of 2); p = 0.95.
- 13 • Inouye\* (1999) reported that the median length of stay was 7 days in the  
14 intervention group and 6.5 days in the control group; this was not a  
15 significant difference (p = 0.95).
- 16 • The Wong\*\* (2005) study reported that the median length of stay was 10  
17 days (2-44) in the intervention group and 8 days (3-41) in the control  
18 group; this was not a significant difference.
- 19 • The Harari\*\*\* (2007a) study reported a median length of stay of 10.0  
20 days (range 4-26) and 14.5 (2-80) days for the intervention and control  
21 groups respectively (this was not a significant difference; p=0.058).

22

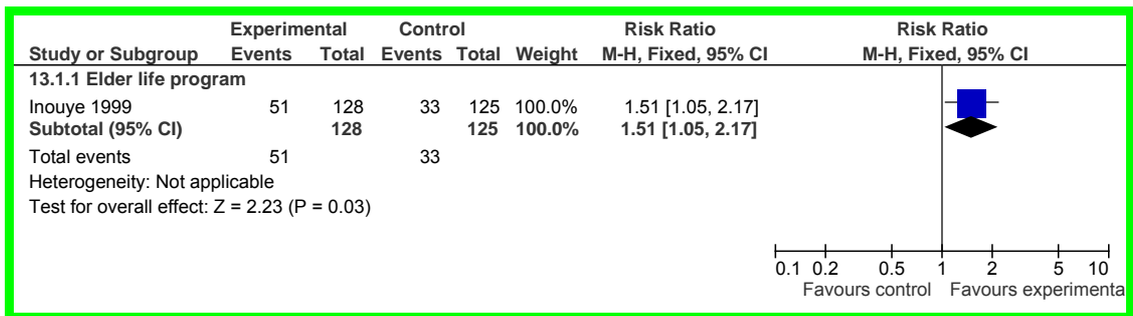
1 **9.17.1.6 Cognitive impairment**

2 The Inouye\* (1999) study reported an adjusted orientation score (10 items on  
 3 the MMSE) at reassessment (day 5 or at discharge if earlier); adjustment was for  
 4 the patients' baseline score. We note that all patients received the cognitive  
 5 impairment protocol once daily and those with an MMSE score below 20 or an  
 6 orientation score below 8 received the protocol 3 times daily (advanced  
 7 protocol); results were only reported for 253 of the original 852 patients (as  
 8 two groups) – we assume this included the patients receiving the advanced  
 9 protocol and their matched pairs in the control group. There were significantly  
 10 more patients who had improved by 2 points on the MMSE at 5 days or at  
 11 discharge (figure 9.15).

12 There was no significant difference in MMSE score in 580 patients (i.e. more than  
 13 20% missing data) at 6 months follow up in the Bogardus\* (2003) study:  
 14 adjusted mean difference -0.3 (95%CI -0.7 to 0.1) on a scale of 0-23. This  
 15 study reported the MMSE score for all patients available, regardless of whether  
 16 they had the advanced protocol.

17  
 18

Figure 9.15: improvement in cognitive impairment at 5 days or discharge



19  
 20  
 21  
 22  
 23

One low quality RCT (Landefeld\*\* 1995) reported no significant difference (p = 0.3) in MMSE scores (0 to 21) between the intervention (17.3) and control (17.7) groups for patients surviving to hospital discharge.

24

25 **9.17.1.7 Number of patients discharged to new long-term care placement**

26 One low quality RCT (Landefeld\*\* 1995) reported that, of the patients admitted  
 27 from private homes who survived to discharge, significantly fewer patients in the  
 28 intervention group were discharged to new long-term care (figure 9.16); RR  
 29 0.64 (95% CI 0.46 to 0.90) which corresponds to a number needed to treat of  
 30 13 (95% CI 8 to 50), for a control group rate of 22%.

31 In addition, two studies (Marcantonio 2001; Wanich\*\*1992) presented  
 32 percentages of patients discharged to institutional settings (e.g. nursing home,  
 33 rehab hospital); however, it was not clear how many of the patients were in  
 34 long-term care settings at baseline.

36

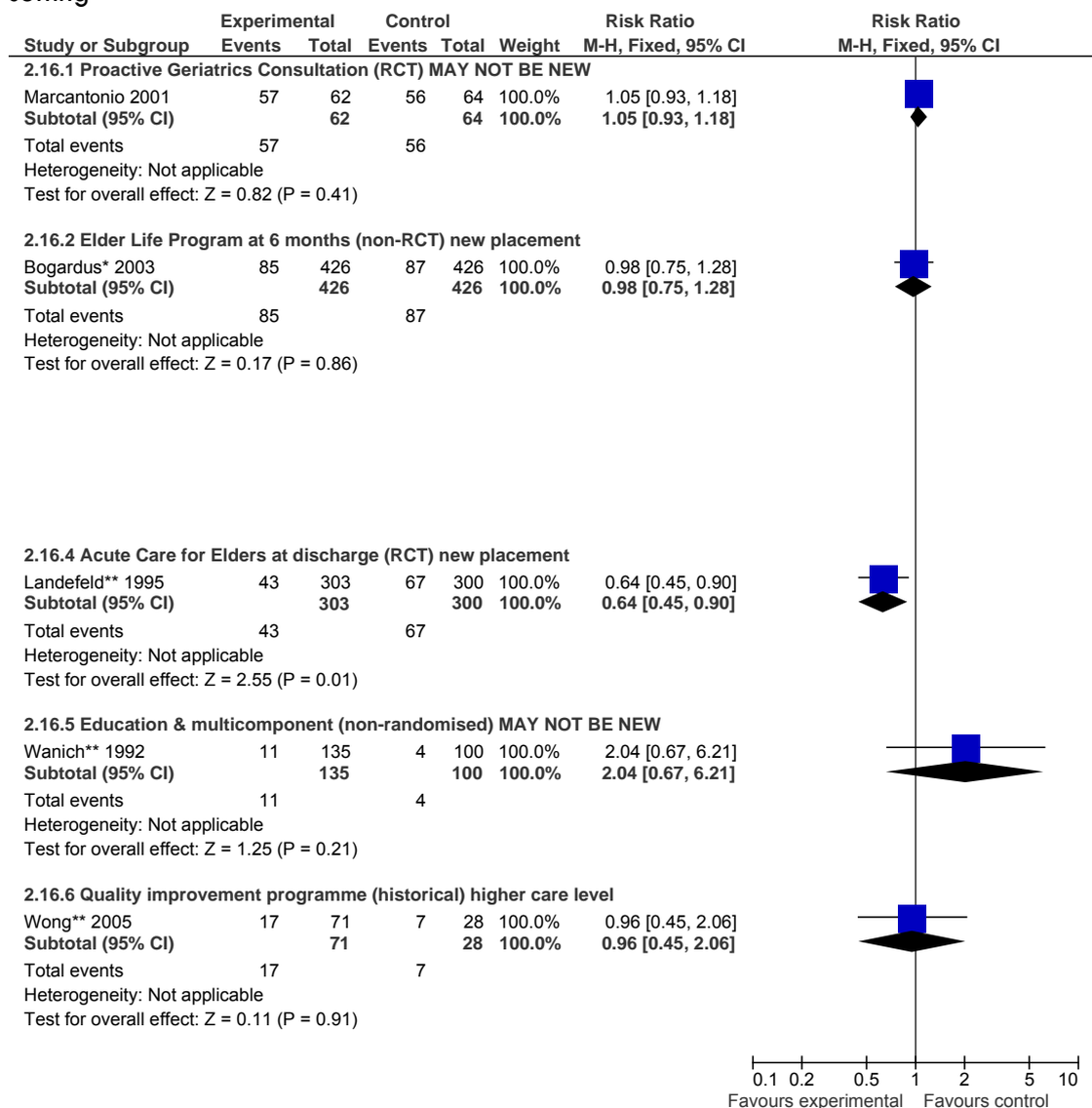
1 In a non-randomised study (Wong\*\* 2005), no significant difference in the  
 2 number of patients discharged to higher level care was found between the  
 3 intervention and control groups (figure 9.16); RR 0.96 (95% CI 0.45, 2.06).

4

5 The Bogardus\* (2003) study reported the number of patients with a new long-  
 6 term placement at 6 months follow up of the Inouye\* (1999) study. The  
 7 denominators used were the number of patients in the original study. There was  
 8 no significant difference between interventions.

9

10 Figure 9.16: number of patients discharged to a new institutional  
 11 setting



12

13

1 **9.17.1.8 Mortality**

2 Two low quality RCTs (Landefeld\*\* 1995; Lundström\*\* 2005) and four non-  
3 randomised studies reported on mortality (Harari\*\*\* 2007a; Inouye\*  
4 1999/Bogardus\* 2003; Wanich\*\*1992; Wong\*\* 2005).

5 The Inouye\* (1999) non-randomised study reported mortality during the  
6 hospitalisation period and the Bogardus\* (2003) study reported mortality  
7 between hospital admission and 6 months follow up. In the latter case, the  
8 denominators used were the number of patients in the original study. There was  
9 no significant difference between interventions, but the confidence interval was  
10 consistent with significant benefit and significant harm.

11 The Lundström\*\* (2005) study reported on mortality but only in patients with  
12 delirium. They found that mortality was less *in delirious patients* who received the  
13 intervention, than in delirious patients who received usual care (2/63 (3.2%)  
14 compared to 9/62 (14.5%),  $p=0.03$ ).

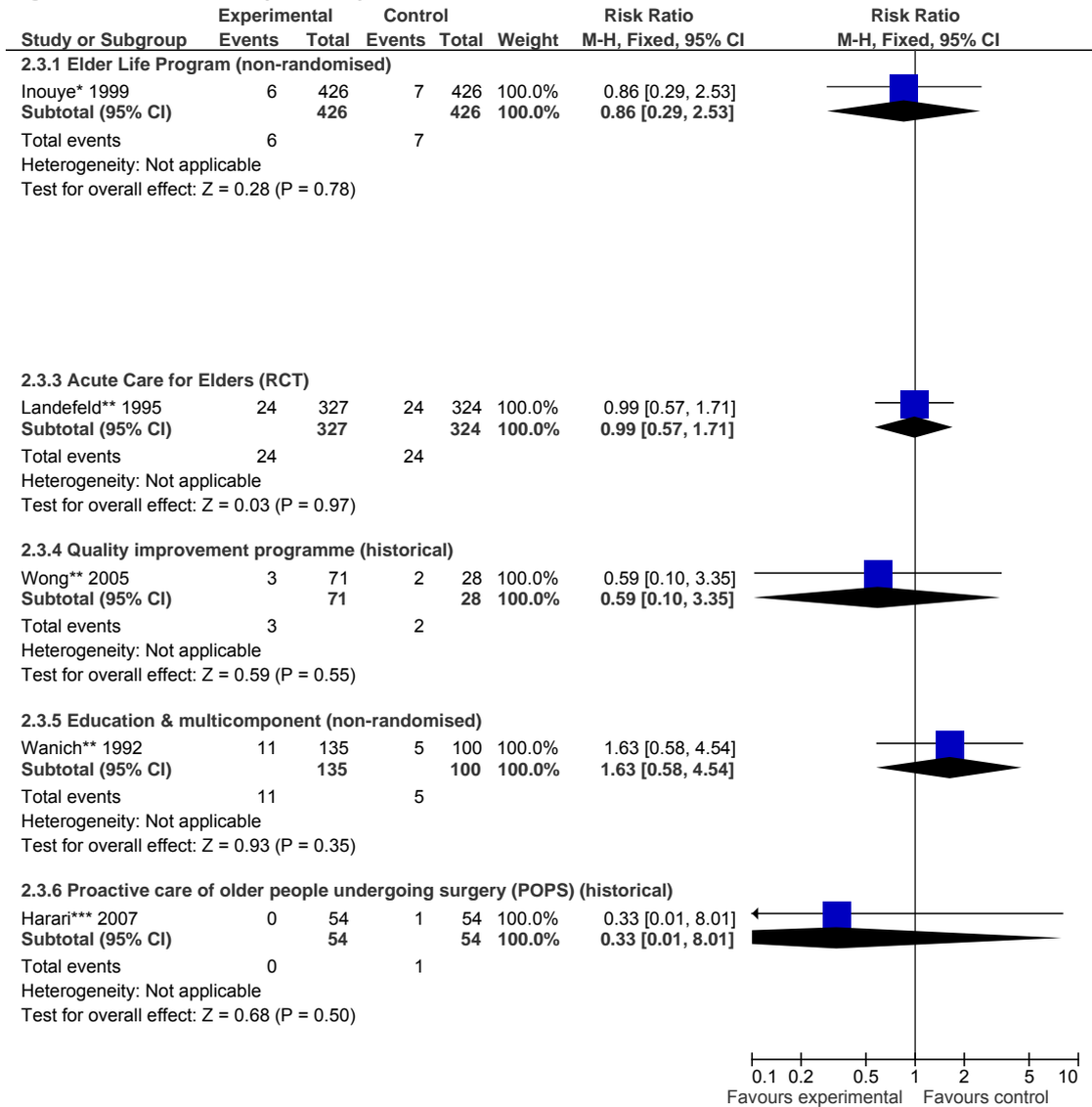
15 In Harari\*\*\* (2007a), the figures reflect the number of patients who died within  
16 30 days of surgery. The Landefeld\*\* (1995) also reported the number of deaths  
17 post discharge and up to 3 months and we used these data to calculate the  
18 number of deaths between admission and 3 months.

19 Overall none of the studies showed an effect on mortality, but often the CIs were  
20 wide and the results imprecise (figures 9.17 and 9.18).

21

22

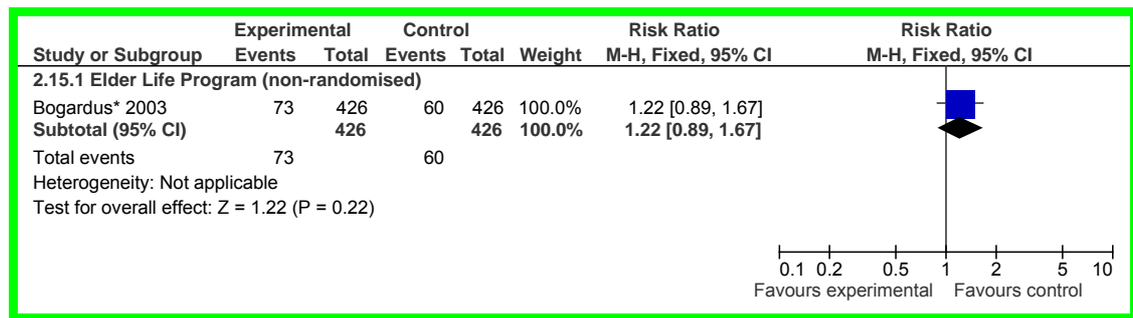
1 Figure 9.17: mortality in hospital



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4

1 Figure 9.18: mortality at up to 6 months follow up

2  
34 **9.17.1.9 Activities of daily living**

5 Three non-randomised studies evaluated ADL (Inouye\* 1999/Bogardus\* 2003;  
6 Landefeld\*\* 1995; Wanich\*\*1992); figure 9.19. The Lundström\*\* (2005) study  
7 also examined the patients using the Katz ADL scale but no results were  
8 reported.

9 The Inouye\* (1999) study reported an adjusted Katz ADL score, on a scale of  
10 0–14 (low scores indicate functional impairment), at reassessment (day 5 or at  
11 discharge if earlier); adjustment was for their baseline score. Although the study  
12 reported that standard deviations were given, this did not agree with the p  
13 value reported and it was assumed that the SDs were standard errors.  
14 Accordingly we calculated standard deviations. There was no significant  
15 difference between interventions (figure 9.20); MD 0.40 (95%CI -0.43, 1.23) on  
16 a scale of 0 to 14. There was no significant difference in the number whose  
17 immobility improved by 2 points but this result was imprecise (figure 9.19). We  
18 note that all patients had ambulation where possible and additional measures  
19 were provided when patients were non-ambulatory, Results were only reported  
20 for 194/852 patients.

21

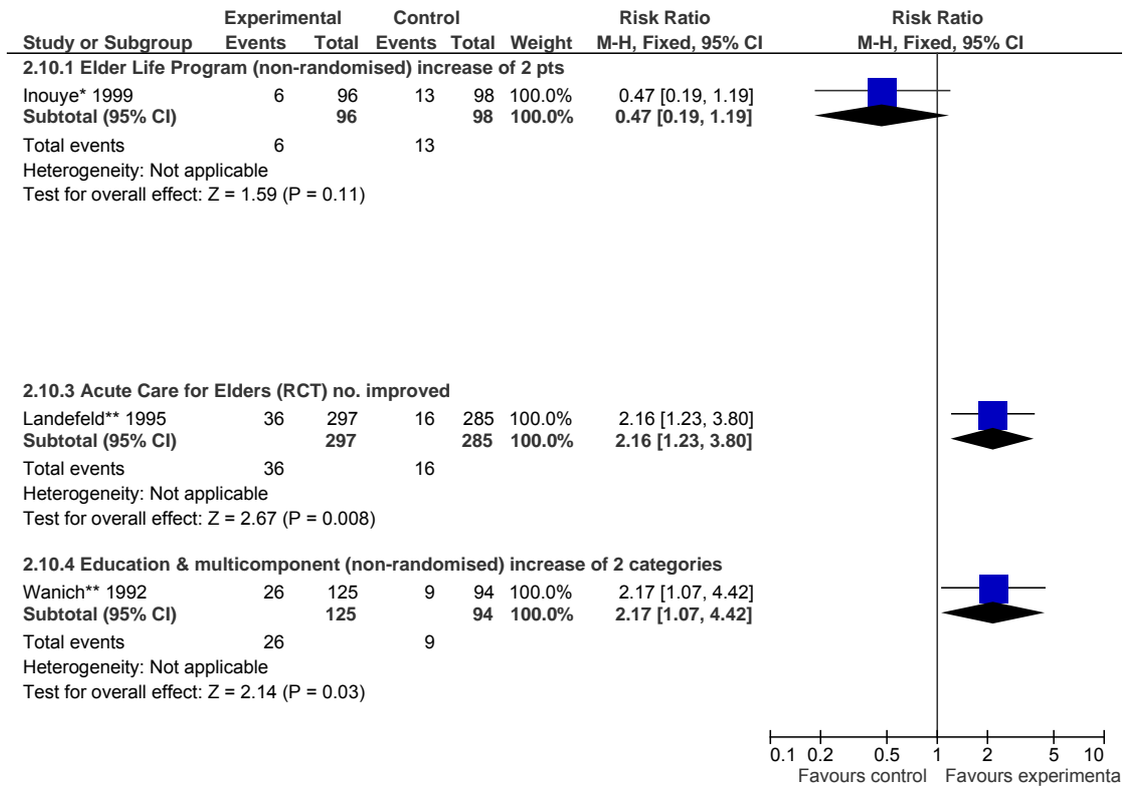
22 In Wanich\*\* (1992) a change in functional status was determined as an increase  
23 or decrease in two or more levels of function (e.g. Katz level C to E or C to A).  
24 By comparing the proportion of patients who were 'better', 'same' and 'worse',  
25 more patients in the intervention group had improved functional status and fewer  
26 had deteriorated in function compared to patients in the control group (p=0.02).  
27 The Wanich (1992) study also carried out a multiple logistic regression analysis  
28 to take into account baseline differences; the adjusted odds ratio was still  
29 significant; OR 3.29 (95%CI 1.26 to 8.17).

30

31 Landefeld\*\* (1995) also reported on the change from admission to discharge in  
32 the number of basic activities performed independently (using the Katz index);  
33 the authors reported the number of patients with improved or much improved  
34 levels of function (figure 9.19) and the mean number of basic activities that could  
35 be performed at discharge (up to 5); this was 3.6 and 3.3 for the intervention  
36 and control groups respectively, which was of borderline significance (p = 0.05).

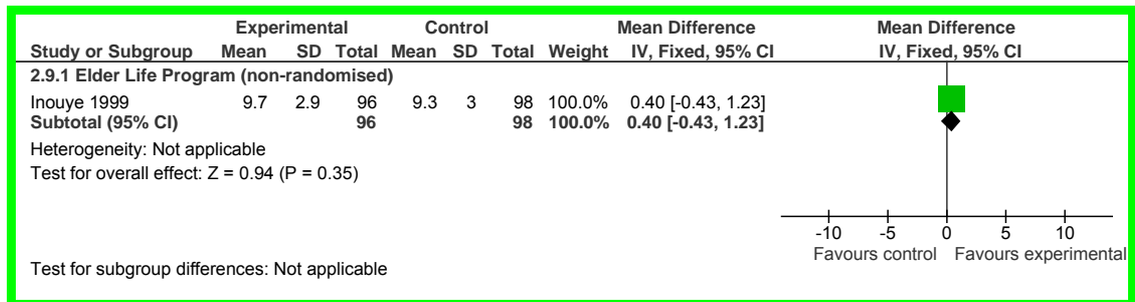
37

1 Figure 9.19: number of patients with an improvement in ADL



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3

4 Figure 9.20: adjusted ADL score



5  
6

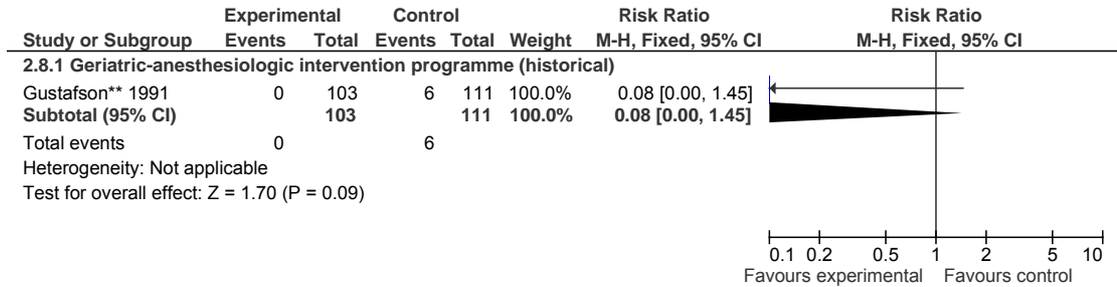
7 **9.17.1.10 Post-discharge follow up**

8 There was no significant difference in ADL score in 704 patients at 6 months  
 9 follow up in the Bogardus\* (2003) study: adjusted mean difference 0.1 (95%CI –  
 10 0.2 to 0.4) on a scale of 0–14. There was also no significant difference in the  
 11 mean number of basic activities that could be performed in the 3 months after  
 12 discharge in the Landefeld\*\* (1995) study; this was 4.0 and 3.8 for the  
 13 intervention and control groups respectively, (p = 0.3).

1 **9.17.1.11 Severe falls**

2 One study (Gustafson\*\* 1991) reported the number of people with severe falls.  
 3 The confidence interval was too wide to determine if there was a difference  
 4 between interventions (figure 9.21).

5  
 6 Figure 9.21: number of patients with severe falls

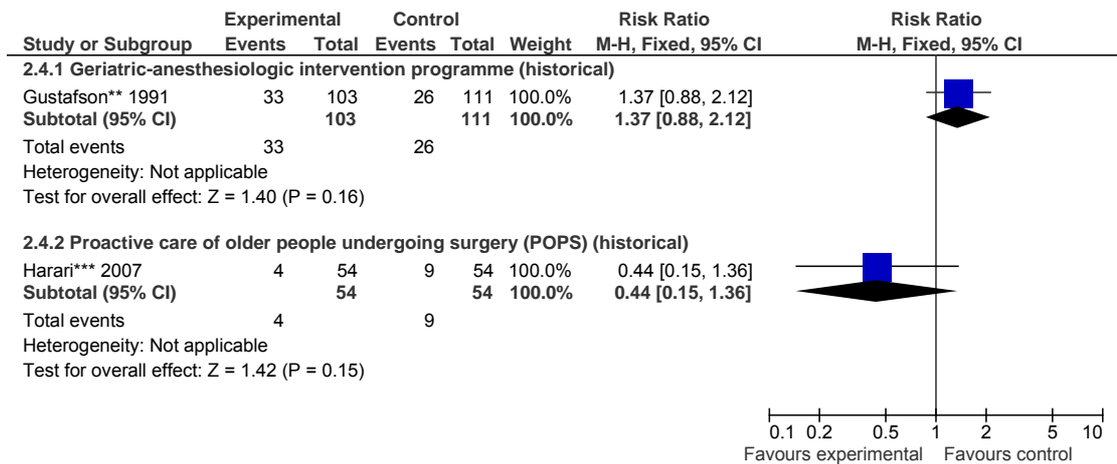


7  
 8 **9.17.1.12 Infections**

9 **Urinary tract infections (figure 9.22)**

10 Two studies (Gustafson\*\* 1991; Harari\*\*\* 2007a) reported the number of  
 11 patients with urinary infections). There was no significant difference between the  
 12 intervention and control studies in the number of patients with urinary tract  
 13 infections, although the results were imprecise in the Gustafson\*\* (1991) study  
 14 and very imprecise in the Harari\*\*\* (2007a) study.

15  
 16 Figure 9.22: urinary tract infections

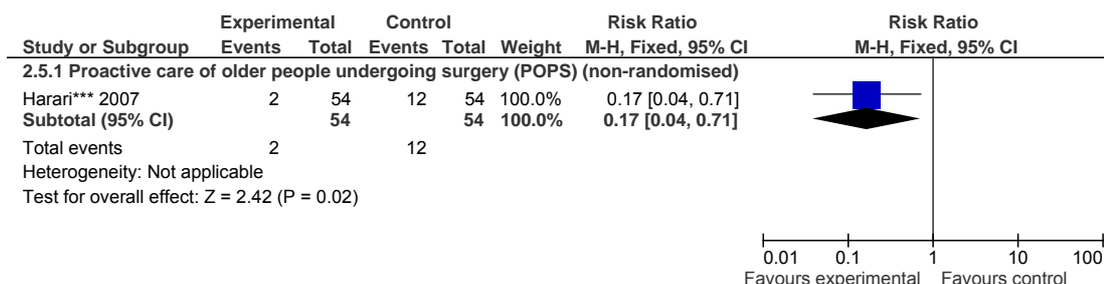


### 1 **Wound infection (figure 9.23)**

2 One study (Harari\*\*\* 2007a) reported the number of patients with wound  
3 infections. There was a clinically significant difference but there was imprecision  
4 in this small study.

5

6 Figure 9.23: wound infections



7

8 NB scale 0.01 to 100

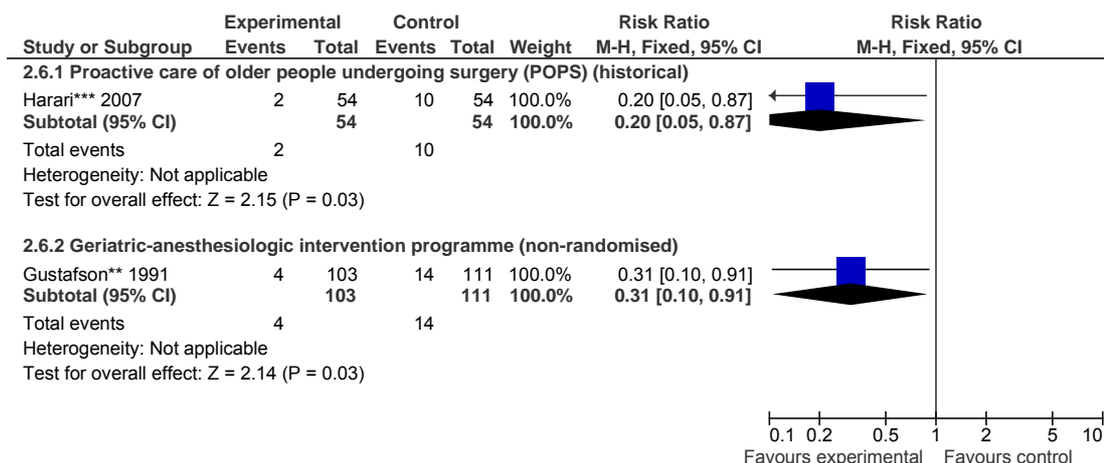
9

### 10 **9.17.1.13 Pressure ulcers (figure 9.24)**

11 Two non-randomised studies (Gustafson\*\* 1991; Harari\*\*\* 2007a) reported the  
12 number of people with pressure ulcers. There was a significant difference  
13 between interventions in both studies, but the results are imprecise.

14

15 Figure 9.24: pressure ulcers



16

### 17 **9.17.1.14 Sensory impairment**

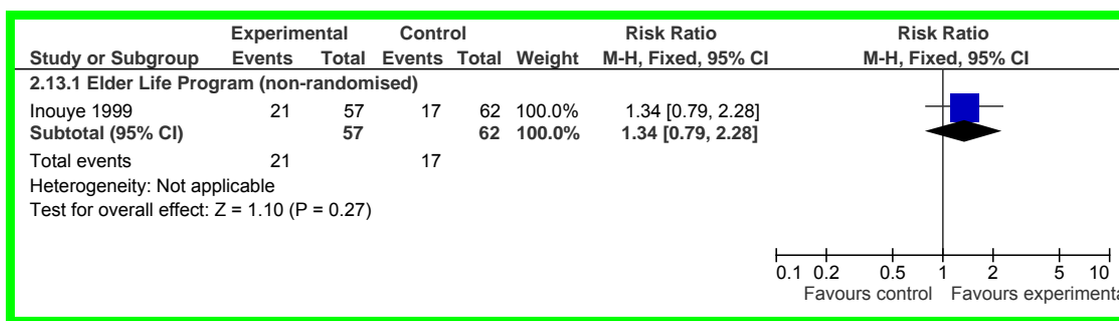
#### 18 **Visual impairment**

19 The Inouye\* (1999) study reported the number of patients with early vision  
20 correction at reassessment (day 5 or at discharge if earlier). There was no

1 significant difference between interventions (figure 9.25); RR 1.34 (95%CI 0.79  
 2 to 2.28), but the results are imprecise. We note that only patients who had a  
 3 visual acuity of less than 20/70 on binocular near vision testing received the  
 4 vision protocol; results were only reported for 119/852 patients.

5

6 Figure 9.25: early vision correction at reassessment (day 5 or at discharge if  
 7 earlier)

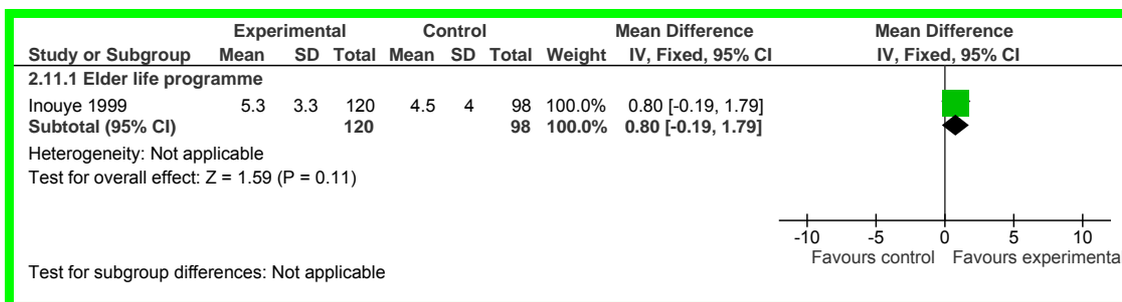


8  
9

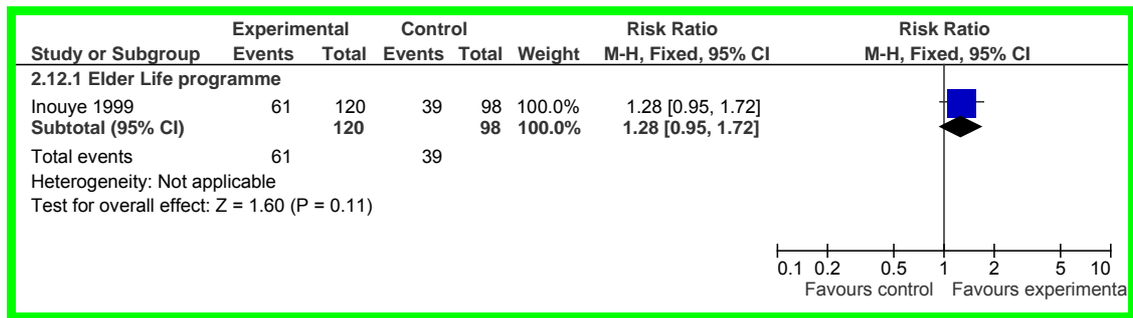
10 **Hearing impairment**

11 The Inouye\* (1999) study reported an adjusted Whisper test score at  
 12 reassessment (day 5 or at discharge if earlier); adjustment was for the patients'  
 13 baseline score. Although the study reported that standard deviations were  
 14 given, this did not agree with the p value reported and it was assumed that the  
 15 SDs were standard errors. Accordingly we recalculated standard deviations.  
 16 There was no significant difference between interventions (figure 9.26); MD 0.80  
 17 (95%CI -0.19, 1.79) on a scale of 0 to 12 (good hearing). There was no  
 18 significant difference in the number whose score improved by 1 point (figure  
 19 9.27). We note that only patients who had a Whisper test score below 7  
 20 received the protocol once daily; results were only reported for 218/852  
 21 patients.

22 Figure 9.26: whisper test



24  
25  
26  
27 **Figure 9.27: whisper test – number of patients with improvement by one**  
 28 **point**



1  
2  
3

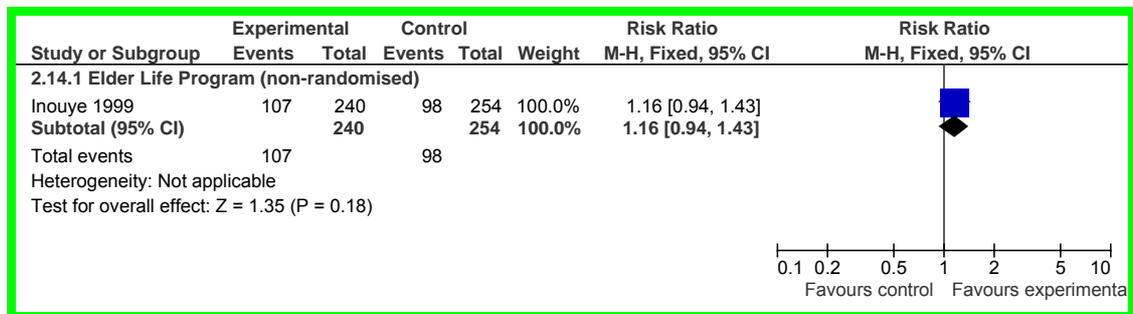
#### 4 9.17.1.15 Dehydration

5 Two non-randomised studies reported on dehydration (Harari\*\*\* 2007; Inouye\*  
6 1999).

7 The Inouye\* (1999) study reported the number of patients assessed to be  
8 improved by 5 points for the adjusted ratio of blood urea nitrogen to creatinine  
9 at reassessment; adjustment was for the patients' baseline score. There was no  
10 significant difference in the number who were assessed to be improved (figure  
11 9.28) although the results are imprecise. We note that only patients who had a  
12 ratio of blood urea nitrogen to creatinine of at least 18 received the protocol;  
13 results were only reported for 494/852 patients.

14  
15

Figure 9.28: number of patients with improvement in dehydration

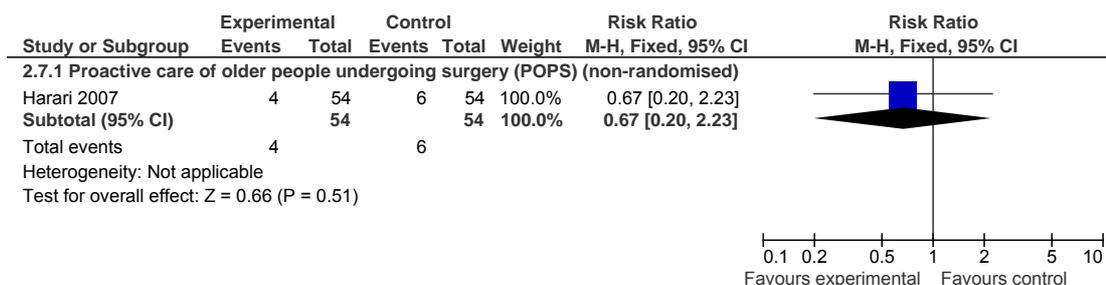


16  
17  
18

19 The Harari\*\*\* (2007a) study reported the number of patients with dehydration  
20 (figure 9.29); the CI was very wide and consistent with both important benefits  
21 and important harms.

22

23 Figure 9.29: number of patients with dehydration



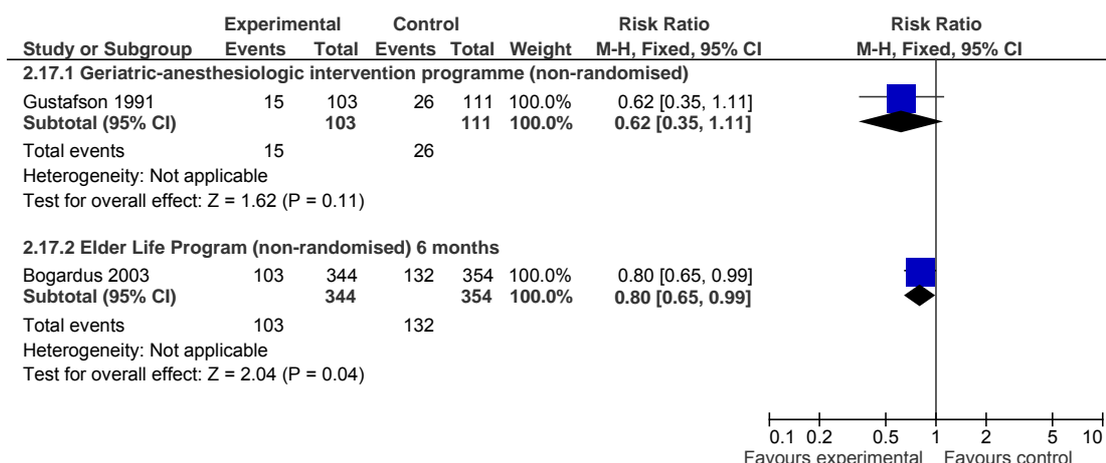
1  
2

3 **9.17.1.16 Urinary incontinence (Figure 9.30)**

4 Two studies investigated urinary incontinence (Gustafson\*\* 1991; Bogardus\*  
5 2003/Inouye\* 1999). There was no significant difference between the  
6 intervention and control studies in the number of patients with urinary infections in  
7 Gustafson\*\* 1991, but the 6 months follow up of the Inouye\* (1999) study  
8 showed a significant difference in the number of people with incontinence  
9 compared with the usual care group. Both studies showed imprecision.

10  
11

Figure 9.30: urinary incontinence



12  
13

14 **9.17.1.17 Adherence**

15 One study (Inouye\* 1999) reported the overall rate of adherence to all  
16 protocols (87%) and the rate of adherence to individual protocols: orientation  
17 96%; vision 92%; hearing 92%; therapeutic activities 86%; early mobilisation  
18 84%; volume repletion 81% and non-pharmacological sleep 71%. No adverse  
19 effects were associated with the intervention protocols. The Marcantonio (2001)  
20 study reported an overall adherence to recommendations of 77%, and the  
21 Wong\*\* (2005) study reported 90%.

22

23 **9.18 Clinical evidence statements**

24 There is **low quality evidence** to show the following results for a multicomponent  
25 intervention based on targeting 6 modifiable risk factors (cognitive impairment,

1 sleep deprivation, immobility, vision impairment, hearing impairment,  
 2 dehydration), with training (Inouye 1999) in patients at high or intermediate risk  
 3 of delirium:

- 4 • A significant reduction in the incidence of delirium; RR 0.66 (95%CI 0.46 to  
 5 0.95)
- 6 • A significant reduction in the total number of days of delirium amongst all  
 7 patients in the group (105 versus 161 days)
- 8 • A significant difference in the number with urinary incontinence after 6  
 9 months follow up; RR 0.80 (95%CI 0.65 to 0.99)
- 10
- 11 • No significant difference in:
  - 12 ○ the incidence of delirium after 6 months follow up; the evidence  
 13 was **very low quality** for this outcome
  - 14 ○ the MMSE score after 6 months follow up
  - 15 ○ delirium severity
  - 16 ○ the median length of stay in hospital
  - 17 ○ the number of patients with a new long-term care placement
  - 18 ○ the number of patients who died, either during the hospitalisation  
 19 period or in the time between hospital admission and 6 months  
 20 follow up; the evidence for hospitalised patients was **very low**  
 21 **quality**
- 22

23 There is **low quality evidence** to show the following results for a multicomponent  
 24 intervention based on targeting 6 modifiable risk factors with training (Inouye  
 25 1999) in subgroups of patients who were targeted to receive the part of the  
 26 multicomponent intervention appropriate to that outcome (the proportion  
 27 receiving the targeted component is given in brackets)

- 28 • A significant increase in the number of patients with an improvement of 2  
 29 points on their MMSE score after 5 days or at discharge if earlier  
 30 (253/852)
- 31 • No significant difference in the number of patients:
  - 32 ○ with an improvement in activities of daily living (194/852)
  - 33 ○ with early vision correction at reassessment (day 5 or at  
 34 discharge if earlier) (119/852)
  - 35 ○ whose hearing improved at reassessment (day 5 or at discharge  
 36 if earlier) (218/852)
  - 37 ○ whose dehydration improved at reassessment (day 5 or at  
 38 discharge if earlier) (494/852)
- 39

1 There is **moderate quality evidence** to show the following results in patients  
2 undergoing surgery for hip fracture (i.e. higher risk), and receiving a  
3 multicomponent intervention based on targeting 7 modifiable risk factors  
4 (orientation, dehydration, sensory impairment, immobility, environmental  
5 modifications and medication management) following consultation with a  
6 geriatrician preoperatively (Marcantonio 2001) showed the following results:

- 7 • A borderline significant reduction in the incidence of delirium; RR 0.65  
8 (95%CI 0.42 to 1.00)
- 9 • No significant difference in the:
  - 10 ○ mean duration of delirium per episode; this is an indirect outcome  
11 measure
  - 12 ○ median length of stay in hospital
  - 13 ○ number of patients discharged to long-term care (it was unclear if  
14 this was a new placement)

15  
16 There was **very low quality evidence** for the effectiveness of an intervention  
17 consisting of an education programme for staff and reorganisation of nursing  
18 and medical care, such that the patients received patient centred care, rather  
19 than task allocated care. Results for this study (Lundström 2005) showed:

- 20 • A significant reduction in the:
  - 21 ○ incidence of delirium; RR 0.51 (95%CI 0.31 to 0.86); this was  
22 low quality evidence because the outcome assessors were blinded
  - 23 ○ mean length of stay in hospital, although the data were skewed

24  
25 The remaining evidence is from studies with a poor quality study design  
26 (Gustafson 1991, Harari 2007a, Wanich 1992, Wong 2005) or from a low  
27 quality RCT that did not record the incidence of delirium as an outcome measure  
28 (Landefeld 1995).

29 For the outcome, incidence of delirium, there is very low quality evidence to  
30 suggest that the following interventions may have potential to reduce the  
31 incidence of delirium in hospital patients:

- 32 • Multidisciplinary team, pre- and post-operative assessment and targeting  
33 of identified issues including pain management, early mobilisation,  
34 nutrition, and early detection and treatment of medical complications  
35 (Harari 2007a). There is much uncertainty around this result
- 36 • Geriatric-anaesthesiologic intervention programme, including pre- and  
37 postoperative assessment by specialist in geriatric and internal medicine  
38 (Gustafson 1991)
- 39 • Plan-do-study-act programme, including staff education and geriatric team  
40 assessments to address 12 modifiable risk factors (Wong 2005)

41

1 There is very low quality evidence to suggest that the following intervention did  
2 not have a significant effect on the incidence of delirium: education of staff and  
3 assessment by geriatricians to address 6 modifiable risk factors (Wanich 1992)  
4

5

## 6 **9.19 Health economic evidence**

### 7 **9.19.1 Multi-component interventions for the prevention of delirium in** 8 **a hospital setting**

9

10 One economic evaluation study was included as evidence (Rizzo 2001). This was  
11 a non-randomised study of 70 year old patients with no evidence of delirium but  
12 who had intermediate or high risk of delirium. It was conducted in the USA in  
13 2001 with the following objectives:

- 14 • to determine the impact of the multi-component intervention strategy on  
15 total hospital costs, average daily costs, and length of stay,
- 16 • to estimate the impact of the multi-component intervention on specific  
17 hospital cost components,
- 18 • to describe the intervention costs associated with the intervention strategy,  
19 and
- 20 • to combine the results of cost and effectiveness analyses to assess the cost-  
21 effectiveness of the intervention strategy.

22

23 Patients in the intervention group were those who met the inclusion criteria of  
24 being 70 years and older with no evidence of delirium but had intermediate or  
25 high risk of delirium. Control patients were prospectively selected and matched  
26 on age, gender, and baseline delirium risk. The intervention group received the  
27 multi-component intervention (Hospital Elder Life Program) strategy which  
28 consisted of interventions targeted toward six delirium risk factors (cognitive  
29 impairment, sleep deprivation, immobility, visual impairment, hearing impairment  
30 and dehydration). The core interventions included orienting communication,  
31 therapeutic activities, sleep enhancement strategies, exercise and mobilisation,  
32 provision of vision and hearing aids, and oral volume repletion for dehydration.  
33 Others included geriatric nursing assessment and interdisciplinary rounds. The  
34 control arm did receive usual hospital care.

35 The cost of the intervention was based on personnel and equipment costs during  
36 the three year study period. The total personnel and equipment costs over this  
37 period were \$252,885 and \$257,385 respectively. The non-intervention costs in  
38 the intervention and usual care groups were reported as \$6,484 and \$7,300  
39 respectively. The additional cost of the intervention was \$592 per patient  
40 (standard error, se=21). Unit cost and resources use were reported and the  
41 perspective of the analysis was third party (hospital healthcare provider). The

1 multi-component intervention was estimated to result in cost savings (excluding  
2 intervention costs) of \$831 for intermediate risk patients after multivariate  
3 adjustment for confounding variables but there was no significant difference for  
4 the high risk group. In the intermediate delirium risk patients the net cost saving  
5 attributable to the intervention was \$99 if intervention costs were included. This  
6 was statistically insignificant after multivariate adjustment. The intervention had a  
7 statistically significant cost increase of \$1,308 in high risk patients.

8 The overall incidence of delirium was 9.9% and 15.0% in the intervention and  
9 control groups respectively. The incidence of delirium in the intermediate risk  
10 group was 6.5% with intervention and 11.7% without intervention. In the high risk  
11 group, it was 18.5% and 23.5% respectively. Incidence of delirium was based  
12 on CAM, MMSE and digital span test. A mortality rate of 1% and 2% were  
13 reported in the respective groups. Costs were not assessed from a UK NHS and  
14 PSS perspective. The measure of health benefit from the intervention was not in  
15 QALY units. The results of this study were judged to be not applicable to the  
16 guideline population.

17

## 18 **9.20 Health economic evidence statements**

19

20 The results of the economic model (chapter 16) showed the following:

- 21 • The use of two multi-component targeted interventions was cost effective in:
  - 22 ○ elderly patients at intermediate or high risk of delirium and who
  - 23 ○ were admitted to the general medicine service.
  - 24 ○ elderly patients who were admitted emergently for surgical
  - 25 ○ repair of hip fracture.
- 26 • These findings were robust as the interventions remained cost-effective
- 27 ○ after a series of sensitivity analyses were conducted.

28

29

30

1

2

Table 9.1: multicomponent interventions for the prevention of delirium

Study	Multi-disciplinary team	Education intervention	Care methods	assessment of patients	orientation	Dehydration nutrition	Sleep	Sensory impairment improvement	Early mobilisation	Environmental modifications	Medication management	Pain management	Other
Lundström (2005)	No; mainly nursing care	staff education on Ass; PTD: NPI; Med. Monthly guidance for staff	Patient-allocation care , with individualised care	yes: via education	only via education	No	No	No	No	No	only via education	no	No
Inouye (1999): Elder Life Program	Yes: N, Physio, G, TRS, V	yes: trained team; performance evaluated quarterly	not changed	Yes in order to determine risk factors addressed	Yes: schedule / name board; reorienting communication	Yes for those with dehydration: early recognition of dehydration and volume repletion	Yes: non-pharmacological sleep protocol; sleep-enhancement protocol	Yes for visually impaired and hearing impaired people	yes	yes: unit-wide noise reduction strategies	No	no	cognitively stimulating activities (e.g. discussion of current events)
Gustafson (1991): Geriatric-anesthesiologic intervention programme	Yes; nursing, anaesthetist and geriatrician care	No	not changed; task oriented	pre- and postop by geriatrician	No	No	No	No	No	No	individualised thrombosis prophylaxis	no	O2 therapy from admission; phenylephrine for low systolic bp; surgical policy
Harari 2007a: Proactive care of older people undergoing surgery (POPS)	Yes: N, Physio, G, OT, SW	Yes: patients preop (N, Ex, RT, PM); staff postop (TMC, EM, PM, BBF, N, DP)	no change	preop planning and postop review by geriatrician and nurse; targetting issues identified	No	Yes: nutrition	No	No	Yes	No	early detection and treatment of medical complications	Yes	discharge planning

Study	Multi-disciplinary team	Education intervention	Care methods	assessment of patients	orientation	Dehydration nutrition	Sleep	Sensory impairment improvement	Early mobilisation	Environmental modifications	Medication management	Pain management	Other
Landefeld (1995); Acute Care for Elders programme	yes: daily visits (D, N, SW, Diet, Physio, VNL)	No	patient centred care	Yes: daily assessment by nurses of physical, cognitive and psychosocial function; daily review of medical care	Yes: large clock, calendar	yes nutrition (no details)	yes (no details)	No	yes (no details)	Yes: specially designed environmt (carpeting, handrails, uncluttered hallways, elevated toilet seats, door levers)	yes: minimise medications (e.g. sedative-hypnotic agents)	no	minimise effects of procedures (e.g. catheterisation); discharge planning
Wannich (1992):	yes: for discharge planning (N, Physio, OT, SW, Diet)	Yes: staff (Ass, SI, Mob, En); families (RC, O, En)	Not stated	Yes: assessment and management plans recorded on charts and shared with staff and families	Yes (e.g. day of week, current events, updated calendars in every room)	No	No	Yes for visually impaired and hearing impaired people only (glasses and hearing aid + encouragement to use them)	yes	Yes: lighting to decrease sensory deprivation; night lights	assess medics contributing to delerium, e.g. neuroleptics, antidepressants, narcotic analgesics, sedative-hypnotics, and use discouraged	no	discharge planning; Communication: clear and slow, with repetition
Wong (2005)	Yes: project team supervised programme (N, G, Ph, D, QI, A, Diet)	Yes: staff on PTD, POD, Ass, MMD	not changed	Yes for identification of needs	Yes: clock, calendar	Yes: nutrition (including properly fitting dentures); maintenance of fluid/electrolyte imbalance	No	Yes: sensory stimuli - glasses, hearing aid	yes	Yes: soft lighting, not putting delirious patients in same room	treatment of major complications; stop unnecess benzodiazepines, antihistamines, anticholinergics	Yes	regulation of bladder / bowel function; O2; tmt of agitated delirium
Marcantonio (2001): Proactive geriatrics consultation	No consultation with geriatrician	no	Not stated	Yes: consultation with geriatrician preop / within 24 h postop. Geriatrician daily visits during hospitalisation => target recs made	Yes: clock, calendar	Yes: nutrition (including properly fitting dentures); maintenance of fluid/electrolyte imbalance; treat dehydration/ overload	No	Yes: sensory stimuli - glasses, hearing aid	yes	Yes: soft lighting, use of radio/tape recorder - not rec for any patient though	treatment of major complications; stop benzodiazepines, antihistamines, anticholinergics; eliminate medicn redundancies; tmt to raise bp	Yes	regulation of bladder / bowel function; O2; tmt of agitated delirium

Key: N = nurses; Physio = physiotherapists; OT = occupational therapists; D = doctor (generally); G = geriatrician; SW = social worker; TRS = therapeutic recreation specialist; V = volunteer; A = anaesthetist; QI = member of the quality improvement unit; Ph = pharmacist, Diet = dietitian / nutritionalist; VNL = visiting nurse liaison; Ass = assessment; PTD = prevention and treatment of delirium; CD = training on cognitive impairment; POD = prevalence and outcome of delirium; NPI = nurse patient interaction; N = nutrition; MMD = medication management of delirium; Ex = exercise; RT = relaxation therapy; PM = pain management; TMC = treatment of medical complications; EM = early mobilisation; PM = pain management; BBF = bowel bladder function; DP = discharge planning).

# 1 10 Pharmacological prevention

## 2 10.1 Clinical introduction

3 The serious nature of delirium and its consequences makes all methods of  
4 prevention important to establish. Pharmacological agents are a recognised  
5 cause of delirium and so the use of these agents for prevention needs to be  
6 approached cautiously. Antipsychotic, benzodiazepines, acetylcholinesterase  
7 inhibitor classes of drugs in particular, and products that influence the immune  
8 system, may prove useful, based on early evidence from small studies, or from a  
9 theoretical perspective.

10  
11 People at risk of delirium are already vulnerable to the adverse effects of  
12 pharmacological products. It will be essential to establish the efficacy and risks  
13 of preventative drug treatment from well conducted clinical trials before they  
14 might be considered for routine use in clinical practice.

15

## 16 10 A) Prevention in hospital

17

### 18 10.2 Description of studies

19 Ten papers were evaluated for inclusion. Two studies were excluded because  
20 there were fewer than 20 patients in one the comparison groups (Sampson  
21 2007; Dautzenberg 2004). Reasons for exclusion are reported in Appendix G.  
22 Two Cochrane Reviews were identified (Lonergan 2007; Siddiqi 2007) and  
23 updated. Six RCTs (Aizawa 2002; Gamberini 2009; Kalisvaart 2005; Kaneko  
24 1999; Liptzin 2005; Prakanrattana 2007) were included.

25

#### 26 10.2.1 Study Design

27 Two studies reported receiving funding from the pharmaceutical industry (Liptzin  
28 2005; Gamberini 2009), one study (Prakanrattana 2007) reported the study  
29 was supported by a hospital research grant that does not appear to be  
30 associated with industry, one study reported no funding was received (Kalisvaart  
31 2005), and two did not state if the study was funded (Aizawa 2002; Kaneko  
32 1999).

33

34 None of the studies were conducted in the UK. One study (Liptzin 2005) was  
35 conducted in the USA, one in The Netherlands (Kalisvaart 2005), one in

1 Switzerland (Gamberini 2009), two in Japan (Aizawa 2002; Kaneko 1999) and  
2 one in Thailand (Prakanrattana 2007).

3 Study duration was reported in four studies (Aizawa 2002: 7 days; Gamberini  
4 2009: 6 days postoperatively; Kalisvaart 2005: varied to a maximum of six  
5 days depending on the onset of delirium; Liptzin 2005: 28 days).

6  
7

## 8 10.2.2 Population

9 One study included fewer than 50 patients (Aizawa 2002: n=42), two studies  
10 included fewer than 100 patients (Kaneko 1999: n=80; Liptzin 2005: n=90);  
11 two studies included 100 or more patients (Gamberini 2009: n=120;  
12 Prakanrattana 2007: n=129) and one study was larger, and included 430  
13 patients (Kalisvaart 2005).

14 All of the studies were conducted in hospital settings in patients undergoing  
15 surgery. The type of surgery included resection for gastric or colorectal cancer  
16 (Aizawa 2002); hip surgery for acute fractures or hip replacements (Kalisvaart  
17 2005); gastrointestinal surgery (Kaneko 1999); total joint replacement surgery  
18 of the knee or hip (Liptzin 2005); cardiac surgery with cardiopulmonary bypass  
19 (Prakanrattana 2007), cardiac surgery (Gamberini 2009). The Kaneko (1999)  
20 study reported that all patients were admitted to an ICU before the scheduled  
21 surgery.

22 The age range across the studies was 51 years to 90 years. All studies included  
23 men and women. The patients' ethnicity was described as being 95% white and  
24 5% other in one study (Liptzin 2005) and was not reported in the remaining  
25 studies.

26 Cognitive status was not reported in two studies (Aizawa 2002; Prakanrattana  
27 2007), one study (Liptzin 2005) reported that at baseline patients did not have  
28 dementia, and one study (Gamberini 2009) reported that patients with an  
29 MMSE score of less than 15 were excluded. Three studies reported that the  
30 method used to assess dementia was the Mini Mental State Examination (MMSE)  
31 (Gamberini 2009; Kalisvaart 2005; Liptzin 2005). The reported MMSE scores  
32 indicated that at least some of the patients had dementia. One study did not  
33 report the method used for the assessment of dementia (Kaneko (1999).

34 One study reported the risk of postoperative delirium (Kalisvaart 2005). In this  
35 study, 84% of the patients had an intermediate risk for postoperative delirium  
36 and 16% had a high risk for postoperative delirium (as based on four predictive  
37 risk factors not specifically described); low risk patients were excluded. Patients  
38 with delirium at hospital admission were excluded from the study.

39 The Kalisvaart (2005) study also described their patients as having light  
40 dehydration.

41

1 **10.2.3 Interventions**

2

3 **10.2.3.1 Acetylcholinesterase**

4 One study (Liptzin 2005) investigated the acetylcholinesterase inhibitor,  
5 donepezil.

- 6 • 5–10 mg donepezil per day.

7 One study (Gamberini 2009) investigated the acetylcholinesterase inhibitor,  
8 rivastigmine

- 9 • 1.5 mg oral rivastigmine three times per day every 8 hours, starting on the  
10 evening preceding surgery and continuing until the sixth postoperative  
11 day; each patient received 22 doses in total.

12

13 **10.2.3.2 Atypical antipsychotics**

14 One study (Prakanrattana 2007) investigated the atypical antipsychotic,  
15 risperidone.

- 16 • 1 mg (orally disintegrating tablet) sublingually as a one-off dose when  
17 patients started to wake up in the ICU.

18

19 **10.2.3.3 Typical antipsychotics**

20 Two studies (Kalisvaart 2005; Kaneko 1999) investigated the typical  
21 antipsychotic drug haloperidol. The interventions included:

- 22 • 1.5 mg haloperidol tablet three times per day, starting on hospital  
23 admission and continued until 3 days after surgery; a maximum delay  
24 from admission of 72 hours was permitted before surgery (Kalisvaart  
25 2005)

- 26 • 5 mg intravenous haloperidol once per day, starting on the first  
27 postoperative day (Kaneko 1999)

28

29 **10.2.3.4 Benzodiazepines**

30 One study (Aizawa 2002) investigated the use of a 'Delirium Free Protocol  
31 (DFP)' which was designed to address the risk factor of insomnia. The DFP  
32 included:

- 33 • a combination of two benzodiazepines with pethidine: (diazepam 0.1  
34 mg/kg per day intramuscularly given at 20.00h and a drip infusion of  
35 flunitrazepam 0.04 mg/kg) and pethidine 1 mg/kg (both given from

1 20.00 to 04.00h), for the first 3 days postoperatively, starting on the  
2 day of the operation.

- 3 • The GDG expressed concern that the method of delivery of the drug (IM  
4 diazepam), and the addition of pethidine made the effect of  
5 benzodiazepines unclear, the study was addressing symptoms of  
6 improving insomnia, which in turn is a risk factor for delirium; this study  
7 was therefore not considered further.

8

#### 9 **10.2.4 Comparisons**

10 The following comparisons were carried out:

##### 11 **10.2.4.1 Acetylcholinesterase inhibitors**

- 12 • Donepezil versus placebo (Liptzin 2005)
- 13 ○ The intervention was given for 14 days preoperatively and a  
14 further 14 days postoperatively; patients were not admitted to  
15 hospital until the day before surgery.
- 16 ○ The control group received placebo once a day at breakfast,  
17 and again, where symptoms of delirium were experienced, the  
18 placebo dose was doubled.
- 19 • Rivastigmine versus placebo (Gamberini 2009)
- 20 ○ The intervention was given the evening before surgery, three  
21 times per day every 8 hours thereafter until the evening of the  
22 sixth postoperative day.
- 23 ○ The control group was administered the placebo (liquid identical  
24 to rivastigmine solution) following the same dosing scheme.
- 25 ○ If postoperative delirium occurred. patients received haloperidol  
26 (starting with 0.5 mg every 6 to 8h) and lorazepam (1 mg per  
27 day)

28

##### 29 **10.2.4.2 Atypical antipsychotics**

- 30 • Risperidone (orally disintegrating tablet) versus placebo (an antiseptic strip  
31 applied sublingually). The interventions were a one-off dose.  
32 (Prakanrattana 2007)

33

##### 34 **10.2.4.3 Typical antipsychotics**

- 35 • Haloperidol versus placebo
- 36 ○ 1.5 mg haloperidol tablet three times per day, up to 6 days pre  
37 and postoperatively (Kalisvaart 2005)
- 38 - all patients received a proactive geriatric consultation  
39 (geriatric medical attention; enhancement of orientation)

1 and cognition; sensory and mobility improving advice;  
2 attention to pain and sleeping problems; extra attention to  
3 food and fluid intake; patient, family and nursing staff  
4 education). This study also gave the patients haloperidol  
5 and/or lorazepam 3 times a day if postoperative delirium  
6 occurred.

- 7 ○ 5 mg intravenous haloperidol once per day, 5 day intervention  
8 period postoperatively (Kaneko 1999)

9  
10 Concurrent medications were not reported in three studies (Liptzin 2005;  
11 Kalisvaart 2005; Kaneko 1999). Comorbidities were not reported in three  
12 studies (Kalisvaart 2005; Kaneko 1999; Liptzin 2005). One study (Prakanrattana  
13 2007) reported that 67% of the patients were suffering from coexisting diseases  
14 including hypertension, diabetes mellitus, cerebrovascular accident, renal failure,  
15 or atrial fibrillation and another study (Gamberini 2009) reported that patients  
16 had arterial hypertension (78%) and were being treated for diabetes mellitus  
17 (7%) and for chronic pulmonary obstructive disease (4%).

### 18 19 **10.3 Methodological quality**

20 The Liptzin (2005) study reported that initially 1038 patients were contacted  
21 and 732 were not followed up or refused to participate. The remaining 306  
22 were contacted 2–3 weeks before surgery and underwent screening. From these,  
23 90 patients were randomised, although 10 were not operated on and the results  
24 are based upon 80 patients. The study reported there were no significant  
25 differences between the randomized patients and the non participants, in  
26 relation to age, gender, ethnicity, and site of operation (knee or hip joint  
27 surgery).

28 The method of sequence generation was adequate in three studies (computer  
29 generated blocks of 20: Gamberini 2009; computer-generated sequence:  
30 Kalisvaart 2005; Prakanrattana 2007). Sequence generation was not reported  
31 in two studies (Kaneko 1999; Liptzin 2005).

32 Allocation concealment was partially met in all of the studies. Gamberini (2009)  
33 reported that optically identical solutions in identical bottles were delivered by  
34 the hospital pharmacy, labelled with a number. Kalisvaart (2005) used identical  
35 containers prepackaged by a hospital pharmacist, which were sequentially  
36 assigned; Kaneko (1999) used sealed envelopes. In the Liptzin (2005) study the  
37 patients were randomised by the research pharmacist, but no further details  
38 were given, and in the Prakanrattana (2007) study, a concealed envelope was  
39 used.

40 Four studies (Gamberini 2009; Kalisvaart 2005; Liptzin 2005; Prakanrattana  
41 2007) were described as double-blind (Kalisvaart 2005: blinding was checked  
42 by interviewing the study assessors). Although in the Prakanrattana (2007) study  
43 the patients' placebo was an antiseptic strip rather than tablet, the authors

1 stated that the assessors were blind to treatment. The Kaneko (1999) study did  
2 not report on blinding, although a placebo was used.

3 An *a priori* sample size calculation was reported in three studies (Kalisvaart  
4 2005; Liptzin 2005; Prakanrattana 2007). The Gamberini (2009) study  
5 reported that a sample size of 120 was required to detect a relative risk  
6 reduction of 50%, with 80% power at a 5% significance level. One study  
7 (Kalisvaart 2005) reported a sample size of 206 patients per group was  
8 required to detect a 13% decrease in risk with 80% power at a 5% significance  
9 level. The sample sizes included in this study (n= 430), slightly exceeded this  
10 sample size estimate. The Liptzin (2005) study reported that a sample of 80 was  
11 required to have an 80% power to detect a difference of 22% in the study  
12 groups at a one-sided significance level of 5% assuming a delirium rate of 44%  
13 in the placebo group. Another study (Prakanrattana 2007) required a sample  
14 size of 63 per group to detect a 30% reduction in risk with 90% power at a 5%  
15 significance level; 63 patients per group were recruited and completed the  
16 study.

17 All studies demonstrated baseline comparability.

18 The Kalisvaart (2005) study reported no significant differences in mean age,  
19 proportion of males to females, mini-mental examination scores, visual acuity,  
20 health scores, geriatric depression scores, Barthel Index, or baseline risk of  
21 delirium between treatment and control groups. The Kaneko (1999) study  
22 reported no differences in the proportion of males to females by group, pre-  
23 existing diseases, preoperative medicines, duration of operation and anesthesia.  
24 They did observe that fewer patients in the haloperidol group had premorbid  
25 cognitive impairment (5% versus 10% in the placebo group), but the difference  
26 was not statistically significant. In the Liptzin (2005) study patients were  
27 comparable at baseline for age, gender, ethnicity, the surgeon who operated,  
28 the joint operated on and the MMSE questionnaire and clock-drawing test scores.  
29 The Prakanrattana (2007), study demonstrated baseline comparability between  
30 intervention groups for age, proportion of males to females, weight, New York  
31 Heart Association functional class, coexisting disease, type of operation  
32 (coronary artery bypass graft, valve or others), anaesthesia time,  
33 cardiopulmonary bypass time, and aortic cross-clamp time. In the Gamberini  
34 (2009) study patients were comparable for age, gender, baseline MMSE,  
35 baseline clock-drawing test scores, pre-existing diseases, type of operation  
36 (CABG, valve repair).

37 One study (Prakanrattana 2007) reported no missing participants; all patients  
38 were included in the analysis.

39 Three studies (Gamberini 2009; Kalisvaart 2005; Kaneko 1999) reported  
40 acceptable missing levels of data (that is less than 20%).

41 • The Gamberini (2009) study reported there was missing data for 25%  
42 (15/61) and 24% (14/59), in the intervention and control groups  
43 respectively. The study reported that only patients who were not  
44 assessed with CAM within 6 days after surgery (4/61: 3/59) were  
45 excluded from the analysis; however, the authors reported that an  
46 intention to treat analysis was carried out.

1           • In the Kaneko (1999) study 5% (2/40) in the intervention group and 0%  
2           in control group were missing, and the authors analysed all available  
3           participants in their analyses (n = 78).

4           • In the Kalisvaart (2005) study, 5% (11/212) were lost to follow-up in the  
5           treatment group and 11% (24/218) were lost to follow-up in the  
6           placebo group. However the authors analysed all patients who were  
7           randomised (ITT analysis).

8  
9           One study (Liptzin 2005) had inadequate levels of missing data (more than 20%  
10          missing data in each group). Originally 90 patients were included in the study,  
11          but ten patients were not included in the final analyses because they were not  
12          operated on, or took no further part in the analysis; the groups to which they  
13          were assigned were not reported. Of the remaining 80 patients, a further  
14          11/39 (28%) and 11/41 (27%) did not complete the study. A per protocol  
15          analysis was reported based on the 80 patients, although it was not clear what  
16          was assumed about the missing data.

17          Methods to assess concordance were partially reported in Kalisvaart (2005).  
18          They stated that clinical staff recorded the level of adherence to the intervention,  
19          but it was not stated how this was done. Concordance was determined by  
20          patients keeping records of their medication usage, and this was assessed by a  
21          research assistant (Liptzin 2005). Methods to assess concordance were not  
22          reported in the remaining studies.

23  
24          The method of delirium assessment was:

25           • **adequate** in three studies (Kalisvaart 2005; Liptzin 2005; Prakanrattana  
26           2007)

- 27           ○ One study used the DSM-IV criteria (Liptzin 2005)
- 28           ○ One study used the CAM and DSM-IV criteria (Kalisvaart 2005)
- 29           ○ One study used the CAM-ICU instrument (Prakanrattana (2007)

30

31           • **partially adequate** in one study (Gamberini 2009). The Gamberini (2009)  
32           study used the CAM instrument in both the surgical and ICU setting.

33

34          Method of delirium assessment was unclear in one study (Kaneko 1999). The  
35          DSM-IV and DSM III-R criteria were used for 'psychotic diagnoses' and also  
36          stated that delirium was 'clinically diagnosed'. Data were collected from the  
37          patients and nursing charts on the fifth day after surgery; it was not clear if the  
38          charts were used to record delirium.

39          One study (Kalisvaart 2005) assessed severity using the DRS-R-98 [range 0 (no  
40          severity) to high 45 (high severity)], MMSE, and the Digit Span test [assessment  
41          of attention, range 0 (no attention) to 42 (good attention)].

1  
2 All studies evaluated the incidence of delirium as a primary outcome. Secondary  
3 outcomes were: severity of delirium (Kalisvaart 2005), duration of delirium  
4 (Gamberini 2009; Kalisvaart 2005; Kaneko 1999; Liptzin 2005) and adverse  
5 events (Kalisvaart 2005; Kaneko 1999), length of hospital stay (Gamberini  
6 2009; Kalisvaart 2005; Liptzin 2005; Prakanrattana 2007), length of ICU stay  
7 (Gamberini 2009; Prakanrattana 2007), and sleep-wakefulness rhythm (Kaneko  
8 1999).

9  
10 Overall two studies were considered to have a higher risk of bias for the  
11 following reasons:

- 12 • The method of measurement of delirium was unclear (Kaneko (1999).
- 13 • Inadequate levels of missing data [over 20%] (Liptzin 2005)

14  
15 The use of rescue medication in the Kalisvaart (2005) study may have led to  
16 confounding for the following outcomes: duration of delirium, severity of delirium  
17 and length of stay.

## 18 19 **10.4 Results**

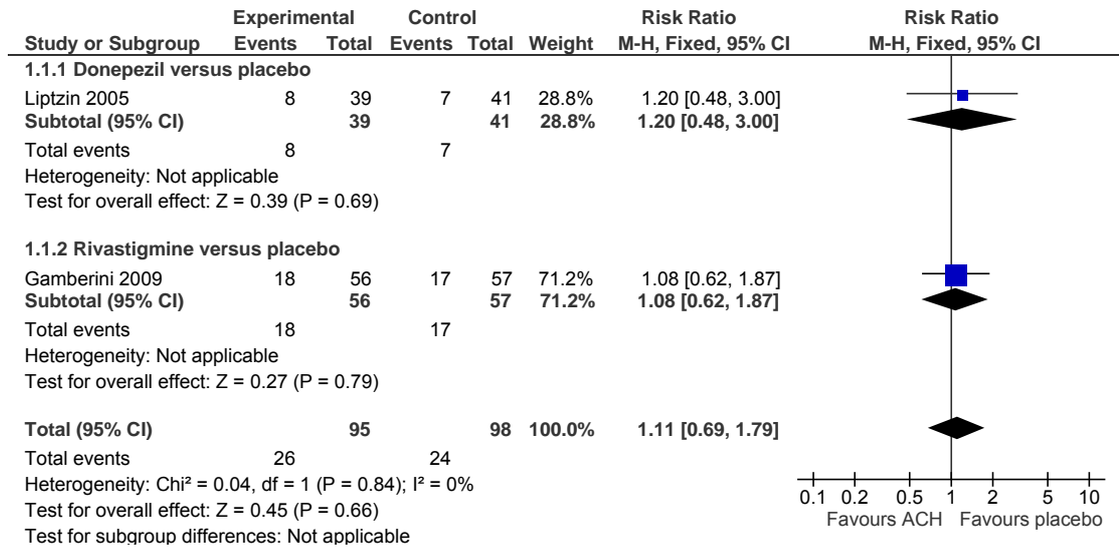
### 20 **10.4.1 Acetylcholinesterase inhibitor versus placebo**

#### 21 **10.4.1.1 Acetylcholinesterase inhibitor versus placebo**

##### 22 23 1. Incidence of postoperative delirium (endpoint 28 days)

24 Meta-analysis of two studies (Gamberini 2009; Liptzin 2005) with 193 patients,  
25 comparing acetylcholinesterase (ACH) with placebo showed no significant  
26 difference in the incidence of delirium between the groups (Figure 10.1); RR  
27 1.11 (95% CI 0.69 to 1.79); although the results are very imprecise.

1 Figure 10.1: number of patients with delirium



2

3

4 **2. Duration of postoperative delirium**

5 Two studies (Gamberini 2009; Liptzin 2005) reported the duration of  
6 postoperative delirium.

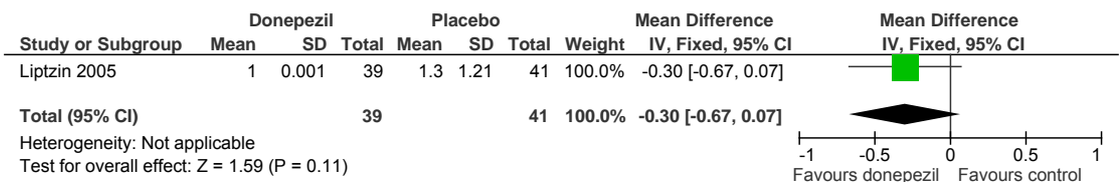
7 The Gamberini (2009) study compared rivastigmine versus placebo, in 113  
8 patients and reported there was no difference in the duration of delirium. The  
9 results from this study are not shown on the forest plot because study reported  
10 values for the median and range. The reported median and range were as  
11 follows: 2.5 days (range 1 to 5) and 2 days (range 1 to 6) for the rivastigmine  
12 and placebo groups respectively (reported p value= 0.3).

13 The remaining study (Liptzin 2005) comparing donepezil with placebo in 80  
14 patients found no significant difference in the duration of postoperative delirium  
15 (end point) (figure 10.2); mean difference (MD) -0.30 days (95%CI -0.67 to  
16 0.07), for a placebo group duration of 1.3 days; the results are imprecise. The  
17 standard deviation in the donepezil group was stated to be zero, but for the  
18 purposes of analysis this was assumed to be 0.001.

19

20

Figure 10.2: duration of delirium



21

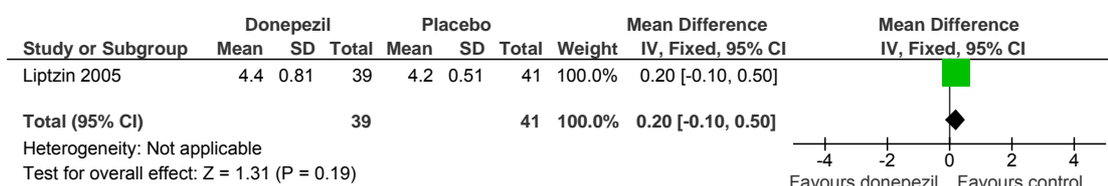
22

1 3. Length of hospital stay

2 Two studies (Gamberini 2009; Liptzin 2009) reported the length of stay. The  
 3 Gamberini (2009) study reported the median and range and the results for this  
 4 study are not shown on the forest plot. The (Gamberini 2009) study comparing  
 5 rivastigmine versus placebo in 113 patients reported there was no difference in  
 6 the length of hospital stay; the median and range was 13 days (range 7 to 39)  
 7 for both the rivastigmine and placebo groups respectively (reported p value =  
 8 0.3).

9 One study (Liptzin 2005) comparing donepezil with placebo in 80 patients  
 10 found no significant difference in the length of hospital stay(endpoint 28 days)  
 11 between the groups (figure 10.3); MD 0.20 days (95%CI -0.10 to 0.50). There  
 12 was imprecision because of the small sample size.

13  
 14 Figure 10.3: length of hospital stay



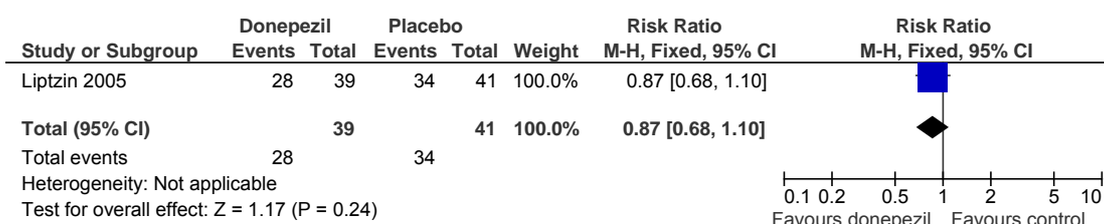
15  
 16  
 17 4. Length of ICU stay

18 One study (Gamberini 2009) comparing rivastigmine versus placebo in 113  
 19 patients reported there was no difference in the length of ICU stay; the median  
 20 and range were as follows: 2 days (range 2 to 7) and 2 days (range 2 to 6)  
 21 for the rivastigmine and placebo groups respectively (reported p value: 0.9).

22 5. Number of patients discharged to a rehabilitation facility (endpoint 28 days)

23 Analysis of one study comparing donepezil with placebo in 80 patients found no  
 24 significant difference between the groups for the number of patients discharged  
 25 to 'a rehabilitation facility', but it was not clear what this facility was (figure  
 26 10.4); RR 0.87 (95%CI 0.68 to 1.10). There was some imprecision in this  
 27 outcome.

28  
 29 Figure 10.4: discharge to rehabilitation facility



30  
 31  
 32  
 33 6. Use of rescue medications

1 The Gamberini (2009) study reported the use of haloperidol and lorazepam  
2 rescue medications. 32% and 30% of the patients receiving rivastigmine and  
3 placebo respectively were given haloperidol ( $p=0.9$ ). 61% and 68%, of the  
4 patients receiving rivastigmine and placebo, respectively were given lorazepam;  
5  $p=0.3$ ). There were no significant differences between the two groups in the  
6 number of patients who received the rescue medications.

7  
8  
9

## 10 10.4.2 Typical antipsychotics

### 11 10.4.2.1 Typical antipsychotics versus placebo

12

#### 13 1. Incidence of postoperative delirium

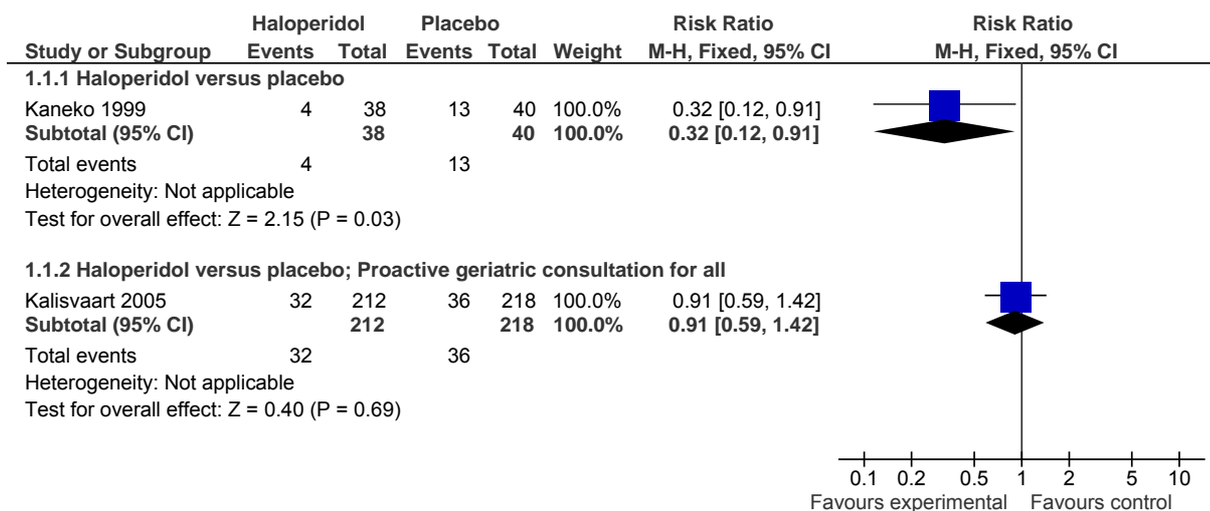
14 Two studies (Kaalisvaart 2005; Kaneko 1999) reported the use of haloperidol  
15 versus placebo on incidence of postoperative delirium. The Kaalisvaart (2005)  
16 study reported that all patients received a proactive geriatric consultation, thus  
17 the study was investigating the adjunctive effect of haloperidol. Therefore, these  
18 two studies are reported separately on the forest plots (figure 10.6)

- 19 • One study (Kalisvaart 2005) with 440 patients showed no significant  
20 difference in the incidence of postoperative delirium; RR 0.91 (95% CI  
21 0.59 to 1.42).
- 22 • The Kaneko (1999) study with 78 patients showed a small significant effect  
23 [0.32 (95% CI 0.12 to 0.91)]. We note this study was at higher risk of  
24 bias.

25

26 Figure 10.6: number of patients with postoperative delirium

27  
28



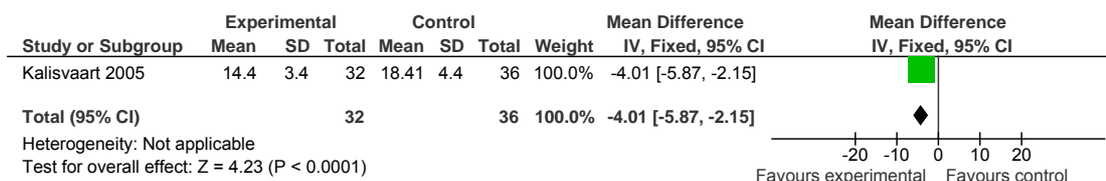
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7

**2. Severity of delirium**

Two studies (Kalisvaart 2005; Kaneko 1999) evaluated the severity of delirium, and only Kalisvaart (2005) presented data for analysis. In 78 patients who had delirium, Kalisvaart (2005) used the highest value obtained during delirium, on the DRS-R-98 scale, (maximum value on this scale is 39) to assess the severity of delirium. The analysis demonstrates a significant effect in favour of haloperidol: MD -4.01 (95% CI -5.87 to -2.15; figure 10.7). It is noted that the severity of delirium may have been confounded by the use of rescue medication.

The Kaneko (1999) study reported that the postoperative delirium was more severe in the placebo group (no data or statistical analyses were presented).

Figure 10.7: severity of delirium scores



**3. Duration of delirium**

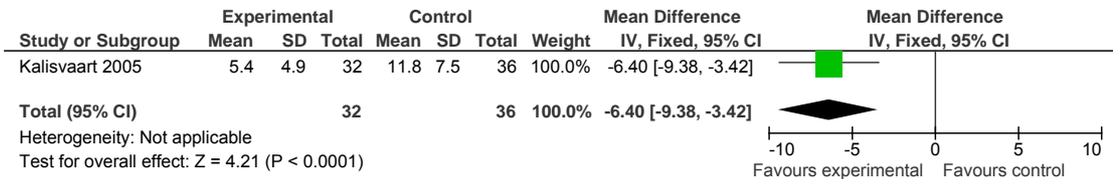
Two studies (Kalisvaart 2005; Kaneko 1999) evaluated the duration of delirium, and only Kalisvaart (2005) presented data for analysis. The analysis demonstrates that patients who received haloperidol, had, on average, significantly fewer days of delirium (of those who had delirium): MD -6.40 (95% CI -9.38 to -3.42; figure 10.8). It is noted that the duration of delirium may have been confounded by the use of rescue medication and that results were reported only for those with delirium. We also note that the distribution for the duration of delirium is skewed for both the intervention and placebo groups (mean values less than twice the standard deviation). The Kaneko (1999) study

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1 reported that the duration of postoperative delirium was longer in the placebo  
2 group (no data or statistical analyses were presented).

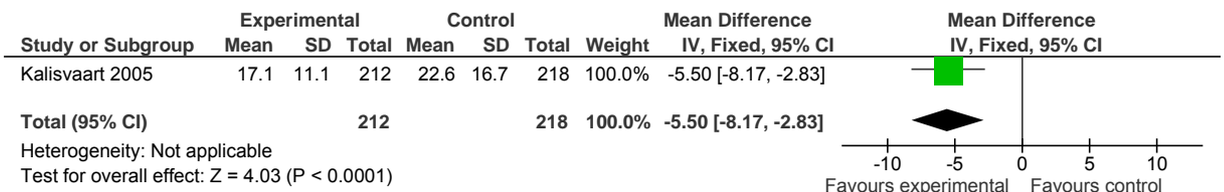
3  
4 Figure 10.8: duration of delirium



5  
6  
7 4. Length of hospital stay

8 The Kalisvaart (2005) study demonstrated that the number of days spent in  
9 hospital was significantly shorter in patients who received haloperidol compared  
10 to patients who received placebo in addition to the proactive geriatric  
11 consultation; MD  $-5.50$  ( $-8.17$  to  $-2.83$ ; figure 10.9). The study included the  
12 results for hospital length of stay in a table that was stated to apply to patients  
13 with delirium only. However, we have assumed this should refer to all patients;  
14 we also note that the summary statistics are incorrectly noted in the table in the  
15 report (the upper confidence limit is lower than the mean). Furthermore, the  
16 distribution for length of stay is skewed for both intervention and placebo  
17 groups.

18  
19 Figure 10.9: length of hospital stay

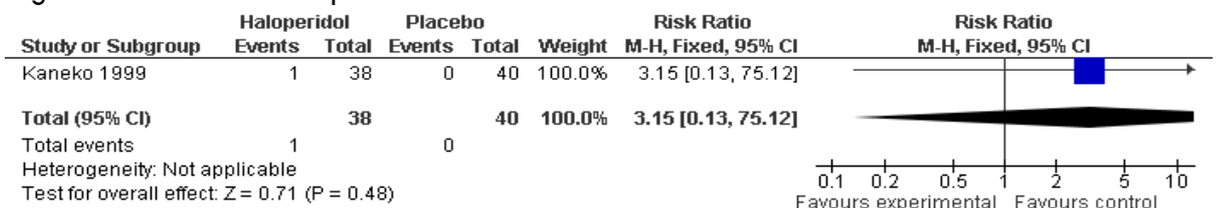


20  
21 NB: Scale -10 to 10

22 5. Adverse events

23 Two studies (Kalisvaart 2005; Kaneko 1999) evaluated adverse events.  
24 Kalisvaart (2005) reported that there were no drug-related side effects. Only  
25 Kaneko (1999) presented data for analyses; they observed that one patient in  
26 the treatment group developed transient tachycardia. The results are very  
27 imprecise (figure 10.10).

28  
29 Figure 10.10: number of patients with adverse events



1

2 **10.4.3 Atypical antipsychotics**

3 **10.4.3.1 Atypical antipsychotics versus placebo**

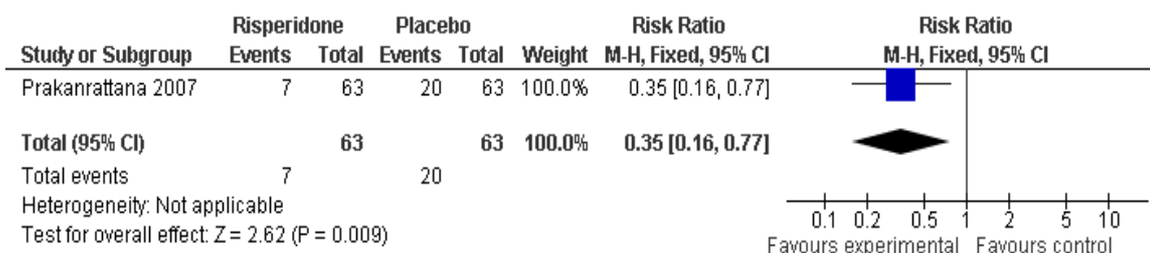
4 1. Incidence of delirium

5 In the Prakanratta (2007) study, delirium was recorded twice daily in the ICU  
6 and once daily on discharge from the ICU. The study reported results as  
7 percentages, so we calculated the number of patients with delirium.

8 In one study (Prakanrattana 2007) comparing risperidone with placebo in 126  
9 patients, there were significantly fewer patients with delirium in the risperidone  
10 group compared with placebo, although the result was imprecise (figure 10.11);  
11 RR 0.35 (95%CI 0.16 to 0.77) which corresponds to a number needed to treat  
12 of 5 (95%CI 3 to 14), for a control group rate of 32%. The authors reported  
13 that all episodes of delirium occurred within the first three postoperative days.

14

15 Figure 10.1: number of patients with delirium



16

17

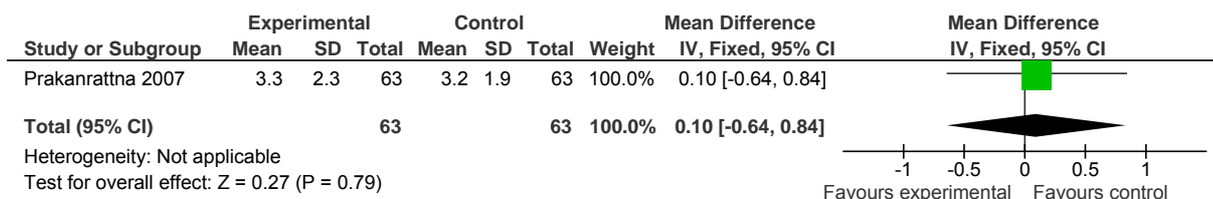
18 2. Length of ICU stay

19 There was no significant difference between the treatment groups for the number  
20 of days spent in ICU; MD 0.10 (95% CI -0.64 to 0.84; figure 10.12). The  
21 results are very imprecise. (clinically important difference: 0.5 days)

22

23 Figure 10.12: length of ICU stay

24



25

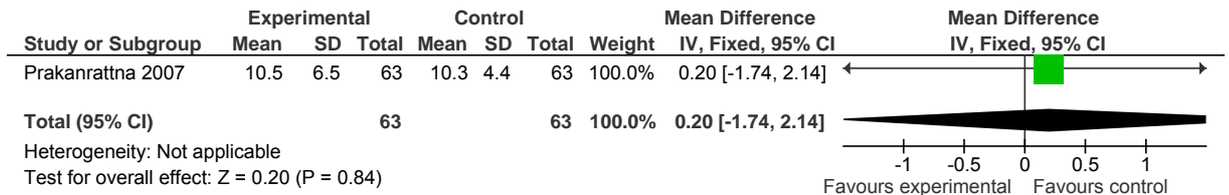
26

27

3. Length of hospital stay

1 There was no significant difference between the treatment groups for the number  
 2 of days spent in hospital; MD 0.20 (95% CI -1.66 to 2.06; figure 10.13). The  
 3 results are very imprecise.

4  
 5 Figure 10.13: length of hospital stay



7  
 8  
 9  
 10  
 11  
 12 **10.5 Clinical evidence statements**

13 Refer to Appendix K for the GRADE profile.

14  
 15 **10.5.1 Acetylcholinesterase inhibitor versus placebo**

- 16 • Meta-analysis of 2 RCTs comparing acetylcholinesterase with placebo  
 17 showed:
- 18 ○ no significant effect on the incidence of delirium (very low quality)

19 **10.5.1.1 Donepezil versus placebo**

- 20 • 1 RCT comparing donepezil with placebo showed:
- 21 ○ no significant effect on the length of hospital stay and the number  
 22 of patients discharged to a rehabilitation facility (low quality)

23  
 24  
 25 **10.5.2 Typical antipsychotics**

26 **10.5.2.1 Haloperidol versus placebo**

- 27 • 1 RCT comparing haloperidol with placebo as an adjunct to a proactive  
 28 geriatric consultation (non-pharmacological intervention) showed:
- 29 ○ no significant effect on the incidence of postoperative delirium  
 30 (low quality).
  - 31 ○ a significantly lower severity of delirium and fewer days of  
 32 delirium in favour of the haloperidol group ( low quality)

- 1                   ○ a significantly shorter length of hospital stay in patients who  
2                   received haloperidol (low quality)
- 3                   • 1 RCT comparing haloperidol with placebo showed:
- 4                   ○ no significant effect on the incidence of postoperative delirium  
5                   (low quality)
- 6                   ○ no difference between the groups for the number of adverse  
7                   events (transient tachycardia); (insufficient evidence)
- 8

### 9    **10.5.2.2 Atypical antipsychotics versus placebo**

- 10                  • 1 RCT conducted in ICU, comparing risperidone with placebo showed:
- 11                  ○ a lower incidence of delirium in patients receiving risperidone  
12                  (moderate quality).
- 13                  • 1 RCT comparing risperidone with placebo showed:
- 14                  ○ no significant difference between the groups for length of stay in  
15                  ICU and hospital. (low quality)
- 16
- 17

## 18   **10.6 Health economic evidence**

### 19   **10.6.1                  Pharmacological interventions for the prevention of delirium in** 20                  **a hospital setting**

21                  One economic evaluation study was included as evidence (Bracco 2007). This  
22                  was a non-randomised clinical trial of 1293 patients who underwent cardiac  
23                  surgery in Canada. The objective was to examine outcomes and use of intensive  
24                  care resources for a cohort of consecutive patients who underwent cardiac  
25                  surgery with or without thoracic epidural anaesthesia. The intervention group  
26                  received thoracic epidural anaesthesia for cardiac surgery. The control group  
27                  did not receive thoracic epidural anaesthesia. Detailed description of  
28                  intervention and control strategies is given in Appendix J (table J1). The  
29                  intervention shortened ventilation time and the length of stay in the ICU by 9.6  
30                  hours and 12.7 hours respectively after adjusting for type of surgery in a  
31                  multivariate analysis. This reduction decreased the ICU and mechanical  
32                  ventilation costs by US\$2700 and US\$700 respectively, per patient. The  
33                  additional cost of thoracic epidural use was given as US\$82. Post-operative  
34                  delirium complication rate was reported as 24/506 in the intervention arm, and  
35                  20/787 in the control arm. This was measured using CAM-ICU scale. A relative  
36                  risk of 0.3 was reported. Intensive care unit mortality rate of 2/506 was also  
37                  reported in the intervention arm and 14/787, in the control arm. A multivariate  
38                  analysis for mortality was not statistically significant. Cost data was taken from  
39                  the literature and QALY estimates were not reported. The study sample was not  
40                  randomised and there was no sensitivity analysis on variables whose values will  
41                  probably be uncertain. The results are not directly applicable and should be  
42                  cautiously interpreted.

1

## 2 **10 B) Prevention in long-term care:**

### 3 **acetylcholinesterase inhibitors**

#### 4 **10.7 Description of studies**

5 One paper was evaluated for inclusion Moretti (2004). The study was an RCT.

6

##### 7 **10.7.1 Study Design**

8 The RCT was conducted in Italy in a community based setting; this was treated as  
9 an indirect setting for long-term care. Patients without reliable carers were  
10 excluded from the trial. The funding source was not reported. Two hundred and  
11 forty six patients were randomised; the unit of randomisation was the patient.

12

##### 13 **10.7.2 Population**

14 The patients all had an MMSE score of at least 14, indicating patients had mild  
15 to moderate dementia. All patients met the DSM-IV criteria for dementia.  
16 Patients also satisfied the criteria for probable vascular dementia, or multi-  
17 infarct dementia with the NINDS-AIREN criteria (National Institute of  
18 Neurological Disorders and Stroke and Association Internationale pour la  
19 Recherché et l'Enseignement en Neurosciences). Their ages ranged from 65–80  
20 years with a mean age of 76 years. One hundred and sixteen men and 130  
21 women were included in the study, although 12 patients died during the study  
22 and four refused to participate; all data were based on the remaining groups of  
23 115 in the rivastigmine group and 115 in the aspirin group. All were ambulatory  
24 outpatients living in the community. Their delirium risk was not stated in the study.  
25 The comorbidity was vascular dementia, although other comorbidities were  
26 implied because of the drugs patients were taking; patients with previous  
27 psychiatric illness or central nervous system disorders or alcoholism were  
28 excluded.

1

2 **10.7.3 Interventions**

3 The included study investigated rivastigmine, a cholinesterase inhibitor,  
4 compared with cardio-aspirin (considered as usual care). Participants were  
5 ambulatory outpatients and were given the interventions for 2 years after  
6 randomisation. Rivastigmine was titrated to the higher dose after the first 16  
7 weeks. The interventions included:

- 8 • 3–6 mg/day rivastigmine
- 9 • aspirin 100 mg/day

10 It was assumed that the cardio-aspirin was representing usual care and was not  
11 an active intervention.

12

13 **10.7.4 Comparisons**

14 The following comparison was carried out:

- 15 • rivastigmine versus cardio-aspirin for 2 years (Moretti 2004)

16

17 The patients were allowed to continue taking their existing drug therapies, anti-  
18 hypertensives, anti-dyslipidemic, anti-diabetic drugs, diuretics, bronchodilators.

19

20 Patients received benzodiazepines or neuroleptic drugs during delirium, which  
21 were significantly less in the intervention group. This may have led to  
22 confounding for some outcomes, but would serve to underestimate the size of the  
23 effect.

24

25 **10.8 Methodological quality**

26 The methods of sequence generation and allocation concealment were not  
27 described, although the patients were matched for age and education level. It  
28 was not reported if all eligible patients were recruited.

29

30 The study did not report whether patients and investigators were blinded to  
31 treatment allocation. An *a priori* sample size calculation was not reported.

32

33 Originally 246 patients were included in the study, but 16 were not included in  
34 the final analyses (7% missing data; 12 patients died during the follow up and

four refused to participate in the follow up). The groups to which they were assigned were not reported. The remaining 230 patients completed the two year follow up. Patients were found to be comparable at baseline on the following scales: BEHAVE-AD (Behavioural Pathology in Alzheimer's Disease Rating); Clinical Dementia Rating; and the Cumulative Illness Rating Scale. Concordance was monitored by care givers, who controlled the intake of drugs.

Delirium was assessed using the Confusion Assessment Method (CAM).

Overall, the study may have been at a higher risk of bias because allocation concealment and blinding were unclear; appear to have a higher potential for bias, although the differential use of rescue medication may have led to confounding for some outcomes.

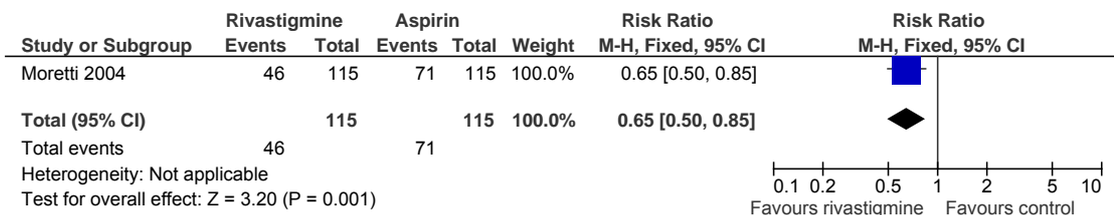
## 10.9 Results

### 10.9.1 Rivastigmine versus usual care (aspirin)

#### 10.9.1.1 Incidence of delirium (endpoint 2 years)

Analysis of one study in 230 patients showed that the incidence of delirium was significantly lower in the rivastigmine group compared with usual care (figure 10.14); RR 0.65 (95%CI 0.50 to 0.85), which corresponds to a number needed to treat of 5 (95%CI 4 to 12), for a control group rate of 62%. The result was imprecise.

Figure 10.14: incidence of delirium

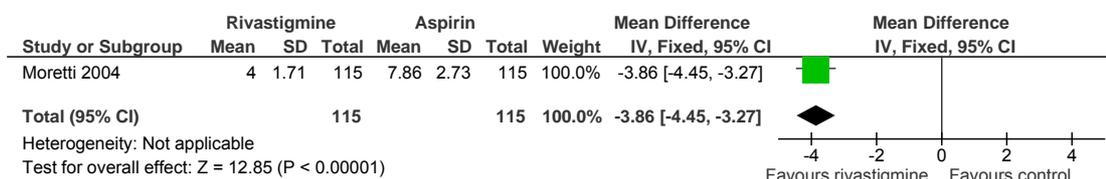


#### 10.9.1.2 Duration of delirium

Analysis of one study in 230 patients showed that the duration of delirium was significantly shorter in the rivastigmine group compared with usual care (figure 10.15a); MD -3.86 days (95%CI -4.44 to -3.28), for a control group duration of 7.86 days. It was unclear whether the duration of delirium was reported just for those who had delirium or was a mean across all patients: the paper

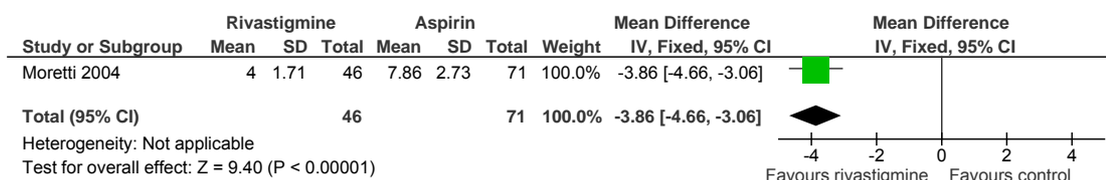
1 describes 'the main duration of the delirium'. In addition, the different standard  
 2 deviations across the groups, indicates the mean may just be for those with  
 3 delirium. Figure 10.15b shows the analysis with this assumption; the only  
 4 difference is a slightly wider CI; MD -3.86 days (95%CI -4.66 to -3.06).

5  
 6 Figure 10.15a: duration of delirium (all patients)



7  
 8  
 9

10 Figure 10.15b: duration of delirium assuming mean is across those with delirium

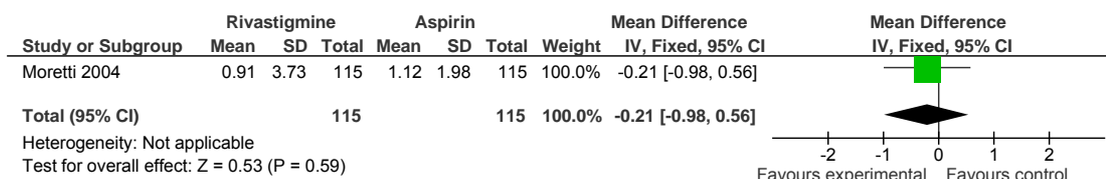


11  
 12

13 **10.9.1.3 Cognitive impairment**

14 The study assessed global performance using the Clinical Dementia Rating (scale  
 15 0–3), and reported the change from baseline at 12 months. Analysis of 230  
 16 patients showed there was no significant difference between the groups,  
 17 although the table in Moretti (2004) stated the difference was significant (figure  
 18 10.16);

19  
 20 Figure 10.16: cognitive impairment (Clinical Dementia Rating change scores)



21  
 22  
 23  
 24

25 **10.9.1.4 Behavioural disturbance (change score at 1 year)**

26 Analysis of one study in 230 patients showed that behavioural disturbance was  
 27 significantly lower in the rivastigmine group compared with usual care (figure  
 28 10.17a). The study used the BEHAVE-AD to assess individual behavioural items  
 29 on this scale (delusions, hallucinations, activity alterations, aggressiveness,  
 30 anxiety/phobia, sleep disturbances, affective disturbances, anxiety). All  
 31 individual items were stated to be statistically significant, with the exception of

delusions. The overall score showed a statistically significant mean difference, favouring the intervention; MD  $-39.66$  (95%CI  $-40.06$  to  $-39.26$ ). This seems to be a very narrow CI, even for a change score from baseline, but if these were standard errors, rather than standard deviations (despite what was reported in the text), the standard deviations would be rather large for the intervention group (figure 10.17b). The assumption of a standard error gave a large significant mean difference of  $-39.66$  (95% CI  $-43.91$  to  $-35.41$ ), favouring the intervention group.

Figure 10.17a: BEHAVE-AD scale change scores

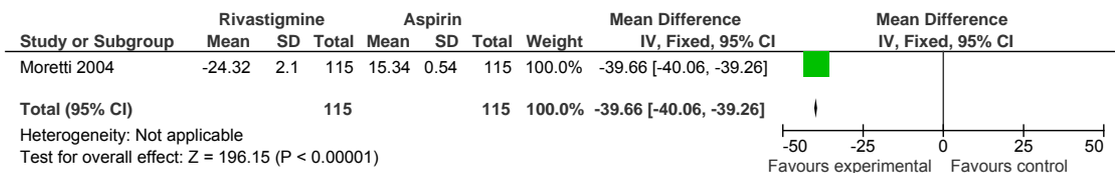
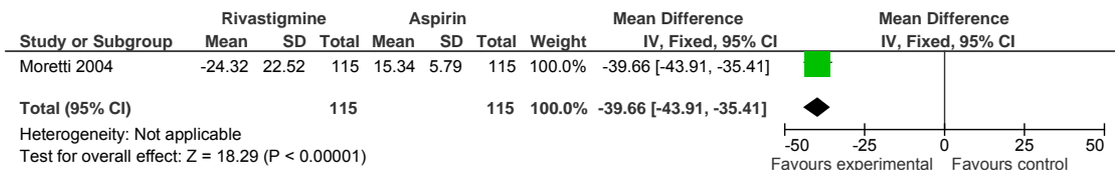


Figure 10.17b: BEHAVE-AD overall change scores



## 10.10 Evidence summary

- 1 RCT comparing rivastigmine with usual care (indirect evidence) showed that:
  - the rivastigmine group had a significantly lower incidence of delirium.
  - the rivastigmine group had significantly fewer days of delirium.
  - the rivastigmine group had significantly lower behaviour disturbances (change score at 1 year).
  - at 12 months there was no significant difference between the groups for change in cognitive impairment from baseline.  
(very low quality)

# 1 11 Adverse effects

2

## 3 11.1 Background

4 A wide variety of pharmacological interventions are available for the prevention  
5 and treatment of delirium. The drugs have varying pharmacological actions, and  
6 patients may potentially be troubled by a wide spectrum of adverse effects  
7 depending on the agent administered.

8

9 In making rational treatment choices, healthcare professionals need to carefully  
10 weigh up evidence on the anticipated benefits against that of any relevant  
11 concerns about the safety and tolerability. There are two important aspects in a  
12 review of adverse effects data for drugs in delirium:

- 13 • Evaluation of comparative data among different drugs can help healthcare  
14 professionals arrive at a treatment decision for a particular agent based  
15 on whether the safety profile (nature and frequency of adverse effects)  
16 is more, or less, acceptable than the other available agents.
- 17 • Healthcare professionals should be aware of the most important adverse  
18 effects that can arise after giving the therapy so that they can take  
19 appropriate measures to detect and minimize the risk from adverse  
20 effects

21

22 In most illnesses, patients are given adverse effects information to guide their  
23 choice of treatment and to enable them to seek medical attention for any  
24 untoward symptoms. However, patients receiving treatment for delirium may  
25 have little say in the matter, and have to rely on the actions of the healthcare  
26 professional. As such the onus is on the healthcare professional to make the  
27 appropriate decisions and to institute relevant monitoring and precautionary  
28 measures.

29

30 While some details on adverse effects have been covered in the parallel  
31 efficacy reviews of delirium, there is limited information on the specific adverse  
32 effects. It is also unclear whether the classes of drugs differ in their safety and  
33 tolerability profile.

34

### 35 11.1.1 Objective:

36 To determine what specific adverse effects may arise from drug therapy for  
37 prevention or treatment of delirium.

1

## 2 **11.2 Selection criteria**

3 The selection criteria described in the general methodology section were used,  
4 but some were specific to the evaluation of adverse effects and are reported in  
5 the following sections.

6

### 7 **11.2.1 Types of studies**

8

9 We did not apply any specific inclusion criteria based on study design; however,  
10 we aimed to exclude:

- 11 • Published case reports and case series of specific adverse events, as there  
12 is a large degree of publication bias stemming from authors' and editors'  
13 decisions favouring manuscripts covering esoteric or interesting patients.  
14 Such cases are unlikely to be representative of the general patient  
15 population
- 16 • Cross-over studies, as it is impossible to discriminate between events that  
17 arise as a complication of the first (previous) treatment, or as events  
18 resulting from the present therapy (carry-over).
- 19 • Small studies with fewer than 20 patients exposed to the intervention of  
20 interest, as such studies are unlikely to be able to detect any important  
21 adverse effects, and may lead to falsely reassuring findings that no  
22 safety problems were identified.

23

### 24 **11.2.2 Types of participants**

- 25 • Adults (18 years and over)
- 26 • Patients requiring treatment for delirium or being given treatment to  
27 prevent delirium
- 28 • Not end-of-life patients or patients with primarily psychiatric disorders such  
29 as schizophrenia, bipolar disorder or other psychoses.

30

31 Following GDG advice and post-hoc evidence from an indirect population was  
32 included in order to investigate stroke as an adverse event. The GDG extended  
33 the population to include older patients and those with dementia.

### 34 **11.2.3 Interventions of interest**

- 35 • Typical antipsychotics: haloperidol
- 36 • Atypical antipsychotics: risperidone, olanzapine, quetiapine, amisulpride

- 1           • Benzodiazepines: diazepam, flunitrazepam
- 2           • Cholinesterase inhibitors: donepezil, rivastigmine
- 3           • 5-HT3 antagonists: ondansetron

4

5           Duration of intervention: any

6

7   **11.2.4     Comparators**

8           For controlled studies, we accepted comparisons of any of the above agents  
9           versus placebo or no treatment. We also included studies that directly compared  
10          two or more agents from the above list of interventions. However, we excluded  
11          studies if the relevant intervention was tested against an active comparator that  
12          was not on the list of included drugs, as it would then be impossible to draw any  
13          valid conclusions on the relative safety profile of the agent of interest (safer or  
14          more harmful than an intervention of unknown effect).

15

16   **11.2.5     Outcomes**

17          All outcomes reported within the categories of 'adverse effects, side effects,  
18          adverse events, complications, safety, or tolerability'.

19

20   **11.2.6     Assessment of Validity of Adverse Effects Data**

21

22          The methods for assessing validity were based on recommendations of chapter  
23          14 of the Cochrane Handbook of Systematic Reviews. This focuses on two major  
24          factors:

- 25           • How thorough were the methods used in monitoring adverse effects?
- 26           • How complete or detailed was the reporting?

27

28          In view of this, the following parameters were recorded:

- 29           • What methods (if any) did the studies stipulate for the specific assessment  
30           of adverse effects?
- 31           • Did the investigators prespecify any possible adverse events that they  
32           were particularly looking out for?
- 33           • What categories of adverse effects were reported?

34

35

36

## 1 11.3 Identification of studies

2 Articles that had already been retrieved for the efficacy reviews were  
3 considered and reference lists were checked to identify specific articles on  
4 adverse effects.

5

6 A total of 170 full text articles were screened, with 18 studies fulfilling the  
7 inclusion criteria.

8 However, we had to make further exclusions due to no adverse effects data  
9 being extractable. Three eligible studies failed to mention anything about  
10 adverse effects and were not evaluated any further. (Hu 2006: olanzapine,  
11 haloperidol and control; Liu 2004: risperidone; Moretti 2004: rivastigmine).

12

13 Adverse effects data were extracted from 15 included papers.

14 Following GDG advice, indirect evidence was obtained from three further  
15 studies.

16

### 17 11.3.1 Study Design

18 The following types of studies were included in the adverse effects analysis:

- 19 • Direct head to head comparison of two antipsychotic agents: 1 RCT (Lee  
20 2005), 1 quasi-randomised study (Skrobik 2004), 1 prospective cohort  
21 study (Gill 2005\*; with retrospective elements), and 2 retrospective  
22 cohort studies (Herrmann 2004\*; Miyaji 2007).
- 23 • Typical antipsychotic: haloperidol, 2 placebo controlled RCTs (Kalisvaart  
24 2005; Kaneko 1999); typical antipsychotics generally, 1 retrospective  
25 cohort study (Douglas\* 2008)
- 26 • Atypical antipsychotics: 6 studies consisting of 1 RCT (Prakanratta 2007), 3  
27 open trials without control arms (Breitbart 2002; Kim 2001; Pae 2004),  
28 and 3 observational studies (Douglas 2008\*; Parellada 2004).
- 29 • Benzodiazepines: diazepam, flunitrazepam: no studies met the eligibility  
30 criteria. One study (Aizawa 2002) that was included in the efficacy  
31 review had to be excluded as the intervention involved three agents –  
32 diazepam, flunitrazepam and pethidine, and it would not have been  
33 possible to tell if any adverse effects were due to the benzodiazepine or  
34 the pethidine.
- 35 • Cholinesterase inhibitors: donepezil, rivastigmine. One double blind  
36 placebo controlled RCT (Liptzin 2005).

- 1           • 5-HT3 antagonists: ondansetron – one open trial without control arm  
2                    (Bayindir 2000)

3           \* indicates studies in an indirect population  
4  
5

### 6   **11.3.2    Population**

7           The studies looked at a wide range of participants, but for the most part were in  
8           patients undergoing surgery or admission to intensive care. Three of the studies  
9           (Douglas 2008\*; Gill 2005\*; Herrmann 2004\*) reported on stroke adverse  
10          events associated with antipsychotics in older patients, who were likely to be at  
11          risk of delirium.

12

### 13   **11.3.3    Intervention and Comparisons**

14          There was a diverse range of interventions, and associated comparator agents  
15          across the trials.

16

### 17   **11.3.4    Assessment and Reporting of Adverse Effects**

18          A diverse range of methods were used, with the most well-defined ones being  
19          scales for assessing extrapyramidal signs and symptoms. It is not clear though  
20          how useful such scales are in postoperative or intensive care patients, in contrast  
21          to ambulant psychiatric patients.

22

## 23   **11.4 Results**

24          The interventions, comparators and populations were extremely varied, as was  
25          the reporting of adverse effects outcomes. Descriptive summaries are given in  
26          Appendix D.

27

### 28   **11.4.1    Direct comparison of active agents**

29          Five studies (Gill 2005\*; Herrmann 2004\*; Lee 2004; Miyaji 2007; Skrobik  
30          2004) reported direct comparisons between two antipsychotic agents.

31

32          Extrapyramidal adverse effects were the main focus of three studies (Lee 2004;  
33          Miyaji 2007; Skrobik 2004), with one study (Skrobik 2004) describing specific  
34          efforts to “carefully record” such events. Two studies reported specifically on  
35          stroke as an adverse event (Gill 2005\*; Herrmann 2004\*). One study was in  
36          older adults (mean age 81.7 years) (Herrmann 2004) and one study was in  
37          older adults with dementia (mean age 82.6 years) (Gill 2005\*).

1

2 No extrapyramidal events were found in the Lee (2004) study, but both Miyaji  
3 (2007) and Skrobik (2004) studies described a higher incidence of  
4 extrapyramidal effects with haloperidol as compared to quetiapine, and  
5 olanzapine respectively. However the Miyaji (2007) study was retrospective  
6 while Skrobik (2004) was quasi-randomised, and neither study had any blinding  
7 and are thus subject to bias from investigators who may favour the new atypical  
8 antipsychotics when recording the extrapyramidal effects.

9

10 While the ascertainment of mortality is less subjective, the baseline differences in  
11 populations receiving the interventions in the Miyaji (2007) study makes it  
12 difficult to draw any reliable conclusions, simply because the more severely ill  
13 patients may have received parenteral haloperidol.

14

15 Two studies carried out multivariate analyses (Gill 2005\*; Herrmann 2004\*). The  
16 Gill (2005) study did not take into account confounders such as smoking history,  
17 presence and severity of hypertension, lipid status and specific valvular heart  
18 conditions. Similarly the Herrman (2004) study did not take into consideration  
19 smoking or obesity. The most commonly prescribed antipsychotic was risperidone  
20 in both studies (Gill 2005\*: 76%; Herrmann 2004\*: 61%)

21

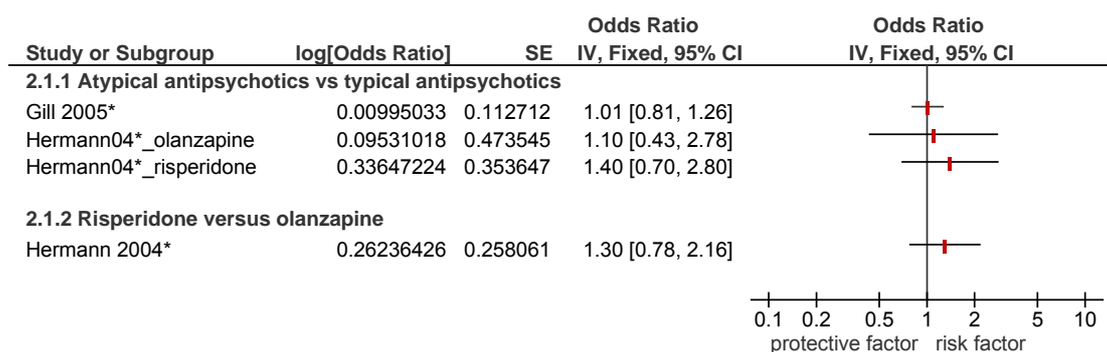
22 The Gill (2005)\* study reported that in older patients with dementia there is no  
23 significant difference in the effects of atypical antipsychotics compared with  
24 typical antipsychotics.

25

26 The Herrmann (2004)\* study reported results separately for olanzapine and  
27 risperidone compared with typical antipsychotics. There was no significant  
28 effect of olanzapine [RR 1.1 (95% CI 0.4 to 2.3)] or risperidone [RR 1.4 (95% CI  
29 0.7 to 2.8)] on the incidence of stroke. A head to head comparison (risperidone  
30 versus olanzapine) showed no difference in effect [RR 1.3 (95% CI 0.8 to 2.2);  
31 figure 11.1.

32

1 Figure 11.1: antipsychotics as a risk factor for stroke



2  
3  
4  
5

6 **11.4.2 Results of specific classes of interventions versus no treatment or**  
7 **placebo**

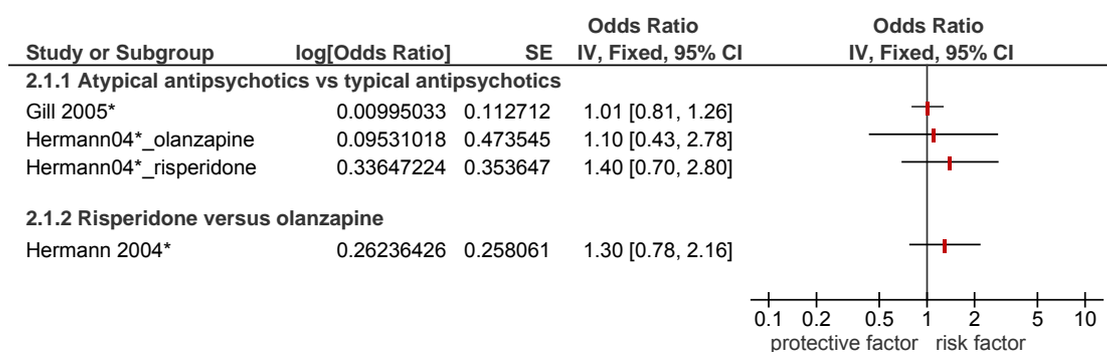
8 **11.4.2.1 Typical and atypical antipsychotics**

9 One retrospective cohort study (Douglas 2008\*) was an intra-patient study  
10 comparing periods of antipsychotic use and non-use in older adults (indirect  
11 population). Median age when first exposed to any antipsychotic drug was 80  
12 years. The study reported on the risk of stroke in patients presenting with first  
13 ever stroke (at least 12 months after initial registration on the UK general  
14 practice database). The most commonly prescribed atypical antipsychotic was  
15 risperidone (81%), followed by olanzapine (18%), amisulpride and quetiapine  
16 (4% in each group).

17 Exposure to any of the antipsychotics was a significant risk factor for stroke [RR  
18 1.73 (95% CI 1.60 1.87)]. When typical and atypical antipsychotics were  
19 analysed separately, a significant effect was observed (figure 11.2).

20  
21

Figure 11.2: antipsychotics as a risk factor for stroke



22  
23

24 **11.4.2.2 Haloperidol**

1 There were two included RCTs, both covering the use of haloperidol in  
2 postoperative patients. (Kalisvaart 2005, Kaneko 1999)

3

4 Both trials reported on active measures to detect adverse effects, with frequent  
5 clinical assessments. Haloperidol use in this setting appeared to be relatively  
6 safe with no excess of withdrawals from adverse events compared to control,  
7 and no extrapyramidal effects.

8

#### 9 **11.4.2.3 Atypical antipsychotics**

10 For risperidone, we identified one RCT (Prakanratta 2007) and one  
11 observational study (Parellada 2004). There were two open uncontrolled trials  
12 of olanzapine (Breitbart 2002, Kim 2001), and one of quetiapine (Pae 2004).

13 Both the risperidone studies looked for specific adverse effects but did not show  
14 any clear trend for harm.

15 One olanzapine study (Breitbart 2002) used clinical methods to evaluate  
16 adverse effects, and this showed sedation to be a problem necessitating dosage  
17 reduction.

18 The remaining two studies (Kim 2001, Pae 2004) did not mention any specific  
19 monitoring for adverse effects, and data were sparse.

20

21

#### 22 **11.4.2.4 Cholinesterase inhibitors**

23 One study (Liptzin 2005) which was a randomised double-blind controlled trial  
24 of donepezil was identified. Despite methodological strengths elsewhere, this  
25 study did not describe any specific monitoring of adverse effects, and did not  
26 provide numerical data, even though there was a statement about equivalent  
27 rates of adverse effects between drug and placebo. Moreover, adherence to  
28 treatment was poor, and as such, no conclusions can be drawn on the relative  
29 safety of donepezil.

30

#### 31 **11.4.2.5 5-HT<sub>3</sub> antagonists**

32 One study (Bayindir 2000) which was an open-label uncontrolled evaluation of  
33 ondansetron in postoperative patients was identified. The authors did not state  
34 what, if any monitoring was used for detecting adverse effects, and it is difficult  
35 to have any confidence in their conclusions that the therapy was safe, without  
36 any apparent side effects.

1

2 **11.4.3 Limitations of the results**

3 The paucity of the reported adverse effects data is a major limitation here. Most  
4 of the investigators appear to have focused on extrapyramidal effects, and  
5 omitted to consider or discuss the possibility of other adverse events. Another  
6 important weakness here is that patients with delirium are unable to accurately  
7 complain of any untoward symptoms, and thus adverse events may have been  
8 missed by the clinicians. The heterogeneous data on haloperidol are of interest  
9 here, as this may possibly reflect susceptibility to bias in the unblinded studies  
10 that found an excess of extrapyramidal symptoms, when compared to newer  
11 atypical agents. The data on extrapyramidal effects and mortality should be  
12 judged cautiously, given that higher quality randomized controlled trials with  
13 thorough adverse effects monitoring have failed to replicate these findings.

14 All three studies (Douglas 2004\*; Gill 2005\*; Herrmann 2004\*) reporting on the  
15 incidence of stroke and antipsychotic use attempted to take into account known  
16 confounders, but each had limitation; the Gill (2005)\* may have been higher  
17 quality because it was prospective but was solely in patients with dementia and  
18 the results may therefore not be generalisable.

19

20

21 **11.5 Clinical evidence statements**

22

23

- There is moderate quality evidence in a large:

24

- retrospective cohort study that antipsychotics have a significant effect on the incidence of stroke in patients who have a median exposure time of 0.37 years. This is indirect evidence for patients who receive antipsychotics for delirium, who will have the drugs for much shorter periods.

25

26

27

28

29

- mixed prospective-retrospective cohort study in patients with dementia to suggest there is no significant difference in the effects of typical and atypical antipsychotics compared head to head.

30

31

32

33

- retrospective cohort study to suggest that there is no significant difference between risperidone and olanzapine as risk factors for stroke in patients who received drugs for at least 30 days.

34

35

# 12 Diagnosis: accuracy of diagnostic tests in different clinical settings

## 12.1 Clinical Introduction

Delirium is common but is frequently unrecognised by doctors and nurses despite the fact that it can be life-threatening and lead to serious preventable complications. Unfortunately there is no simple quick test for delirium comparable to the ECG or Troponin test in myocardial infarction. The reference standard for diagnosis is a careful clinical assessment using the DSM-IV criteria at the bedside but this takes time and needs clinical expertise. There are however many screening tests available and these are reviewed in this section. Clinical suspicion should be high in any patient with a sudden change of behaviour or mental state especially in older patients with dementia, severe illness or fracture neck of femur. Early identification of patients with delirium and patients at increased risk is an essential first step in improving the management and outcome for this serious condition.

16  
17

### 12.1.1 Clinical Question:

What are the practical diagnostic tests to identify patients with delirium in different clinical settings?

21

### 12.1.2 Primary objective of the review

To determine the accuracy of various diagnostic tests in diagnosing delirium in patients in hospital and long-term care.

25

### 12.1.3 Inclusion criteria

The following inclusion criteria were used for this review:

28

#### 12.1.3.1 Patients

Adult patients in hospital; studies were stratified by setting (hospitals, long-term care and ICU).

32

#### 12.1.3.2 Prior tests

No prior tests were undertaken

34

35

1 **12.1.3.3 The target condition**

2 Delirium  
3

4 **12.1.3.4 The index test and who executes the test**

5 • Hospital:

- 6 ○ Abbreviated Mental test (AMT); any personnel can do this;
- 7 ○ Clock-drawing test; can be used by untrained nurses or  
8 volunteers;
- 9 ○ Confusion Assessment Method [long version] (CAM); trained  
10 healthcare professionals;
- 11 ○ Confusion Assessment Method [long version] (CAM); trained  
12 healthcare professionals;
- 13 ○ DRS-R-98; trained healthcare professional;
- 14 ○ Mini Mental State Examination (MMSE) or other cognitive  
15 assessment instrument; trained healthcare professional.

16

17 • ICU:

- 18 ○ CAM-ICU and RASS (together)

19

20

21 **12.1.3.5 The reference standard**

22 DSM-IV or ICD-10 applied by trained specialists  
23

24 **12.1.3.6 Sensitivity analyses**

25 Sensitivity analyses were carried out to address QUADAS quality items  
26

27 **12.1.3.7 Subgroup analyses**

28 For this review, we stratified the data according to the setting (hospital, ICU,  
29 long-term care), and considered the following subgroups in order to investigate  
30 heterogeneity

31 • ethnicity

32 • whether English is the first language

33 • writing ability

34 • patients with and without dementia/cognitive impairment

1

## 2 12.2 Characteristics of included studies

3 Thirty-four reports were identified as being potentially relevant. Fourteen were  
4 excluded and these are listed in Appendix G, along with reasons for exclusion.  
5 20 reports of 18 studies were included (Andrew 2009; Cole 2003; Ely 2001;  
6 Ely 2001b; Fabbri 2001; Gonzalez 2004; Hestermann 2009; Laurila 2002;  
7 Laurila 2003; Laurila 2004; Lin 2004; Monette 2001; Ni Chonchubhair 1995;  
8 O'Keeffe 2005; Pompei 1995; Radtke 2008; Rockwood 1994; Rolfson 1999b;  
9 Yates 2009; Zou 1998). One study (Laurila 2003) had more than one report  
10 (Laurila 2003 and 2004); hereafter, these studies are referred to by their first  
11 named reports, but separately in the results section. One study (Vreeswijk 2009)  
12 was identified in the update search and was not analysed in depth as it did not  
13 add substantially to the body of evidence.

14

15 One study (Laurila 2002) may have included some of the same patients as those  
16 included in the Laurila (2003) study. The study enrollment period or the time  
17 period when assessments were carried out was not reported in the Laurila  
18 (2002) study. However, as the setting was limited to hospitals only in the 2002  
19 study and as the other study (Laurila 2004) included hospital and long-term care  
20 setting, the results are reported separately.

21

22 The Cole (2003) study reported a secondary analysis of information collected in  
23 what the authors reported as a randomised controlled trial of management of  
24 delirium and a prospective study of prognosis of delirium which included non  
25 delirious patients [references were not provided for either study in the text].

26

27 Study size ranged from fewer than 50 patients in two studies (Ely 2001b: n=38;  
28 Hestermann 2009: n=39), between 50 and 100 patients in six studies (Ely 2001:  
29 n=96; Fabbri 2001: n=100; Laurila 2002: n=81; Rolfson 1999b: n=71; Yates  
30 2009: n=62; Zou 1998: n=87), between 100 and 1000 in ten studies (Andrew  
31 2009: n=145; Cole 2003: n=322; Gonzalez 2004: n=153; Laurila 2003:  
32 n=425; Lin 2004: n=109; Monette 2001: n=110; Ni Chonchubhair 1995:  
33 n=100; O'Keeffe 2005: n=165; Radtke 2008: n=154; Rockwood 1994:  
34 n=434) and one study recruited over 1000 patients (Pompei 1995: n=1168).

35

### 36 12.2.1 Study design

37 There were 20 included reports, all of which were studies of diagnostic test  
38 accuracy. Most studies had a cross-sectional design, but the Cole (2003) study,  
39 which reported a secondary analysis of data collected in an RCT and  
40 prospective study, appeared to be a case-control study; one set of patients  
41 were included if they had a score of 3 or more on the Short Portable Mental  
42 Status Questionnaire (SPMSQ) or if their nursing notes indicated symptoms of  
43 delirium and who met the DSM III-R criteria for delirium. The other set of included  
44 patients were people free of delirium, selected following screening for delirium;  
45 the study reported that the selection of non delirious patients in the study took  
46 into account the patients' age and initial cognitive impairment status (SPSMQ  
47 score <3).

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The studies were conducted in different settings:

- Fifteen studies were carried out in hospital (Andrew 2009; Cole 2003; Fabbri 2001; Gonzalez 2004; Hestermann 2009; Laurila 2002; Monette 2001; Ni Chonchubhair 1995; O'Keeffe 2005; Pompei 1995; Radtke 2008; Rockwood 1994; Rolfson 1999b; Yates 2009; Zou 1998);
  - The Andrew (2009) study included 73% [106/145] inpatients and the remainder were outpatients; 15/39 of the outpatients (10% overall) were seen at home
  - The Gonzalez (2004) study reported excluding patients in psychiatric wards.
- Three studies were conducted in an ICU setting (Ely 2001; Ely 2001b; Lin 2004);
- One study was conducted in both hospital and long-term care settings (Laurila 2003).

Two studies were carried out in the UK (Ni Chonchubhair 1995; Yates 2009) and the rest were conducted in: Ireland (O'Keeffe 2005); the USA (Ely 2001; Ely 2001b; Pompei 1995); Canada (Andrew 2009; Cole 2003; Monette 2001; Rockwood 1994; Rolfson 1999b; Zou 1998); Finland (Laurila 2002; Laurila 2003); Germany (Hestermann 2009; Radtke 2008); Spain (Gonzalez 2004); Brazil (Fabbri 2001); and China (Lin 2004).

## 24 12.2.2 Population

25 The inclusion and exclusion criteria for each of the studies are shown in  
26 Appendices D and G.

27  
28 Rates of delirium ranged from 14% (Radtke 2008) to 64% (Zou 1998) in the  
29 hospital setting; 86% (Ely 2001; Ely 2001b) in the ICU setting; and 25%  
30 (Laurila 2003) in the mixed setting (hospital and nursing home wards).

31  
32 Where reported, the mean age of the participants in the studies was mostly  
33 above 65 years but varied as follows:

- mean age above 65 years (Andrew 2009; Cole 2003; Fabbri 2001; Gonzalez 2004; Hestermann 2009; Inouye 2005; Laurila 2003; Lin 2004; Monette 2001; Ni Chonchubhair 1995; O'Keeffe 2005; Pompei 1995; Rolfson 1999b; Yates 2009; Zou 1998)
  - Five studies were in much older patients: mean age over 80 years (Andrew 2009; Cole 2003; Hestermann 2009; Laurila 2003; Zou 1998)
- mean age below 65 years (Ely 2001; Ely 2001b; Radtke 2008)

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Eight studies had a lower limit to age for inclusion: the Monette (2001) study reported that patients were eligible for enrollment if their age was 66 years and over; five studies (Gonzalez 2004; Ni Chonchubhair 1995; O'Keeffe 2005;

1 Pompei 1995; Zou 1999) included patients over 65 years; and two studies  
2 (Laurila 2002; Laurila 2003) excluded patients younger than 70 years.

3  
4 The studies varied in the proportion of patients with dementia/cognitive  
5 impairment:

- 6 • Patients with dementia were excluded in one study (Lin 2004);
- 7 • The Ely (2001b) study reported patients with a history of severe dementia  
8 were excluded, however, patients with suspected dementia (29%) were  
9 identified following enrollment;
- 10 • One study (Ely 2001: 12.5% ) reported that less than 20% of the patients  
11 had suspected dementia;
- 12 • Five studies (Andrew 2009: 40%; Cole 2003: 29%; Gonzalez 2004: 50%;  
13 O’Keeffe 1997: 22%; Pompei 1995: 21%) reported between 20 and  
14 50% of the patients had dementia;
- 15 • Three studies (Hestermann 2009: 84.6% ; Laurila 2003: 64%; Monette  
16 2001: 53%) reported over 50% of the patients had dementia;
- 17 • One study (Yates 2009) reported the mean MMSE scores for delirium and  
18 non delirium groups (4.64 versus 14.94; p=0.003); the scores indicate  
19 that the included patients in this study were likely to be severely  
20 cognitively impaired.
- 21 • Four studies did not report dementia status (Fabbri 2001; Ni Chonchubhair  
22 1995; Radtke 2008; Zou 1998).
- 23 • One study (Rolfson 1999b) reported that patients were ‘highly selected  
24 with a low proportion of dementia’. Patients were undergoing coronary  
25 artery bypass graft surgery.

26  
27 The studies varied in their inclusion or otherwise of non-English speaking people.  
28 None of the studies reported if English was the first language. Five studies (Ely  
29 2001; Ely 2001b; Inouye 2005; Pompei 1995; Rolfson 2005) reported  
30 excluding patients who did not speak English; two studies (Cole 2003; Monette  
31 2001) reported excluding patients who did not speak English or French and one  
32 study (Radtke 2008) conducted in Germany reported excluding patients who  
33 did not speak the local language. Four studies reported the validation of the  
34 translated CAM instrument into: Portuguese (Fabbri 2001); Chinese (Lin 2004);  
35 Spanish (Gonzalez 2004); Hestermann (German). One study (Laurila 2002)  
36 reported using a previously validated, Finnish version of the CAM instrument. For  
37 the translation studies we have assumed English was not the first language.

38  
39 Ethnicity was reported in six studies (Ely 2001; Ely 2001b; Fabbri 2001; Inouye  
40 2005; O’Keeffe 2005; Pompei 1995); with three studies reporting the majority  
41 of the patients were white (Ely 2001; Ely 2001b; O’Keeffe 2005); European  
42 descent (Fabbri 2001), and one study (Pompei 1995) reporting that 29% of the  
43 patients were African-American.

44  
45 One study (Fabbri 2001) reported that 32% of the patients included in the study  
46 were unable to read or write fluently.

1

2 **12.2.3 Index tests**

3 A range of index tests were described:

- 4 • Abbreviated Mental Test (AMT); serial test (comparison of day before  
5 surgery and 3 day postoperatively) (Ni Chonchubhair 1995);
- 6 ○ A 10 item questionnaire (scale score range: 0 to 10, with a score  
7 less than 6 indicative of dementia);
- 8 • Confusion Assessment Method (CAM):
- 9 ○ CAM (short version: Laurila 2002; Monette 2001; Pompei 1995;  
10 Radtke 2008)
- 11 - The CAM short version assesses on the following 3 criteria;  
12 acute onset and fluctuating course and inattention *and*  
13 disorganised thinking or altered level of consciousness.
- 14 ○ CAM (long version: Cole 2003; Yates 2009; Zou 1998)
- 15 - The CAM long version assesses on the following 10  
16 criteria: acute onset, inattention, disorganised thinking,  
17 altered level of consciousness, disorientation, memory  
18 impairment, perceptual disturbances, psychomotor  
19 agitation, psychomotor retardation, and altered sleep-  
20 wake cycle
- 21 ○ CAM (type of version unclear: Rockwood 1994; Rolfson 1999b);
- 22 ○ CAM translations (Fabbri\* 2001 [Portuguese]; Gonzalez\* 2004  
23 [Spanish]; Hestermann\* 2009 [German]; Laurila\* 2002 [Finnish];  
24 (translations are indicated by an asterisk in the rest of this  
25 document)
- 26 - Three studies reported a translation of the short version  
27 (Gonzalez\* 2004; Hestermann\* 2009; Laurila\* 2002) and  
28 the other study (Fabbri 2002\*) reported a translation of  
29 the long version.
- 30 • Confusion Assessment Method (ICU) (CAM-ICU) (Ely 2001; Ely 2001b);
- 31 ○ The CAM-ICU assess on the presence or absence of the following  
32 features: acute onset or fluctuation course and inattention and  
33 either disorganised thinking or altered level of consciousness;
- 34 ○ Both studies reported the Attention Screening Examinations (ASE)  
35 scores, with Ely (2001b) reporting that the ASE was used to assess  
36 the 'inattention' feature of CAM-ICU. The Ely (2001b) study  
37 reported that the Vigilance A Random Letter Test which is part of  
38 the ASE was performed selectively in visually impaired patients.  
39 The Ely (2001) study reported that patient's delirium status was  
40 assessed with RASS when they were alert.
- 41 ○ CAM-ICU translations: (Lin\* 2004: Chinese)
- 42 - The study reported patients were followed up daily with  
43 the Glasgow Coma Scale and the RASS for assessment of  
44 acute onset of mental status changes or fluctuation course.

- 1           • Clock-drawing test (Rolfson 1999b);
- 2           ○ The clock-drawing test is an instrument used for screening of
- 3           cognitive disorders. The test can be administered in three formats:
- 4           in the free-drawn method, the patient is asked to draw a clock
- 5           from memory; in the pre-drawn method, the patient is presented
- 6           with a circular contour and is expected to draw in the numbers on
- 7           the clock face; or in the third method the patient is asked only to
- 8           set the hands at a fixed time on a pre-drawn clock, complete with
- 9           contour and numbers.
- 10          ○ The Rolfson (1999b) study did not report the clock-drawing test
- 11          format. The study reported a score of 6 or less was considered
- 12          abnormal (range: 1 to 10, with 10 being error-free).
- 13          • Mini Mental State Examination (MMSE) (Rolfson 1999b; O’Keeffe 2005);
- 14          ○ The MMSE is a test that is used to screen for cognitive impairment.
- 15          (range 0 to 30);
- 16          ○ Score of 23 or less was considered to be indicative of cognitive
- 17          impairment (Rolfson 1999b)
- 18          ○ Serial change in MMSE score; change in score between day 1
- 19          and day 6 (O’Keeffe 2005)
- 20                  - The study reported using a version of the MMSE that was
- 21                  previously adapted and validated for use in an Irish
- 22                  population.
- 23          • Delirium Index (DI) (Cole 2003);
- 24          ○ An instrument designed to be used in conjunction with the MMSE,
- 25          for the measurement of severity of symptoms of delirium based
- 26          solely on observation of the patients. Patients are assessed on the
- 27          following seven domains: inattention, disorganised thinking,
- 28          altered level of consciousness, disorientation, memory impairment,
- 29          perceptual disturbances, and motor disturbances. Score range
- 30          from 0 to 21, with 21 points indicating maximum severity.
- 31          • DRS-R-98 (Andrew 2009);
- 32          ○ The revised version of the DRS, allows assessment for both
- 33          diagnosis of delirium and severity of delirium. This 16-item scale
- 34          includes 3 ‘diagnostic items’ (temporal onset, fluctuation and
- 35          physical disorder) and 13 ‘severity symptoms’ (attention,
- 36          orientation, memory [short and long-term], sleep-wake cycle
- 37          disturbances, perceptual disturbances and hallucinations,
- 38          delusions, lability of affect, language, thought process
- 39          abnormalities, and motor agitation or retardation). Scores range
- 40          from 0 to 44, and patients with a score of at least or over 17.75
- 41          points were screened as positive for delirium.
- 42          • Chart assessment (Rolfson 1999b);
- 43          ○ Documentation of delirium or its symptoms in the health records
- 44          by physicians and nurses

- 1                   ○ A retrospective review of the records by non study physicians and  
2                   nurses were conducted for terms [including 'delirium' , 'confusion',  
3                   'acute confusion', 'toxic psychosis' and 'metabolic  
4                   encephalopathy'] and themes [features of delirium, for e.g. acute  
5                   onset, altered mental status, hallucinations, memory impairment]  
6                   that suggested the recognition of delirium; Results for this index  
7                   test will not considered as the GDG considered retrospective  
8                   chart review to be an inadequate method of delirium assessment.

9  
10                   Most studies reported that the patients received only one index test; the  
11                   exceptions were four reports of five studies (Cole 2003: CAM; DI; DSMIII;  
12                   DSMIII-R; ICD-10; Laurila\* 2003: DSM-III-R; DSM-III; ICD-10; Rolfson 1999b:  
13                   CAM; MMSE; clock-drawing test; Chart assessment).

14  
15                   Three other studies (Andrew 2009; Pompei 1995; Radkte 2008) reported  
16                   patients received other index tests that were not considered within this review  
17                   (Andrew 2009: Delirium Symptom Interview (DSI); Pompei 1995: Digit Span Test,  
18                   Vigilance 'A' Test, Clinical Assessment of Confusion (CAC); Radkte 2008: Delirium  
19                   Detection Score (DDS); Nursing Delirium Screening Scale (Nu-DESC))  
20

#### 21                   **12.2.4                   Reference standard (and index tests with which they were** 22                   **compared)**

23                   Although the GDG specified that the reference standard was to be DSM-IV or  
24                   ICD-10, a number of studies compared tests only with the reference standard of  
25                   DSM IIIIR or DSM III. The GDG ruled that this was acceptable, especially for the  
26                   purpose of comparing different index tests.  
27

28                   The reference standards were carried out in different ways:

##### 29                   • DSM-IV

- 30                   ○ Five studies (Ely 2001; Ely 2001b; Gonzalez\* 2004;  
31                   Hestermann\* 2009; Lin\* 2004) reported the DSM-IV criteria for  
32                   delirium was applied following clinical interview, family and/or  
33                   nurse interviews, medical records and/or mental status records.  
34                   ○ Two studies (Ely 2001; Ely 2001b) reported patients were  
35                   assessed as either normal, delirious, stupor or comatose using  
36                   DSM-IV or standardised definition of stupor and coma.  
37                   ○ Two studies (Radtko 2008; Yates 2009) reported that the  
38                   presence of delirium was determined using the DSM-IV criteria  
39                   and did not provide further information.  
40                   ○ One study (Laurila\* 2002) reported the criteria addressed in the  
41                   DSM-IV were operationalised in one questionnaire which also  
42                   addressed the criteria in other classification systems (DSM-III-R,  
43                   DSM-III, ICD10).

##### 44 45                   • ICD-10

- 1                   ○ One study (Laurila\* 2002) reported the criteria addressed in the  
2 ICD-10 were operationalised in one questionnaire which also  
3 addressed the criteria in other classification systems (DSM-IV,  
4 DSM-III-R, DSM-III).
- 5                   ● DSM III R
- 6                   ○ In the Cole (2003) study, a nurse gave CAM to patients with a  
7 SPMSQ score  $\geq 3$  or delirium symptoms in the nursing notes; then  
8 the 10 CAM symptoms of delirium appeared to be used to  
9 determine the reference standard.
- 10                  ○ One study (Laurila\* 2002) reported the criteria addressed in the  
11 DSM-III-R were operationalised in one questionnaire with  
12 addressed in other classification systems (DSM-IV, DSM-III, ICD-  
13 10).
- 14                  ● CAM and Clinician interview
- 15                  ○ One study (O’Keeffe 2005) had an experienced consultant  
16 geriatrician interview the patients using the CAM (short version)
- 17                  ● Consensus diagnosis
- 18                  ○ In the Zou (1998) study, the study team comprised of two  
19 geriatric psychiatrists, research fellow and a nurse clinician  
20 arrived at a consensus diagnosis using a nominal group method  
21 based on the following: results reported by the nurse for the  
22 CAM, SPMSQ, chart review; one assessment by a psychiatrist  
23 based on chart review and clinical examination; and independent  
24 assessment by each member of the team indicating the presence  
25 or absence of the five DSM-IV criteria for delirium (both ‘definite’  
26 cases, requiring five criteria and ‘probable’ cases, requiring four  
27 of the five were included.).
- 28
- 29                  Where reported, the reference standard was mainly carried out by  
30 geriatricians or psychiatrists, with the exception of three studies (Pompei 1995:  
31 assessed by geriatricians and a geriatric nurse specialist; Yates 2009: junior  
32 medical doctor; Zou 1998; consensus diagnosis included a nurse’s CAM findings).  
33
- 34                  Two studies compared different diagnostic criteria. In each of these comparisons  
35 the patients were given the same questionnaire/interview and the criteria were  
36 deduced from the symptoms reported
- 37                  ● DSM-III-R versus DSM-IV (Cole 2003; Laurila\* 2003) ; the test was carried  
38 out by a:
- 39                  ○ geriatrician in the hospital setting, and a nurse’s interview and  
40 notes were used to arrive at an assessment for the long-term care  
41 setting (Laurila\* 2003)
- 42                  ○ nurse (Cole 2003).
- 43                  ● DSM III versus DSM-IV (Laurila\* 2003) ; the test was carried out by :

- 1                   ○ geriatrician in the hospital setting, and a nurse's interview and  
2 notes were used to arrive at an assessment for the long-term care  
3 setting (Laurila\* 2003)
- 4                   • ICD-10 versus DSM-IV (Laurila\* 2003); the test carried out by:
- 5                   ○ geriatrician in the hospital setting and a nurse's interview and  
6 notes were used to arrive at an assessment for the long-term care  
7 setting
- 8                   • DSM-III versus DSM-III-R (Cole 2003); the test was carried out by :
- 9                   ○ nurse (Cole 2003).
- 10                  • ICD-10 versus DSM-III-R (Cole 2003) the test was carried out by :
- 11                  ○ nurse (Cole 2003).

12  
13 The following tests were compared with the different reference standards:

- 14                  • Reference standard DSM-IV
- 15                  ○ CAM: short version (Gonzalez\* 2004; Hestermann\* 2009;  
16 Laurila\* 2002; Radtke 2008); the test was carried out by a:
- 17                   - geriatrician (Fabbri\* 2001; Laurila\* 2002);
- 18                   - general physician or psychiatrist (Gonzalez\* 2004);
- 19                   - psycho gerontologist and a resident (Hestermann\* 2009);
- 20                   - trained assessor (Radtke 2008).
- 21                  ○ CAM: long version (Fabbri\* 2001; Yates 2009
- 22                   - geriatrician (Fabbri\* 2001)
- 23                   - one of two junior medical doctors (Yates 2009)
- 24                  –
- 25                  ○ CAM-ICU (Ely 2001; Ely 2001b; Lin\* 2004); the test was carried  
26 out by:
- 27                   - two nurses (Ely 2001; Ely 2001b) and an intensivist (Ely  
28 2001b).
- 29                   - a research assistant (Lin\* 2004).
- 30                  ○ DRS-R-98 (Andrew 2009);
- 31                   - Test was carried out by either a geriatrician or a resident.
- 32
- 33                  • Reference standard ICD 10
- 34                  ○ CAM: short version (Laurila\* 2002);
- 35                   - Test was carried out by a geriatrician
- 36
- 37                  • Reference standard DSM IIIR

- 1                   ○ CAM : short version (Laurila\* 2002; Pompei 1995); the test was  
2                   carried out by:
- 3                         - a geriatrician (Laurila\* 2002)  
4                         - a research assistant (Pompei 1995)
- 5                   ○ CAM: long version (Cole 2003; Rockwood 1994; Rolfson 1999b);  
6                   the test was carried out by:
- 7                         - a nurse (Cole 2003)
- 8                   ○ CAM: type of version unclear (Rockwood 1994; Rolfson 1999b)
- 9                         - the study physician (Rockwood 1994)  
10                        - both physician (first 41 patients) and trained research  
11                        nurses (second 30 patients) (Rolfson 1999b).
- 12                   ○ MMSE (Rolfson 1999b);
- 13                        - Unclear whether a physician or nurse carried out the  
14                        assessment.
- 15                   ○ Clock-drawing test (Rolfson 1999b);
- 16                        - Unclear whether a physician or nurse carried out the  
17                        assessment.
- 18                   ○ Delirium Index (DI) (Cole 2003)
- 19                        - Test carried out by a trained research assistant
- 20
- 21                   • Reference standard DSM III
- 22                        ○ AMT (Ni Chonchubhair 1995);
- 23                        - For the reference standard, the study reported that a  
24                        single experienced physician examined patients using the  
25                        Delirium Assessment Scale and determined which patients  
26                        had delirium according to the DSMIII criteria
- 27                        - Unclear who carried out the test.
- 28                   ○ CAM: short version (Laurila\* 2002);
- 29                        - Test carried out by a geriatrician.  
30                        - Reference standard Consensus diagnosis;
- 31                   ○ CAM: long version (Zou 1998);
- 32                   • Test carried out by a nurse.
- 33
- 34                   Additionally, two studies compared different index tests, using CAM (carried out  
35                   by a geriatrician) as a reference standard. These studies are included for  
36                   completeness, but should be considered indirect comparisons for studies of  
37                   diagnostic test accuracy
- 38                   • Reference standard CAM (short version)

- CAM test carried out by one of three lay interviewers. The team of lay interviewers included a nurse without prior research experience, a nurse with some experience as a research interviewer and one research assistant without a nursing degree but with experience as a research interviewer (Monette 2001);
- Reference standard CAM (long version) and Clinician interview
  - MMSE test carried out by one of two trained registrars in geriatric and general internal medicine (O'Keeffe 2005);

### 12.2.5 Outcomes

Methods of reporting outcomes varied:

- One study reported raw data to enable calculation of diagnostic test accuracy, and 2 x 2 tables were constructed (Laurila\* 2003);
- In ten studies the raw data were back-calculated from accuracy measures (Andrew 2009; Cole 2003; Ely 2001; Gonzalez\* 2004; Lin\* 2004; O'Keeffe 2005; Pompei 1995; Radtke 2008; Rockwood 1994; Yates 2009);
- in six studies both the raw data and accuracy measures were reported (Fabbri\* 2001; Laurila\* 2002; Monette 2001; Ni Chonchubhair 1995; Rolfson 1999b; Zou 1998);
- In one study (Ely 2001b), the raw data were obtained by an estimation process in order to reproduce the reported accuracy parameters.

In the Rockwood (1994) study limited raw data was reported. We estimated the number of patients who were delirious and non delirious by assuming the 52 patients (who were referred to the study physician) were roughly equally spread between the two groups.

One study (Laurila\* 2004), provided insufficient raw data and we were unable to calculate accuracy measures.

### 12.3 Methodological quality of included studies

The methodological quality was assessed (Appendix E) using QUADAS criteria.

Most of the studies used a reference standard that was likely to classify the target condition correctly. Two studies (Monette 2001: CAM assessment by geriatrician; O'Keeffe 1997: CAM and clinical interview) used the CAM as the reference standard. In one study (Andrew 2009) it was unclear who performed the assessment.

Generally the studies reported the availability of additional clinical data, for example MMSE scores or other measures indicative of cognitive impairment or

1 dementia, medical records or notes from interviews with family/carers were  
2 available when patients were assessed.

3  
4 Overall, most studies briefly reported the execution of the index test and  
5 reference standard, with the exception of four studies which provided detailed  
6 information on the tests and/or the method of assessments (Ely 2001; Ely 2001b;  
7 Gonzalez\* 2004: index test; Laurila\* 2002). One study (Radtke 2008) reported  
8 that patients were assessed only once in the recovery room and length of stay  
9 ranged between 22 minutes to 147 minutes.

10  
11  
12 None of the studies reported intermediate or uninterpretable results.  
13 Withdrawals (18%: 35/200) in one study (O’Keeffe 2005) were due to deaths,  
14 early discharge or error. Two studies reported missing data (Andrew 2009: 1%,  
15 values were replaced with the mid-range score; Pompei 1995: 0.9% missing  
16 data and were excluded from the analysis);

17  
18 In addition to the above quality issues, the following studies were found to be at  
19 risk of bias on the following criteria:

- 20 • Spectrum bias (Andrew 2009; Cole 2003; Monette 2001; Radtke 2008;  
21 Rolfson 1999b)
- 22 ○ Following first stage CAM assessment by the nurse, patients were  
23 selected from those classified as having probable delirium and no  
24 delirium; the CAM negative group had a higher proportion of  
25 cognitively impaired people (Monette 2001)
  - 26 ○ 30% of the patients were outpatients, of whom 10% were  
27 assessed at home. (Andrew 2009)
  - 28 ○ Case control study in which two groups of patients with and  
29 without delirium were selected (Cole 2003)
  - 30 ○ Patients were in the recovery room following general  
31 anaesthesia. The GDG considered the ordinary version of CAM to  
32 be inappropriate for this environment (Radtke 2008)
  - 33 ○ Patients were undergoing CABG surgery and had a low  
34 proportion with dementia (Rolfson 1999b)
- 35
- 36 • Disease progression bias (Andrew 2009; Inouye 2005; Ni Chonchubhair  
37 1995; O’Keeffe 2005; Rockwood 1994; Rolfson 1999b; Yates 2009;  
38 Zou 1998)
- 39 ○ The authors reported that the index and reference tests were not  
40 necessarily done on the same day, which given the fluctuating  
41 course of delirium, is a limitation. (Andrew 2009);
  - 42 ○ The study reported that reference standard assessment was within  
43 the same day (O’Keeffe 2005)
  - 44 ○ The study reported that the time between assessments varied  
45 between 30 min and 8 hours (Zou 1998)

- 1                   ○ Time period was not reported so the studies were downgraded  
2                   for this quality criterion (Ni Chonchubhair 1995; Rockwood 1994;  
3                   Rolfson 1999b; Yates 2009).  
4
- 5                   ● Partial verification bias (Cole 2003; Pompei 1995)
- 6                   ○ Reference standard appeared to be given only to patients with  
7                   SPMSQ score  $\geq 3$  or delirium symptoms in notes (Cole 2003)
- 8                   ○ Only the patients with an acute change in mental status  
9                   (61%:263/432) were referred to clinician for reference standard  
10                  assessment (Pompei 1995)  
11
- 12                  ● Review bias (Andrew 2009; Cole 2003; Laurila\* 2003; Monette 2001;  
13                  Rockwood 1994; Rolfson 1999b; Yates 2009; Zou 1998)
- 14                  ○ Two studies used the same data for both the reference standard  
15                  and index test and it was very likely that there was review bias  
16                  (Cole 2003; Laurila\* 2003)
- 17                  ○ One study included the index test as part of the reference  
18                  standard; results for DSM-IV as a separate reference standard  
19                  were not reported (Zou 1998)
- 20                  ○ One study had the index and reference tests carried out by the  
21                  same person (Rockwood 1994)
- 22                  ○ One study may have had the index and reference tests carried  
23                  out by the same person/people (Yates 2009)
- 24                  ○ It was unclear whether the index test was interpreted without the  
25                  knowledge of the reference standard, as the nurse [conducting  
26                  the index test] observed the geriatrician [reference standard]  
27                  (Monette 2001)
- 28                  ○ In the Rolfson (1999b) study the CAM assessments were  
29                  administered by a physician [41/71 patients] and a nurse  
30                  administered the CAM for the remaining patients; the same  
31                  physician assessed the reference standard (but the other tests  
32                  were not carried out by the same people)
- 33                  ○ For the rest of the above studies it was unclear whether the  
34                  reference standard was interpreted with the knowledge of the  
35                  result of the index test so studies were downgraded for this  
36                  quality criterion  
37
- 38                  ● Incorporation bias (Cole 2003; Laurila\* 2003; Zou 1998)
- 39                  ○ The index test [CAM administered by the nurse] was part of the  
40                  reference standard [consensus diagnosis] (Zou 1998)
- 41                  ○ The index tests and reference tests were based on the same data  
42                  (Cole 2003; Laurila\* 2003)  
43

Overall, nine studies were considered as potentially or at risk of bias (Andrew 2009; Cole 2003 (all comparisons); Laurila\* 2003 (all comparisons); Monette 2001; Pompei 1995; Rockwood 1994; Rolfson 1999b (for CAM only); Yates 2009; Zou 1998). These studies were considered in sensitivity analyses.

## 12.4 Results – hospital setting

The purpose of the tests examined is to identify delirium, possibly to be used as a screening tool. The GDG stated that they were most interested in a test that had high sensitivity and would 'rule in' patients with delirium. We examined the sensitivity, specificity, positive likelihood ratio and the pre and post test probabilities.

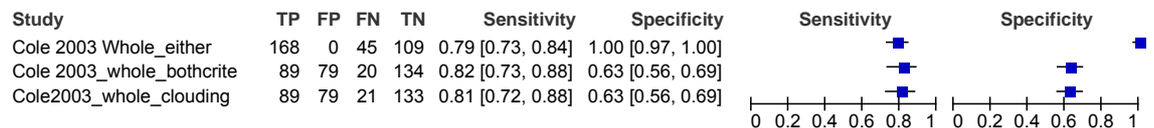
### 12.4.1 Comparison of diagnostic criteria (table 12.1)

One low quality, case control study (Cole 2003) compared different diagnostic criteria; raw data were calculated from the accuracy measures.

#### 12.4.1.1 DSM-III-R versus DSM-IV

One low quality, case control study (Cole 2003) compared DSM-III-R with DSM-IV using the same symptoms to determine both test results, and considered the effect on sensitivity and specificity in relation to criterion A from the DSM-III-R and the DSM-IV (inattention versus clouding of consciousness). The test showed moderate sensitivity: 79%; specificity: 100% when *either* inattention or clouding of consciousness criterion was used. However, when the required criterion was *both* inattention and clouding of consciousness, the sensitivity showed a slight improvement [82%], however, the specificity was compromised [63%] and similar results were reported [sensitivity: 81%; specificity: 63%] when only the clouding of consciousness was the required criterion (figure 12.1).

Figure 12.1: forest plot of DSM-III-R diagnostic test with DSM-IV as a reference standard in a hospital setting

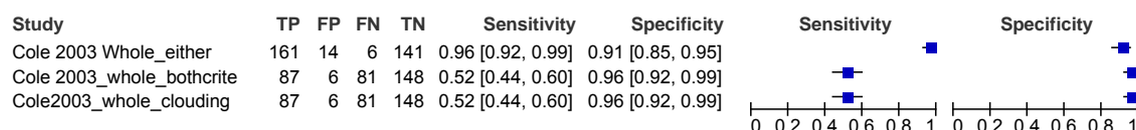


#### 12.4.1.2 DSM III versus DSM-III-R

One low quality, case control study (Cole 2003) compared DSM-III with DSM-III-R and considered the effect on sensitivity and specificity in relation to criterion A (inattention versus clouding of consciousness). The test showed high sensitivity [96%] and specificity [91%] when *either* inattention or clouding of consciousness

1 criterion was used. However, when the required criterion was *both* inattention  
 2 and clouding of consciousness, the sensitivity was compromised [52%], however,  
 3 the specificity slightly improved [96%] and similar results were reported  
 4 [sensitivity: 52%; specificity: 96%] when only the clouding of consciousness was  
 5 the required criterion (figure 12.2).

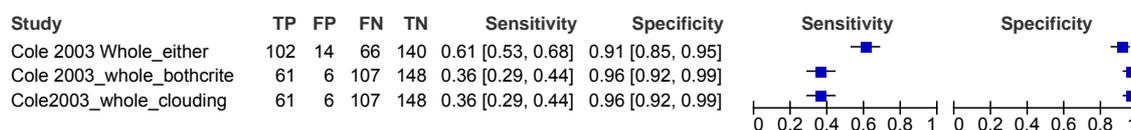
6  
 7  
 8 Figure 12.2: forest plot of DSM-III diagnostic test with DSM-III-R as a reference  
 9 standard in a hospital setting  
 10



11  
 12  
 13  
 14 **12.4.1.3 ICD-10 versus DSM-III-R**

15 One low quality, case control study (Cole 2003) compared ICD-10 with DSM-III-  
 16 R and considered the effect on sensitivity and specificity in relation to criterion A  
 17 (inattention versus clouding of consciousness). The test showed moderate  
 18 sensitivity: 61%; specificity: 91% when *either* inattention or clouding of  
 19 consciousness criterion was used. However, when the required criterion was *both*  
 20 inattention and clouding of consciousness, the sensitivity was low [36%], however,  
 21 the specificity slightly improved [96%] and similar results were reported  
 22 [sensitivity: 36%; specificity: 96%] when only the clouding of consciousness was  
 23 the required criterion (figure 12.3).  
 24

25  
 26  
 27 Figure 12.3: forest plot of ICD-10 diagnostic test with DSM-III-R as a reference  
 28 standard in a hospital setting  
 29



30  
 31  
 32 The DSM-III-R compared with DSM-IV showed moderate sensitivity and a high  
 33 post predictive value (PPV) (which is the proportion of patients with a positive  
 34 test who have the target condition) indicating the DSM-III-R is inclusive. Of the  
 35 two diagnostic tests (DSMIII and ICD-10) compared with DSM-III-R, the ICD-10  
 36 was least inclusive.  
 37

1 Table 12.1: diagnostic test accuracy statistics for different reference standards

Study name	Comments	Test operator	Sensitivity	Specificity	PPV	LR+	Pre-test probability	Post-test probability
Cole 2003	DSM-III-R vs DSM-IV; criterion A: either inattention or clouding of consciousness	Nurse	79.23	100	100	NA	65.84	100
Cole 2003	DSMIII vs DSM-III-R; criterion A: either inattention or clouding of consciousness	Nurse	96.4	90.9	92.1	10.67	6.83	43.9
Cole 2003	ICD10 vs DSM-III-R; criterion A: either inattention or clouding of consciousness	Nurse	60.71	90.92	87.9	6.68	52.17	87.9

2

3 **12.4.1.4 CAM (short version) versus different diagnostic criteria**

4 One moderate quality study (Laurila\* 2002) compared the CAM index test  
5 (short version) with different reference standards. The CAM test, which is based  
6 on the DSM-III-R criteria, showed a moderate sensitivity (80% to 85%) and  
7 specificity (63.4% to 83.7%) against the reference standards. The CAM had the  
8 most concordance with the DSM-IV [sensitivity: 81.3% and specificity: 83.7%]  
9 and was the least concordant with the ICD-10 [sensitivity: 80% and specificity:  
10 63.4%]; table 12.2.

11  
1213 Table 12.2: diagnostic test accuracy statistics for CAM for different reference  
14 standards

CAM index test (short version)	Study name	Comments	Test operator	Sensitivity	Specificity	PPV	LR+	Pre-test probability	Post-test probability
DSM-IV	Laurila * 2002	CAM vs DSM-IV	Geriatrician	81.3	83.7	76.0	5.0	39.5	76.5
ICD-10	Laurila * 2002	CAM vs ICD-10	Geriatrician	80.0	63.4	24.0	2.2	12.3	23.5
DSM IIIIR	Laurila * 2002	CAM vs DSMIII-R	Geriatrician	81.0	71.7	50.0	2.9	25.9	50.0
DSM III	Laurila * 2002	CAM vs DSMIII	Geriatrician	85.0	72.1	50.0	3.1	24.7	50.0

15

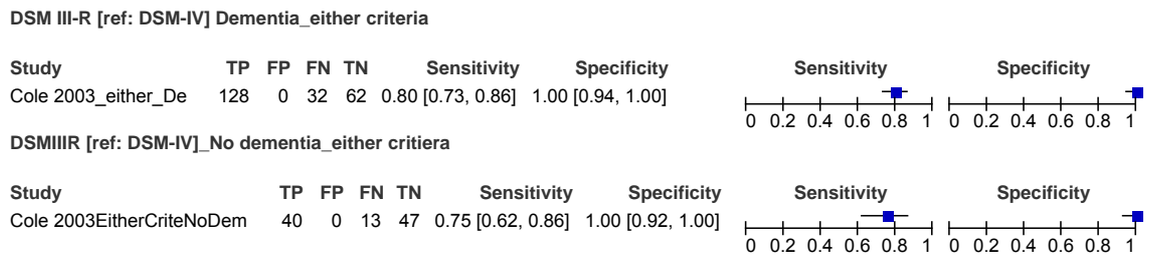
1 **12.4.1.5 Subgroup analysis by dementia or no dementia**

2 The Cole (2003) study reported separately the accuracy measures for different  
 3 diagnostic criteria in patients with and without dementia. Dementia was  
 4 diagnosed with the IQCODE.  
 5

6 **12.4.1.6 DSM-III-R versus DSM-IV**

7 The DSM-III-R instrument (compared with DSM-IV) shows a slightly higher  
 8 sensitivity in people with dementia [80%] than in people without dementia  
 9 [range: 75%] when the criterion A is interpreted as either clouding of  
 10 consciousness or inattention. A forest plot of sensitivity and specificity is shown in  
 11 figure 12.4, but we note that the study used both tests to interpret the same  
 12 symptoms.  
 13  
 14

15 Figure 12.4: forest plot of DSM-III-R compared with DSM-IV in a hospital setting-  
 16 subgroup analyses

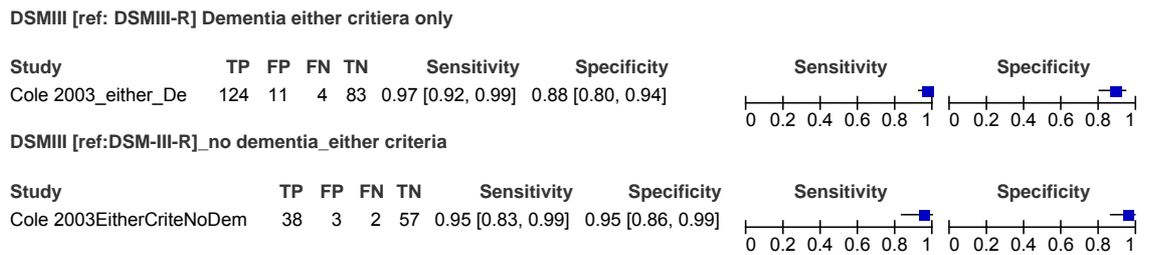


17  
 18  
 19

20 **12.4.1.7 DSM-III versus DSM-III-R**

21 The DSM-III instrument (compared with DSM III-R) shows a high sensitivity and the  
 22 ability of the test to rule in those with delirium is high and this is the case whether  
 23 the patients have dementia [sensitivity: 97%] or not [sensitivity: 95%]; figure  
 24 12.5. The reported results are for criterion A being interpreted as either  
 25 clouding of consciousness or inattention.  
 26

27 Figure 12.5: forest plot of DSM-III-R compared with DSM-III-R in a hospital  
 28 setting - subgroup analyses



29  
 30

### 1 12.4.1.8 ICD10 versus DSM-III-R

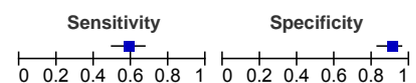
2 The ICD-10 instrument (compared with DSM III-R) showed a fairly low sensitivity  
3 and this is the case for patients with dementia [sensitivity: 59%] or for patients  
4 without dementia [sensitivity: 68%]; figure 12.6. The reported results are for  
5 criterion A being interpreted as either clouding of consciousness or inattention.  
6

7 Figure 12.6: forest plot of ICD-10 compared with DSM-III-R in a hospital setting-  
8 subgroup analyses

9

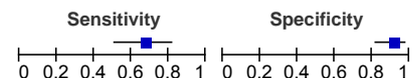
ICD 10 [ref: DSM-III-R] Dementia \_either criteria

Study	TP	FP	FN	TN	Sensitivity	Specificity
Cole 2003_either_De	75	9	53	85	0.59 [0.50, 0.67]	0.90 [0.83, 0.96]



ICD10 [ref:DSM-III-R] No dementia \_either criteria

Study	TP	FP	FN	TN	Sensitivity	Specificity
Cole 2003EitherCriteNoDem	27	5	13	55	0.68 [0.51, 0.81]	0.92 [0.82, 0.97]



10  
11

12

### 13 12.4.2 Diagnostic test accuracy (DSM-IV as the reference standard)

14 Seven studies compared index tests with DSM-IV as the reference standard: four  
15 investigated CAM short version (Gonzalez\* 2004; Hestermann\*2009; Laurila\*  
16 2002; Radtke 2008); two studies investigated CAM long version [Fabbri\* 2001;  
17 Yates 2009 (low)]; and one study investigated the DRS-R-98 [(Andrew 2009  
18 (low)].

19

20 A forest plot of sensitivity and specificity is shown in figure 12.7. The GDG  
21 agreed that the CAM long version, which assessed for 10 symptoms (acute onset,  
22 inattention, disorganised thinking, altered level of consciousness, disorientation,  
23 memory impairment, perceptual disturbances, psychomotor agitation,  
24 psychomotor retardation) and the CAM short version, which assessed for 3  
25 symptoms (acute onset, inattention, disorganised thinking or altered level of  
26 consciousness) of delirium, should be treated separately and these are reported  
27 as subgroups. The diagnostic test accuracy statistics are summarised in table  
28 12.3.

29

#### 30 12.4.2.1 DRS-R-98

31 One low quality study (Andrew 2009) assessed the DRS-R-98 with DSM-IV  
32 showed a moderate specificity and fairly low sensitivity [sensitivity: 56%;  
33 specificity: 82%]. The study included patients with dementia (40%), had a high  
34 proportion of inpatients (73%), with high comorbidity [mean co-morbidity count  
35 7.1 (SD 2.7)]. The study also examined a sub-sample of patients with underlying  
36 dementia, which had a sensitivity of 59% and a specificity of 67%. The study

1 reported that the assessors of the index test had varying expertise and did not  
2 have extensive training in the use of the instrument; the study showed a  
3 moderate inter-rater reliability ( $k=0.76$ ).

4  
5 The number of patients identified with the DRS-R-98 instrument as delirious have  
6 a small likelihood of being delirious [likelihood ratio: 3.17]. However, the results  
7 are based on one low quality study so some uncertainty exists on DRS-R-98  
8 utility as a screening instrument for delirium.  
9

#### 10 **12.4.2.2 CAM**

11 Of the six studies [Fabbri\* 2001; Gonzalez\* 2004; Hestermann\* 2009; Laurila\*  
12 2002; Radtke 2008; Yates 2009 (low)] comparing CAM, we note that four of  
13 these (Fabbri\* 2001; Gonzalez\* 2004; Hestermann\* 2009; Laurila\* 2002) used  
14 a foreign language version of the CAM: Portuguese, Spanish, German, and  
15 Finnish respectively. The Gonzalez\* (2004) study reported that in order to  
16 further assess the onset and course of the mental status changes and to evaluate  
17 thinking and attention, items from the Spanish version of the MMSE were included  
18 in the interview – so this study was considered as an adaptation study.  
19

20 Two of the studies (Fabbri\* 2001; Hestermann\* 2009) reported that the  
21 instrument was translated and back translated and in the other two studies  
22 (Gonzalez\* 2004; Laurila\* 2002) the final version of the instrument was based  
23 on expert panel consensus.  
24

25 In all of the studies, the CAM was rated by a physician, with the exception of the  
26 Yates (2008) study, where a trained assessor administered the instrument (CAM  
27 long version).  
28

29 For the CAM short version, the sensitivity ranged from 43% to 90% and the  
30 specificity from 84% to 100%. The positive predictive value ranged from: 76%  
31 to 100% and likelihood ratio ranged from: 5.0 to 28.5.

32 There was heterogeneity, particularly for sensitivity and some variation in the  
33 specificity. Heterogeneity was considered in terms of the following factors:  
34 language and type of patients. As noted earlier, assessment was carried out with  
35 a foreign language version of the CAM in three studies (Gonzalez\* 2004;  
36 Hestermann\* 2009; Laurila\* 2002). We note that the Radtke (2008) study,  
37 conducted in Germany, reported patients who did not speak the local language  
38 were excluded; however, it was unclear if the CAM instrument was a version  
39 translated into the local language.

40 In terms of type of patients included in the study, we note the Radtke (2008)  
41 study was the only study which included patients with a mean age below 65  
42 years (mean [range]: 54.5 years [25.4 to 80.8]) and the study included patients  
43 who were in the recovery following general anaesthesia. The GDG considered  
44 whether the ordinary version of CAM to be inappropriate for this environment.  
45

46 The type of patients included, the setting and the inappropriate measure for this  
47 setting may account for the low sensitivity [43%] observed in the Radtke (2008)  
48 study.  
49

1 For the CAM long version, the sensitivity ranged from 91% to 94% and the  
2 specificity was 96%. We note the Yates (2009) study was poor quality.

3  
4 The CAM instrument when compared with DSM-IV as the reference standard,  
5 was able to detect delirium and the likelihood of patients having delirium when  
6 CAM had identified patients as being delirious is high.  
7

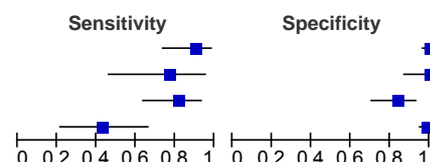
8  
9 Table 12.3: diagnostic test accuracy statistics for DSM-IV as the reference  
10 standard

DSM-IV	Study name	Comments	Test operator	Sensitivity	Specificity	PPV	LR+	Pre-test probability
CAM Long version	Fabbri* 2001	CAM [geriatrician] vs DSMIV [psychiatrist]	Geriatrician	94.1	96.4	84.0	26.0	17.0
	Yates 2009	CAM vs DSM-IV	Study physician	90.90	96.10	83.00	23.2	17.7
CAM Short version	Gonzalez * 2004	CAM vs DSMIV	General Physician or Psychiatrist	90.0	100.0	100.0	NA	24.4
	Hesterma nn * 2009	CAM [rater 1= psychogerontologist] vs DSM-IV[consensus]	Psychologist / Gerontologist and Resident	76.9	96.2	91.0	20	33.3
	Hesterma nn* 2009	CAM [rater2= internal resident in geriatric medicine] vs DSM-IV[consensus]	Psychologist/G erontologist and Resident	76.9	100.0	100.0	NA	33.3
	Laurila* 2002	CAM vs DSM-IV	Geriatrician	81.3	83.7	76.0	5.0	39.5
	Radtke 2008	CAM vs DSM-IV	Trained assessor (trained by psychiatrist)	42.9	98.5	82.0	28.5	13.6
DRS-R-98	Andrew 2009	Index: DRS-R98 Ref: 'clinically diagnosed delirium'=DSMIV	Geriatrician/ Resident	56.40	82.20	66.00	3.2	37.9

1 Figure 12.7: forest plot of index tests compared with DSM-IV in a hospital  
 2 setting

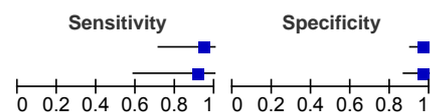
CAM short version [ref: DSMIV]

Study	TP	FP	FN	TN	Sensitivity	Specificity
Gonzalez 2004	27	0	3	93	0.90 [0.73, 0.98]	1.00 [0.96, 1.00]
Hestermann 2009_Rater 2	10	0	3	26	0.77 [0.46, 0.95]	1.00 [0.87, 1.00]
Laurila 2002	26	8	6	41	0.81 [0.64, 0.93]	0.84 [0.70, 0.93]
Radtke 2008	9	2	12	131	0.43 [0.22, 0.66]	0.98 [0.95, 1.00]



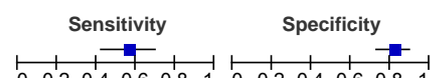
CAM long version [ref: DSMIV]

Study	TP	FP	FN	TN	Sensitivity	Specificity
Fabbri 2001	16	3	1	80	0.94 [0.71, 1.00]	0.96 [0.90, 0.99]
Yates 2009	10	2	1	49	0.91 [0.59, 1.00]	0.96 [0.87, 1.00]



DRS-R-98 [Ref: DSM-IV]

Study	TP	FP	FN	TN	Sensitivity	Specificity
Andrew 2009	31	16	24	74	0.56 [0.42, 0.70]	0.82 [0.73, 0.89]



3

4

5

6 **12.4.2.3 Subgroup analyses by dementia or no dementia**

7

8 Subgroup analyses for DRS-R-98 compared with DSM-IV

9 One low quality study (Andrew 2009) reported subgroup analyses for patients  
 10 with and without dementia for the DRS-R-98 test compared with DSM-IV as  
 11 reference standard.

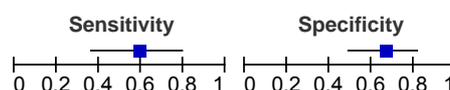
12  
 13 Dementia was diagnosed with DSM-IV and the number of patients with dementia  
 14 and underlying dementia with superimposed delirium was 58. The study showed  
 15 low sensitivity and specificity, 59% and 67%, respectively (figure 12.8). We  
 16 note that this study was considered low quality.

17

18

19 Figure 12.8: forest plot of DRS-R-98 compared with DSM-IV in a hospital  
 20 setting- subgroup analysis

Study	TP	FP	FN	TN	Sensitivity	Specificity
Andrew 2009	13	12	9	24	0.59 [0.36, 0.79]	0.67 [0.49, 0.81]



21

22

23 Subgroup analyses for CAM (short version) compared with DSM-IV

24 One moderate quality study (Gonzalez\* 2004) reported the diagnostic  
 25 accuracy measures for the CAM test (short version) compared with DSM-IV as  
 26 reference in people with and without dementia. Dementia was diagnosed on the  
 27 basis of DSM-IV criteria, medical records, MMSE rating, and interviews with  
 28 relatives. The study did not provide the number of patients diagnosed with  
 29 delirium for the subgroups so we were unable to back-calculate the raw data.

The Spanish translation of the CAM (short version) showed a slightly lower sensitivity in people with dementia [sensitivity: 87%] compared to people without dementia [sensitivity: 93%]; the specificity was similar for both groups [100%].

### 12.4.3 ICD-10 as reference standard

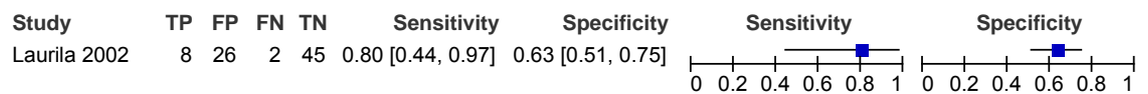
One moderate quality study (Laurila\* 2002) compared CAM (short version) with ICD-10 as a reference standard. We note that in this study, four reference standards [DSM-IV, DSM-III-R, DSM-III, and ICD-10] were operationalised in one questionnaire. The index test was a previously validated foreign language [Finnish] version of the CAM, which was developed by consensus.

The forest plot showing the specificity and sensitivity is shown in figure 12.9. The CAM (short version) showed moderate sensitivity [80%] with the ICD-10 classification, however, the specificity was fairly low [63%].

Although the positive predictive value is 24%, the negative predictive value is 96% which indicates that a negative result on the CAM test is able to exclude delirium. The low positive likelihood ratio of 2.18 indicating that a patient identified with delirium using the CAM instrument for assessment is 2.18 more likely to be delirious than non delirious.

As shown earlier in section 12.4.1.3, the ICD-10 diagnostic criteria (compared with DSM-III-R), performs poorly in relation to specificity and may have some limitations as a reference standard.

Figure 12.9: forest plot of CAM compared with ICD-10 in a hospital setting-subgroup analysis



### 12.4.4 DSM-III-R as the reference standard

Two studies compared CAM short version with DSM-III-R (Laurila\* 2002; Pompei 1995 (low)); one study compared CAM long version with DSM-III-R (Cole 2003 (low)); and type of version was unclear in two studies (Rockwood 1994 (low); Rolfson 1999b (partly low)). One study (Rolfson 1999b) also gave the patients other index tests compared with DSM-III-R [MMSE; clock-drawing test] – the study quality was considered to be moderate for these tests.

1 A forest plot of sensitivity and specificity is shown in figure 12.10. Results for the  
2 CAM short and long versions are reported as subgroups. The diagnostic test  
3 accuracy statistics are summarised in table 12.4.

4 The low quality Cole (2003) study also reported classification of delirium by  
5 number of symptoms for the CAM and DI; this is reported separately under  
6 section X.4.4.5. In Figure 11, for the Cole (2003) study, the values for more than  
7 6 symptoms and more than 4 symptoms are used respectively. We note that the  
8 same data were used for the CAM and reference standard, but a separate test  
9 was carried out for the DI, so the CAM results are likely to be more biased.

10

#### 11 **12.4.4.1 CAM**

12 Two studies compared CAM short version with DSM-III-R (Laurila\* 2002; Pompei  
13 1995 (low)); one study compared CAM long version with DSM-III-R (Cole 2003  
14 (low)); and type of version was unclear in two studies (Rockwood 1994 (low);  
15 Rolfson 1999b (partly low)).

16  
17 The Cole (2003) study used the CAM (long version) to determine 10 symptoms  
18 which were used for the reference standard. The study reported the sensitivity  
19 and specificity (for more than 6 symptoms) for patients with dementia or without  
20 dementia. The sensitivity and the specificity was 98% and 76% for patients with  
21 dementia and 95% and 83% for patients without dementia. We note this was a  
22 case control study; therefore the sensitivity and specificity are likely to be  
23 overestimated.

24  
25  
26 The two studies (Laurila\* 2002; Pompei 1995 (low)) comparing CAM short  
27 version with DSM-III-R showed sensitivity ranging from 46% to 81% and  
28 specificity ranging from 72% to 92%. A sensitivity analysis was carried out  
29 excluding the low quality studies. Considering the remaining study (Laurila\*  
30 2002), which was of moderate quality, the CAM showed an 81% sensitivity and  
31 72% specificity compared with DSM-III-R. The positive predictive accuracy was  
32 50% and the negative predictive value was 91%, indicating that a negative  
33 result on the CAM instrument will accurately exclude delirium. The likelihood ratio  
34 is 2.86, which suggests a not particularly strong test.

35  
36  
37 In two studies (Rockwood 1994 (low); Rolfson 1999b (low)) the type of version  
38 used was unclear. The Rolfson (1999) study reported that the CAM and  
39 reference standard were carried out by the same physician for 41 patients and  
40 by different assessors for the next 30 patients: for the latter, assessment was by  
41 nurses, and these results are considered to be low quality. The results are  
42 reported separately for the two groups.

43  
44 The Rockwood (1994) study reported the sensitivity [64%] and specificity [93%],  
45 however, there was insufficient information and we were unable to calculate the  
46 raw data from the reported accuracy measures, although a rough estimate was  
47 obtained by assuming the 52 patients were roughly equally spread between  
48 delirium positive and delirium negative; the study is not included in the forest  
49 plot.

1

2 **12.4.4.2 Clock- drawing and MMSE tests**

3 Both the MMSE and the clock-drawing test index tests were administered on the  
4 day prior to surgery and on the fourth day postoperative day in the Rolfson  
5 (1999) study; results were reported for the latter time. The MMSE showed a low  
6 sensitivity, 35%, a small positive likelihood ratio of 1.9. It was unclear in the  
7 study how many patients had impaired communication which would not allow the  
8 MMSE to be administered (albeit patients with coma before day 4 were  
9 excluded).

10

11 The clock-drawing test showed a very low sensitivity of 9%, and a positive  
12 likelihood ratio of 4.2. It was unclear whether patients had been assessed with  
13 impaired writing ability at baseline as the administration of this index test in such  
14 population would be limited.

15

16 **12.4.4.3 Test comparison**

17 Overall, the CAM performed better than the MMSE or the clock-drawing tests;  
18 although this is based on different studies and there was variation in the index  
19 and reference test assessors.

20

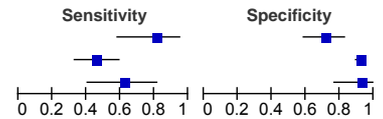
1 Table 12.4: index test compared with DSM-III-R (the pale blue shading indicates  
2 moderate quality studies)

DSM-III-R	Study name	Comments	Test operator	Sensitivity	Specificity	PPV	LR+	Pre-test probability	Post-test probability
CAM Long Version	Cole 2003	CAM >6 symptoms vs DSM III-R for patients with dementia	Nurse	97.7	75.0	84.0	4.0	57.7	84.5
	Cole 2003	CAM >6 symptoms vs DSM III-R for patients without dementia	Nurse	95.0	83.3	79.0	5.7	40.0	79.2
CAM Short Version	Laurila* 2002	CAM vs DSMIII-R	Geriatrician	81.0	71.7	50.0	2.9	25.9	50.0
	Pompei 1995	CAM vs DSMIIIR without 4 patients for whom no results	Research Assistant	45.9	92.1	49.0	5.8	14.3	49.1
CAM type of version unclear	Rockwood 1994	CAM vs DSMIIIR raw data estimated based on sensitivity and specificity	Study physician	63.0	93.0	88.2	8.75	46.15	88.2
	Rolfson 1999b	CAM nurse	Nurse	12.5	100.0	100.0	NA	26.7	100.0
	Rolfson 1999b	CAM [physician] vs DSM III-R [geriatrician]	Physician	69.6	100.0	100.0	NA	32.4	100.0
MMSE	Rolfson 1999b	MMSE vs DSM III-R	Nurse/physician	34.8	81.2	47.0	1.9	32.4	47.0
Clock Drawing	Rolfson 1999b	Clock-drawing test vs DSM III-R	Nurse/physician	8.7	97.9	67.0	4.2	32.4	66.7

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6 Figure 12.10: forest plot of index test compared with DSM-III-R

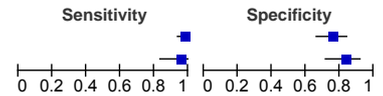
## CAM short version [ref: DSM III-R]

Study	TP	FP	FN	TN	Sensitivity	Specificity
Laurila 2002	17	17	4	43	0.81 [0.58, 0.95]	0.72 [0.59, 0.83]
Pompei 1995	28	28	33	338	0.46 [0.33, 0.59]	0.92 [0.89, 0.95]
Rockwood 1994	15	2	9	26	0.63 [0.41, 0.81]	0.93 [0.76, 0.99]



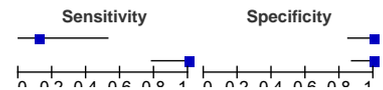
## CAM long version [ref: DSM-III-R]

Study	TP	FP	FN	TN	Sensitivity	Specificity
Cole 2003 Dementia more than 6 smptoms	125	24	3	75	0.98 [0.93, 1.00]	0.76 [0.66, 0.84]
Cole 2003 No dementia; more than 6 symptoms	38	10	2	50	0.95 [0.83, 0.99]	0.83 [0.71, 0.92]



## CAM [type of version unclear] [ref: DSM-III-R]

Study	TP	FP	FN	TN	Sensitivity	Specificity
Rolfson 1999b_nurseassess	1	0	7	22	0.13 [0.00, 0.53]	1.00 [0.85, 1.00]
Rolfson 1999b_physician	15	0	0	26	1.00 [0.78, 1.00]	1.00 [0.87, 1.00]

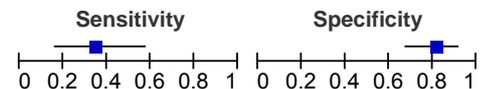


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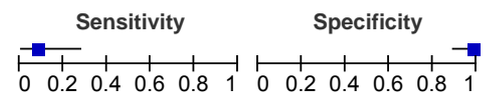
## MMSE [ref: DSM-III-R]

Study	TP	FP	FN	TN	Sensitivity	Specificity
Rolfson 1999b	8	9	15	39	0.35 [0.16, 0.57]	0.81 [0.67, 0.91]



## Clock drawing test [ref: DSM-IIIR]

Study	TP	FP	FN	TN	Sensitivity	Specificity
Rolfson 1999b	2	1	21	47	0.09 [0.01, 0.28]	0.98 [0.89, 1.00]



3

4

## 5 12.4.4.4 Subgroup analyses

6 One low quality study (Pompei 1995) reported subgroup analyses for patients  
 7 (21%: 96/438) with impaired cognitive status on admission. Cognitive status was  
 8 assessed with the MMSE (range 0 to 30); with varying cut-off points adjusted for  
 9 education level (score less than 21 was indicative of cognitive impairment for  
 10 those with less than a high school; score less than 23 points was indicative of  
 11 cognitive impairment for those with high school experience; and score less than  
 12 24 points was indicative of cognitive impairment for those with college  
 13 education).

14  
 15 The study showed moderate/low sensitivity and specificity, 54% and 79%,  
 16 respectively and a likelihood ratio of 2.6. The CAM's ability to screen patients  
 17 with delirium when presented with underlying cognitive impairment was  
 18 moderately compromised; however, we note that this study was of low quality.

19  
 20 The Cole (2003) study reported the sensitivity and specificity for patients with  
 21 dementia [69%: n=222/322; sensitivity: 100.0%; specificity: 96.8%] and those  
 22 without dementia [31%: n=100/322; sensitivity: 100.0%; specificity: 98.3%].  
 23 We note that this study was low quality and the same symptoms were used to  
 24 determine the index test and reference standard results.  
 25

1 **12.4.4.5 Within group comparisons**

2 One study (Cole 2003) separately compared the CAM (long version) and the  
 3 Delirium Index (DI) with the DSM-III-R to identify the sensitivity and specificity of  
 4 number of symptoms of delirium, irrespective of the type of symptoms. We note  
 5 that this was a low quality case control study and that the same data were used  
 6 for the CAM and the reference standard, but a separate test was carried out for  
 7 the DI. This makes a direct comparison between CAM and DI unreliable (figure  
 8 12.11)

9  
 10 As shown in figure 12.12, the ROC plot that explores the effect of varying  
 11 thresholds on sensitivity and specificity in a single study, the presence of 6 or  
 12 more number of symptoms of delirium on the CAM (long version) compared with  
 13 the DSM-III-R criteria was considered the best threshold point. This cut-off point  
 14 was similar for patients with and without dementia.

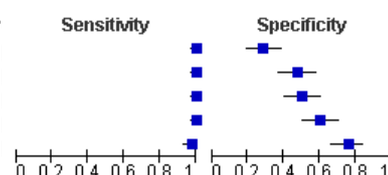
15  
 16 We note this is a poor quality study and the same symptoms were used to  
 17 determine the index test and reference standard results.

18  
 19 On the Delirium Index instrument, the presence of 4 or more symptoms and 3 or  
 20 more symptoms showed the best sensitivity and specificity in patients with and  
 21 without dementia, respectively.

22  
 23  
 24 Figure 12.11: forest plot of number of symptoms in index tests compared with  
 25 DSMIII-R as the reference standard in a hospital setting

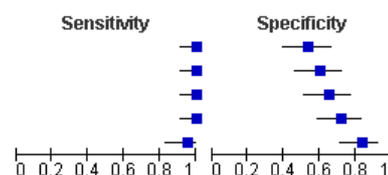
**CAM [number of symptoms] [ref: DSMIII-R] patients with dementia**

Study	TP	FP	FN	TN	Sensitivity	Specificity
Cole 2003 Dementia >2	128	67	0	27	1.00 [0.97, 1.00]	0.29 [0.20, 0.39]
Cole 2003 Dementia >3	128	52	0	47	1.00 [0.97, 1.00]	0.47 [0.37, 0.58]
Cole 2003 Dementia >4	128	49	0	50	1.00 [0.97, 1.00]	0.51 [0.40, 0.61]
Cole 2003 Dementia >5	128	39	0	60	1.00 [0.97, 1.00]	0.61 [0.50, 0.70]
Cole 2003 Dementia more than 6 smptoms	125	24	3	75	0.98 [0.93, 1.00]	0.76 [0.66, 0.84]



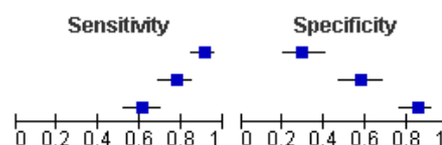
**CAM [number of symptoms] [ref: DSMIII-R] no dementia**

Study	TP	FP	FN	TN	Sensitivity	Specificity
Cole 2003 No dementia >2	40	28	0	32	1.00 [0.91, 1.00]	0.53 [0.40, 0.66]
Cole 2003 No dementia >3	40	24	0	36	1.00 [0.91, 1.00]	0.60 [0.47, 0.72]
Cole 2003 No dementia >4	40	21	0	39	1.00 [0.91, 1.00]	0.65 [0.52, 0.77]
Cole 2003 No dementia >5	40	17	0	43	1.00 [0.91, 1.00]	0.72 [0.59, 0.83]
Cole 2003 No dementia >6sy	38	10	2	50	0.95 [0.83, 0.99]	0.83 [0.71, 0.92]



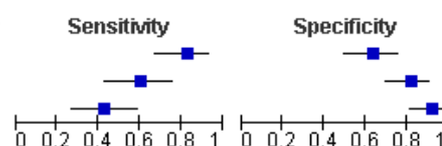
**DI- dementia**

Study	TP	FP	FN	TN	Sensitivity	Specificity
Cole 2003 Dementia >2	116	66	12	28	0.91 [0.84, 0.95]	0.30 [0.21, 0.40]
Cole 2003 Dementia >3	99	40	29	54	0.77 [0.69, 0.84]	0.57 [0.47, 0.68]
Cole 2003 Dementia >4	78	14	50	80	0.61 [0.52, 0.69]	0.85 [0.76, 0.92]



**DI- no dementia**

Study	TP	FP	FN	TN	Sensitivity	Specificity
Cole 2003 No dementia >2	33	22	7	38	0.82 [0.67, 0.93]	0.63 [0.50, 0.75]
Cole 2003 No dementia >3	24	11	16	49	0.60 [0.43, 0.75]	0.82 [0.70, 0.90]
Cole 2003 No dementia >4	17	5	23	55	0.42 [0.27, 0.59]	0.92 [0.82, 0.97]

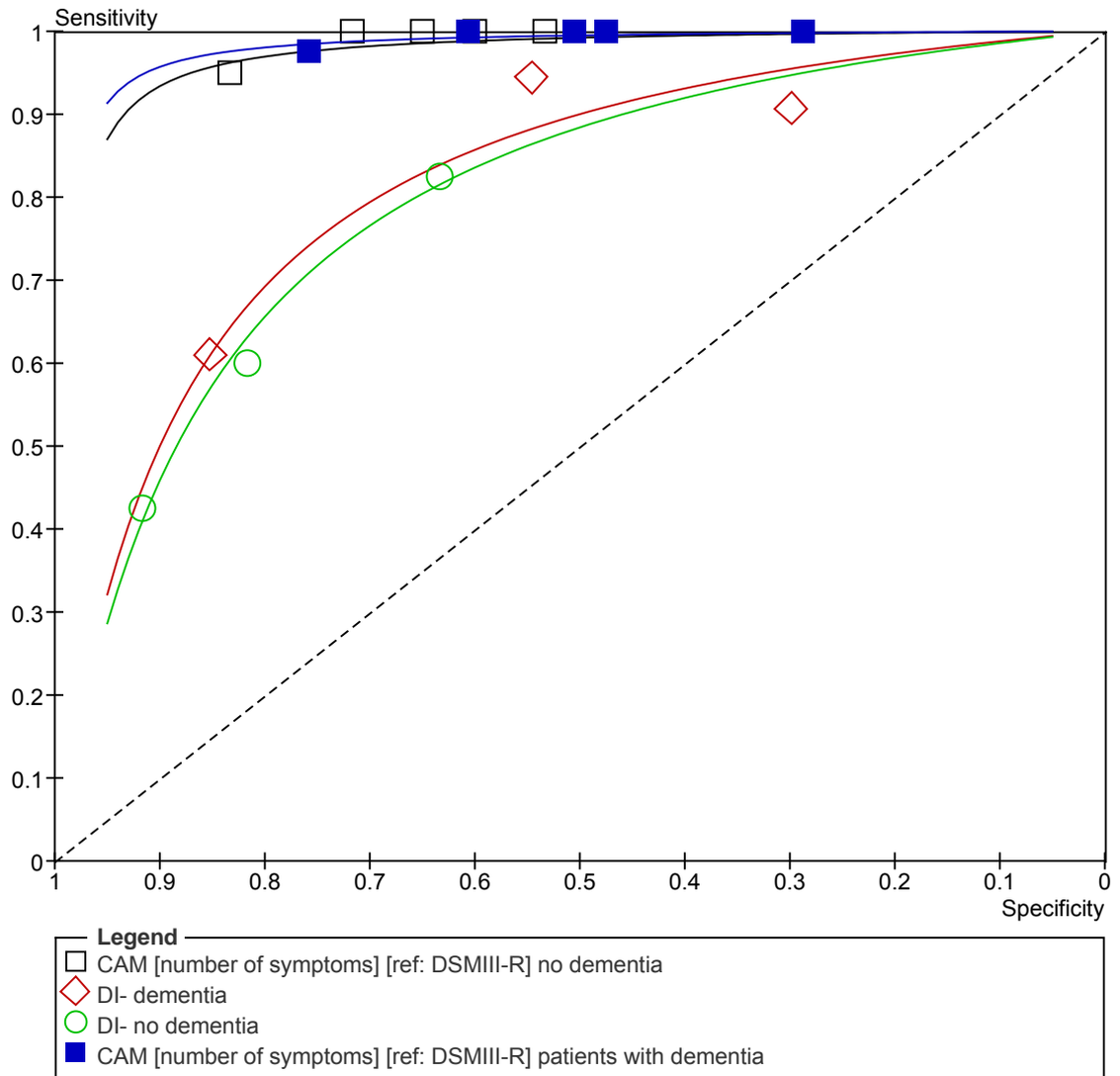


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2 Figure 12.12: ROC plot of effects of varying threshold for CAM and DI  
 3 compared with DSM-III-R



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#### 7 **12.4.4.6 DSM III as the reference standard**

8 Two studies (Laurila\* 2002; Ni Chonchubhair 1995) reported an index test  
 9 compared with DSM III as the reference standard. A forest plot of sensitivity and  
 10 specificity is shown in figure 12.13, and the diagnostic test accuracy statistics are  
 11 summarised in table 12.5.

12

#### 13 **12.4.4.7 AMT serial test**

One study (Ni Chonchubhair 1995) compared the change in AMT scores using the Delirium Assessment Scale to determine delirium according to the DSM III criteria. A 2 point decrease between preoperative and postoperative AMT score showed high sensitivity and specificity, 93% and 84%, respectively. A 3 point decline in AMT scores showed a lower sensitivity [67%] and higher specificity [95%].

The ROC curve (figure 12.15), shows a 2 point change threshold performs better.

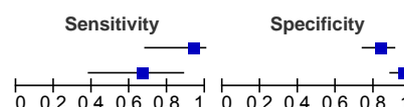
#### 12.4.4.8 CAM

One study (Laurila\* 2002) comparing CAM (short version) with DSM-III showed a moderate sensitivity and specificity [85% and 82%, respectively]. The ability of the instrument to exclude the condition is still high [94%]; but the positive likelihood ratio is low [3.05].

Figure 12.13: forest plot of index tests with DSM-III as the reference standard in a hospital setting

AMT [ref: DSMIII]- decline in score 2 points & 3 points

Study	TP	FP	FN	TN	Sensitivity	Specificity
Ni Chonchubhair 1995-2 pt	14	14	1	71	0.93 [0.68, 1.00]	0.84 [0.74, 0.91]
Ni Chonchubhair 1995-3pt	10	4	5	81	0.67 [0.38, 0.88]	0.95 [0.88, 0.99]



CAM [ref: DSM III]

Study	TP	FP	FN	TN	Sensitivity	Specificity
Laurila 2002	17	17	3	44	0.85 [0.62, 0.97]	0.72 [0.59, 0.83]

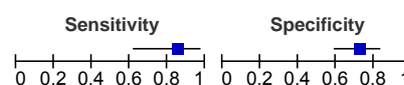
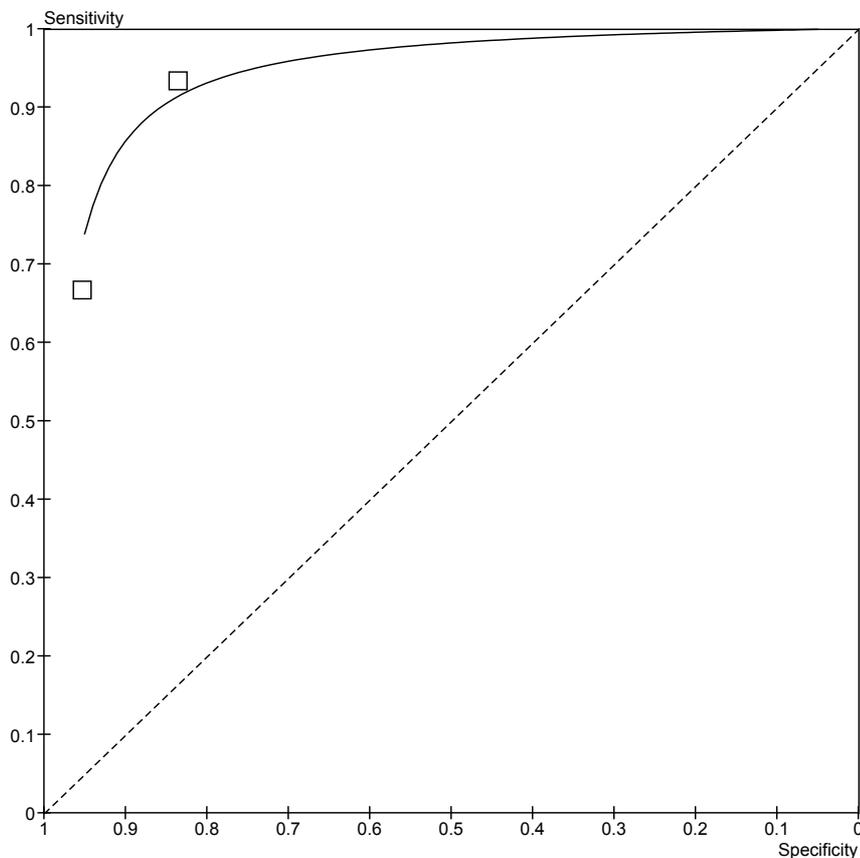


Table 12.5: index test compared with DSM-III-R

DSM-III	Study name	Comments	Test operator	Sensitivity	Specificity	PPV	LR+	Pre-test probability	Post-test probability
CAM short version	Laurila *2002		Geriatrician	85.00	72.10	50.00	3.05	24.70	50.00
AMT	Ni Chonchubhair 1995	Cut off at decline of 3 points or more	Not stated / unclear	66.70	95.30	71.00	14.17	15.00	71.40
AMT	Ni Chonchubhair 1995	Cut off at decline of 2 points or more	Not stated / unclear	93.30	83.50	50.00	5.67	15.00	50.00

1 Figure 12.14: ROC curve - AMT

2  
34 **12.4.5 Consensus diagnosis as a reference standard**

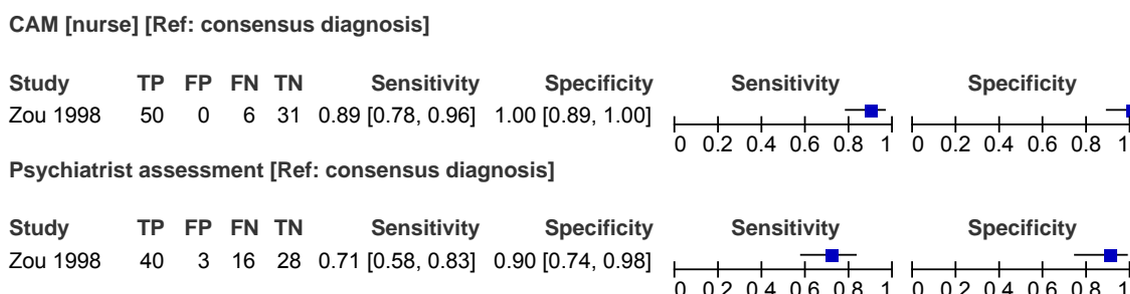
5 One low quality study (Zou 1998) reported separately the sensitivity and  
 6 specificity for two index tests [nurse assessed CAM (long version) and psychiatrist  
 7 assessment] compared with a reference standard (expert consensus diagnosis);  
 8 the expert group comprised two geriatric psychiatrists, a research fellow and a  
 9 nurse. The consensus diagnosis was comprised of the following: psychiatrist's  
 10 findings from a chart review and clinical examination; each professional's  
 11 independent assessment on the presence or absence of delirium  
 12 based on the psychiatrist's application of the DSM-IV criteria and the nurse's  
 13 findings from the CAM and chart review. The forest plot of the sensitivity and  
 14 specificity is shown in figure 12.15. The nurse's CAM rating showed a higher  
 15 sensitivity [89%] than the psychiatrist diagnosis [71%]. The authors attributed  
 16 this partly to the fact the nurse had more opportunities to observe and reassess  
 17 the patient, as opposed to the psychiatrist who assessed the patient only once.

18  
 19 The results from the study should be treated with caution as this was considered  
 20 a low quality study.  
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Figure 12.15: forest plot of index test compared with consensus diagnosis as the reference standard in a hospital setting



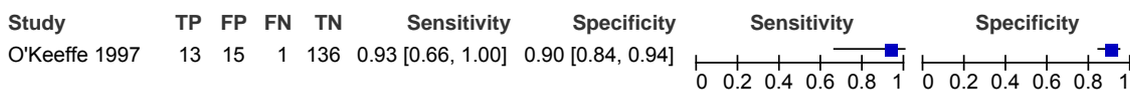
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**12.4.6 CAM (short version) and expert interviewer as the reference standard; MMSE serial test**

One study (O’Keeffe 2005) examined the change in the MMSE scale between day 1 and day 6 of hospitalisation, to identify the best determinant for detecting the development and resolution of delirium. The diagnosis of delirium was with the CAM (short version) instrument and clinician interview.

The study found, for the detection of delirium, a decline of 2 or more points was the best determinant. The sensitivity and specificity were 93% and 90% respectively (figure 12.16). There was some uncertainty with the raw data which were back calculated from the diagnostic accuracy measures. The diagnostic test accuracy statistics are summarised in table 12.6.

Figure 12.16: forest plot of index test compared with CAM (short version) and clinical interview as the reference standard



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Table 12.6: index test compared with CAM (short version) and clinician interview

CAM + interview by experienced clinician	Study name	Comments	Test operator	Sensitivity	Specificity	PPV	LR+	Pre-test probability	Post-test probability
MMSE (serial change)	O’Keeffe 2005	Some uncertainty with the raw	Trained assessor	92.90	90.10	46.00	8.9	8.48	46.40

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**12.4.7 Comparison of different assessors for CAM (short version)**

One low quality study (Monette 2001) compared CAM (short version) assessment by a lay interviewer with a geriatrician; there was no reference standard in this study. The team of lay interviewers included a nurse without prior research experience, a nurse with some experience as a research interviewer or an experienced research assistant without a nursing degree but with experience as a research interviewer.

**12.4.7.1 Subgroup analyses by dementia or no dementia**

The low quality Monette (2001) study presented results by those with possible or suspected dementia or no dementia. High sensitivity was observed for the two subgroups, but the lower specificity [78%] observed in the possible dementia group was attributed to a suggested weakness in CAM's (short version) ability to exclude those with underlying cognitive impairment. However, we note that this is a low quality study, so that results should be treated with caution (figure 12.17). The diagnostic test accuracy statistics are summarised in table 12.7.

Figure 12.17: forest plot CAM (lay person) compared with CAM (geriatrician) - subgroup analyses

CAM (geriatrician) vs CAM (lay interviewer)-dementia



CAM(geriatrician) vs CAM (lay interviewer)- no dementia



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Table 12.7: CAM (lay person) compared with CAM (geriatrician)

CAM short version	Study name	Comments	Test operator	Sensitivity	Specificity	PPV	LR+	Pre-test probability	Post-test probability
	Monette 2001	CAM for patients with possible or probable	Trained assessor (trained by psychiatrist)	96.40	78.30	84.00	4.4357	54.90	84.40

dementia

Monette 2001	no dementia	Trained assessor (trained by psychiatrist)	94.70	95.00	90.00	18.947	38.80	92.30
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2 **12.5 Results: ICU setting**

3 **12.5.1 Diagnostic test accuracy (DSM-IV as the reference standard)**

4 **12.5.1.1 CAM-ICU**

5 Three moderate to high quality studies (Ely 2001; Ely 2001b; Lin\* 2004)  
6 compared CAM-ICU with DSM-IV.

7

8 A forest plot of sensitivity and specificity is shown in figure 12.17, and diagnostic  
9 test accuracy statistics are summarised in table 12.8.

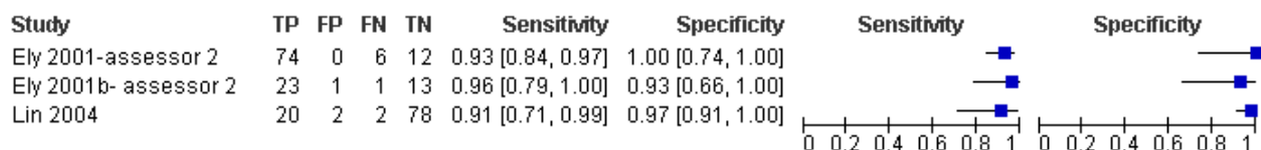
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11 The remaining studies were of good quality and showed a high sensitivity  
12 [range: 91% to 96%] and specificity [93% to 100%]. The likelihood ratio  
13 ranged from 13.42 to 36.36, showing a high likelihood that a patient found to  
14 be delirious based on the CAM-ICU, is delirious.

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16

17 Figure 12.18: forest plot of CAM-ICU index test with DSM-IV as reference  
18 standard in an ICU setting



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22 Table 12.8: diagnostic test accuracy statistics for CAM-ICU

CAM-ICU	Study name	Comments	Test operator	Sensitivity	Specificity	PPV	LR+	Pre-test probability	Post-test probability
	Ely 2001	CAM-ICU [Nurse 2] vs DSM-IV	Nurse	93.00	100.00	100.00	NA	14.13	100
	Ely 2001b	CAM-ICU [Nurse 2] vs DSMIV	Nurse	96.00.	93.00	96.00	13.42	63.20	95.80
	Lin 2004	CAM-ICU [Chinese] [Assessor 1] vs DSMIV [psychiatrist]	Research Assistant	90.90	97.50	91.00	36.364	21.60	90.90

23

1

2 **12.5.1.2 Subgroup analyses by dementia or no dementia**

3 Two studies (Ely 2001; Ely 2001b) reported subgroup analyses by dementia  
 4 status. The number of patients with suspected dementia was 12.5% [12/96] and  
 5 28.9% [11/38], respectively in the two studies. In both studies suspected  
 6 dementia was defined as: the delirium expert rating of having dementia, a  
 7 Blessed Dementia Rating Scale score of at least 3, or a rating by a surrogate of  
 8 at least 3 of out of 5 as 'possibly having dementia'.  
 9

10 The diagnostic test accuracy statistics are summarised in table 12.9.  
 11

12 Both studies reported 100% sensitivity and 100% specificity for patients with  
 13 suspected dementia. However, the 95% confidence interval around these values  
 14 was 56% to 100% for both the sensitivity and specificity in the Ely (2001b)  
 15 study for all three raters and 63% to 100% (nurse 1; nurse 2: 95% CI 66% to  
 16 100%) for sensitivity and 40% to 100% for the specificity (nurse 1; nurse 2:  
 17 95% CI 3% to 100%) in the Ely (2001) study. The number of patients within this  
 18 subgroup analysis in both studies is small (Ely 2001: n=12; Ely 2001b: n=11)  
 19 and the authors suggested that the criteria for identifying patients with suspected  
 20 dementia was liberal.  
 21  
 22

23 Table 12.9: diagnostic test accuracy statistics for CAM-ICU - dementia subgroup

CAM-ICU	Study name	Comments	test operator	sensitivity	specificity
	Ely 2001	CAM-ICU [Nurse 1] vs DSMIV; suspected dementia (n=12)	Nurse 1	100.00	100.00
	Ely 2001	CAM-ICU [Nurse 2] vs DSM-IV Suspected dementia (n=12)	Nurse 2	100.00	100.00
	Ely 2001	CAM-ICU [Nurse 1] vs DSMIV; not suspected dementia (n=84)	Nurse 1	98.00	100.00
	Ely 2001	CAM-ICU [Nurse 2] vs DSM-IV not suspected dementia (n=84)	Nurse 2	100.00	91.00
	Ely 2001b	CAM-ICU [Nurse 1] vs DSMIV Suspected dementia (n=11)	Nurse 1	100.00	100.00
	Ely 2001b	CAM-ICU [Nurse 2] vs DSMIV Suspected dementia (n=11)	Nurse 2	100.00	100.00
	Ely 2001b	CAM-ICU [Intensivist] vs DSMIV Suspected dementia (n=11)	Intensivist	100.00	100.00

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### 12.6 Results: mixed setting

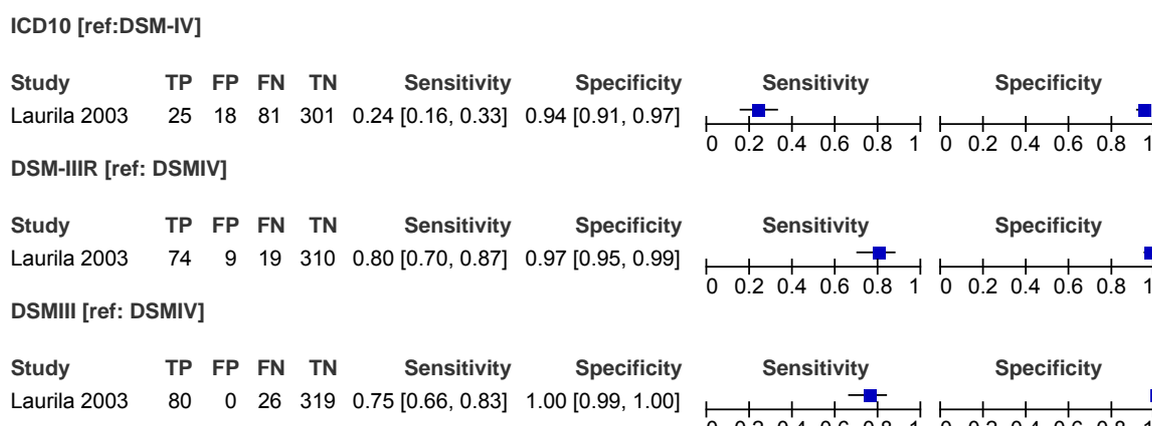
#### 12.6.1 Comparison of diagnostic criterion tools [DSM-IV as the reference standard].

One low quality study (Laurila\* 2003) and one report of that study (Laurila\* 2004) compared three sets of diagnostic criteria in the same patients, using the same data: DSM-III-R; DSM-III and ICD-10 with DSM-IV, in both hospital wards and nursing homes. The study operationalised the clinical and research criteria of the ICD-10 and the criteria from the DSM-IV, DSM-III-R, and DSM-III into one questionnaire. The Laurila\* (2004) study reported a subgroup analysis (see section 12.6.1.1.).

The forest plot of sensitivity and specificity is shown in figure 12.19 and diagnostic test accuracy statistics are summarised in table 12.10.

The ICD-10 showed the lowest sensitivity [24%], whilst the DSM-III-R showed the highest sensitivity [78%]. All three tests showed high specificity. The study reported that the DSM-IV criteria were the most inclusive in the hospital [34.8% of the patients were considered to be delirious], and the DSM-III-R criteria were the most inclusive in the nursing homes [14.4% of the patients were considered to be delirious].

Figure 12.19: forest plot of ICD-10, DSM-III-R and DSM-III compared with DSM-IV; mixed setting (hospital and long-term care)



1 Table 12.10: diagnostic test accuracy statistics for diagnostic criterion tools;  
2 mixed setting

Study name	Comments	Test operator	Sensitivity	Specificity	PPV	LR+	Pre-test probability	Post-test probability
Laurila* 2003	ICD10 vs DSMIV	Geriatrician [hospital]/Nurse [LTC]	40.60	100.00	100.00	NA	24.90	100.00
Laurila* 2003	DSM IIIR vs DSMIV	Geriatrician [hospital]/Nurse [LTC]	79.57	97.18	89.00	28.20	24.94	90.3
Laurila* 2003	DSMIII vs DSMIV	Geriatrician [hospital]/Nurse [LTC]	75.50	100.00	100.00	NA	24.90	100.00

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### 5 12.6.1.1 Subgroup analyses

6 One report (Laurila\* 2004) of the low quality Laurila\* (2003) study reported  
7 the number of patients with and without dementia diagnosed with delirium with  
8 three index tests. Dementia diagnosis was based on the consensus diagnosis of  
9 three geriatricians based on the following information: prior dementia diagnoses,  
10 Clinical Dementia Rating Scale, operationalised criteria according to the DSM-IV,  
11 nurses and/or caregivers' interviews and the results of the brain CT (computed  
12 tomography)/MRI (magnetic resonance imaging) and prior MMSE scores, where  
13 available. The number of patients diagnosed with and without dementia were as  
14 follows: ICD-10: 15% [38/255]: 2.9% [ 5/170]; DSM-III-R: 23% [58/255]:  
15 13% [22/170]; DSM III: 23% [58/255]:13% [22/170] in comparison with DSM-  
16 IV (26% :[66/255]: 24% [40/170]) as the reference standard. However, there  
17 was insufficient information so we were unable to construct 2x2 tables and  
18 report on the sensitivity and specificity of these results.  
19

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## 21 12.7 Clinical evidence statements

22 The GDG's view was that the CAM short version is widely used in practise whilst  
23 the long version was used for research purposes Therefore, the evidence  
24 summary for the CAM short version are reported here.

25

### 26 12.7.1 Hospital setting

- 27 • There is moderate quality evidence to show that:
  - 28 ○ the CAM test (short version) has the most agreement with the
  - 29 DSM-IV criteria for delirium, followed by the DSM-III and DSM-III-
  - 30 R, and is in least agreement with the ICD-10 criteria for delirium.

- 1                   ○ the CAM test (short version) compared with the DSM-IV has a  
2                   moderate ability as a screening instrument for delirium.
- 3                   ○ the MMSE test compared with the DSM-III-R has a low ability as a  
4                   screening instrument for delirium.
- 5                   ○ the clock-drawing test compared with the DSM-III-R has a low  
6                   ability as a screening instrument for delirium.
- 7
- 8                   • There is low quality evidence to show that:
- 9                   ○ the DSM-III-R criteria for delirium shows a moderate agreement  
10                  with the DSM-IV criteria for delirium; same symptoms were used  
11                  to determine both test results.
- 12                  ○ the ICD-10 criteria for delirium are less inclusive than the DSM III  
13                  criteria, when compared with the DSM-III-R criteria for delirium.
- 14                  ○ the DRS-R-98 test compared with the DSM-IV has a fairly low  
15                  ability to moderate as a screening instrument for delirium.
- 16                  ○ the CAM test (short version) compared with the DSM-III-R has a  
17                  low ability to screen patients with delirium with underlying  
18                  cognitive impairment.
- 19                  ○ the presence of 6 or more symptoms of delirium on the CAM test  
20                  compared with the DSM-III-R criteria is considered the best  
21                  threshold point, irrespective of dementia status. We note the  
22                  study was of poor quality and the same symptoms were used to  
23                  determine the index test and reference standard results.
- 24                  ○ the presence of 4 symptoms of delirium on the Delirium Index test  
25                  compared with the DSM-III-R criteria is considered the best  
26                  threshold point in patients with dementia,
- 27                  ○ the presence of 3 or more symptoms of delirium on the Delirium  
28                  Index test compared with the DSM-III-R criteria is considered the  
29                  best threshold point in patients without dementia,
- 30

### 31   12.7.2            **ICU setting**

- 32                   • There is moderate to high quality evidence to show that the CAM-ICU test  
33                   compared with the DSM-IV, has a moderate ability as a screening  
34                   instrument for delirium, irrespective of dementia status.
- 35

### 36   12.7.3            **Mixed setting (hospital and long-term care)**

- 37                   • There is moderate quality evidence to show that the:
- 38                   ○ DSM-III-R criteria is the most inclusive followed by the DSM-III  
39                   criteria compared with the DSM-IV criteria for delirium.
- 40                   ○ ICD-10 criteria to be the least inclusive compared with the DSM-  
41                   IV criteria for delirium.

1  
2

# 1 **13 Non-pharmacological treatment:**

## 2 **multicomponent interventions for**

### 3 **treatment of delirium in a hospital**

#### 4 **setting**

5

#### 6 **13.1 Clinical introduction**

7 Despite the advances in medical science over the last three decades, mortality  
8 and morbidity from delirium have remained unchanged and health costs for this  
9 syndrome remain high. Current management of delirium relies on early  
10 recognition, elimination or correction of underlying causal factors and general  
11 symptomatic and supportive measures. However, there is much uncertainty about  
12 the effectiveness of various interventions.

13 Early recognition and investigation of delirium is challenging and studies have  
14 repeatedly shown that delirium is missed in two-thirds of patients in hospitals.  
15 Moreover, delirium often has multi-factorial causes and multiple potential  
16 consequences. This has led to suggestions that multi-component interventions,  
17 including non-pharmacological interventions might be appropriate for the  
18 treatment of delirium, and several such interventions have been investigated.

19

#### 20 **13.2 Description of studies**

21 Nine papers were evaluated for inclusion. Three studies were excluded and  
22 listed in Appendix G with reasons for exclusion. Seven reports of six studies  
23 were included: three (Cole 1994; Cole 2002; Pitkala 2006; Pitkala 2008) that  
24 reported randomised controlled trials (RCTs); and three (Milisen 2001; Naughton  
25 2005; Rahkonen 2001) that reported prospective studies with historical control  
26 groups. One study (Pitkala 2006) had more than one report (Pitkala 2006 and  
27 Pitkala 2008); hereafter these studies are referred to by the first name reports,  
28 but separately in the results section.

29

##### 30 **13.2.1 Study Design**

31 The unit of randomisation in the RCTs was at patient level. In one of the historical  
32 controlled trial (Naughton 2005), eligible patients were enrolled at two different  
33 time periods. The Naughton (2005) study considered three groups of patients:  
34 those studied in the pre-intervention and two groups after the intervention had  
35 ceased – these patients were studied 4 and 9 months after the initial education  
36 phase of the intervention was completed.

37

1 No studies were conducted in the UK. One study was conducted in the USA  
2 (Naughton 2005); two studies were carried out in Canada (Cole 1994; Cole  
3 2002), two in Finland (Pitkala 2006; Rahkonen 2001) and one in Belgium  
4 (Milisen 2001).

5  
6 Five studies were funded by non-industry sources (Cole 1994; Cole 2002;  
7 Pitkala 2006; Milisen 2001; Naughton 2005) and one did not specify the source  
8 of funding (Rahkonen 2001).

9  
10 One included study had fewer than 100 patients (Cole 1994: n=88), three  
11 studies had more than 100 but fewer than 200 patients (Milisen 2001: n=120;  
12 Pitkala 2006: n=174; Rahkonen 2001: n= 102) and two studies enrolled more  
13 than 200 patients (Cole 2002: n=227 Naughton 2005: n = 374).  
14

### 15 13.2.2 Population

16 All studies took place in a hospital setting; the intervention in the Rahkonen  
17 (2001) study continued after discharge from hospital as it involved support for  
18 the patient over 3 years; Patients were all admitted to medical wards, with the  
19 exception of one study (Milisen 2001). Patients were included in each of the  
20 studies if they had delirium: this was based on screening with CAM, apart from  
21 the Rahkonen (2001) study which specified that the diagnosis was based on  
22 DSM-III-R but did not specify that CAM was used. In the Pitkala (2006) study,  
23 patients found to be positive on CAM screening had their diagnosis confirmed by  
24 a physician using DSM-IV criteria.  
25

26  
27 The Naughton (2005) study reported that for all patients admitted to the Acute  
28 Geriatric Unit (AGU) one criterion for admission was cognitive impairment (score  
29 less than 25 on the MMSE).  
30

31 Some patients had dementia in the studies, (Cole 1994; Cole 1992; Pitkala  
32 2006) ranging from 10% to 58% of participants, except in the Rahkonen (2001)  
33 study, where patients with dementia were excluded.  
34

35 Method of assessment of dementia varied and the following methods were  
36 reported:

- 37 • SPSMQ; scale scores range from: 0 to 10, from no impairment to severe;  
38 score of 5 or more indicative of moderate to severe cognitive  
39 impairment) (Cole 1994)
- 40 • IQCODE (Cole 2002);
- 41 • Medical record data for the diagnosis of preexisting dementia (Milisen  
42 2001)
- 43 • Clinical Dementia Rating Scale (CDR; scale scores range from 0.5 to 3,  
44 from very mild to severe dementia), DSM-IV criteria for dementia or  
45 diagnosis by specialist using standard diagnostic tests (no further details  
46 were given) (Pitkala 2006).

1  
2 The mean age across the studies was 81 to 85.5 years; the studies had a mixed  
3 gender population with a majority of females (Cole 1994: 65%; Cole 2002:  
4 54%; Milisen 2001: 81%; Naughton 2005: 63%; Pitkala 2006: 74%; Rahkonen  
5 2001: 90%). Ethnicity was not reported in any of the studies.  
6

### 7 **13.2.3 Interventions**

8 The included studies investigated multicomponent interventions in a hospital (or  
9 hospital plus community in the case of Rahkonen 2001) setting for the treatment  
10 of delirium (table 13.1).

#### 11 12 **13.2.3.1 Nursing intervention protocol (Cole 1994, Cole 2002),**

13 This intervention comprised of a multidisciplinary team consisting of geriatricians  
14 and liaison nurse.

- 15 • consultation by a geriatrician or geriatric psychiatrist (completed within 24  
16 hours after referral)
- 17 • follow-up by a liaison nurse
  - 18 ○ follow up included daily visits during the patients' stay (up to a  
19 maximum of 8 weeks), liaising with family members, recording  
20 information on patient's mental status and discuss management with  
21 the patient's nurses with the use of the protocol
  - 22 ○ assess compliance with consultant recommendations. Where  
23 appropriate, the nurse discussed management problems with the  
24 geriatrician or geriatric psychiatrist and where necessary patient  
25 was reassessed by the specialists.
- 26 • the intervention protocol targeted the following risk factors:
  - 27 ○ environment (not having excessive, inadequate or ambiguous  
28 sensory input, medication not interrupting sleep, presenting one  
29 stimulus or task at a time);
  - 30 ○ orientation (room should have a clock, calendar, and chart of the  
31 day's schedule; evaluate need for glasses, hearing aid,  
32 interpreter)
  - 33 ○ familiarity (objects from home, same staff, family members  
34 staying with patient, discussion of familiar areas of interest),
  - 35 ○ communication (clear, slow, simple, repetitive, facing patient,  
36 warm, firm kindness, address patient by name, identify self,  
37 encourage verbal expression)
  - 38 ○ activities (avoid physical restraint, allow movement, encourage  
39 self care and personal activities).

40  
41 The intervention in the later trial (Cole 2002) was described as more intensive  
42 than in the earlier study (Cole 1994) and the following components were added  
43 to the intervention:

- 1 • consultant not only assessed initially but also followed up the patients;
- 2 • the study nurse visited the patient 5 days per week;
- 3 • the intervention team (2 geriatric psychiatrists, 2 geriatric internists and the
- 4 study nurse) met after every 8 to 10 patients were enrolled to discuss
- 5 delirium management problems; and
- 6 • the study investigator met the nurse weekly to discuss problems of
- 7 diagnosis, enrollment and interventions.

8

### 9 **13.2.3.2 Multicomponent geriatric intervention (Pitkala 2006)**

10 Patients received a comprehensive geriatric assessment, which included history  
11 taking, interview with caregiver, physical examination, assessment of cognition  
12 and physical functioning, screening for depression, nutrition, and medication  
13 review.

14 Other aspects of the intervention included:

- 15 • recognising delirium and any underlying conditions
- 16 • orientation (with calendars, clocks, photographs)
- 17 • physiotherapy
- 18 • general geriatric interventions (calcium and vitamin D supplements;
- 19 nutritional supplements for those at risk of malnutrition or malnourished;
- 20 hip protectors)
- 21 • comprehensive discharge planning (including consultation of a social
- 22 worker, occupational therapist's home visit, involvement of caregivers).
- 23 • medical management (avoiding neuroleptics; administering atypical
- 24 antipsychotics for hyperactive/psychotic symptoms; use of cholinesterase
- 25 inhibitors if patient's cognition did not improve to MMSE score above 23).

26

27 The intervention group received significantly more atypical antipsychotic drugs  
28 than the control group (69.0% versus 29.9%,  $p < 0.001$ ), more  
29 acetylcholinesterase inhibitors (58.6% versus 9.2%,  $p < 0.001$ ), vitamin D and  
30 calcium supplements (77.0% versus 9.2%,  $p < 0.001$ ), nutritional supplements  
31 (92.0% versus 0.0%,  $p < 0.001$ ) and fewer conventional neuroleptics (8.0%  
32 versus 23.0%,  $p = 0.006$ ).

33

### 34 **13.2.3.3 Nurse-led interdisciplinary intervention (Milisen 2001)**

35 This intervention involved nurse education to identify high-risk patients which  
36 included:

- 37 • education: a poster was developed to educate all nurses on the essential
- 38 aspects of delirium, depression and dementia. This poster included the
- 39 core symptoms of delirium according to the CAM criteria, comparative

- 1 features and differences between delirium, dementia and depression and  
2 the relevance of correct and early recognition of delirium;
- 3 • systematic screening of cognitive function using the NEECHAM Confusion  
4 Scale following training;
- 5 • pain management: scheduled pain medication to provide effective post-  
6 operative pain control; and
- 7 • consultative service: access to a resource nurses who were given training in  
8 identifying patients by a geriatric nurse specialist in the identification and  
9 management of older hip-fracture patients. If necessary, the resource  
10 nurses could consult with a geriatric nurse specialist or psycho  
11 geriatrician; resource nursed to help the primary nurses in implementing  
12 appropriate antidelirium interventions.
- 13 • the nurses were provided with 'A nursing guide for the evaluation of causes  
14 of delirium in elderly hospitalised patients' (as reported in Milisen 1998).  
15 The guide advised a nurse to report to the attending physician of any  
16 changes in patient's status on the following: medication, pain, hypoxemia,  
17 dehydration, electrolyte and metabolic disturbances, and infection. The  
18 interventions are briefly described below:
- 19 ○ medication: to be vigilant of polypharmacy, especially  
20 anticholinergics, antiparkinsonian drugs, histamine H<sub>2</sub>-receptor  
21 antagonists;
  - 22 ○ pain: inquire systematically about pain; observe verbal and  
23 nonverbal expressions; use of as many possible analgesics based  
24 on nonopioid drug (e.g. paracetamol) and where required  
25 minimum dose of opioids combined with non opioid drug;
  - 26 ○ hypoxemia: monitor abnormalities in rate, depth and quality of  
27 respiration, cyanosis, PO<sub>2</sub> ≤ 32; administer oxygen as ordered;  
28 determine source of hypoxia; low respiration (<10 l/min) due to  
29 opioid intoxication; consult attending physician for treatment with  
30 naloxone as antidote; in patients undergoing surgery: monitor  
31 hypothermia and postoperative shivering; maintain optimal  
32 patient temperature by applying warming [fluids and blood;  
33 gowns and blankets; humidified oxygen]; be alert for nocturnal  
34 desaturation during the first 3 days postoperatively and  
35 especially in obese patients; administer 2 l of O<sub>2</sub> (unless  
36 contraindicated);
  - 37 ○ dehydration: encourage patient to drink water regularly and  
38 when necessary prepare for blood or fluid replacement;
  - 39 ○ electrolyte and metabolic disturbances: monitor abnormalities of  
40 blood and urine chemistry; give frequent small meals and add  
41 nutritional supplements, such as calorie/protein rich drink;
  - 42 ○ infection: be alert for urinary tract, respiratory, mouth and feet  
43 infections; stimulate patient for adequate water intake (2 l/day)  
44 (unless contraindicated); observe for abrupt onset for fever  
45 (rectal temperature >100°F) and apply cooling techniques as  
46 needed.
- 47

#### 1 **13.2.3.4 Systematic intervention (Rahkonen 2001).**

2 The intervention consisted of a case manager (nurse specialist) and an annual one-  
3 week rehabilitation period at a Brain Research and Rehabilitation Centre.

4 Patient's rehabilitation team included the study physician, the nurse specialist,  
5 physiotherapist, neuropsychologist and occupation therapist.

- 6 • a nurse specialist trained in geriatrics and care of the elderly acted as the  
7 case manager. Patients received continuous and systematic support  
8 provided by the case manager with responsibility in supporting the  
9 patients during community care through out the 3 year follow-up acting  
10 as a counsellor and advocate and in the rehabilitation unit (as the  
11 primary care nurse);

- 12 • care in the community: arranged in consultation with relatives and health  
13 and social care services, and continuity of care was achieved with  
14 regular follow-ups, including in-home visits and 'phone calls by the case  
15 manager. Study physician was also available for consultation and  
16 medical care throughout the follow up; and

- 17 • rehabilitation period: individually structured physiotherapy once or twice  
18 daily; mobility and other special aides for daily living (e.g. hearing aids  
19 and special shoes) were arranged when needed; patients were  
20 encouraged to participate in occupational therapy and free-time events.

21

#### 22 **13.2.3.5 Education and management intervention (Naughton 2005)**

23 The intervention was designed to improve the recognition of delirium in medically  
24 ill older adults evaluated in the emergency department [ED triaged these  
25 patients with delirium specifically to the acute geriatric unit (AGU)]. This was  
26 achieved by addressing the following factors:

- 27 • education:

- 28 ○ The charting procedures in ED were changed and physicians were  
29 reminded to evaluate adults aged 75 years and older for  
30 cognitive impairment and delirium and direct the admission to the  
31 AGU. Nurses and physicians were trained to triage patients using  
32 yes/no answers to four questions from the history and mental  
33 status examination. A study nurse periodically reported the  
34 proportion of older adults correctly admitted to the AGU from  
35 the ED.

- 36 • the education component for the AGU nurses (provided by geriatricians  
37 and geriatric nurse) involved:

- 38 ○ educating on prevalence and outcome of delirium;
- 39 ○ sensitivity training on cognitive impairment;
- 40 ○ training on methods of mental status assessment;
- 41 ○ guidelines on medication management of cognitive impairment  
42 and delirium.

- 1                   ○ small group consensus process used to develop assessment and  
2                   charting procedures; and
- 3                   ○ AGU physicians were provided with information on cognitive  
4                   impairment and delirium in the elderly, recommended mental status  
5                   assessment procedures, and review of the intervention guidelines.
- 6                   ● treating underlying medical factors;
- 7                   ● treating precipitating factors (removing precipitating medications;  
8                   addressing immobility);
- 9                   ● providing family support;
- 10                  ● using non-pharmacological support for: physically non aggressive  
11                  behaviour and episodes triggered with ADL care;
- 12                  ● medication management: reduce the use of psychotropic medications  
13                  (benzodiazepines and anticholinergics); consider using synergistic agents  
14                  such as neuroleptics or antidepressants that supplement behaviour  
15                  treatment; sleep medication: trazadone 50 to 100 mg; zolpidem: 5 mg;
- 16                  ● fewer patients in the AGU received benzodiazepines (22.6% compared  
17                  with 30.9% at baseline); antihistamines (6% compared with 15.5%;  
18                  p<0.02); increased use of antidepressants (22.7% compared with 10%  
19                  at baseline; p<0.02); and neuroleptics (27.4% compared with 10.9% at  
20                  baseline; p<.01)
- 21                  ● simplifying pain regimen (minimise p.r.n.); and
- 22                  ● environmental stimuli: addressing problems with environmental stimuli for  
23                  example, noise, sleep disruption, disruptive room mate,

24  
25  
26  
27  
28

None of the studies included more than two study arms, and the comparator in all studies was 'usual medical care' (no further details given).

#### 29   **13.2.4                   Comparisons**

30                  The following comparison was carried out:

- 31                  ● Multicomponent intervention versus usual care.
- 32                         ○ Two RCTs followed patients up to 8 weeks (Cole 1994, Cole  
33                         2002) and one followed patients up to 1 year (Pitkala 2006). Of  
34                         the non-RCTs, one study followed patients up to 12 days (Milisen  
35                         2001), 2 months (Naughton 2005) and 3 years (Rahkonen 2001).

36  
37

Two studies (Naughton 2005; Pitkala 2006) reported concurrent medications:

- 38                  ● opiates (42.7%); benzodiazepines (30.9%); antihistamines (15.5%);  
39                  antidepressants (10.0%); neuroleptics (10.9%)
- 40                  ● conventional neuroleptics (22%); atypical antipsychotics (14%) and  
41                  cholinesterase inhibitors (6%) (Pitkala 2006).

42

### 1 13.2.5 Outcome measures

2

3

The following primary and secondary outcome measures were reported:

4

5

- primary outcomes:

6

- complete response (Pitkala 2006 RCT; Naughton 2005 non RCT)

7

- duration of delirium (Milisen 2001 non RCT)

8

9

- secondary outcomes:

10

- cognitive impairment (Cole 1994; Pitkala 2006)

11

- length of stay (Cole 1994; Cole 2002)

12

- health related quality of life (Pitkala 2008)

13

- discharge (higher dependency: Cole 1994; Cole 2002; long-term care: Pitkala 2006)

14

15

- days in new long-term care (non RCT: Rahkonen 2001)

16

- mortality (RCTs: Cole 1994; Cole 2002; Pitkala 2006; non RCT: Rahkonen 2001)

17

18

19

## 20 13.3 Methodological quality

21

### 13.3.1 RCTs

22

The method of sequence generation was adequate in two RCTs in which a computer-generated sequence was employed (Cole 2002, Pitkala 2006), and was not stated in one RCT (Cole 1994).

23

24

25

26

One RCT reported adequate allocation concealment - central randomisation with details of a retained schedule (Pitkala 2006). One RCT was partially adequate (with independent allocation but no further details, Cole 2002). In the third RCT, allocation concealment was not stated (Cole 1994).

27

28

29

30

31

Outcome assessors were stated to be blinded in two RCTs (Cole 1994, Cole 2002) and this was not stated in the other RCT (Pitkala 2006). Patients were not blinded in any of the RCTs.

32

33

34

35

Two RCTs reported an *a priori* sample size calculation. One RCT (Cole 1994) reported that a sample of 30 or more was required for 80% power to detect a difference of at least 1 SD in the change in the measures used ( $p=0.05$ ). One RCT (Pitkala 2006) reported that 58 to 91 patients per group were needed to show a 20% difference in the combined endpoint (discharge to permanent institutional care or death) with 80% power ( $p=0.05$ ). The third RCT did not report a sample size calculation (Cole 2002).

36

37

38

39

40

41

1  
2 All three RCTs included in the review demonstrated baseline comparability of the  
3 groups on measures such as age, gender and baseline scores measuring delirium  
4 or mental state.  
5

6 All RCTs used an intention to treat analysis for at least some outcome measures.  
7 One RCT reported no missing data in either group (Pitkala 2006). In one RCT  
8 (Cole 2002), 7 patients withdrew in the intervention group (6.2%) versus 2  
9 (1.8%) in the control group. In the third RCT (Cole 1994), 33% of patients died  
10 in the intervention group versus 37% in the control group; mean scores for some  
11 of the outcome measures SPMSQ and Crichton Geriatric Behavioural Rating  
12 Scale [CGBRS] were given for surviving patients only (i.e. fewer than 70% of the  
13 number randomised), although all patients were included in some outcome  
14 measures (length of stay, discharge to new long-term care, mortality).  
15

16 Overall, one RCT was considered to have the potential for bias (Cole 1994). This  
17 study did not state randomisation or allocation concealment methods, and some  
18 outcome measures had missing data due to patients who had died (Cole 1994).  
19 This study was considered in sensitivity analyses.  
20

### 21 13.3.2 Non-RCTs

22 In the Rakhonen (2001) study, the control group was formed by matching pairs  
23 of patients on age and gender from patients fulfilling the inclusion criteria from  
24 the earlier time period; in the remaining two studies patients were not  
25 individually matched but the groups were comparable on age and gender. The  
26 Milisen (2001) study reported that the non intervention cohort had significantly  
27 greater comorbid conditions (e.g. cardiac, vascular and abdominal problems).  
28

29 One study reported that the investigator was blinded to the data of the main  
30 outcome measure of the study in the control patients (Rahkonen 2001:  
31 information was collected from registers for the control patients) and unclear in  
32 the other two studies.  
33

34 One study (Rahkonen 2001) reported not all eligible patients were included  
35 (10%) and it was unclear in the other two studies.  
36

37 Overall, we considered the three non-RCT studies to be of weak quality because  
38 of the study design.  
39  
40

## 41 13.4 Results

### 42 13.4.1 Multicomponent intervention versus usual care

#### 43 13.4.1.1 Primary outcomes of the review:

##### 44 Duration of delirium

45 Only one study reported the duration of delirium (Milisen 2001). This was  
46 significantly shorter in the intervention cohort (median = 1 day, interquartile

1 range [IQR] = 1) compared with the non-intervention cohort (median = 4 days,  
2 IQR = 5.5, p=0.03, Mann-Whitney U test).

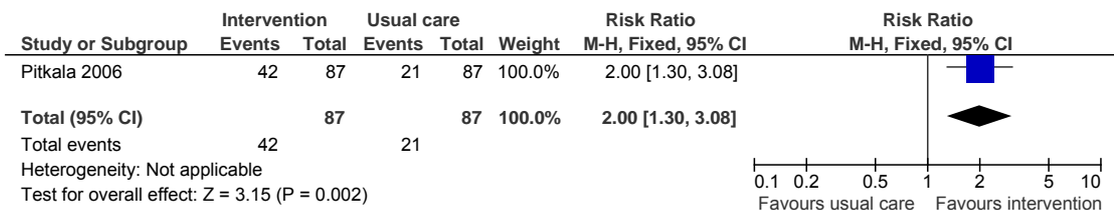
### 3 Number of patients recovered from delirium (complete response)

4 Two RCTs (Cole 2002; Pitkala 2005) reported complete response. The Pitkala  
5 (2006) study defined the response rate as a permanent improvement of at least  
6 4 points on the MDAS (severity of delirium scored 0 to 30, with 30 being the  
7 worst) at 8 days; although no data or references were supplied to justify the use  
8 of this score as the measure for improvement, and the GDG considered this to be  
9 a weak measure of complete response.

10  
11  
12 Cole (2002) reported the number of patients with an improvement in cognitive  
13 status, as defined by the MMSE, during the hospital stay (mean length of stay 19  
14 days). "Improvement" was defined as an increase in MMSE of 2 or more points;  
15 with no decrease below baseline plus 2 points thereafter. If the MMSE score at  
16 baseline was 27 or more, improvement was no decrease below 27; MMSE  
17 ranges from 0=poor to 30=excellent; a score of 23 or less indicates cognitive  
18 impairment) or 'not improved'. The GDG decided that 'the number improved'  
19 was an unsatisfactory definition of recovery from delirium, so the study was not  
20 included in the analysis for this outcome.

21  
22 In the Pitkala (2006) study, the intervention significantly increased the number of  
23 patients who had recovered from delirium at 8 days after admission (RR 2.00,  
24 95% CI 1.30 to 3.08) This corresponds to a number needed to treat of 5 (95%  
25 CI 3 to 10); figure 13.1. The GDG debated whether a change of 4 points on the  
26 MDAS scale would clearly show improvement and considered that any  
27 conclusions drawn from the Pitkala (2006) study should be treated with caution.

28  
29  
30  
31 Figure 13.1: number of patients with complete response.



33

### 34 **13.4.1.2 Secondary outcomes of the review:**

#### 35 Cognitive impairment

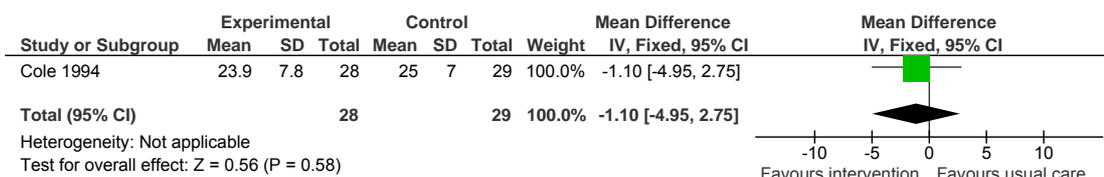
1 Three studies (Cole 1994; Milisen 2001; Pitkala 2006) reported cognitive  
 2 impairment.

3  
 4 The Cole (1994) study reported scores for the SPSMQ, a 10-item questionnaire  
 5 that evaluates orientation, memory and concentration (0=no impairment to  
 6 10=severe impairment) at 8 weeks. There was no difference between the  
 7 intervention and usual care groups (figure 13.2), although the result is imprecise.  
 8

9 The Pitkala (2006) study measured cognitive impairment with the MMSE at 6  
 10 months (Pitkala 2006). The study reported a mean score of 18.4 in the  
 11 intervention group versus 15.8 in the usual care group, but no standard  
 12 deviations were given ( $p=0.047$  for repeated measures analysis of variance  
 13 (ANOVA); baseline scores used as covariates). This was just significant.  
 14

15 The Milisen (2001) study reported the mean MMSE scores for the delirious  
 16 patients in the intervention group and the non intervention group (mean MMSE  
 17 scores: intervention group (delirious): 15.5; non intervention group (delirious):  
 18 9.5); the study reported that although the intervention group showed a higher  
 19 overall cognitive function this difference was not statistically significant;  $p$   
 20 values or standard deviations were not reported.  
 21  
 22

23 Figure 13.2: cognitive impairment



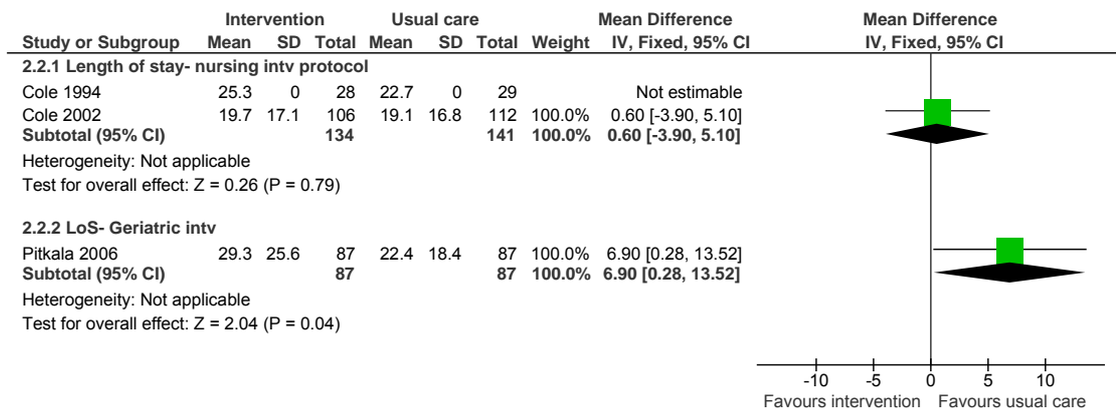
24  
 25  
 26  
 27  
 28 Length of stay

29 Length of hospital stay was reported by all three RCTs (Cole 1994; Cole 2002;  
 30 Pitkala 2006). The result for the Pitkala (2006) study is presented as a subgroup  
 31 as the intervention differed from the other two studies (Cole 1994; Cole 2002).  
 32

33 The Cole (1994) study did not report standard deviations, so the study's  
 34 contribution to the meta-analysis of the two studies was not estimable. There was  
 35 no significant difference between intervention and usual care groups in Cole  
 36 (2002), although the result is imprecise (figure 13.3).  
 37

38 In the Pitkala (2006), length of stay appeared shorter in the usual care group.  
 39 We note that the distribution of lengths of stay was skewed (median 21 days in  
 40 the intervention group, range 2 to 110 days; median 16 in the usual care group,  
 41 range 1 to 90 days; mean 29.3 days, SD 25.6 in intervention group and mean  
 42 22.4 days, SD 18.4 in control group; means are less than twice SD so data likely  
 43 to be skewed). The result is imprecise.  
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 45

46 Figure 13.3: length of stay



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Two non RCTs also reported length of hospital stay (Milisen 2001; Naughton 2005). The Milisen (2001) study reported a median of 13.5 days (IQR 3.75 days) for the intervention cohort and 14 days (IQR 5 days,  $p=0.6$ ) for the non-intervention cohort. The Naughton (2005) study reported that following intervention, a mean of 3.3 days was saved in length of stay following each episode of delirium.

#### Discharge to long-term care

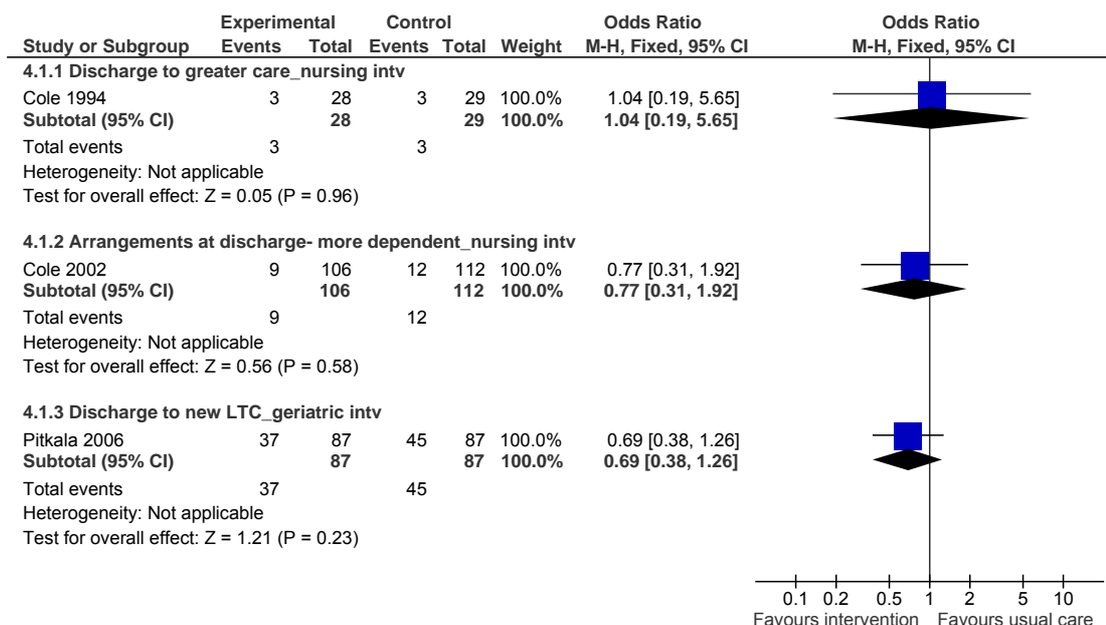
All three RCTs reported discharge of patients who had become more dependent since their admission. Two studies reported that patients were discharged at a greater level of dependency: Cole (1994) reported the percentage of patients discharged required more care (numbers were calculated as the proportion of patients remaining alive at the end of the study); Cole (2002) reported that living arrangements were arranged hierarchically from least dependent (e.g. home alone) to most dependent (e.g. nursing home); living arrangements at discharge were compared with those at admission and were rated as more dependent, same, or less dependent.

The Pitkala (2006) study reported the number of patients discharged to permanent institutional care, and these represented new admissions to such care as patients already in permanent institutional care at admission were excluded from the study.

The results are presented as subgroups in figure 13.4. There was no significant difference in effect of the intervention on discharge to higher care or to new long-term care, although the results for all three studies are imprecise.

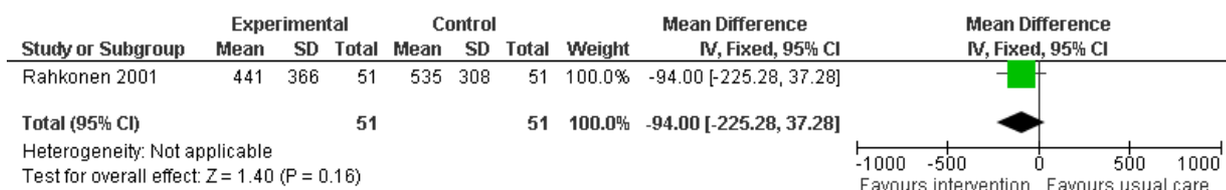
Excluding the Cole (1994) study due to its possible bias would not materially alter the results.

Figure 13.4: discharge to higher dependency or to new long-term care (RCTs)



The Rahkonen (2001) study reported the duration of long-term care in the three years of the study. This was a mean of 441 days (SD 366) in the intervention group compared with 535 days (SD 308) in the control group (figure 13.5). The mean age was comparable (82.1 years in both groups) and the study excluded patients with confirmed or suspected dementia, however, individuals with mild cognitive impairment were included.

Figure 13.5: number of days in new long-term care (non RCT)



NB: Scale -1000 to 1000

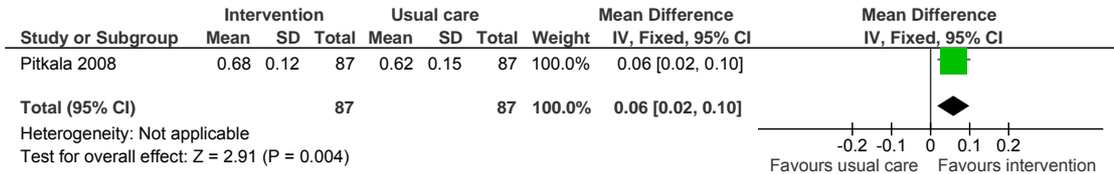
Health related quality of life (HRQoL)

One report (Pitkala 2008) of the Pitkala (2006) study reported health related quality of life along the following dimensions: mobility, vision, hearing, breathing, sleeping, eating, speech, elimination, usual activities, mental function, discomfort and symptoms, depression, distress, and vitality. Patients were assessed with the 15D questionnaire at baseline and discharge [range 0 (poor HRQoL) to 1 (excellent HRQoL)].

There was a small significantly higher HRQoL for the intervention group (MD 0.06 (95% CI 0.02 to 0.10); figure 13.6. The study reported that there were significant differences for the intervention and usual care group on the following dimensions on the 15D questionnaire: mental function corresponding to cognition and alertness (p<0.001), usual activities corresponding to functioning in activities

1 of daily living ( $p < 0.001$ ), vitality ( $p = 0.004$ ), depression ( $p = 0.044$ ), and speech  
 2 ( $p = 0.024$ ).  
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Figure 13.6: improvement in HRQoL



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 8 NB: Scale -0.2 to 0.2

### 9 10 11 Mortality

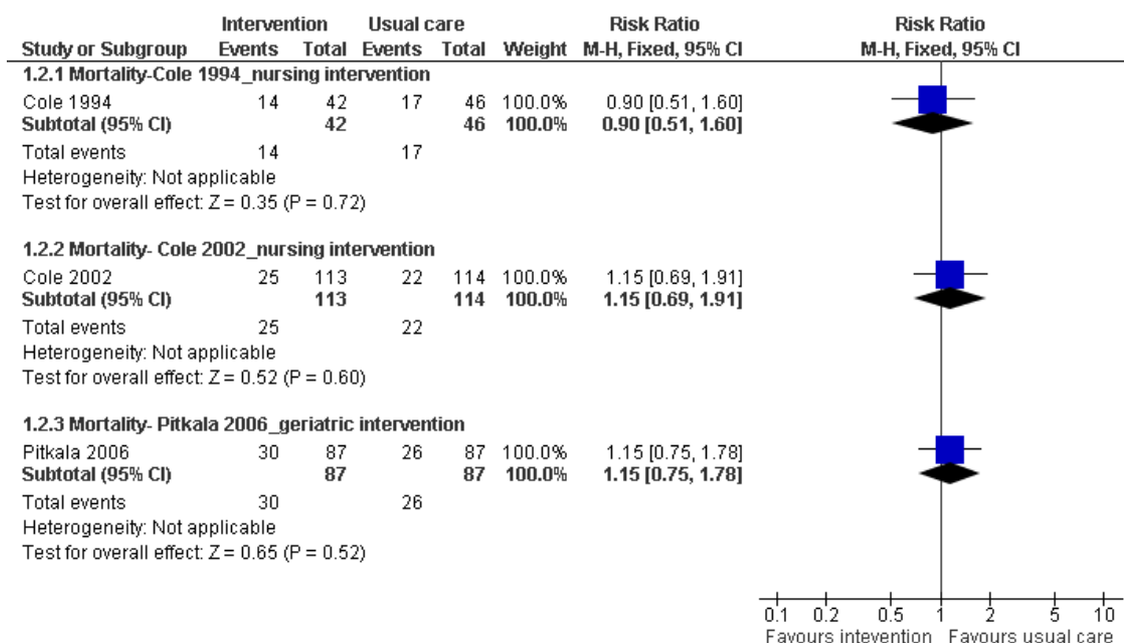
12 Three RCTs (Cole 1994; Cole 2002; Pitkala 2006) and one non-RCT (Rahkonen  
 13 2001) evaluated the number of patients who died: two RCTs at 8 weeks (Cole  
 14 1994; Cole 2002) and the other RCT at 1 year (Pitkala 2006) and the non-RCT  
 15 at 3 years (Rahkonen 2001).  
 16

17 The Cole (1994) study reported that overall 35% (31/88) patients died in 8  
 18 weeks (33% [14/42] and 37% [17/46] deaths occurring in the intervention and  
 19 control groups, respectively); the causes of death were not given.  
 20

21 The Cole (2002) study reported that overall 21% (47/227) of patients died  
 22 (22% [25/113] and 19% [22/114] deaths occurring in the intervention and  
 23 control groups, respectively); and the Pitkala (2006) study reported that overall  
 24 32% (56/174) patients died over 1 year (34% [30/87] and 30% [26/87]  
 25 deaths occurring in the intervention and control groups, respectively); the causes  
 26 of death were not reported in either study.  
 27

28 There was no significant difference between the interventions and usual care in  
 29 the mortality rates, but the results were very imprecise (figure 13.7).  
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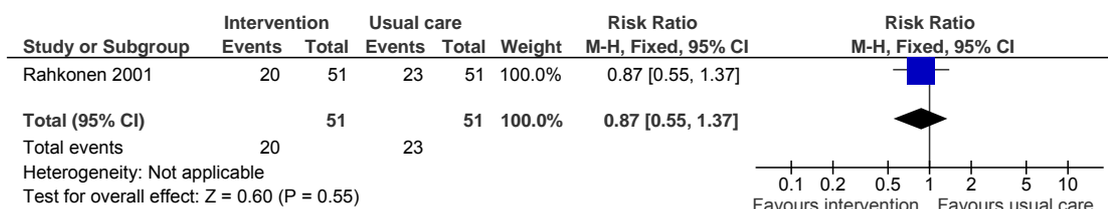
Figure 13.7: mortality (RCTs only)



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The non-RCT study (Rahkonen2001) reported that during the three-year follow up, a total of 42% (43/102) patients died, the causes of death were not reported (figure 13.8).

Figure 13.8: mortality (non-RCT)



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### 12 13.5 Clinical evidence statements

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- There is very low quality evidence which showed that a multicomponent intervention targeting six modifiable risk factors (orientation, sleep, sensory impairment improvement, early mobilisation, environmental, medication) following a consultation with a geriatrician or geriatric psychiatrist and follow up by a liaison nurse showed no significant difference in:
  - cognitive impairment (measured at 8 weeks). However, there is much uncertainty around this result
  - the number of patients discharged with a greater level of dependency; there is much uncertainty around this result
  - mortality rates at 8 weeks; there is uncertainty around this result

(Cole 1994)

- 1           • There is very low quality evidence which showed that a multicomponent  
2           intervention targeting six modifiable risk factors (orientation, sleep,  
3           sensory impairment improvement, early mobilisation, environmental,  
4           medication) followed by an assessment and follow up by a geriatrician  
5           or geriatric psychiatrist and follow up by a liaison nurse showed no  
6           significant difference in:
- 7           ○ the number of patients discharged to a 'more dependent' level  
8           of care; there is some uncertainty around this result
- 9           ○ mortality rates at 8 weeks; there is some uncertainty around this  
10          result
- 11          (Cole 2002)
- 12
- 13          • There is very low quality evidence to show that a multicomponent  
14          intervention targeting three modifiable risk factors (dehydration/nutrition,  
15          pain management, medication management) with training showed:
- 16          ○ significantly shorter duration of delirium in patients in the  
17          intervention group
- 18          ○ no significant difference in the median length of stay in hospital
- 19          (Milisen 2001)
- 20
- 21          • There is moderate quality evidence to show that a multicomponent geriatric  
22          intervention based on targeting four modifiable risk factors (orientation,  
23          dehydration/nutrition, early mobilisation, medication management) with  
24          comprehensive geriatric assessment showed:
- 25          ○ a significant number of patients recovered from delirium (at 8  
26          days) in the intervention group; however, there is much uncertainty  
27          around this result
- 28          ○ a borderline significant difference showing a lower level of  
29          cognitive impairment at 6 months for the intervention group
- 30          ○ a significant difference showing a decreased length of stay in the  
31          usual care group; however there is much uncertainty around this  
32          result
- 33          ○ no significant difference in the number of patients discharged to  
34          long-term care; there is much uncertainty around this result.
- 35          (Pitkala 2006)
- 36          ○ a small significant improvement in the health related quality of  
37          life (mental function, daily functioning, depression, vitality, and  
38          speech) for the intervention group at discharge
- 39          (Pitkala 2008)
- 40
- 41          • There is very low quality evidence to show a multicomponent intervention  
42          targeting two modifiable risk factors (orientation, early mobilisation) with

- 1 training, continuous nursing support and annual one-week visits to a  
2 rehabilitation unit showed no significant difference in:
- 3 ○ the stay in long-term care over the duration of the study (3 years)  
4 ; there is much uncertainty around this result
  - 5 ○ mortality rates at 3 years
- 6 (Rahkonen 1998)  
7  
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## 9 13.6 Health economic evidence

### 10 13.6.1 Multi-component interventions for the treatment of delirium in a 11 hospital setting

12 One economic evaluation study was included as evidence (Pitkala 2008). This  
13 was a Finnish RCT of 174 consecutive delirium patients aged above 69 years  
14 who were admitted to the general medicine services and whose life expectancy  
15 was predicted to be above 6 months. The study aimed at assessing the effects of  
16 multi-component geriatric treatment on costs of care and HRQoL in delirious in-  
17 patients. Patients in the intervention group received a comprehensive geriatric  
18 assessment at baseline for good detection of delirium, as well as careful  
19 diagnosis of the underlying etiological conditions. They received atypical  
20 antipsychotics if necessary and effective general treatments were implemented  
21 for all patients. After the acute phase of delirium, all patients not recovering  
22 from impaired cognition underwent detailed diagnostics for dementia and  
23 thereafter, received acetyl cholinesterase inhibitors. Patients in the comparator  
24 arm received usual care and this was not exactly described.

25 The average cost per patient in the intervention arm was €19,737 while the  
26 average cost per patient in the usual care arm was €19,557. The extra cost  
27 attributable to intervention was €446 per patient. This included the cost of  
28 atypical antipsychotics, acetylcholinesterase inhibitors, vitamin D-calcium  
29 supplements, hip protectors, and nutritional supplements. Average unit costs in  
30 Finland were used. Health related quality of life was measured using the 15D  
31 questionnaire but the question on sexual activity was omitted. Subjective health  
32 was assessed using an ordinal scale at discharge. An unadjusted mortality rate  
33 of 35% and 30% were reported in the intervention and usual care groups  
34 respectively. The patient's measure of health status was 0.68 and 0.62 in the  
35 intervention and control groups respectively. The dimensions of HRQoL showing  
36 significant differences favouring intervention were mental function, usual  
37 activities, vitality, depression and speech.

38 The results of this study could be used to estimate the cost per unit of  
39 improvement in health status of delirium patients. However, patient's measure of  
40 health status was based on 15D which elicited health status scores from a Finnish  
41 general population. It was reported only at the point of discharge from  
42 hospitalisation for delirium and quality-adjusted life years were not reported.  
43 Furthermore, there was no sensitivity analysis to test the effect of the  
44 uncertainties surrounding the cost and health outcome measures. Costs were not  
45 assessed from a UK NHS and PSS perspective. The results of this study were  
46 judged to be not directly applicable to this guideline.

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1 Table 13.1: multicomponent interventions for the treatment of delirium

Study	Multi-disciplinary team	Education intervention	Treatment intervention	Care methods	assessment	orientation	Dehydration nutrition	Sleep	Sensory impairment improvement	Early mobilisation	Pain management	Environmental modifications	Medication management	Other (including communication, discharge planning)
Cole (1994)	Yes: consultation by geriatrician or geriatric psychiatrist & follow up by liaison nurse	No	daily visits & management by protocol	Daily visits from liaison nurse	Yes: consultation by geriatrician or geriatric psychiatrist	Yes: clock, calendar, chart of day's schedule. Verbal reminders of time, day & place. Assess the need for glasses, hearing aid, foreign language interpreters. Keep the pt in the same surroundings; Familiarity [familiar possession from home, request family members to stay w/pt, discuss familiar areas of interest, same staff members to care for the pt]	No	Yes: medication rounds not interrupting sleep	Yes: hearing aid	Yes: encouraging self-care	No	Yes: not excess, inadequate or ambiguous sensory input. Present one stimulus or task at a time. Determine if pt prefers radio or TV	Yes: medication rounds not interrupting sleep	Yes: communication (clear, facing patient, frequently address the pt by name and convey identifying info )
Cole (2002)	Yes: consultation and follow up by geriatrician or geriatric psychiatrist & follow up by liaison nurse	No	daily visits & management by protocol	Daily visits from liaison nurse	Yes: consultation by geriatrician or geriatric psychiatrist	Yes: clock, calendar, glasses, hearing aid, foreign language interpreters, familiarity (objects from home, same staff)	No	Yes: medication rounds not interrupting sleep	Yes: hearing aid	Yes: encouraging self-care	No	Yes: not excess sensory input	Yes: medication rounds not interrupting sleep	Yes: communication (clear, facing patient, frequently address pt by name and convey identifying info)
Milisen (2001)	Yes: access to resource nurses/consultants	Education posters for nurses on core symptoms of delirium according to Cam, features and difference btw delirium, dementia, & depression and the relevance of correct and early recognition of delirium; All nurses trained in using the NEECHEM Confusion	nurse education; screening; antidelirium intervention; access to resource nurses/consultants; scheduled pain medication	Not changed, nurse	Usual nurse (Resource nurses verified regular staff nurses assessments)	No	Yes: additional nutrition supplements (e.g. calorie/protein rich drink) for those with malnutrition (especially, vit B deficiency and low serum albumin)	No	No	No	Yes: scheduled pain medication upto 5th postop day. From 6th day postop pain meds given as requested by pt or on basis of judgement of the primary nurse, inquire systematically and observe for	No	Yes: scheduled pain medication	Not stated
Naughton (2005)		Yes: staff on POD, CD, Ass, MMD	not changed	Yes: small group process and audit	No	No	yes: pharma and non-pharma (latter implied)	No	yes: immobility treated	Yes: noise, disruptive room mate, sleep disturbance	Reduce use of psychotropic medications; use of neuroleptics	Yes	non-pharma mgt	Not stated
Pitkala (2006)	Yes: nurses, physician/geriatrician, physiotherapy, social worker, occupational therapist		recognise delirium & underlying conditions; assessment & treatment (e.g. nutrition, review drugs), avoid neuroleptics; orientation, physiotherapy, Calcium/Vitamin D/other supplements, hip protectors, screen for treatable causes, cholinesterase inhibitor, discharge plan	Not changed, nurse	Comprehensive geriatric assessment; recognition of delirium and underlying conditions	Yes: calendar, clocks, photos	Yes: nutritional supplements for those at risk of malnutrition or malnourished; calcium and vitamin D supplements	No	No	Physiotherapy	No	Hip protectors	review drugs, avoid neuroleptics & administer atypical antipsychotics for hyperactive/psychotic symptoms, use cholinesterase inhibitors	Lab tests and scans for treatable causes of dementia; screening for depression; comprehensive discharge planning (consultation of social worker, OT home visit, discharge planning w/caregivers)
Rahkonen 2001	yes; nurse specialist, physician, physiotherapist, neuropsychologist, OT. Speech therapist	No	Continuous & systemic support by a nurse specialist and one rehabilitation period (for 1 wk) pt received individual recd physiotherapy.	in-home visits and phone contacts for 3 years	medical examinations; nurse conducted dementia,	No	No	No	Special aids for daily living (e.g. hearing aids) arranged for the rehab week	Physiotherapy; Special aids for daily living (e.g. special shoes) arranged when	No		No	Community care plan arranged with pt, relative and professional from district social & health

2

# 1 14 Pharmacological treatment

## 2 14.1 Clinical introduction

3 Delirium is characterised by a range of symptoms that can cause distress,  
4 behaviour disturbance and place people at risk. Medications are used in clinical  
5 practice to manage these symptoms though the evidence base remains limited.  
6 Pharmacological agents that alter the course of delirium or control particular  
7 symptoms will need to demonstrate safety as well as effectiveness but would be  
8 a valuable development in treatment.

9

10 The pathophysiology of delirium is complex and people with delirium may have  
11 serious physical illness that complicates the use of drug treatment. Should drugs  
12 be given routinely or for selected symptoms? If selected symptoms then for which  
13 symptoms? Does the clinical context alter decisions about drug treatments?  
14 Would all people receive them or those at risk? These are questions for which  
15 answers are needed.

16

## 17 14.2 Description of studies

18 Twenty-three papers were evaluated for inclusion. Two Cochrane Reviews  
19 (Lonergeren 2009; Overshott 2008) were identified and updated. Sixteen studies  
20 were excluded: eight because there were fewer than 20 patients in each arm  
21 (Aakerlund 1994; Breitbart 1996; Han 2004; Horikawa 2003; Kim 2003;  
22 Maneeton 2007; Mittal 2004; Sasaki 2003); two because there were fewer  
23 than 20 patients in one arm (Nakamura 1995; Nakamura 1997); one because  
24 delirium was induced by morphine (Morita 2005). Three studies had a before  
25 and after study design (Bayindir 2001; Ikezawa 2008; Paradella 2004); and  
26 one was excluded because one of the interventions was not licensed in the UK for  
27 any indication (Pandharipande 2007; dexmedetomidine versus lorazepam). One  
28 other study was excluded because it was not a primary study (Appendix G).

29

30 Three studies were included that had randomised (Hu 2006; Lee 2005); or  
31 quasi-randomised designs (Skrobik 2004).

32

33 Two non-randomised comparative studies comparing typical and atypical  
34 antipsychotics were also included initially, because their comparator for  
35 haloperidol was risperidone, rather than olanzapine (which was used in the  
36 RCTs). Both had retrospective comparative designs, in which patients were  
37 selected from records (Liu 2004; Miyaji 2007). In the Liu (2004) study, patients  
38 were treated at the clinician's choice; in the other (Miyaji 2007), allocation was  
39 presumed to be by clinician choice but this was not stated. In the Liu (2004)  
40 study, there was a large difference in age between the risperidone and

1 haloperidol groups (risperidone 68 years, range 40–85 years; haloperidol 50  
2 years, range 15–77 years). In the Miyaji (2007) study, the participants in the  
3 injection haloperidol group were significantly younger than those in the other two  
4 groups (median 69 years versus 73 years).

5  
6 In view of these methodological limitations, the GDG decided to exclude these  
7 two studies from the review, and to rely on the class effect for the comparison  
8 between typical and atypical antipsychotics. Therefore these two non-  
9 randomised studies were not considered further except for the adverse effects  
10 review (chapter 11).

11  
12 Thus the efficacy review focuses on three studies (Hu 2006; Lee 2005; Skrobik  
13 2004).

#### 15 **14.2.1 Study Design**

16 None of the studies were conducted in the UK. One study was carried out in  
17 China (Hu 2006); one in Korea (Lee 2005) and one in Canada (Skrobik 2004).

18 One study (Skrobik 2004) received some funding from Eli Lilly and the other two  
19 studies did not state a funding source.

20 One study had fewer than 50 patients (Lee 2005, n = 40). One study had more  
21 than 50, but fewer than 100 patients (Skrobik 2004, n = 77) and the other  
22 study had more than 100 patients (Hu 2006, n = 180).

#### 24 **14.2.2 Population**

25 One study (Skrobik 2004) was in an ICU, in which the patients were mostly  
26 surgical (48 elective operations; 21 urgent operations; 4 medical patients), and  
27 patients were treated within 2 hours of the diagnosis of delirium.

28 The two other studies had patients in a non-ICU hospital setting. In the Hu (2006)  
29 study, the type of ward was not stated, but the patients had 'senile delirium' due  
30 to metabolic (n = 68), toxic (n = 47), structural (n = 25) or infectious (n = 35)  
31 causes; the duration of delirium was reported to be between 30 minutes and 17  
32 days. In the Lee (2005) study, patients had been referred to a psychiatric  
33 consultation service from departments of neurosurgery, internal medicine,  
34 neurology and rehabilitation medicine: those who had immediately recovered  
35 from a major operation were excluded.

36 Different methods were used to diagnose delirium, however, all the studies used  
37 the DSM-IV criteria in some form: in the ICU study (Skrobik 2004), patients were  
38 screened using the ICU-Delirium Screening Checklist, then if they scored 4 or

1 more (or had a clinical diagnosis of delirium); this was confirmed using DSM-IV  
2 criteria. In the Hu (2006) study, patients were assessed using the DSM-IV criteria.  
3 They also had to have a total score on the Delirium Rating Scale of 12 or more,  
4 and a clinical global impression scale: severity of illness (CGI-SI) score of 4 or  
5 more. In the Lee (2005) study, patients meeting the criteria for delirium were  
6 diagnosed using the DSM-IV criteria and evaluated using the Delirium Rating  
7 Scale-Revised-98 (DRS-R-98). This includes a 16-item scale to diagnose delirium  
8 and the 13-item severity subscale.

9 None of the studies reported whether the patients had dementia or cognitive  
10 impairment, although the Lee (2005) study excluded patients who had a  
11 previous history of a 'psychiatric disorder'.

12  
13 The age range across the studies was 42 to 99 years, with the mean age  
14 ranging from 61 to 74 years. All studies included men and women. Ethnicity was  
15 not reported.

### 16 17 **14.2.3 Interventions**

18 The included studies investigated the following drugs: typical antipsychotics  
19 (haloperidol) and atypical antipsychotics (amisulpride, olanzapine, and  
20 quetiapine) in the treatment of delirium in a hospital setting. The interventions  
21 were:

#### 22 • Haloperidol

- 23 ○ orally or by enteral tube: given within 2 h of the diagnosis of  
24 delirium, initially 2.5–5 mg every 8 hours (patients over 60 years  
25 0.5–1 mg) then titrated based on clinical judgement for up to 5  
26 days (Skrobik 2004)
- 27 ○ intramuscular injection 2.5–10 mg per day, depending on  
28 response; the effect was observed for one week; delirium had  
29 occurred from 30 min to 17 days (Hu 2006)

#### 30 • Olanzapine

- 31 ○ orally or by enteral tube: given within 2 h of the diagnosis of  
32 delirium, initially 5 mg per day (patients over 60 years 2.5 mg)  
33 then titrated based on clinical judgement for up to 5 days  
34 (Skrobik 2004)
- 35 ○ orally or sublingually initial dose 1.25–2.5 mg then adjusted,  
36 depending on response, to 1.25–20 mg per day; the effect was  
37 observed for one week; delirium had occurred from 30 min to 17  
38 days (Hu 2006)

#### 39 • Amisulpride

- 40 ○ 50–800 mg/day (initial dose mean: 156.4 mg/day (SD 97.5));  
41 the dose was flexible according to clinicians preferences and  
42 experience; it was unclear when the drug was administered  
43 following the diagnosis of delirium; treatment was administered  
44 until the CGI score reached 2 or less; mean duration of  
45 stabilisation was 6.3 (SD 4.4) days (Lee 2005)

- 1           • Quetiapine
- 2                 ○ 50–300 mg/day (initial dose mean: 113 mg/day (SD 85.5)); the
- 3                 dose was flexible according to clinicians preferences and
- 4                 experience; it was unclear when the drug was administered
- 5                 following the diagnosis of delirium; treatment was administered
- 6                 until the CGI score reached 2 or less; mean duration of
- 7                 stabilisation was 7.4 (SD 4.1) days (Lee 2005)

8

#### 9   **14.2.4           Comparisons**

10           The following comparisons were carried out:

- 11           • Typical antipsychotic (haloperidol) versus no treatment (Hu 2006)
- 12                 ○ all patients also had ‘somatic treatment aiming at delirium’
- 13           • Atypical antipsychotic (olanzapine) versus no treatment (Hu 2006)
- 14                 ○ all patients also had ‘somatic treatment aiming at delirium’
- 15           • Comparison of two drugs in the same class (atypical antipsychotics)
- 16                 ○ Amisulpride versus Quetiapine (Lee 2005)
- 17           • Comparison of two drug classes
- 18                 ○ Typical antipsychotic (haloperidol) versus atypical antipsychotic
- 19                 (olanzapine) (Hu 2006; Skrobik 2004)
- 20                         - all patients in Hu (2006) also had ‘somatic treatment
- 21                         aiming at delirium’

22

23           One study (Skrobik 2004) reported that the patients received concurrent

24           benzodiazepines and fentanyl for analgesia; some patients also received other

25           sedatives; there was no significant difference between interventions for these

26           concurrent drugs or in the amount of rescue IV haloperidol used. The Hu (2006)

27           study reported that all patients received ‘somatic treatment for delirium’; and the

28           Lee (2005) study reported that other antipsychotics or benzodiazepines were not

29           allowed.

30

### 31   **14.3 Methodological quality**

#### 32   **14.3.1           Randomised and quasi-randomised studies**

33           The method of sequence generation was inadequate in the quasi-randomised

34           study (Skrobik 2004), in which the patients were allocated on an even/odd day

35           basis, and allocation concealment was also judged inadequate because the

36           sequence was likely to be known in advance. The methods of sequence

37           generation and allocation concealment were not stated in either of the two RCTs

38           (Hu 2006; Lee 2005).

39

1 In the Skrobik (2004) study, outcomes were assessed by a clinician or research  
2 nurse blinded to the allocation; it was unclear whether patients were blinded, but  
3 this was unlikely because the frequency of dosing was different. In the other two  
4 studies, it was unclear whether assessors were blinded, and it was also unclear if  
5 the patients in the Lee (2005) study were blinded. In Hu (2006) it was unlikely  
6 that the patients were blinded because of the nature of the interventions (no  
7 placebos and different routes of administration for the active drugs).

8  
9 None of the studies reported an *a priori* sample size calculation.

10  
11 In the Skrobik (2004) study, patients were comparable on gender, weight and  
12 APACHE score, but those on haloperidol were significantly younger. In the Lee  
13 (2005) study, there were no significant differences between the groups on age,  
14 gender, baseline DRS-R-98 and CGI scores. In the Hu (2006) study, there were  
15 no significant differences between the groups on age, gender, pre-treatment  
16 severity of mental symptoms or causes of delirium.

17  
18 Two studies had less than 20% missing data in either group (Hu 2006; Skrobik  
19 2004). One study (Lee 2005) had more than 20% missing data: 5/20 (25%)  
20 dropped out from the quetiapine group and 4/20 (20%) from the amisulpride  
21 group; only patients who completed the study were included in the analysis. In  
22 the Skrobik (2004) study, patients were analysed according to their allocation  
23 group; and the Hu (2006) study, carried out an available case analysis.

24  
25 All the studies used an adequate method of delirium assessment at baseline  
26 [DSM-IV; ICDSC (Skrobik 2004)] and used an adequate method of assessment to  
27 evaluate delirium following treatment (Hu 2006: DRS; Lee 2005: DRS-R-98,  
28 administered by a trained psychologist; Skrobik 2004: Delirium Index (DI) scale  
29 administered by a trained clinician).

30  
31 Two studies (Hu 2006; Lee 2005) also used the CGI scale to evaluate treatment  
32 effects. The GDG noted that the CGI scale is not a direct measure of delirium  
33 and needs to be interpreted accordingly.

34  
35 Overall, the Skrobik (2004) study was considered to be at high risk of bias  
36 because there was inadequate allocation concealment, the patients were not  
37 blinded and there was a significant difference in patient age. In addition, the  
38 patients received rescue medication which may have confounded the outcome  
39 measures. The other two studies also had some potential for bias because the  
40 patients were unlikely to be blinded (Hu 2006), and because of more than 20%  
41 missing data in one group (Lee 2005).

## 42 43 **14.4 Results**

### 44 **A) TYPICAL ANTIPSYCHOTICS VERSUS PLACEBO/NO TREATMENT**

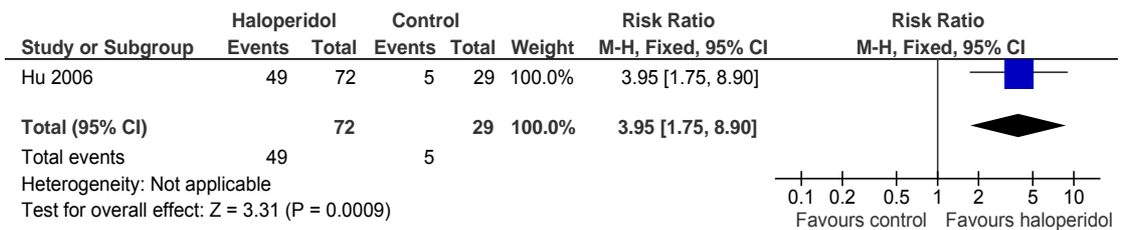
45 One RCT (Hu 2006) compared a typical antipsychotic (haloperidol) versus a no  
46 treatment control.

#### 47 48 **14.4.1 Primary Outcomes**

1 **14.4.1.1 Complete response**

2 One study Hu (2006) in 101 patients reported the measure of recovery from  
 3 delirium as 'symptoms alleviated or disappeared completely' on the global  
 4 improvement item of the CGI (CGI-GI) at 7 days. The analysis showed a  
 5 significant improvement of delirium in the haloperidol group compared to the  
 6 control group, although the result is imprecise (figure 14.1); RR 3.95 (95% CI  
 7 1.75 to 8.90). This corresponds to an NNT of 2 (95% CI 2 to 3) for a control  
 8 group rate of 17%.

10 Figure 14.1: complete response



13 **14.4.1.2 Duration of delirium**

14 The Hu (2006) study reported the 'time to take effect', the mean number of days  
 15 for the drug to take into effect, in responders only. The GDG considered that  
 16 these results were potentially biased and did not consider 'time to take effect'  
 17 was an adequate proxy/surrogate outcome for duration of delirium. Therefore  
 18 the results are not reported.

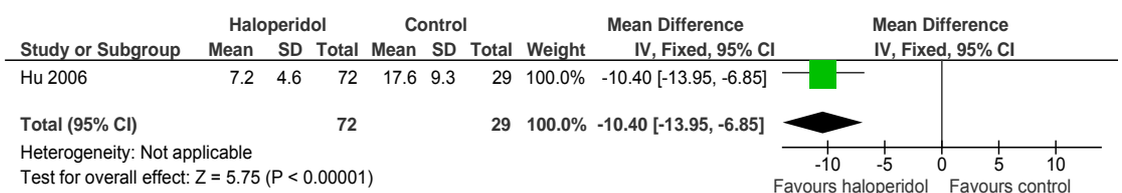
20 **14.4.2 Secondary Outcomes**

21 **14.4.2.1 Severity of delirium**

22 The Hu (2006) study reported scores on the DRS (0 to 32 scale) following  
 23 treatment. The severity of delirium assessed at the seventh day of treatment was  
 24 significantly lower in the haloperidol group (figure 14.2); MD: -10.40 (95% CI -  
 25 13.95 to -6.85) for a control group severity score of 17.6.

27 This study also reported the scores on the CGI-SI. These were 1.79 (SD 1.12) for  
 28 haloperidol and 3.97 (SD 1.76) for the control group. The GDG stated this scale  
 29 is not a direct measure of delirium and should be interpreted accordingly.

31 Figure 14.2: severity of delirium



## B) ATYPICAL ANTIPSYCHOTICS VERSUS PLACEBO/NO TREATMENT

One RCT (Hu 2006) compared an atypical antipsychotic (olanzapine) versus a no treatment control.

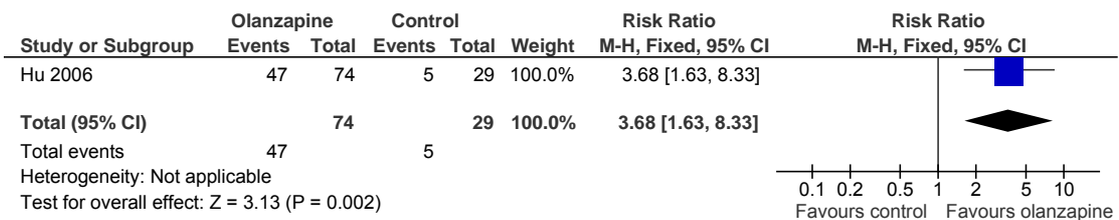
### 14.4.3 Primary outcomes:

#### 14.4.3.1 Recovery from delirium (complete response); figure 14.3

One study Hu (2006) with 103 patients reported the 'symptoms alleviated or disappeared completely' on the global improvement item of the CGI-GI scale at 7 days.

The analysis showed a significant improvement of delirium in the olanzapine group compared to the control group, but the result is imprecise; RR 3.68 (95% CI 1.63 to 8.33). This corresponds to an NNT of 3 (95% CI 2 to 4) for a control group rate of 17%.

Figure 14.3: complete response



#### 14.4.3.2 Duration of delirium

The Hu (2006) study reported the 'time to take effect', in responders only, but again this outcome was considered to be biased and are not reported here.

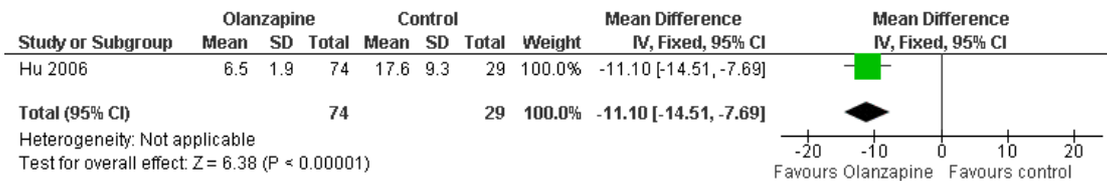
### 14.4.4 Secondary outcomes

#### 14.4.4.1 Severity of delirium

One study (Hu 2006) in 103 patients reported scores on the DRS (0 to 32 scale). There was a large significant difference between the treatments on this measure; mean difference: -11.10 (95% CI -7.69 to -14.51) for a control group severity score of 17.6 (figure 14.4).

This study also reported the scores on the CGI-SI. These were 2.05 (SD 0.99) for olanzapine and 3.97 (SD 1.76) for the control group. The GDG stated this scale is not a direct measure of delirium and should be interpreted accordingly.

1 Figure 14.4: severity of delirium



2

3 NB: Scale -20 to 20

4

5 **C) ATYPICAL ANTIPSYCHOTIC 1 VERSUS ATYPICAL ANTIPSYCHOTIC 2**

6 **14.4.5 Amisulpride versus Quetiapine**

7 One study (Lee 2005) compared two atypical antipsychotic drugs. It is noted  
8 that the study size was very small (40 patients randomised, but only 31  
9 analysed) and that this study cannot be expected to determine a difference  
10 between two active interventions.

11

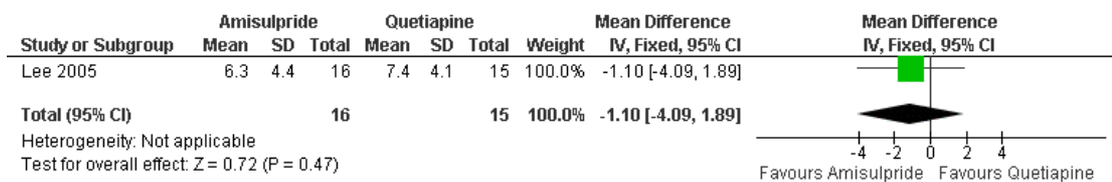
12 **14.4.6 Primary outcome**

13 **14.4.6.1 Duration of delirium**

14 One study (Lee 2005) with 31 patients reported the mean 'duration of  
15 stabilisation', which was the time for the patients to reach recovery from delirium;  
16 there was no significant difference between groups; but the result is imprecise;  
17 MD: -1.10 days (95%CI -4.09 to 1.89), for a duration of 7.4 days for the  
18 quetiapine group (figure 14.5).

19

20 Figure 14.5: duration of delirium



21  
22

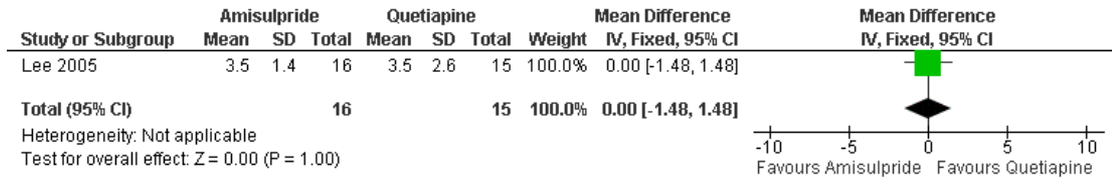
23 **14.4.7 Secondary outcomes**

24 **14.4.7.1 Severity of delirium**

25 One study (Lee 2005) with 31 patients reported scores on the DRS-R-98 (0 to  
26 39 scale); there was no significant difference between the treatments; mean  
27 difference 0.00 (95%CI -1.48 to 1.48) for a severity score of 3.5 in the  
28 quetiapine group (figure 14.6).

1  
2

Figure 14.6: severity of delirium

3  
45 **14.4.7.2 Adverse effects**

6 The Lee (2005) study reported that there were no serious adverse events  
7 observed, such as acute dystonia and dyskinesia, but there were very few  
8 patients in this study.

9

10

11

**D) TYPICAL ANTIPSYCHOTICS VERSUS ATYPICAL ANTIPSYCHOTICS**

12

13

One RCT (Hu 2006) and one quasi randomised study compared a typical  
antipsychotic (haloperidol) versus an atypical antipsychotic (olanzapine).

14

14.4.8 **Primary outcomes**

15

14.4.9 **Complete response**

16

17

18

19

Both randomised/quasi-randomised studies evaluated a measure of recovery  
from delirium, although these were reported differently and neither constituted a  
direct outcome measure (Hu 2006; Skrobik 2004).

20

21

22

Hu (2006) reported the 'symptoms alleviated or disappeared completely on the  
global improvement item of the clinical global impression scale' at 7 days.

23

24

25

26

27

28

29

Skrobik (2004) reported the numbers of patients requiring rescue IV haloperidol  
on day 1 (19/45 patients on haloperidol and 10/28 on olanzapine) and the  
numbers for subsequent days (4/45 haloperidol and 1/28 olanzapine). This was  
converted into the numbers not requiring rescue medication (by subtraction), i.e.  
22/45 on haloperidol and 17/28 on olanzapine. This was assumed to be an  
approximation to a complete response to study treatment.

30

31

32

33

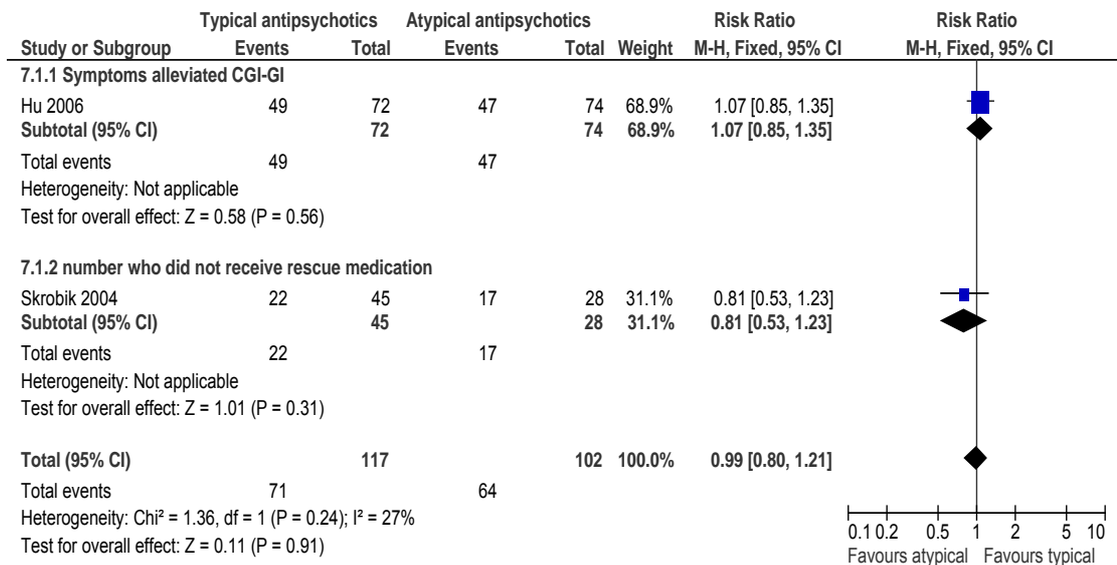
34

35

36

Meta-analysis of the two studies in 219 patients did not demonstrate a  
significant difference between the treatments (figure 14.7); RR 0.99 (95% CI  
0.80 to 1.21). There was insignificant heterogeneity between studies ( $I^2 = 27%$ ;  
 $p = 0.24$ ). In the absence of the Skrobik (2004) study, which was at much higher  
risk of bias, there was no significant difference between interventions; RR 1.07  
(95%CI 0.85 to 1.35).

1 Figure 14.7: complete response



2  
3  
4

5 **14.4.10 Duration of delirium**

6 The Hu (2006) study reported the ‘time to take effect’, in responders only, but  
7 again this outcome was considered to be biased and are not reported here.

8

9 **14.5 Secondary outcomes**

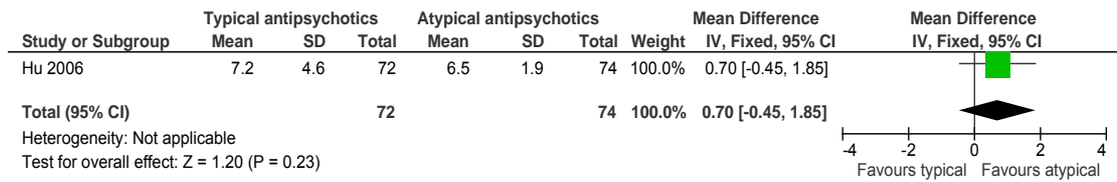
10 **14.5.1 Severity of delirium**

11 One study (Hu 2006) reported scores on the DRS (0 to 32 scale; figure 14.8),  
12 which showed no significant difference between the treatments on this measure;  
13 mean difference 0.70 (95% CI -0.45 to 1.85) for a control group severity of  
14 6.5 units.

15  
16 This study also reported the scores on the CGI-SI. These were 1.79 (SD 1.12) for  
17 haloperidol and 2.05 (SD 0.99) for olanzapine. The GDG considered the CGI  
18 scale was likely to be less specific for measuring delirium symptoms than the DRS.

19  
20

1 Figure 14.8: severity of delirium

2  
3

NB: Scale -4 to 4

4

5

The quasi-randomised study (Skrobik 2004) reported the mean daily delirium index scores on a graph. The mean daily delirium index scores at day 5 were haloperidol 4.85, olanzapine 5.40, mean difference 0.55; there was no significant difference between interventions ( $p=0.83$ ). It is noted that these data were likely to be confounded by the use of rescue IV haloperidol medication, predominantly on the first day in around a third of the patients in each group.

9

10

11

12

13 **14.6 Clinical evidence statements**

14

Refer to Appendix K for the GRADE profile.

15

16 **14.6.1 Typical antipsychotics versus placebo/no treatment**

17

- There is moderate quality evidence from one RCT showing a:

18

- significant improvement of delirium in the haloperidol group compared with no treatment at 7 days. There is some uncertainty around this result.

19

20

21

- significantly lower severity of delirium in the haloperidol group compared with no treatment (an indirect measure of delirium was used).

22

23

24

25 **14.6.2 Atypical antipsychotics versus placebo/no treatment**

26

- There is moderate quality evidence from one RCT showing a:

27

- significant recovery from delirium in favour of the olanzapine group compared with no treatment at 7 days. There is much uncertainty with this result.

28

29

30

- significantly lower severity of delirium in the olanzapine group compared with no treatment (an indirect measure of delirium was used).

31

32

33

34

### 1 14.6.3 Atypical antipsychotic 1 versus atypical antipsychotic 2

- 2 • There is very low quality evidence from one RCT showing no significant  
3 difference in the duration of delirium between amisulpride and  
4 quetiapine groups. There is some uncertainty with this result.
- 5 • There is low quality evidence from one RCT showing no significant  
6 difference in the severity of delirium between amisulpride and  
7 quetiapine groups.

8

### 9 14.6.4 Typical antipsychotics versus atypical antipsychotics

- 10 • There is low quality evidence from a meta-analysis of two studies [one RCT  
11 and one quasi-RCT] showing no significant difference in recovery from  
12 delirium between the haloperidol and olanzapine groups.
- 13 • There is moderate quality evidence from one RCT showing no significant  
14 difference in the severity of delirium between the haloperidol and the  
15 olanzapine groups (an indirect measure of delirium was used).

16

17

## 18 14.7 Health economic evidence statements

19

20 The results of the economic model (chapter 16) showed the following:

- 21 • The use of haloperidol and olanzapine was cost-effective in the treatment  
22 of delirium in the hospital. This finding was robust as the interventions  
23 remained cost-effective after a series of sensitivity analyses were  
24 conducted.
- 25 • Haloperidol was more cost-effective than olanzapine in the treatment of  
26 delirium in the hospital. However, there was a wide uncertainty around  
27 the incremental cost-effectiveness of haloperidol compared to  
28 olanzapine.

29

30

# 1 15 What information is useful for people 2 with delirium and their carers?

## 3 4 15.1 Clinical Introduction

5  
6 Delirium can be a distressing experience for affected individuals, family  
7 caregivers and professionals. The symptoms can be complex and full or partial  
8 recall after the episode has resolved is common. Sometimes this can result in  
9 unpleasant “flashback” episodes. Information and education to improve  
10 understanding of delirium and its effects might help to improve outcomes from  
11 the condition.

## 12 13 15.2 Description of studies

14  
15 Twenty four studies were ordered for this review. Fourteen studies were  
16 excluded.

17  
18 One included study was UK based. There were four Swedish studies, two  
19 studies conducted in the USA and one each in Australia, Canada and Finland.

20  
21 One non randomised control trial was reviewed and nine qualitative studies  
22 were critically appraised using the NICE qualitative methodology checklist. Two  
23 of the included studies used a phenomenologic approach, one study used a  
24 hermeneutic approach and another study used a combined phenomenologic-  
25 hermeneutic approach. There were three studies which employed content  
26 analysis to elicit categories and themes based upon patient interviews and one  
27 further study which used an interview questionnaire to obtain subjective  
28 responses of family carers to the experience of delirium.

29  
30 One of the included studies described an information giving intervention in a  
31 hospice setting. Although people receiving end-of-life care are excluded from  
32 the guideline, this was the only comparative study identified and the only study  
33 which assessed the actual development and implementation of a delirium  
34 educational tool for family caregivers. It was considered that the information in  
35 this study could be imputed to other settings.

## 36 37 15.3 Results

38  
39 Owens & Hutelmyer (1981) conducted a non randomised control trial among 64  
40 adults having cardiac surgery. The study tested the hypothesis that patients who  
41 are educated pre-operatively about the possibility of unusual sensory or  
42 cognitive experiences will not have such experiences postoperatively or will feel  
43 comfortable or in control of the experiences if they occur. Patients were  
44 assigned on a consecutive admission basis to either the intervention or control  
45 group. The staff did not discuss the psychological aspects of postoperative care  
46 with any participants. The investigator discussed the possibility of memory loss,

1 inability to concentrate, inability to recognise familiar objects or persons and the  
2 possibility of seeing or hearing things that could not be explained or were not  
3 really there with the experimental group only. Post-operative interviews were  
4 conducted on days 4-8. Of the 32 patients in the control group, 25 reported at  
5 least one unusual experience. In the experimental group, 19 patients reported  
6 such experiences. The difference was not statistically significant. When the  
7 groups were compared as to whether they felt comfortable or in control during  
8 an unusual experience, the experimental group was significantly ( $p < .05$ ) more  
9 comfortable.

10  
11 Margarey and McCutcheon (2005) interviewed eight patients who had  
12 experienced hallucinations during an ICU admission. Most of these patients  
13 remembered the nurses talking to them even if they did not recall the ICU  
14 environment. Reassurance and comfort from the nurses was important to patients,  
15 particularly reassurance that the experience of delirium is common and that they  
16 were not going mad. The presence of family members was associated with the  
17 beginning of recovery. The authors of this study suggest that post ICU clinics to  
18 allow patients to discuss the experience of delirium and post ICU visits so that  
19 patients can put their experiences into context may be useful.

20  
21 Duppils and Wikblad (2006) interviewed 15 patients who had undergone hip-  
22 related surgery and experienced delirium during their hospital stay. Difficulty in  
23 communication was identified as one of the risk factors in delirium. Patients  
24 complained that the nurses talked 'about' them, not 'to' them. Nurses were  
25 encouraged to try to understand the patients thought and experiences in order  
26 to communicate information in a therapeutic manner.

27  
28 Nineteen patients who had been ventilated and stayed at least 36 hours in the  
29 ICU were interviewed by Granberg et al (1998) about one week after  
30 discharge and again 4-8 weeks after their discharge from the ICU. Patients  
31 described their first feelings and memories after delirium. Relatives provided a  
32 lifeline to reality. Patients were very sensitive to the attitude and behaviour of  
33 staff. They also reported the effort to regain control over their bodies. Patient  
34 reaction to the equipment of ICU which is unfamiliar and uncomfortable and limits  
35 mobility resulted in fear and tension. Caring nurses could provide rest from a  
36 state of prolonged tension and engender a feeling of security by helping with  
37 orientation to the surroundings and providing a sense of 'We are with you.' It  
38 was important for patients to know that unreal experiences are common and that  
39 their intellectual capacity would not be impaired. They appreciated nurses who  
40 would explain equipment and procedures and who understood that they needed  
41 help to regain control over their bodies.

42  
43  
44 Heleena Laitinen (1996) conducted a study of 10 postoperative intensive care  
45 coronary artery by-pass patients. Implications for nursing practice were  
46 highlighted, particularly understanding and acceptance. Being aware of space  
47 and time gives patients more confidence for coping with being in the ICU.  
48 Consciousness of space and time presumes that events and stimuli in the  
49 environment are constantly being explained to the patient in a sensitive manner.  
50

1 Ewa Stenwall et al (2008) interviewed seven geriatric patients who had  
2 experienced acute confusional state (ACS; delirium). Patients stated that gaining  
3 knowledge about what was happening and what was planned evoked feelings  
4 of safety.

5  
6 Good communication occurs through the senses. Relatives can inform carers which  
7 sense the patient prefers and which sense is less efficient.

8  
9 Another study by Stenwell et al (2008) explored the experience of relatives of  
10 patients with delirium. The conclusions of this study with regard to information  
11 giving were as follows:

- 12 • Relatives need information about acute confusional state (delirium) to  
13 alleviate their insecurity about interactions with the patient and to aid  
14 their understanding of the patient's behaviour which will allow trust to  
15 develop. It is necessary to inform relatives of the short term nature of  
16 ACS and the need to have support and advice from professionals on how  
17 to communicate.
- 18 • Relative's knowledge of the patient should be used to inform the  
19 communication style of carers with that individual. Communication must  
20 be responsive to the individual encounter.

21  
22 Fourteen elders participated in a phenomenologic study describing the  
23 experience of delirium patients (McCurren & Cronin, 2003). Three themes were  
24 identified:

- 25 • Being in the confusion event
- 26 • Responding to confusion
- 27 • Dealing with confusion

28  
29 The latter theme involved the responses of family, staff and the patient. Among  
30 the interventions which helped with delirium included explanations from nurses  
31 which helped to reassure patients and families. Anticipatory explanations for  
32 surgical patients were also identified as helpful.

33  
34 Another interpretative phenomenological analysis of nine patients (Harding  
35 2008) aimed to understand the delirium experience of older people after  
36 reparative hip surgery. Semi structured interviews were conducted and two  
37 primary themes were identified:

- 38 • Struggling to understand the experience of delirium
- 39 • Strategies used in discussing delirium

40  
41 Based upon an in-depth analysis of the experiences and concerns of the  
42 participants the authors suggested the following:

- 43 • Providing information for patients and relatives (e.g. in a leaflet) to help  
44 them understand delirium

- Training healthcare staff to help facilitate open discussions with patients about their delirious symptoms and supervision to help staff better understand and manage their own anxieties.

A psycho-educational intervention was implemented in a palliative care hospice to help family caregivers cope with delirium and eventually to contribute to early detection (Gagnon et al, 2002). Phase 1 of this study aimed to develop the framework of an optimal psycho-educational intervention about delirium through focus group discussion. Phase 2 was the development of a brochure to be used as part of the psycho-educational intervention and Phase 3 included the implementation and evaluation of the intervention by comparing 58 family who received 'usual care' and 66 families who received explanations by nursing staff and a brochure on delirium. The delirium brochure included the symptoms of delirium, the cause of delirium, staff actions when a patient has delirium and how to behave with a patient with delirium.

Those who received the intervention felt more competent in making decisions than those in the usual care group ( $p=0.006$ ) and the majority felt that all family caregivers should be informed on the risk of delirium ( $p<0.009$ ).

#### 15.4 Clinical evidence statements

Overall, the studies on information giving to patients employ a variety of qualitative methods, with typically small numbers of participants in each study. Papers on information giving address the needs of patients, professional staff and family carers and identify needs throughout the delirium continuum from pre-delirium, to the delirium experience itself and finally to the post-delirium state. The following recommendations for information sharing appear in the literature:

- Patients need insight into the experience of delirium to promote their understanding and to decrease fear. Pre-op information and a visit to the ICU are recommended.
- Nurses require insight into the patient experience in order to promote empathy.
- As relatives provide a link with reality and can facilitate communication, they require anticipatory information about the risk for delirium.
- Post-delirium it is recommended to offer the opportunity to discuss the experience and provide reassurance. Post-extubation time in the ICU or a post-ICU visit may help a patient understand his/her experience.

# 1 16 Cost-effectiveness analysis

## 2 16.1 Introduction

3

4 The occurrence of delirium has been shown in a systematic review to result in  
5 adverse consequences (section 8). The adverse consequences could lead to a  
6 reduction in patients' health-related quality of life, HRQoL, and the expenditure  
7 of the resources of the NHS or PSS. It will therefore be useful to know the cost-  
8 effectiveness of prevention and treatment interventions for delirium.

9

10 We searched the literature for existing cost-effectiveness results that could  
11 reliably inform the guideline recommendations and we identified four papers  
12 (Rizzo et al 2001, Pitkala et al 2008, Bracco et al 2007, Robinson et al 2002).  
13 However, none of them were felt to be directly applicable to the guideline  
14 population. It therefore became necessary to develop an original economic  
15 evaluation model to determine the cost-effectiveness of strategies for the  
16 prevention and treatment of delirium in different care settings. As described  
17 above in the general cost-effectiveness method section (section 2.6), the model  
18 was constructed for prevention and treatment interventions in hospital care  
19 setting.

20

### 21 16.1.1 Interventions

22 There were a number of interventions strategies included in the systematic review  
23 of prevention and treatment interventions (chapters 9, 10, 13 and 14). However,  
24 after considering the existing evidence, the GDG wanted more information on  
25 the cost-effectiveness of two multi-component prevention interventions and two  
26 pharmacological treatment interventions. They advised that these should be  
27 evaluated in the economic model. The two multi-component prevention  
28 interventions were those included in the Inouye et al study (1999) and  
29 Marcantonio et al study (2001). The two pharmacological treatment interventions  
30 were those in Hu et al (2006). These studies have been described fully (chapters  
31 9 and 14).

32

33 Study participants in the Inouye et al (1999) study were consecutive patients  
34 admitted to the general medicine service in the non-intensive care section  
35 between March 1995 and March 1998. Patients were at least 70 years old,  
36 had no delirium at the time of admission, and were at intermediate or high risk  
37 for delirium at base line. There were 852 patients in the study and half of the  
38 sample received the multi-component targeted intervention, Elder Life Program.  
39 They received standard protocols for the management of six risk factors for  
40 delirium namely, cognitive impairment, sleep deprivation, immobility, visual  
41 impairment, hearing impairment, and dehydration. Geriatric nursing assessment  
42 and interdisciplinary rounds were other program interventions targeted towards

1 the risk factors. Patients in the usual care group received standard hospital  
2 services in the general-medicine unit.

3  
4 Study participants in the Marcantonio et al study were 65 years old or older  
5 patients and were admitted non-electively for surgical repair of hip fracture.  
6 Patients in the intervention group received proactive geriatric consultation, which  
7 began preoperatively or within 24 hours of surgery. They received targeted  
8 recommendations based on a structured protocol from the geriatrician during the  
9 period of hospitalization. Patients in the control group received usual care. They  
10 received management by the orthopaedics team, including internal medicine  
11 consultants or geriatricians on a reactive rather than proactive basis.

12  
13 The study participants in the Hu et al study were elderly inpatients with senile  
14 delirium selected from September 2001 to September 2003. The enrolled  
15 patients were divided into three groups including two treatment groups and a  
16 control group. Each of the two treatment groups received somatic treatment in  
17 addition to either haloperidol or olanzapine. The control group received only  
18 somatic treatment only.

#### 20 **16.1.2 Population**

21 The model was developed for patients in hospital settings. The two multi-  
22 component interventions were targeted at patients with specific risk factors for  
23 delirium while the treatment interventions were indicated for patients with  
24 delirium. For the prevention interventions, we chose to model the cost-  
25 effectiveness in the trial population rather than extrapolate to other populations  
26 as the patients were selected on the basis of specific risk factors and the  
27 intervention was targeted at modifying those specific risk factors. Therefore the  
28 GDG felt that the efficacy may not translate to other populations. The starting  
29 age used in the model was 79 years. This was based on the mean age reported  
30 in the largest of the three studies above (Inouye et al 1999).

#### 32 **16.1.3 Outcomes**

33 The outcomes of interest for the model were the incremental cost and the  
34 incremental quality-adjusted life years (QALY) gained. These were used to  
35 calculate the incremental cost effectiveness ratio (ICER) and the incremental net  
36 monetary benefit (INMB).

## 1 16.2 The prevention model

2

### 3 16.2.1 The model structure for the prevention interventions

4

#### 5 16.2.1.1 Decision Tree

6

7 The cost-effectiveness model consists of a simple decision tree which captures the  
8 outcomes of economic importance. The outcomes at the end of each branch of the  
9 tree include the adverse consequences of delirium. These outcomes will  
10 negatively impact on patient's health status and will lead to the expenditure of  
11 the resources of the NHS and PSS. The GDG advised that the adverse  
12 consequences to be used in the economic model should include falls, pressure  
13 ulcer, new dementia, new admission to institution, extended stay in the hospital  
14 and fatality. The decision tree was applied to each strategy and was used to  
15 estimate the impact of each strategy on the expected number of delirium cases,  
16 cost and QALYs associated with the adverse consequences. The decision tree is  
17 as shown below in figure 16.1.

18

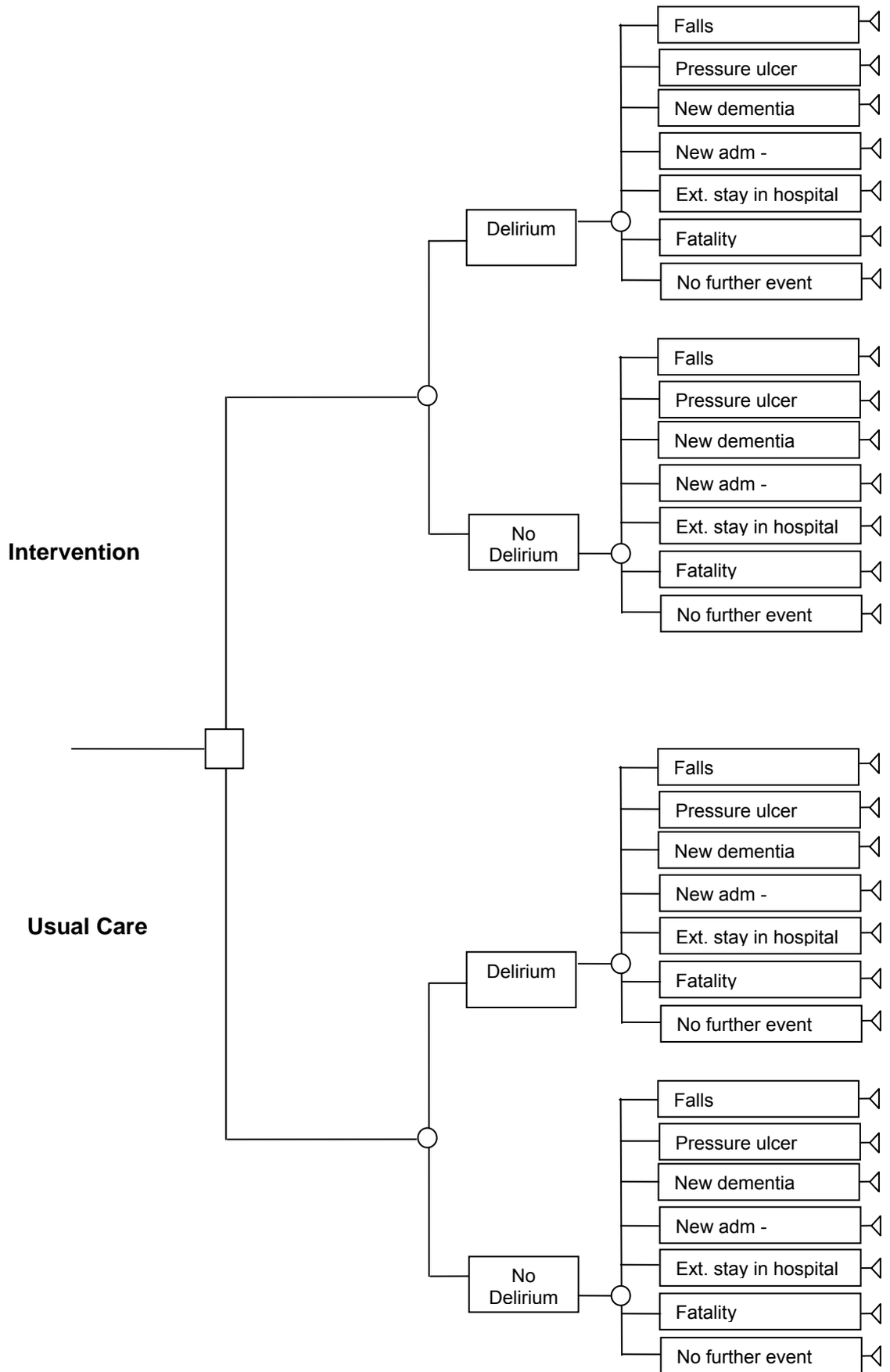
19 Some members of a hypothetical cohort receiving each intervention strategy will  
20 become delirious and others will not. In the usual care strategy, the number that  
21 will become delirious will depend on the baseline risk of delirium in the care  
22 setting. The baseline risk of delirium is the risk of becoming delirious under no  
23 intervention conditions. In the intervention strategy, the number will depend on  
24 the baseline risk as well as the relative risk of becoming delirious if exposed to  
25 the intervention. The relative risk measure here is a measure of the efficacy of  
26 the intervention strategy. It is a ratio of the risk of becoming delirious among  
27 members of a population exposed to an intervention compared with a similar  
28 population that is not exposed to the intervention.

29

30 In non-delirious patients, the number of cases of the adverse consequences will  
31 depend on the baseline risk of the adverse consequence. In delirious patients, it  
32 will depend on the baseline risk as well as the relative risk of experiencing the  
33 adverse consequences if exposed to delirium. The end point of each branch of  
34 the tree implies a particular cost and a particular QALY. The total number of  
35 cases of delirium and the adverse consequences, the associated total cost and  
36 QALYs are summed up for each strategy.

37

1      Figure 16.1: decision tree for prevention intervention strategies



## 1 16.2.2 Baseline Risk

2

### 3 16.2.2.1 Hospital (*intervention in general medicine services*)

4 The baseline risk of delirium in the hospital was taken from a matched controlled  
5 trial in the USA (Inouye et al 1999). The study has been described in the review  
6 of prevention interventions (section 9.15). Study participants were consecutive  
7 patients admitted to the general medicine service in the non-intensive care  
8 section. Patients were at least 70 years old, had no delirium at the time of  
9 admission, and were at intermediate or high risk for delirium at base line. Half  
10 of the sample received the multi-component targeted intervention while the other  
11 half received usual care. Usual care was defined as standard hospital services in  
12 the general-medicine unit. Patients were screened and baseline assessments were  
13 completed within 48 hours after admission. They were subsequently evaluated  
14 daily until discharge with a structured interview consisting of the Digit Span Test,  
15 Mini-Mental State Examination, and Confusion Assessment Method rating. Their  
16 medical records were reviewed after discharge for evidence of delirium, final  
17 diagnosis, medications, laboratory results, and destination after discharge. The  
18 primary outcome of the study was delirium defined according to the Confusion  
19 Assessment Method criteria. The median lengths of stay in the intervention and  
20 usual care groups were 7.0 and 6.5 days respectively. The incidence of delirium  
21 in the usual care group was 15% and this was used in the model as the  
22 probability of delirium in this group of hospitalized patients. In a sensitivity  
23 analysis, we used a lower incidence of delirium of 12.5%, which was the lower  
24 range of incidence reported in the needs assessment review for general medical  
25 patients (chapter 5).

26

### 27 16.2.2.2 Hospital (*intervention in hip fracture surgery*)

28 The baseline risk of delirium in the hospital for this patient group was taken from  
29 a randomised trial in the USA (Marcantonio et al 2001). The trial has been  
30 described elsewhere (section 9.15). Study patients were 65 years old or older  
31 and were admitted non-electively for surgical repair of hip fracture. Patients in  
32 the intervention group received proactive geriatric consultation, which began  
33 preoperatively or within 24 hours of surgery. Patients in the control group  
34 received usual care. The median length of stay in both groups was 5 days and  
35 the cumulative incidence during acute hospitalization was reported as 50% in the  
36 usual care group. This estimate was used as the probability of delirium in this  
37 patient group. In a sensitivity analysis, we used the lower estimate (15%)  
38 reported above for patients in general medicine services.

39

### 1 **16.2.2.3 Dementia**

2 The baseline risk of dementia was taken from a Canadian prospective cohort  
3 study (Rockwood et al 1999). It has been described in the section on the review  
4 of delirium consequences (chapter 8). Study patients were 65 years old or older  
5 and were consecutively admitted to the general medicine services of a tertiary-  
6 care hospital. A study cohort of 203 patients was followed up between June  
7 1994 and August 1995, and dementia incidence as well as death was the  
8 primary outcome. Dementia diagnosis was done to conform to the Canadian  
9 Study of Health and Ageing dementia protocol. Dementia status was evaluated  
10 using the Informant Questionnaire on Cognitive Decline in the Elderly. Interview  
11 was obtained from proxy informants. A screening interview was also done to  
12 evaluate cognition and function. Cognition was done with the Blessed dementia  
13 rating scale while function was done with the Barthel index and the Physical Self-  
14 Maintenance Scale. The incidence of dementia in patients without cognitive  
15 delirium at baseline was reported as 5.6% per year. This baseline probability  
16 was used in the economic model.

17

### 18 **16.2.2.4 Pressure Ulcer**

19 The baseline risk of pressure ulcer was taken from a study that focussed on  
20 reporting the incidence of pressure sores across a NHS Trust hospital (Clark &  
21 Watts 1994). The number of patients admitted to the wards over 52 weeks  
22 were recorded alongside the number of those developing pressure sores. The  
23 severity and anatomical locations of pressure sores were also recorded. The  
24 incidence was monitored across four medical, three surgical and two orthopaedic  
25 wards and a record form was completed weekly. This enabled the identification  
26 of all patients that developed sores during the preceding seven days. The form  
27 also contained details of admissions and discharges from each ward and the  
28 details were obtained weekly. The number of people admitted in the wards as  
29 in-patients between December 1990 and November 1991 was 8935 and 360  
30 patients developed pressure sores. This is equivalent to an incidence of 4.03%  
31 which we used as the baseline probability of pressure ulcer in the model. Some  
32 of the patients may have had delirium and as such 4.03% could be an over-  
33 estimate. We therefore used 1.68% in a sensitivity analysis. The latter estimate  
34 was reported in the O'Keeffe and Lavan study (1997) where two out of 119  
35 non-delirious hospitalised patients acquired pressure sores. The latter study is  
36 briefly described in the next paragraph.

37

### 38 **16.2.2.5 Falls**

39 The baseline risk of falls was taken from a prospective cohort study in Ireland  
40 (O'Keeffe & Lavan, 1997). The study has been described in the section on  
41 review of delirium consequences and it aimed to determine whether delirium is  
42 an independent predictor of adverse outcomes of hospitalization in older  
43 patients. The study population was 225 people admitted as an emergency over  
44 an 18-month period to an acute geriatric unit in a university teaching hospital.  
45 Only those on first admission within the study period were included in the study.  
46 Patients were excluded if they were not admitted to the geriatric unit on the day

1 of admission, if they were admitted electively for investigations, rehabilitation, or  
2 respite care. Those that had severe aphasia or deafness, those that expected to  
3 remain in hospital for less than 48 hours, and those not assessed by a study  
4 doctor within 48 hours of admission were excluded. Patients were interviewed  
5 using the Delirium Assessment scale to elicit the presence and severity of  
6 individual DSM-III (Diagnostic and Statistical Manual, 3<sup>rd</sup> Edition) criteria for  
7 delirium. An initial assessment was done which included administration of an  
8 adapted Folstein Mini-Mental State Examination (MMSE) validated for use in an  
9 Irish population. All study patients were reviewed regularly and discussed with  
10 nursing and residential medical staff. The delirium status of patients was  
11 discussed at the multidisciplinary team meetings, and members of the team other  
12 than the study physicians were not aware of the underlying hypothesis of the  
13 study. Cases of falls, pressure sores, and urinary incontinence were recorded as  
14 hospital-acquired complications according to standardized criteria and were  
15 identified on the basis of interviews with the nursing staff. Pressure sore  
16 corresponds to grade 2 of Shea's classification. The number of patients studied  
17 was 225 and 42% had delirium defined by the DSM-3 criteria. The mean age  
18 of those with and without delirium was 82 years. Sixty eight percent of those  
19 without delirium were female and 16% of those without delirium were admitted  
20 from long-term care. Nine (7%) of the 131 non-delirious patients had falls, and  
21 we have used 7% as the baseline risk of falls in the economic model.

22

#### 23 **16.2.2.6 New admission to institution**

24 We took the baseline risk of new admission to long-term care (LTC) from a  
25 prospective cohort study and it has been described in the section on the review  
26 of delirium consequences (Bourdel-Marchasson et al 2004). The study was  
27 carried out in France with the aim of assessing the effects of delirium on the  
28 institutionalization rate in older patients hospitalized in an acute care geriatric  
29 unit, taking into account other components of frailty. Study participants were  
30 those older than 75 years old who were admitted between July 2000 and June  
31 2001. Patients were excluded from the analyses if they spent less than 3 days in  
32 hospital, died before discharge or were usually living in an institution. The  
33 assessment of delirium was done with CAM within 24 hours following admission  
34 and then every three days during the hospital stay. The outcome considered for  
35 the analyses of study results was admission to a geriatric institution. There were  
36 230 patients who were reported to be symptom free and 40 (17%) of these  
37 were discharged to geriatric institutions. We used 17% as the baseline risk of  
38 new admission to institute.

39

#### 40 **16.2.2.7 Mortality (in hospital)**

41 The baseline risk of in hospital mortality was taken from the O'Keeffe and Lavan  
42 study (1997) described above. It was reported in the study that five percent of  
43 patients without delirium died during hospitalization, and we used this estimate  
44 as the baseline risk of mortality.

1

2 **16.2.2.8 Mortality or new admission to institution**

3 We have assumed that the adverse outcomes on the decision tree are mutually  
4 exclusive. This could potentially lead to double counting and over-estimation of  
5 costs and QALYs as some patients will experience more than one outcome at a  
6 time. The consequences review reported data on the relative risk of “mortality or  
7 new admission to nursing home” in delirious patients and we used this composite  
8 outcome rather than the single outcomes “mortality” and “nursing home  
9 admission” in a sensitivity analysis. This should reduce the double-counting and  
10 over-estimation of costs and QALYs associated with using the single outcomes in  
11 the model. We explored the effect of this sensitivity analysis on the cost-  
12 effectiveness result. This analysis requires an estimate of baseline risk for this  
13 composite outcome.

14

15 The baseline risk of mortality or new admission to institution was taken from a  
16 prospective cohort study in the USA (Marcantonio et al 2000). The study has also  
17 been described in the section on consequences review. The aim was to evaluate  
18 the role of delirium in the natural history of functional recovery after hip fracture  
19 surgery, independent of pre-fracture status. The study data were collected as  
20 part of a randomised trial to test whether proactive acute geriatrics consultation  
21 could prevent delirium after hip fracture repair. The effect of the intervention  
22 could have potentially affected the relationship between delirium and functional  
23 recovery but it was reported that the effect size of the associations did not differ  
24 between the two groups. Study participants were patients aged 65 years or  
25 older who were admitted to an academic tertiary medical centre for primary  
26 surgical repair of hip fracture. Patients with metastatic cancer or other co-morbid  
27 illnesses likely to reduce life expectancy to less than six months were excluded  
28 from the study. Study participants were interviewed daily during the duration of  
29 the hospitalization, including the Mini-Mental State Examination and Delirium  
30 Symptom Interview, and delirium was diagnosed using the Confusion Assessment  
31 Methods algorithms. They or their proxies were further contacted one and six  
32 months after fracture. They underwent interviews similar to those at enrolment to  
33 determine death, persistent delirium, decline in Activities of Daily Living function,  
34 decline in ambulation, or new nursing home placement. It reported the  
35 percentage of non delirious patients who died or were admitted to nursing home  
36 institute one month after hip fracture to be 12% and we have used this as the  
37 baseline risk of this outcome. This estimate is not compatible with the estimates  
38 reported above for new admission to nursing home and mortality but we  
39 recognise that these estimates were generated from studies carried out in  
40 different settings.

41

42 Mortality is defined in the model to be associated with zero cost. The number of  
43 people experiencing “new admission to institution” alone among the number of  
44 people experiencing “mortality or new admission to institution” was estimated by  
45 multiplying the total number of patients that died or were admitted to institute  
46 by 9%. This estimate was taken from the Marcantonio et al study (2000) which  
47 reported that, after one month, only three people died in a sample of 33 people

1 that either died or had new nursing home placement. This was done to obtain an  
2 accurate cost and QALY estimate for this composite outcome.

3

#### 4 **16.2.2.9 Life Expectancy of delirious and non-delirious persons after discharge**

5 The starting age in the model was 79 years. The survival of non-delirious patients  
6 post-discharge was different from that of delirious patients. We took account of  
7 this in the model by using the Kaplan-Meier survival curve reported in the  
8 Rockwood et al study (1999). Of the delirious patients that were followed up for  
9 a median time of 32.5 months, 21% were alive, while 57% of the non-delirious  
10 patients were alive at follow-up. The median survival time was significantly  
11 shorter for those with delirium than for those without. An adjusted hazard ratio of  
12 occurrence of death of 1.71 was reported after adjusting for potential  
13 confounders on the risk of death. We used the data from the survival curve,  
14 fitted an exponential survival function to the data and estimated a baseline  
15 hazard of mortality of 0.007. In the three years after discharge, we applied  
16 these estimates to capture the different survival expectations in the three years  
17 after discharge for patients who have or haven't experienced delirium during  
18 admission. We then applied the same general population mortality rates (Interim  
19 Life Tables for England and Wales, 2005 - 07) to both groups up to age 100.  
20 We estimated a life expectancy of 3.6 years for patients with delirium and 5.4  
21 years for patients without delirium.

22

#### 23 **16.2.2.10 Life expectancies applied in the model for patients in nursing homes and** 24 **patients with new dementia**

##### 25 **Patients staying in nursing home**

26 The data on length of stay in long-term care attributable to delirium was taken  
27 from the results of two large-scale surveys of residential and nursing home  
28 residents in England (Netten et al 2001). They were a longitudinal survey of  
29 eighteen English local authorities and a cross-sectional survey conducted for the  
30 most part in the same authorities as the longitudinal survey. Information about the  
31 circumstances of 2,544 permanent publicly funded admissions from the  
32 authorities to residential and nursing home care was obtained in the longitudinal  
33 survey during a period from mid-October 1995 to mid-January 1996. In the  
34 cross-sectional survey, information about 11,900 residents in the homes was  
35 returned during the autumn of 1996. Cognitive impairment was identified using  
36 items from the Minimum Data Set. This allowed the compilation of the Minimum  
37 Data Set Cognitive Performance Scale. We assumed that the extra time a  
38 delirium patient spends in the long-term care after being transferred from the  
39 hospital will be equivalent to the time a patient with mild cognitive impairment  
40 spends in long-term care. The median length of stay for people with mild  
41 cognitive impairment was 18.9 months and we have assumed in our model that  
42 this is the survival time of patients that stay in long-term care.

1

2 **New dementia**

3 We took data on the life expectancy of a dementia patient from the study on  
4 the costs of dementia in England and Wales in the 21<sup>st</sup> century (McNamee et al  
5 2001).The McNamee et al study (2001) was a Medical Research Council  
6 Cognitive Function and Ageing Study as well as a Resource Implications study. It  
7 provides estimates of formal care cost of dementia based on a population  
8 subgroup identified as cognitively impaired. The diagnosis of dementia was  
9 done using the Geriatric Mental State, and age- and gender-specific prevalence  
10 rates were estimated using data collected in a multi-centre study of four areas  
11 of England and one area in Wales. A sample of 2500 individuals was randomly  
12 selected from Family Health Services Authority or general practice files in the  
13 five centres. This included individuals in long-term hospital care. Life expectancy  
14 with dementia was estimated by applying age- and gender-specific prevalence  
15 rates for dementia to life tables. Cohort specific expectation of life with  
16 dementia was reported for the age groups, 65-69, 70-74, 75-79, 80-84, and  
17 85+ for men and women. The specific life expectancies in years in the respective  
18 age groups for men were 0.7, 0.7, 0.9, 0.9 and 0.8 respectively. It was 1.5, 1.4,  
19 1.8, 1.8 and 1.3 for the respective age groups in women. The population sizes in  
20 these cohorts were reported and we used in the base case analysis a weighted  
21 mean of 1.2 years as the length of time a dementia patient will live. The GDG  
22 suggested that this is rather an underestimate and suggested that the median  
23 estimate in the Dementia UK report (Dementia UK, Full report, 2007; Fitzpatrick  
24 et al 2005) should be used in a sensitivity analysis. The median life expectancies  
25 for individuals with Alzheimer's disease, vascular dementia and mixed dementia  
26 were reported as 7.1, 3.9 and 5.4 years respectively. The estimates were based  
27 on a US cohort study that examined mortality in 3602 participants who were  
28 evaluated for dementia incidence between 1992 and 1999 and followed for  
29 6.5 years. The study was a subset of a larger Cardiovascular Health Study which  
30 recruited participants from Medicare eligibility lists in four US communities.  
31 Participants were to have completed a magnetic resonance imaging and three  
32 Mini-Mental State Exams in order to be eligible for the study. Dementia status  
33 was ascertained using data already collected in the Cardiovascular Health  
34 Study but supplemented with additional data on cognitive measures. The mean  
35 age of those with Alzheimer's disease, vascular dementia and mixed dementia  
36 were 80.1, 78.3 and 79.8 years respectively. We used a life-expectancy of 1.2  
37 years for patients with dementia in the base case which is less than the modelled  
38 life-expectancy for patients without dementia. But in a sensitivity analysis we  
39 assumed that there is no increased risk of mortality due to dementia and  
40 therefore applied the life-expectancy for patients without dementia but taking  
41 into account the effect of delirium on life-expectancy.

42

43 **16.2.3 Relative Risk of the adverse consequences of delirium**

44

45 The relative risk estimate of adverse consequences of delirium was taken from  
46 the review of those consequences in chapter 8 and the estimates we used are  
47 listed in table 16.1 below.

48

1 The risk of new dementia was taken from the study by Rockwood et al (1999).  
2 This was the only study with a moderate quality that was included in the review  
3 for this outcome. It reported an adjusted odds ratio of 5.97 for new dementia  
4 which was assessed over a period of three years. We used relative risk  
5 estimates in the model and converted the reported odds ratio to a relative risk  
6 estimate using the formula,

7

8 
$$RR = (OR) / [(1 - Po) + (Po \times OR)]$$
 (Zhang & Kai 1998)

9

10 where RR is relative risk; OR, the odds ratio; and Po, the incidence rate in the  
11 unexposed population. The annual incidence of dementia among people without  
12 cognitive impairment at baseline was reported as 5.6% per year. We estimated  
13 a relative risk of 4.67 which we used in our economic model.

14

15 We used a similar method to estimate the relative risk of 2.05 for new admission  
16 to institution using an adjusted odds ratio of 2.64 (Bourdel-Marchasson et al  
17 2004). There was a range of studies that reported the risk for this outcome but  
18 the odds ratio of 2.64 was chosen as it used incident delirium to estimate new  
19 admission to long-term care at the point of discharge.

20

21 The risk of falls and pressure ulcer was available from only one study (O’Keeffe  
22 and Lavan, 1997). The study reported an adjusted odd ratio of 2.3 for  
23 developing hospital-acquired complications which included falls and pressure  
24 ulcer. The relative risk of 2.18 for falls and pressure ulcer was estimated using  
25 the combined rate in the non-delirious group for falls and pressure ulcer.

26

27 The adjusted odds ratio of 2.6 for mortality in delirium patients in the hospital  
28 was taken from the O’Keeffe and Lavan study (1997). We estimated a relative  
29 risk of 2.41 which we used in our model. There were other studies that reported  
30 the risk of in-hospital mortality but the GDG advised that it is best to use a UK  
31 study for this outcome. The way we have treated post-discharge mortality has  
32 already been described above.

33

34 Delirium extends hospital length of stay and the additional length of stay used in  
35 the model was estimated from a Kaplan-Meier plot reported in the Holmes and  
36 House study (2000). This study was chosen because it was a UK study and was  
37 judged as being a high quality study for this outcome. We fitted a Weibull  
38 function using a lambda of 0.08 and gamma of 0.87 that were estimated from  
39 the Kaplan-Meier plot on the proportion of people in hospital at different times  
40 of discharge. This was for the people that were reported to be without a  
41 psychiatric diagnosis. The study also reported the result of a Cox Proportional  
42 Hazards model which showed that delirium is associated with a hazard ratio of  
43 0.53 for hospital discharge. We applied this adjusted estimate to fit a Weibull

1 function for the delirious group and estimated the difference in the area  
2 between the two fitted functions. This difference was 16.83 days and was  
3 treated in the model as the additional hospital length of stay due to delirium.

4  
5 The adjusted odds ratio for the composite outcome of "mortality or new nursing  
6 home placement" after one month was reported as 3.0 (Marcantonio et al  
7 2000). We converted this to a relative risk estimate of 2.41 which was used in a  
8 sensitivity analysis in the economic model.

9  
10  
11  
12  
13  
14 Table 16.1: the baseline and relative risks of the adverse consequences of  
15 delirium

16

Adverse consequences	Baseline risk	Source	Odds ratio (95% CI)	Estimated relative risk (95% CI)	Source
New dementia	5.6%	Rockwood et al 1999	5.97 (1.83, 19.54)	4.67 (1.43, 15.29)	Rockwood et al 1999
New admission to institution	17.4%	Bourdel-Marchasson et al 2004	2.64 (0.83, 8.45)	2.05 (0.65, 6.57)	Bourdel-Marchasson et al 2004
Pressure ulcer	4.0%	Clark & Watts 1994	2.30 (1.7, 5.0)	2.18 (1.61, 4.73)	O'Keefe & Lavan 1997
Falls	6.9%	O'Keefe & Lavan 1997			
Mortality	5.0%	O'Keefe & Lavan 1997	2.60 (0.7, 6.2)	2.41 (0.65, 5.74)	O'Keefe & Lavan 1997
Mortality or new admission to institution	12.2%	Marcantonio et al 2000	3.00 (1.1, 8.4)	2.41 (0.88, 6.76)	Marcantonio et al 2000

17  
18  
19 **16.2.4 Efficacy of Interventions**

20 The efficacy of the different intervention strategies has been reported in the  
21 review of multi-component prevention interventions (section 9.15). It was  
22 reported that the use of these interventions by older general medical patients,  
23 who were at intermediate or high risk of delirium, was associated with a relative  
24 risk of delirium of 0.66 (Inouye et al 1999). The use of these interventions in  
25 older patients that underwent hip fracture surgery was reported to result in a  
26 relative risk of delirium of 0.65 (Marcantonio et al 2000). We have applied  
27 these estimates in our economic model.

1

2 **16.2.5 Cost of Adverse Consequences of Delirium**

3

4 **16.2.5.1 Falls (cost)**

5 The cost of falls data came from a randomised, controlled study of the  
6 prevention of fractures in the UK primary care. (Iglesias et al 2008). Eligible  
7 study participants were women aged 70 years and above with one or more risk  
8 factors for hip fracture and a total of 3,314 women were recruited into the  
9 study. The intervention group received daily oral supplementation using 1000mg  
10 calcium with 800 IU cholecalciferol and information leaflet on dietary calcium  
11 intake and prevention of falls (Porthouse et al 2005). The control group received  
12 leaflet only. Data on fracture and fall incidence, in addition to data on HRQoL  
13 and fear of falling, were collected at baseline and every 6 months after that for  
14 a minimum of 2 years and maximum of 42 months.

15

16 A fall and fracture questionnaire was used for resource use data collection and  
17 was administered to 1190 women participating in the prevention study and who  
18 had previously indicated to be willing to be contacted in the future for research  
19 purposes. Participants were asked if they had experienced a fall and / or  
20 fracture in the last 12 months, the number of times they had seen a doctor, GP or  
21 consultant and whether they had been hospitalised for reasons other than a fall  
22 or fracture and for how long, in the same period. Those that had experienced a  
23 fall or a fracture were further asked whether they had been hospitalised and  
24 how long they spent in hospital, the number of times they had seen a doctor or  
25 nurse, whether they had changed residence because of their fall and / or  
26 fracture and for how long. They were asked to describe any treatments that  
27 were specifically prescribed for their fall or fracture over the same period.  
28 Resource use was valued using unit costs from NHS reference cost data, Personal  
29 and Social Services Research Unit (PSSRU) data, as well as the Chartered  
30 Institute of Public Finance and Accountancy (CIPFA) data base. The NHS  
31 reference cost data was used to cost hospital inpatient length of stay as well as  
32 the cost of surgery following hip, wrist, arm and vertebral fractures. The CIPFA  
33 database was used to cost specialist contact visits, and the PSSRU data was used  
34 to cost GP and nurse visits, residential accommodation and the cost of home help.

35

36 The response rate to the questionnaire was 93% and 302 out of 1110  
37 respondents reported falls in the previous 12 months and 62 of those that fell  
38 reported that their fall resulted in a fracture. Falls that did not result in fractures  
39 were generally associated with less resource use. There were 243 falls events  
40 that did not result in fractures and the mean cost was reported as £1,088. The  
41 number of falls that led to fractures was 10 for hip fracture, 7 for wrist fracture,  
42 10 for arm fracture and 2 for vertebral fracture. The cost of falls leading to a  
43 fracture was reported as £15,133; £2,753; £1,863; £1,331; and £3,498 for

1 hip, wrist, arm, vertebral, and other fractures respectively. We used a weighted  
2 estimate of £1875 in our economic model

3

#### 4 **16.2.5.2 Pressure Ulcer (cost)**

5 The cost of pressure ulcer used in our model was taken from a cost study that  
6 aimed to estimate the annual cost of treating pressure ulcers in the UK (Bennett et  
7 al 2004). Treatment protocols which reflect good clinical practice for treating  
8 pressure ulcers of different grades were developed and costs for the daily  
9 resources defined in the protocol were assigned using representative UK NHS  
10 unit costs at 2000 prices. It was assumed that care is provided in a hospital or  
11 long-term care setting and that pressure ulcer patients are not admitted solely  
12 for the care of pressure ulcer. Resources to be used for care include nurse time,  
13 dressings, antibiotics, diagnostic tests, support surfaces and inpatient days where  
14 appropriate. Pressure ulcer was classified in four grades with grade 1 as the  
15 least severe and grade 4, the most severe. The daily costs for the ulcer grades  
16 were estimated for patients whose ulcer would heal normally as well as for  
17 patients whose ulcers were associated with critical colonisation, cellulitis and  
18 osteomyelitis. We assumed that pressure ulcers resulting from delirium are grade  
19 1 pressure ulcers, would heal normally and are not associated with further  
20 complications. This assumption is conservative and is based on the finding that  
21 more complicated pressure ulcers are less common and represent less than 5% of  
22 all cases (Clark 1994). The cost per day for a grade 1 ulcer that heals normally  
23 is £38 and it will take 4.06 weeks on average for this class of ulcer to heal. The  
24 mean time to heal was taken from the same Bennett et al study (2004) and this  
25 estimate was reported to have come from a review of clinical literature. We  
26 therefore used a cost estimate of £1,064, up rated it to a 2007 estimate of  
27 £1364 (£1228.09 to £1499.86) using the inflation indices reported in PSSRU.  
28 The up rated estimate was applied in the model. The GDG suggested that some  
29 of the pressure ulcer cases due to delirium will be grade 4 pressure ulcers that  
30 will heal normally. They advised that the impact of this on the cost-effectiveness  
31 estimates should be investigated. We carried out a deterministic sensitivity  
32 analysis using the cost of grade 4 ulcer that heals normally. This was equivalent  
33 to a 2007 estimate of £9934.99.

34

#### 35 **16.2.5.3 Stay in long-term care (cost)**

36 The cost of long-term care used in the model was estimated from the unit cost of  
37 stay in private nursing homes, private residential care, voluntary residential care  
38 and local authority residential care facility for older people. The care package  
39 costs per permanent residential week in private nursing homes were reported as  
40 £687 (PSSRU 2007). In private, voluntary and local authority residential care  
41 these were reported as £483, £480 and £858 respectively.

42 These unit costs have been estimated to include cost for external services such as  
43 community nursing, GP services as well as personal living expenses. They also  
44 include capital costs for the local authority residential care, and fees for the  
45 private and voluntary residential care. We subtracted £9.20, the cost of  
46 personal living expenses per week, from each unit cost and estimated £655.66,

1 the weighted average of £677.80, £473.80, £470.80 and £848.80, to be the  
2 unit cost of long-term care. The weighting was based on the distribution of  
3 residents, 65 years and older, in care homes in 1996. It was reported that in  
4 nursing homes, local authority, private and voluntary residential homes the  
5 number of residents were 5746, 5476, 2791 and 3664 respectively (Netten et  
6 al 1998).

7  
8 The NHS does not pay towards long-term care for all patients. It was suggested  
9 that only two percent of residents were funded by the NHS and overall, about  
10 70% of the care home population were publicly funded (Netten et al, 1998).  
11 We will consider the effect of this on the cost-effectiveness result by assuming in  
12 a sensitivity analysis that only 70% of the costs of long-term care will be borne  
13 by the NHS and PSS. The length of time a patient spends in the long-term care  
14 has been assumed to be 18.9 months and the source of this estimate is described  
15 above.

#### 16 17 **16.2.5.4 Hospital stay (Unit Cost)**

18 We have used the unit cost estimates per excess day associated with complex  
19 elderly patients. This was reported as unit cost per day for days exceeding the  
20 trim point. We took all the HRG unit costs reported for all Complex Elderly  
21 patients (Hospital Episode Statistics for England. Inpatient statistics, 2007 – 08)  
22 and found a weighted mean of £152. There will be no additional costs on the  
23 basis of inpatient rehabilitation services as the GDG advised that, if at all, only  
24 a small number of delirium patients will need such services.

#### 25 26 27 **16.2.5.5 New Dementia (Cost)**

28 Our cost estimate for dementia was taken from a report of the prevalence and  
29 cost of dementia prepared by the PSSRU and the Institute of Psychiatry  
30 (Dementia UK, The full report, 2007). The cost estimate was based on an  
31 interview of 132 dementia patients and dementia carers, who were referred to  
32 psychiatric services between January 1997 and June 1999. Service use was  
33 measured with a version of the Client Service Receipt Inventory and study  
34 participants were asked for details of accommodation and services during the  
35 past three months. Medication, inpatient and outpatient care, day hospitals, day  
36 centres, community health services, social care and respite care were the services  
37 included in the costing framework. Resource use for the services was valued using  
38 unit cost and estimated costs were inflated to reflect 2005/6 price levels. Cost  
39 of accommodation was based on a weighted average of unit costs for supported  
40 accommodation. Costs were based on only 114 definite cases of dementia, the  
41 study sample was London-based and an adjustment was made to reflect the UK  
42 as a whole. The cost of informal care was also included but we have excluded  
43 such costs here as the cost of informal care is outside the remit of NICE. The

1 annual cost of late onset dementia per person was reported to be £25,472. Of  
2 this, accommodation accounted for 41%, NHS care services 8%, social care  
3 services 15%, and informal care services 36%. We subtracted the cost of  
4 informal care services and arrived at a cost estimate of £16,302 which was used  
5 as the annual cost of new dementia in our economic model. In a sensitivity  
6 analysis, we assumed that the cost of accommodation has been accounted for in  
7 the model, and have also subtracted the cost of accommodation. We estimated  
8 the cost of dementia to include only the cost of NHS services and social care  
9 services and arrived at a cost of £5,859. In the base case analysis, we have  
10 assumed that the life expectancy of a delirium patient is 1.2 years, and we have  
11 increased this in sensitivity analysis. The sources of the life expectancy estimates  
12 are described above.

13

14

#### 15 **16.2.5.6 Mortality (Cost)**

16 We have not accounted for any additional cost resulting from mortality in our  
17 model. We have assumed that the cost associated with mortality has been  
18 incurred in the period up to the point of death, and that this has been captured  
19 in the model in the cost of adverse consequences that would eventually lead to  
20 death.

21

22

### 23 **16.2.6 Utility of Adverse Consequences of Delirium**

24

#### 25 **16.2.6.1 Falls (Utility)**

26 The utility estimate for falls used in the economic model was taken from a Dutch  
27 randomised controlled trial (Hendriks et al, 2008). It was an economic evaluation  
28 that aimed to assess whether a multidisciplinary intervention program would be  
29 preferable to usual care in the Netherlands. The study participants were those  
30 65 years of age or over, and who had visited the accident and emergency  
31 department or general practice cooperative for the consequences of a fall. The  
32 exclusion criteria were inability to speak or understand Dutch, inability to  
33 complete questionnaires or interviews by telephone, cognitive impairment,  
34 admission for more than 4 weeks to a hospital or other institution, being  
35 permanently wheelchair-dependent or bedridden. Follow-up time was 12 months  
36 after baseline. The intervention included medical and occupational-therapy  
37 assessment that aimed to assess and address potential risk factors for fall. In  
38 usual care, medical risks and other risk factors were not systematically recorded  
39 and addressed by hospital physicians, specialists or GPs. Participants responded  
40 to the standard Dutch version of the EQ-5D in self-administered questionnaires at  
41 baseline and after 4 and 12 months. Utility scores for the EQ-5D responses were  
42 estimated using UK based social tariff. The mean age of the 167 participants in  
43 the usual care arm of the trial was 75.2 years. The mean utility at 4 and 12

1 months was reported as 0.72 and 0.71 respectively. The QALYs at the end of  
2 the follow-up was reported as 0.71.

3  
4 In order to estimate the expected lifetime QALY gains for patients who  
5 experience falls we applied a utility multiplier in the first year of a falls'  
6 patient's life. The utility multiplier was estimated as the ratio of the utility of 0.71  
7 reported at the end of the study follow-up and 0.74, the utility of a person  
8 aged 75.2 years old in the UK population. The utility of the population varies by  
9 age and the population utility was derived from an algorithm that was produced  
10 after a re-analysis of data from Kind et al 1998 in Ward et al 2007. In the  
11 model, the starting age is 79 years and the utility multiplier, 0.96 was used to  
12 adjust 0.72, the utility of an average British person aged 79. The QALY gains  
13 for the rest of the patient's life expectancy were estimated from a Markov  
14 survival model from the Life Table. In our estimates, we took account of the three  
15 year differences in survival chances of delirious and non-delirious patients (see  
16 section on mortality after hospital discharge). .

#### 17 18 **16.2.6.2 Pressure Ulcer (Utility)**

19 We did not identify any useful utility data on the HRQoL impact of pressure  
20 ulcer. The life-time expected QALY gain for a person who has experienced a  
21 pressure ulcer was assumed to be equal to the QALY gain of a person without  
22 any adverse consequence of delirium. This was estimated from a Markov survival  
23 analysis from the Life Table and we accounted for the three year differences in  
24 the survival chances of delirious and non-delirious patients (see section on  
25 mortality after hospital discharge). We estimated the expected lifetime QALY  
26 gain of a delirious person as 2.13 and the expected lifetime QALY gain of a  
27 non-delirious person as 3.09.

#### 28 29 **16.2.6.3 Long-term care (utility)**

30 We could not identify a useful study that measured the utility of patients in long-  
31 term care. The GDG advised that the utility of a delirium in long-term care  
32 should be assumed to be equivalent to 0.25, the utility of a patient with severe  
33 dementia (Ekman et al, 2007). The Ekman et al (2007) study aimed to obtain  
34 primary data on community-based health utilities in different stages of mild  
35 cognitive impairment and dementia from a general population sample. It was a  
36 cross-sectional study of subjects aged 45 – 84 years who were randomly  
37 selected in Sweden. A questionnaire was sent to a sample of 1,800 subjects and  
38 a description of the health conditions and how to value them was given. Four  
39 vignettes describing health conditions involving cognitive impairments typical for  
40 the progressive stages of dementia were made using the Clinical Dementia  
41 Rating scale. Mild cognitive impairment was defined as an overall Clinical  
42 Dementia Rating score of 0.5. Valuation of the perceived quality of life in these  
43 stages was carried out using the time trade-off techniques. Respondents were  
44 reported as fairly representative of the general population in terms of age,

1 gender, and employment. The mean age of women and men were 66.4 and  
2 67.1 years respectively and 54.4% of the study sample was women. The mean  
3 utility score for severe dementia was reported as 0.25. This was used as a utility  
4 multiplier in the model. The mean age in the model is 79 years and the utility  
5 multiplier, 0.25 was multiplied with 0.72, the utility of an average British person  
6 aged 79. The adjusted utility of 0.18 was used to estimate the expected lifetime  
7 QALY gains after admission to long-term care.

8

#### 9 **16.2.6.4 Hospital stay (Utility)**

10 We would expect some utility changes for staying in the hospital but the  
11 associated QALY gain will be small because of the short length of stay in  
12 hospital. We have therefore not included the impact of utility changes resulting  
13 from hospital care in our economic model.

14

#### 15 **16.2.6.5 New Dementia (Utility)**

16 The utility score for new dementia was taken from the report by Ekman et al,  
17 2007. This study has been described above in the section on the utility of  
18 patients in long-term care. The mean utility score for mild, moderate and severe  
19 dementia were reported as 0.62, 0.40 and 0.25 respectively. The GDG advised  
20 that we use the utility score reported for moderate dementia. We applied this as  
21 a utility multiplier in the model and estimated a utility of 0.28 which was used to  
22 estimate the expected lifetime QALY gains for this outcome. The life expectancy  
23 used in the base case was 1.2 years and in the sensitivity analysis we used 3.6  
24 years for dementia patients who experienced delirium and 5.4 years for those  
25 who did not experience delirium.

26

#### 27 **16.2.6.6 Mortality (Utility)**

28 We have used zero QALY gain in the event of mortality.

29

30

### 31 **16.2.7 Cost of multi-component Targeted Intervention**

32

#### 33 **16.2.7.1 The use of multi-component targeted intervention in older patients admitted non-** 34 **electively for surgical repair of hip fracture**

35

36 The costing of multi-component targeted intervention in patients admitted for  
37 surgical repair of hip fracture is based on the intervention protocol of a  
38 randomised controlled trial in an orthopaedic surgery service (Marcantonio et al,

1 2001). The trial has been described in the section on the use of multi-component  
2 interventions for delirium prevention (section 9.15). The trial aimed to determine  
3 whether proactive geriatrics consultations can reduce delirium after hip fracture  
4 repair. It was carried out in US patients, 65 years or older, who were admitted  
5 non-electively for surgical repair of hip fracture. All study patients had an intake  
6 assessment that included a patient interview, a proxy interview, and a review of  
7 the medical record. Patients in the intervention group received proactive  
8 geriatric consultation, which began preoperatively or within 24 hours of surgery.  
9 They received targeted recommendations based on a structured protocol from  
10 the geriatrician during the period of hospitalization. Patients in the control group  
11 received usual care. They received management by the orthopaedics team,  
12 including internal medicine consultants or geriatricians on a reactive rather than  
13 proactive basis.

14  
15 The structured protocol used for the recommendations included 10 modules with  
16 each containing two to five specific recommendations (Appendix J).  
17 Recommendations were prioritized and limited to no more than five after the  
18 initial visit by the geriatrician and no more than three after follow-up visits. This  
19 was done to improve adherence. The GDG suggested that the geriatrician and  
20 other NHS personnel would be needed to apply this intervention on patients. It  
21 was suggested that modules one to four, eight, and 10 would be delivered by  
22 doctors. This will require additional 15 minutes of geriatrician's time per patient  
23 per week. The duration of application of this intervention was taken to be  
24 equivalent to the median length of stay of patients with fracture of neck of femur  
25 which was reported as 16 days (HES Online, 2007 – 2008). It will therefore cost  
26 an additional £100 to apply the four modules. The application of modules five  
27 to seven, and module nine were assumed to be part of the routine work for  
28 nurses on pay Band 5. However, additional work and NHS resources would be  
29 expected for applying module 6a and 7b. The additional time for applying  
30 module 6a was suggested to be ten minutes thrice daily per patient while  
31 module 7b would require ten minutes four times daily per patient. The hourly cost  
32 of a nurse pay Band 5, including cost of qualification, is £22 [PSSRU 2007]. The  
33 application of module 6a would cost £11 per patient daily and module 7a  
34 would cost £15 per patient daily. This is equivalent to £176 and £240  
35 respectively over 16 days. The total cost of applying multi-component targeted  
36 intervention to older patients admitted non-electively for surgical repair of hip  
37 fracture would therefore amount to £516.

38  
39  
40 **16.2.7.2 The use of multi-component targeted intervention in consecutive older patients at**  
41 **intermediate or high risk of delirium who were admitted to the general**  
42 **medicine service**

43  
44 The cost estimate for using multi-component targeted intervention in older  
45 patients at intermediate or high risk of delirium who were admitted to the

1 general medicine service was based on a trial of patients aged 70 years or  
2 older who were consecutively admitted to the general medicine service of a  
3 hospital (Inouye et al 1999). This trial has been described in the section on the  
4 use of multi-component interventions for delirium prevention (section 9.15). At the  
5 point of admission, the patients in the trial showed no evidence of patients  
6 having delirium, but they were assessed to be at immediate or high risk of  
7 developing delirium. The study sample was 852 people, including 426 matched  
8 pairs of intervention and control, enrolled in the clinical trial in a hospital  
9 between March, 1995 and March 1998. The trial had three aims namely, to  
10 compare the effectiveness of a multi-component strategy for reducing the risk of  
11 delirium with that of a usual plan of care for hospitalized older patients, to  
12 determine the level of adherence to the intervention protocol, and to measure  
13 the effect of the intervention on the targeted risk factors. Eligible study patients  
14 underwent screening and base line assessments which were completed within 48  
15 hours after admission. Patients in the intervention group received standard  
16 protocols for the management of six risk factors for delirium namely, cognitive  
17 impairment, sleep deprivation, immobility, visual impairment, hearing impairment,  
18 and dehydration (Appendix J). Geriatric nursing assessment and interdisciplinary  
19 rounds were other program interventions targeted towards the risk factors. The  
20 intervention, the Hospital Elder Life Program, was implemented by a trained  
21 team, which consisted of a geriatric nurse-specialist, two specially trained Elder  
22 Life specialists, a certified therapeutic-recreation specialist, a physical-therapy  
23 consultant, a geriatrician, and trained volunteers. Patients in the usual care group  
24 received standard hospital services provided by physicians, nurses, and support  
25 staff. The study reported the total cost of intervention to be \$139,506. The  
26 number of people in the intervention group was 426 and the average cost of  
27 intervention was reported as \$327 per patient. This included staff time spent in  
28 intervention activities, equipment, supplies and consultant costs.

29  
30 It was recommended that the staff required to implement the Hospital Elder Life  
31 Program in 200 to 250 patients per year are one full-time Elder Life Specialist  
32 who also serves as Volunteer Coordinator, one half-time Geriatric Nurse  
33 Specialist, and 0.10 to 0.20 of a full time equivalent geriatrician, who also acts  
34 as a Program Director (Inouye, 2000). We used this time equivalence in our cost  
35 estimation. A description of the duties of each staff is given in Appendix J.  
36 Volunteers play a critical role in the implementation of the program and the  
37 tasks of a volunteer would be carried out by NHS personnel. It was suggested  
38 that a minimum of 21 Volunteers would be required to operate a program of  
39 200 to 250 patients. Each was to serve one shift per week and 3 to 4 hours per  
40 shift. The GDG advised that the pay band for the geriatric nurse specialist would  
41 be Band 6; Elder Life specialist would be Band 5; Geriatrician would be the  
42 annual salary equivalent of an NHS Medical Consultant and the Volunteer would  
43 be Band 2. We applied the Agenda for Change salaries and used the April  
44 2006 scale mid-point. These were used to estimate the unit cost for the Elder Life  
45 Program Staff. We estimated that the personnel cost per patient would be  
46 £370. We assumed that each of the 21 volunteers would work four hours per  
47 week, geriatricians would work 0.15 Full Time Equivalence and the number of  
48 patients that received intervention would be 225 patients.

49

1 Equipment such as computers, telephone and photocopying machines that would  
 2 be needed to implement the program are assumed to be available and would  
 3 not need to be purchased additionally by the NHS. Some of the materials  
 4 needed for implementing the intervention protocol described in the study by  
 5 Inouye et al (1999) are already available to the NHS patient and are used  
 6 during usual care. The additional materials that would need to be purchased are  
 7 listed in Appendix J. They include standard word games and relaxation tapes or  
 8 music. We have assumed that cost of providing instructions by the intervention  
 9 staff will be accounted for through the salary paid to them by the NHS. We  
 10 have not added any additional cost of providing instructions.

11  
 12 We could not find cost data on what the NHS pays for a standard word game  
 13 or relaxation tapes. We have assumed the cost to be £50 each and life  
 14 expectancies of the materials to be 0.5 and 1 year respectively. We have also  
 15 assumed that 10 pieces of relaxation tapes will be required for a multi-  
 16 component targeted intervention program for 225 patients over a year. We  
 17 assumed that 20 pieces of standard word game will be required for the same  
 18 number of patients over the same time period. The additional cost of the  
 19 materials was estimated at £7 per patient.

20  
 21 We have estimated the cost of using multi-component targeted intervention in  
 22 older patients at intermediate or high risk of delirium who were admitted to the  
 23 general medicine service in the NHS as £377. This does not include additional  
 24 training cost as we have assumed that this has already been included as part of  
 25 the time resources required by the Program staff to implement the program. We  
 26 also did not include the cost associated with screening and base line assessment  
 27 at the beginning of the intervention for the same reason. In a sensitivity analysis,  
 28 we assumed that the Geriatric nurse specialist will be on band 7 and the Elder  
 29 Life Specialist, on band 6. This increased the total cost of personnel to £404. This  
 30 was to account for possible additional work load for these two roles.

31  
 32  
 33 A summary of the data inputs used in the model is given below. The baseline and  
 34 relative risk estimates of the adverse consequences have been given above in  
 35 table 16.2.

36  
 37  
 38 Table 16.2: other inputs used in base case analysis in the economic model

Model input	Point Estimate (95% CI)	Source
Baseline risk		
Delirium in hospital (general medicine services)	15.0%	Inouye et al 1999

Delirium in hospital (hip fracture surgery)	50.0%	Marcantonio et al 2000
Unit cost		
New dementia (per year)	£16,302	Dementia UK, The full report, 2007
Stay in long-term care (per week)	£656	PSSRU 2007, Netten et al 1998
Pressure ulcer	£1,364 (£1,228 to £1,500)*	Bennett et al 2004
Falls	£1,875	Iglesias et al 2008
Utility		
New dementia	0.29	Ekman et al, 2007 (reported 0.4 for moderate dementia)
New admission to institution	0.18	Ekman et al, 2007 (reported 0.25 for moderate dementia, GDG suggested it should be used to estimate utility for this outcome)
Falls	0.69	Hendriks et al, 2008 (reported 0.71 after 12 months)
Duration		
Stay in long-term care (months)	18.9	Netten et al 2001
Extended hospital stay (days)	16.83 (9.36, 25.34)	Holmes & House 2000
Life with dementia (years)	1.2	McNamee et al 2001
Intervention Efficacy		
MTI (general medical services)	0.66 (0.46, 0.95)	Inouye et al 1999
MTI (hip fracture surgery)	0.65 (0.42, 1)	Marcantonio et al 2000
Intervention Cost		
MTI (medical services)	£377	Based on study protocol in Inouye et al 1999
MTI (hip fracture surgery)	£511	Based on study protocol in Marcantonio et al 2000

1 \*Reported as mean (+ and – 10%)

2

3

#### 4 **16.2.8 Sensitivity Analyses**

5

##### 6 **16.2.8.1 Deterministic Sensitivity Analyses**

7 In the deterministic analysis we estimated the point estimate for cost, QALYs  
8 gained, ICER and INMB using the base case model structure and point estimates  
9 for model input parameters. We have carried out a series of deterministic  
10 sensitivity analyses (DSA) to explore the uncertainties that relate to the base  
11 case structure.

1

2 The first approach we have taken is to assume that not all the adverse  
3 consequences are important to the model structure. We assumed that each and  
4 only one of the six adverse consequences was the only adverse outcome  
5 associated with delirium. We estimated the INMB after assuming that new  
6 admission to nursing homes was the only adverse outcome to be associated with  
7 delirium. The same was done for mortality, new dementia, falls, pressure ulcer  
8 and extended hospital stay. In another DSA, we included nursing home admission  
9 and mortality as a composite outcome and did not include them as single model  
10 inputs. We explored the cost-effectiveness of interventions in low risk patients  
11 and used 12.5% as the baseline risk of delirium. This was the lower estimate of  
12 the range of delirium incidence reported in the needs assessment review (chapter  
13 5) for general medical patients. We explored the effect of using this lower  
14 estimate for both populations considered by the model (elderly patients at risk  
15 of delirium who were admitted to the general medicine service and patients  
16 undergoing surgical repair of hip fracture).

17

18 In the base case analysis, we have assumed that the life expectancy of delirious  
19 patients to be shorter than that of non-delirious patients. This was due to  
20 difference in post-hospital chances of survival for the two groups. In a DSA we  
21 have assumed that the survival chances for delirious patients are equivalent to  
22 those of non-delirious patients. In another DSA we have assumed the life  
23 expectancy of dementia patients to be 3.6 years and 5.4 years for patients with  
24 and without previous delirium experience respectively.. In the base case, we used  
25 1.2 years regardless of the previous delirium experience. We have assumed in  
26 the base case that patients in long-term care will survive for only 18.9 months. In  
27 a sensitivity analysis, we estimated lifetime QALY gains over a life expectancy  
28 of 3.6 years for those with delirium and 5.4 years for those without delirium.

29

30 The annual cost of dementia was reduced to £5,859. This was to remove  
31 potential double counting of the cost of stay in long-term care as a proportion of  
32 the cost of dementia in the base case was due to stay in long-term care. In  
33 another DSA, we included only 70% of the cost of stay in long-term care, as we  
34 assumed that 100% of this cost will not be funded by the public. Further analyses  
35 were done to explore the impact on the model results of increased cost of  
36 pressure ulcer resulting from grade 4 ulcer that heal normally, and increased  
37 cost of the multi-component targeted interventions resulting from higher pay  
38 Band to the Geriatric Nurse Specialist and Elder Life Specialist.

39

40

#### 41 **16.2.8.2 Probabilistic Sensitivity Analyses**

42

43 In the DSA we used point estimates for the model input parameters. However,  
44 point estimates are subject to uncertainties. We have carried out a probability  
45 sensitivity analysis, PSA, to reflect the uncertainty in the input parameters of the  
model. The results of the PSA show the uncertainty in the primary outcomes of the

1 model that results from the uncertainty in the model inputs. Each of the input  
 2 parameters is assigned a probability distribution which reflects the standard  
 3 error of each parameter estimate.

4  
 5 We randomly selected from each parameter distribution in a simultaneous  
 6 manner and calculated the cost, QALYs, ICERs and INMB. This was repeated  
 7 5000 times to produce 5000 estimates that reflect the uncertainties in the input  
 8 parameters. An average of the estimates was found and the most cost-effective  
 9 strategy is the one with the highest mean INMB. However, the one with the  
 10 highest mean INMB may or may not be the most cost-effective in all the  
 11 simulations. The model parameters, the type of distribution and distribution  
 12 parameters are listed in the table below (table xxx). The model input  
 13 parameters that we did not vary probabilistically are life expectancy of a  
 14 patient with dementia, survival length of time in long-term care, post-discharge  
 15 mortality differences for delirious and non-delirious patients, and the discount  
 16 rate.

17  
 18 Table 16.3: input parameters, type of distribution and distribution parameters  
 19 used in PSA

Parameter	Type of distribution	Point estimate	Distribution parameters	Source
<b>Baseline Risk</b>				
Delirium in Hospital (general medical services)	Beta	15.0%	$\alpha = 64, \beta = 362$	Inouye et al 1999
Delirium in Hospital (hip fracture surgery)	Beta	50.0%	$\alpha = 32, \beta = 32$	Marcantonio et al 2000
Falls	Beta	6.9%	$\alpha = 9, \beta = 122$	O'Keeffe & Lavan 1997
Pressure Ulcer	Beta	4.0%	$\alpha = 360, \beta = 8575$	Clark & Watts 1994
Dementia	Beta	5.6%	$\alpha = 7, \beta = 117$	Rockwood et al 1999
New admission to institution	Beta	17.4%	$\alpha = 40, \beta = 190$	Bourdel-Marchasson et al 2004
In hospital Mortality	Beta	5.0%	$\alpha = 7, \beta = 124$	O'Keeffe & Lavan 1997
Mortality or new admission to institution	Beta	12.2%	$\alpha = 9, \beta = 65$	Marcantonio et al 2000
<b>Post-discharge survival</b>				
Difference in mortality between delirious and non-delirious patients	Lognormal	HR = 1.71	Log (mean) = 0.54, se = 0.26	Rockwood et al 1999

Relative Risk				
Falls and pressure ulcer	Lognormal	RR = 2.18	Log (mean) = 0.78, se = 0.27	O'Keeffe & Lavan 1997
Dementia	Lognormal	RR = 4.67	Log (mean) = 1.54, se = 0.60	Rockwood et al 1999
New admission to institution	Lognormal	RR = 2.05	Log (mean) = 0.72, se = 0.59	Bourdel-Marchasson et al 2004
Mortality	Lognormal	RR = 2.41	Log (mean) = 0.88, se = 0.56	O'Keeffe & Lavan 1997
Mortality or new admission to institution	Lognormal	RR = 2.41	Log (mean) = 0.88, se = 0.52	Marcantonio et al 2000
Cost				
Falls	Gamma	£1,875	Mean = £1,875, se = £239	Iglesias et al 2008
Pressure Ulcer	Gamma	£1,364	Mean = £1,364, se = £69	Bennett et al 2004
Dementia	Gamma	£16,302	Mean = £16,302, se = £2079	Dementia UK, The Full Report, 2007
Extended hospital stay	Gamma	£152	Mean = £152, se = £19	HES England, 2007-08
Stay in long-term care	Gamma	£656	Mean = £656, se = £84	PSSRU 2007
MTI (general medical)	Gamma	£377	Mean = £377, se = £48	Based on recommended protocol and GDG advice
MTI (hip fracture surgery)	Gamma	£511	Mean = £511, se = £65	Based on recommended protocol and GDG advice
Utility				
Falls	Beta	0.71	$\alpha = 249, \beta = 102$	Hendriks et al 2008
Dementia	Beta	0.40	$\alpha = 730, \beta = 1094$	Kman et al 2007
Stay in institution	Beta	0.25	$\alpha = 293, \beta = 880$	Ekman et al 2007
Population utility	Multinomial	Linear relationship	Age-Utility intercept: 1.06;	Based on a re-analysis of data

		with age	Age-Utility gradient: -0.00	from Kind et al 1998 in Ward et al 2007
Duration				
Extended hospital stay	Gamma	16.83	Mean = 16.83, se = 4.08	Holmes and House 2000
Efficacy of MTI intervention				
Relative risk (general medicine services)	Lognormal	0.66	Log (mean) = -0.42, se = 0.19	Inouye et al 1999
Relative risk (hip fracture surgery)	Lognormal	0.65	Log (mean) = -0.43, se = 0.22	Marcantonio et al 2000

1  
2  
3

## 4 16.3 Results

5

### 6 16.3.1 Cost-effectiveness of multi-component targeted prevention 7 interventions in older patients at intermediate or high risk of delirium 8 who were admitted to the general medicine service

9

10 The table below (table xxx) shows the cost-effectiveness model results for the use  
11 of multi-component prevention interventions in patients at immediate or high risk  
12 of delirium and who were admitted to the general medicine service. The result of  
13 the deterministic analysis suggests that this intervention is cost-effective when  
14 compared to usual care and is associated with an INMB of £2,130.

15

16 The result of the PSA suggests that the usual care strategy will cost £13,200 on  
17 average whereas the prevention strategy will cost £12,690. This is the mean  
18 total cost that includes the cost of the adverse consequences and the unit cost of  
19 the intervention itself. The QALY gains associated with both strategies are 2.140  
20 and 2.220 QALYs respectively. The prevention strategy was therefore the  
21 dominant strategy because it reduced cost and increased QALY gains when  
22 compared to the usual care strategy. It was associated with an ICER of -£6,190  
23 per QALY and an INMB of £2,200.

24

25

1 Table 16.4: costs, QALYs and cost-effectiveness of multi-component targeted  
 2 intervention compared to usual care\*

		Usual Care	MTI
Probabilistic	Mean cost	£13,200	£12,690
	Mean QALYs	2.140	2.220
	Incr Cost	N/A	£-520
	Incr QALYs		0.084
	Incr Cost / QALY		£-6,190
	Incr NMB		£2,200
	% of simulations where strategy was most cost-effective	3%	97%
Deterministic	Incr NMB	N/A	£2,130

3 \*Costs and QALYs are mean total costs and QALYs across 5000 PSA simulations

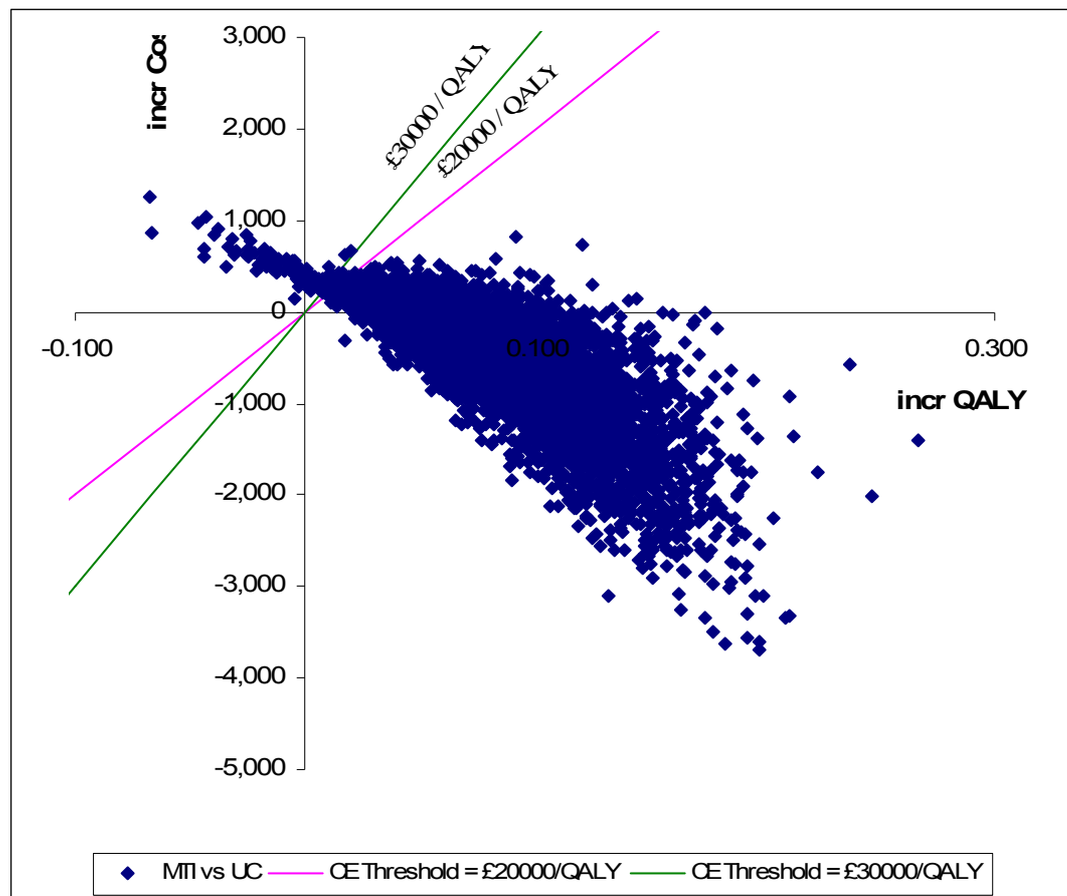
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5

6 At a cost-effectiveness threshold of £20,000 per QALY, the prevention strategy  
 7 was associated with a higher INMB estimate and was more cost-effective in  
 8 96.8% of the simulations that were run in the PSA. In 1.5% of the simulations, the  
 9 intervention strategy increased cost and reduced QALY gains (figure 16.2). The  
 10 INMB was £3,040 at a cost-effectiveness threshold of £30,000 per QALY

11

1 Figure 16.2: cost-effectiveness plane for multi-component targeted intervention  
 2 compared to usual care



3

4

5

6 The results of the one-way deterministic sensitivity analyses are presented in  
 7 table xxx. The use of the prevention intervention remained cost-effective for the  
 8 majority of the DSA. The only exceptions were when we assumed that pressure  
 9 ulcer, falls, in-hospital mortality and extended hospital length of stay were the  
 10 only adverse outcome associated with delirium. In these cases the intervention  
 11 was not cost-effective. The intervention remained cost-effective when we  
 12 excluded the survival difference between delirious and non-delirious cases,  
 13 removed the cost of dementia attributable to stay in long-term care, increased  
 14 the cost of pressure ulcer. The INMB was £2330 when the life expectancy of  
 15 dementia was increased from 1.2 years to 3.6 and 5.4 years for dementia  
 16 patients with and without delirium respectively, An explanation for a higher  
 17 INMB even when the survival implications of dementia are less severe is that the  
 18 additional cost of dementia incurred in additional life years more than off-sets  
 19 the additional health benefits due to increased life expectancy. In further  
 20 analyses, we used the composite outcome of new admission to institution and  
 21 mortality, and assumed that the NHS and PSS would pay only 70% of the cost  
 22 of stay in long-term care but the intervention remained cost-effective.

23

1 Table 16.5: other deterministic sensitivity analyses on the cost-effectiveness of  
 2 multi-component targeted intervention compared to usual care

	<b>Incr NMB (deterministic)</b>
All model parameters (base case)	£2,125
Baseline risk of delirium = 12.5% (base case = 15%)	£1,710
In hospital mortality is the only consequence of delirium	-£140
New dementia is the only consequence of delirium	£440
New admission to nursing home is the only consequence of delirium	£660
Falls is the only consequence of delirium	-£210
Pressure ulcer is the only consequence of delirium	-£370
Extended hospital stay is the only consequence of delirium	-£250
Including 3-year survival difference between delirious and non-delirious patients (as the only adverse outcome in model)	£670
Excluding 3-year survival difference between delirious and non-delirious patients (but including all adverse consequences)	£2009
Excluding the cost of dementia attributable to stay in long-term care (cost of dementia = £5859) (base case = £16,302)	£1994
Life expectancy for dementia patients with previous delirium = 3.6 years, without previous delirium, 5.4 years (base case = 1.2 years)	£2,330
QALY gain for stay in long-term care over life expectancy of 3.6 years for patients with previous delirium and 5.4 years for those without	£2,110
Cost of pressure ulcer using the cost of grade 4 ulcer that heals normally	£2,150
Baseline risk of pressure ulcer = 1.68%	£2,120
Accounted for only 70% of cost of stay in long-term care	£1980
Composite outcome, mortality and new admission to institution	£1980
Increased pay band for Geriatric Nurse (Band 7) and Elder Life Specialist (Band 6)	£2090

3

4 **16.4 Cost-effectiveness of multi-component targeted prevention**  
 5 **interventions in older patients admitted non-electively for surgical**  
 6 **repair of hip fracture**

7

8 The use of multi-component targeted prevent interventions in older patients  
 9 admitted non-electively for surgical repair of hip fracture resulted in an INMB of  
 10 £8070 (table 16.6). In the PSA, the mean total cost of the usual care strategy  
 11 and prevention strategies in this population were estimated as £19,530 and  
 12 £17,040 respectively. The mean QALYs were 1.540 and 1.820 respectively. The

1 intervention strategy reduced cost by £2,490 and increased QALY gain by  
 2 0.290. It therefore dominates the usual care strategy. The ICER and INMB for this  
 3 intervention strategy compared to the usual care strategy were -£8,730 per  
 4 QALY and £8,180 respectively

5

6

7 Table 16.6: costs, QALYs and cost-effectiveness of multi-component targeted  
 8 intervention compared to usual care

		Usual Care	MTI
Probabilistic	Mean cost	£19,530	£17,040
	Mean QALYs	1.540	1.820
	Incr Cost	N/A	-£2,490
	Incr QALYs		0.290
	Incr Cost / QALY		-£8,730
	Incr NMB		£8,180
	% of simulations where strategy was most cost-effective	4%	96%
Deterministic	Incr NMB	N/A	£8,070

9

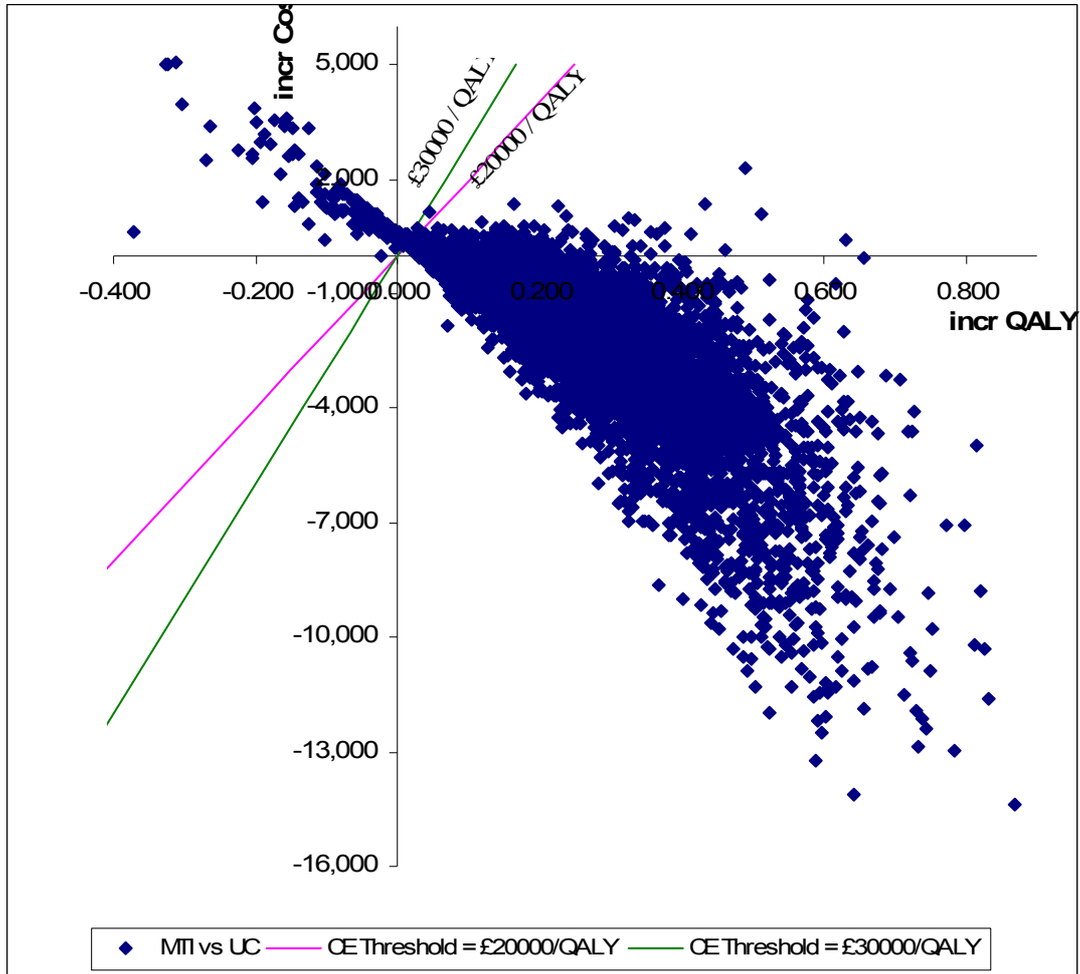
10

11 At a cost-effectiveness threshold of £20,000 per QALY, the prevention strategy  
 12 was more cost-effective in 96.4% of the simulations that were run in the PSA. The  
 13 intervention strategy increased cost and reduced QALY gains in 2.8% of the  
 14 simulations (figure 16.3). The INMB was £11,030 at a cost-effectiveness  
 15 threshold of £30,000 per QALY

16

17

1 Figure 16.3: cost-effectiveness plane for multi-component targeted intervention  
 2 compared to usual care



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7 This intervention strategy remained cost-effective in most of the DSA conducted (table  
 8 16.7). The exceptions were when we assumed that pressure ulcer and extended  
 9 hospital length of stay were the only adverse outcome associated with delirium. The  
 10 intervention was not cost-effective in these cases. When the life expectancy of  
 11 dementia was increased to 3.6 and 5.4 years for dementia patients with and without  
 12 delirium respectively, the INMB was higher than the INMB in base case. In this case, the  
 13 additional cost of dementia incurred in additional life years more than off-sets the  
 14 additional health benefits due to increased life expectancy.

15

1 Table 16.7: other deterministic sensitivity analyses on the cost-effectiveness of  
 2 multi-component targeted intervention compared to usual care

	Incr NMB (deterministic)
All model parameters (base case)	£8,074
Baseline risk of delirium = 12.5% (base case = 50%)	£1,640
In hospital mortality is the only consequence of delirium	£290
New dementia is the only consequence of delirium	£2,270
New admission to nursing home is the only consequence of delirium	£3,060
Falls is the only consequence of delirium	£60
Pressure ulcer is the only consequence of delirium	-£500
Extended hospital stay is the only consequence of delirium	-£62
Including 3-year survival difference between delirious and non-delirious patients (as the only adverse outcome in model)	£3,070
Excluding 3-year survival difference between delirious and non-delirious patients (but including all adverse consequences)	£7,670
Excluding the cost of dementia attributable to stay in long-term care (cost of dementia = £5859) (base case = £16,302)	£7,630
Life expectancy for dementia patients with previous delirium = 3.6 years, without previous delirium, 5.4 years (base case = 1.2 years)	£8,760
QALY gain for stay in long-term care over life expectancy of 3.6 years for patients with previous delirium and 5.4 years for those without	£8,030
Cost of pressure ulcer using the cost of grade 4 ulcer that heals normally	£8,150
Baseline risk of pressure ulcer = 1.68%	£8,070
Accounted for only 70% of cost of stay in long-term care	£7,570
Composite outcome, mortality and new admission to institution	£7,590

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## 7 16.5 THE TREATMENT MODEL

8

### 9 16.5.1 The model structure for the treatment interventions

10

#### 11 16.5.1.1 Decision Tree

12

13 A change in the duration and severity of delirium through treatment will unlikely  
 14 lead to a QALY gain. However, treatment will reduce the cost and QALY loss

1 associated with adverse consequences that will occur in delirious patients. In the  
2 systematic review of the treatment strategies, there were no data on the direct  
3 effect of treatment on the adverse consequences used in the prevention model  
4 above. There were data on intermediate outcomes and we had to use an  
5 intermediate outcome to link the effect of treatment with adverse delirium  
6 consequences. The GDG advised that we use “complete recovery from delirium”  
7 as the intermediate outcome in the model. Data were reported in the adverse  
8 consequences review on the increased risk of nursing home admission and death  
9 for patients without complete recovery.

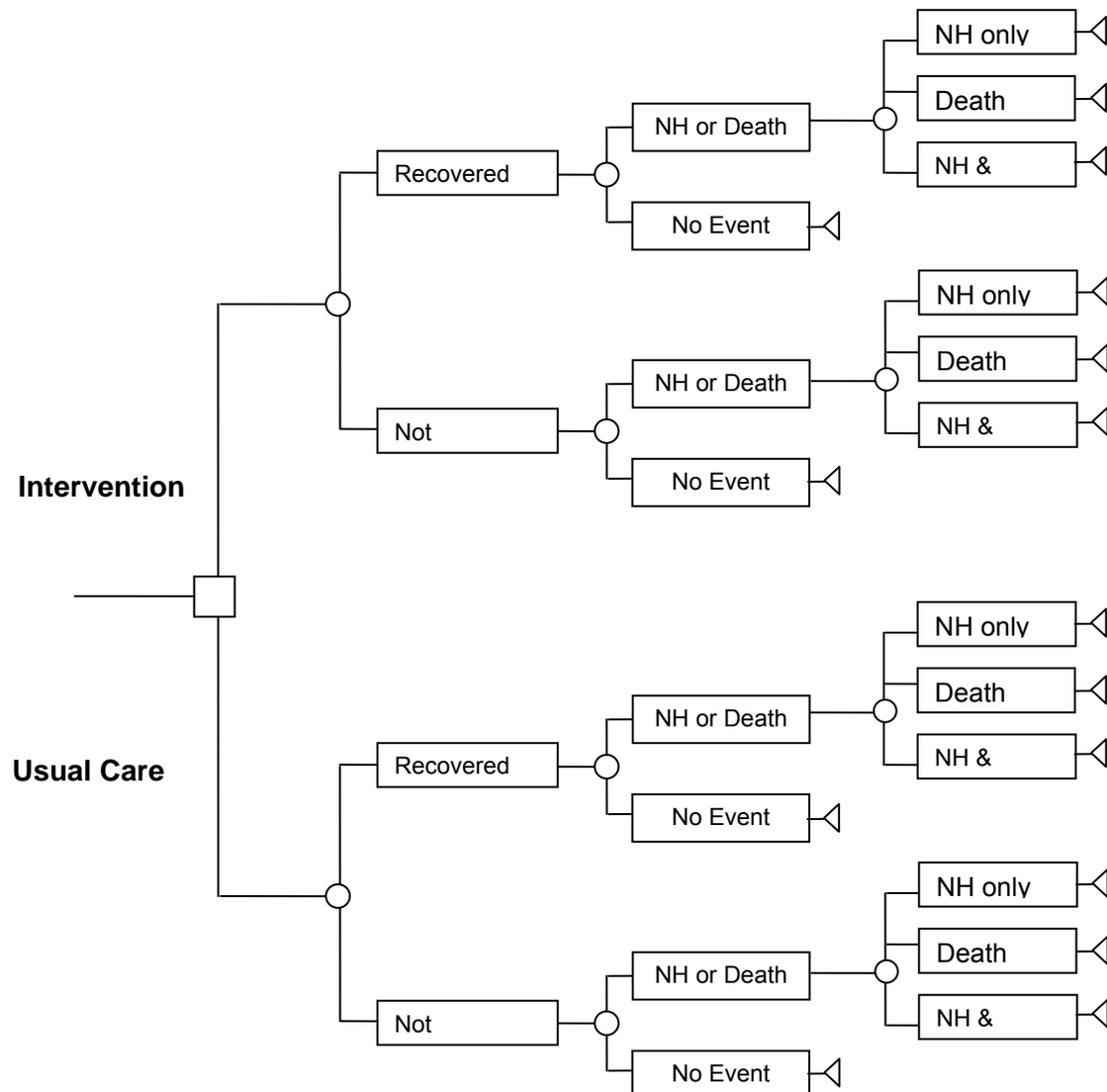
10 The treatment cost-effectiveness model consists of a decision tree (figure 16.3). In  
11 the usual care arm of the tree, the members of a cohort of patients with delirium  
12 will either recover completely or not recover at all. The number of people  
13 recovering will depend on the baseline risk of recovery in a care setting.  
14 Regardless of their recovery status some of them will have no further adverse  
15 event and others will be admitted to the nursing home or will die. Those that  
16 experience further adverse event will either experience admission to nursing  
17 home only, death only or both. The number of people that experience any of  
18 these three outcomes will depend on the baseline risk of these outcomes in the  
19 care setting. In the treatment arm, it will depend on the baseline risks as well as  
20 the relative risk of complete recovery if exposed to the treatment.

21  
22 The GDG advised that we consider the impact of treatment side effects in the  
23 model. A review of the adverse effects of antipsychotic agents suggests that the  
24 only useful evidence for the existence of side effect is for stroke. It was therefore  
25 the only side effect that was considered in the model. We carried out a  
26 sensitivity analysis where stroke was included as one of the branches of the  
27 decision tree.

28  
29 The end of each branch of the tree implies a particular cost and a particular  
30 QALY. The total cost and QALYs are summed up for each strategy.

31  
32

1 Figure 16.3: decision tree for treatment intervention strategies



2

3

4

5 **16.5.2 Absolute Risk Estimates**

6

7 **16.5.2.1 Complete recovery**

8 The baseline risk of complete recovery was taken from the Hu et al study (2006)  
 9 and this study has been described in details in the section on review of hospital  
 10 treatment using pharmacological interventions (chapter 14). It was reported in  
 11 the control arm of the study that five out of a total of 29 people experienced  
 12 complete recovery. We therefore used 17.2% as the baseline risk of complete  
 13 recovery.

14

### 1 **16.5.2.2 Admission to nursing home or death**

2 The baseline risk of “nursing home admission or death” for patients that  
3 recovered as well as those that did not recover were taken from the McAvay et  
4 al study (2006) which has been described in the section on adverse consequences  
5 review (chapter 8). The study compared 1-year institutionalization and mortality  
6 rates of patients who were delirious at discharge, patients whose delirium  
7 resolved by discharge, and patients who were never delirious in the hospital.  
8 Twenty one out of 31 of patients whose delirium resolved experienced “death or  
9 nursing home placement”. An adjusted hazard ratio of 1.73 was reported for  
10 “nursing home admission or mortality” for patients who had delirium at discharge  
11 compared to those whose delirium resolved. We used this adjusted hazard ratio  
12 to estimate the risk of “nursing home admission or mortality” for patients who  
13 had delirium at discharge by assuming that the hazard was constant over time.  
14 This gave a 1 year risk of 85.8%. The McAvay et al study (2006) also reported  
15 data which we used to estimate the proportion of people with death only,  
16 nursing home admission only, and “nursing home admission and death” for  
17 patients whose delirium resolved as well as those whose delirium did not resolve.  
18 For those whose delirium resolved, the proportion of people with nursing home  
19 admission only, death only, and “nursing home admission and death” was  
20 estimated as 61.9%, 33.3% and 4.8% respectively. For those whose delirium  
21 did not resolve, this was estimated as 55.0%, 5.0% and 40.0% respectively.

22

### 23 **16.5.2.3 Stroke**

24 We took the baseline risk of stroke from Wooltorton (2002) who reported an  
25 analysis of drug manufacturer’s trials involving elderly patients with dementia.  
26 Wooltorton (2002) reported that in four placebo-controlled trials lasting one to  
27 three months and involving more than 1200 patients with Alzheimer’s disease or  
28 vascular dementia, cerebrovascular adverse events were twice as common in the  
29 risperidone treated group as in the placebo group. Risperidone is an atypical  
30 antipsychotic and cerebrovascular adverse events were reported to include  
31 stroke and transient ischemic attacks. In the placebo arm, it reported that seven  
32 out of 466 patients experienced this adverse event. We have therefore used  
33 1.5% as the baseline risk of stroke in our model.

34

### 35 **16.5.2.4 Efficacy of Treatment Interventions**

36 The efficacy of different antipsychotic drug treatment interventions has been  
37 reviewed in chapter 14. The two drugs that were identified to be clinically  
38 effective are haloperidol and olanzapine, and we have included only these two  
39 in our model. Haloperidol and olanzapine were estimated to have relative risk  
40 of complete recovery of 3.95 and 3.68 respectively.

41

### 1 **16.5.2.5 Relative risk of stroke as side effect of antipsychotic drugs**

2 The relative risk of stroke following the administration of antipsychotic agents has  
3 been reviewed in chapter 11. We used the data from the Douglas and Smeeth  
4 study (2008) which reported the relative risk of stroke for all antipsychotics  
5 compared to no treatment (RR=1.73); typical antipsychotic compared to no  
6 treatment (RR=1.69); and atypical antipsychotic compared to no treatment  
7 (RR=2.32). In the base case cost-effectiveness analysis we have not included  
8 stroke as a side effect of using antipsychotic agents. In a sensitivity analysis we  
9 have included an increased risk of stroke using the relative risk for all  
10 antipsychotics compared to placebo. In a second sensitivity analysis we have  
11 used the relative risks reported specifically for haloperidol and olanzapine.

12

13

14

## 15 **16.5.3 Cost and QALYs of Outcomes on the decision tree**

16

### 17 **16.5.3.1 Nursing home admission**

18 The estimates of unit cost and duration of stay in long-term care are the same as  
19 the estimates used above in the prevention model. The unit cost of stay in long-  
20 term care is £656 per week and the duration of stay is 18.9 months. The  
21 expected lifetime QALY gain for this outcome has been estimated the same way  
22 it was estimated in the base case of the prevention model.

23

### 24 **16.5.3.2 Death only**

25 The mortality risk was taken from a study (McAvay et al 2006) which reported  
26 this risk in patients followed up for one year post-hospital discharge. We have  
27 assumed that the patient with this outcome will live for six months before death  
28 and we have estimated a QALY again for a 79 year old person who lived for  
29 just six months. We have also assumed that mortality will be associated with zero  
30 cost.

31

### 32 **16.5.3.3 Nursing home admission and death**

33 The cost of this outcome was estimated as a product of the unit cost of stay in  
34 long-term care and the duration of stay. The duration of stay was assumed to be  
35 six months only after which the patient dies. The expected lifetime QALY gain  
36 was estimated in a similar way as it was done in the prevention model. The only  
37 difference is that it was estimated over a period of six months. We used the  
38 same adjusted utility score of 0.18 and the way this was estimated has been  
39 described above.

40

#### 1 **16.5.3.4 Nil Event**

2 For the nil event arm of the decision tree we have assumed that patients will not  
3 experience any death in the first year. Their survival from the second year was  
4 estimated to reflect the increased risk of mortality for persons with delirium.  
5 Adjusted mortality risk was estimated from data from the Rockwood et al study  
6 (1999) and applied in the prevention model for three years. In the treatment  
7 model, we have applied the adjusted increased mortality risk for only 2 years.  
8 The life expectancy of a patient without any event was estimated to be 5.29  
9 years and the QALYs was estimated as 3.24.

10

11

#### 12 **16.5.3.5 Stroke**

##### 13 Cost

14 The cost of stroke in the first year was taken from a cost-effectiveness analysis  
15 that compared different models of stroke care provided in London and  
16 Copenhagen (Grieve et al 2000). In the Copenhagen centre, acute and  
17 rehabilitation unit were combined, and patients could be transferred from the  
18 acute hospital for further inpatient rehabilitation at a separate hospital. In the  
19 London care centre, patients were usually admitted to general wards where they  
20 are treated by general medicine specialist, but could be transferred to a  
21 rehabilitation stroke unit where geriatricians led care. Further rehabilitation as  
22 an inpatient at a separate hospital was not an option. A range of community  
23 services including further rehabilitation and support services were available in  
24 both centres.

25

26 The study participants were first-ever stroke patients and resource use was  
27 recorded one year post stroke. Measurement of resource use took a hospital and  
28 community health perspective and covered primary hospital stay, subsequent  
29 transfer to other hospital, readmissions, institutional care and use of outpatient  
30 and community health services. Data was collected on the use of diagnostic  
31 investigations, the length of stay by ward type, and doctors' and nurses' time  
32 resources. The amount of therapy each patient received was recorded as well as  
33 the length of stay in institutions.

34

35 A standard costing method was reported to have been used in costing inpatient  
36 services. The costs for institutional and community services were based on  
37 interviews undertaken with providers, and the median cost of the item concerned  
38 was used as the unit cost. The cost of a GP consultation came from PSSRU (Netten  
39 & Dennet, 1996) and the same methodology was applied to cost a consultation  
40 in Copenhagen. Disaggregated costs for surgery were not available for the  
41 London centre and were based on costs of surgery in Copenhagen. A factor of  
42 0.74 was used to multiply the costs of surgery in Copenhagen to obtain surgery  
43 costs in London, and the factor was taken from the ratio of costs per hospital day

1 between the centres. Costs were estimated in 1995 local prices but were  
2 converted into dollars using the purchasing power parity index.

3  
4 In the London centre, 358 patients were included in the study but 20 were  
5 excluded from the main analysis because of missing case severity data. Most  
6 patients were admitted to a general medical ward and after an average stay in  
7 the initial area of 8 days, 26% were subsequently transferred to the  
8 rehabilitation stroke unit, and 6% were readmitted to hospital. The mean total  
9 length of all hospital stay in the year following stroke was reported as 35.3  
10 days. On average, there were 3.9 visits to day centre, and the mean length of  
11 days spent in sheltered, residential and nursing homes were 8.1, 8.5 and 16.9  
12 respectively. The mean cost of care in the year following stroke in London was  
13 reported as \$8,825. We converted this to £5,643 using the PPP index for the  
14 year 1995 and up rated the converted estimate to £8,486 using the PSSRU pay  
15 and price indices of 166 for 1995/96 and 256.9 for 2007/08. We applied in  
16 our model £8,486 as the cost of care following first year of stroke.

17  
18 For the cost of care in subsequent years we required information on the life  
19 expectancy of a stroke patient as well as the yearly cost. We took the yearly  
20 cost from the NICE stroke guideline (NICE stroke guideline 2008). Dependent  
21 and independent stroke were reported to cost £11,292 and £876 per patient  
22 per year for subsequent years respectively. These estimates were costs of  
23 inpatient care taken from health technology assessment reports and were largely  
24 determined by calculating total length of hospital stay after stroke and  
25 multiplying by the average cost of inpatient care. We assumed that 62% of the  
26 cases will be independent, 38% will be dependent and the life expectancy of a  
27 stroke patient is 4.7 years (NICE stroke guideline 2008). We estimated the  
28 yearly cost of stroke for subsequent years to be £4827.

### 31 Utility

32 The utility data for stroke was taken from the cross-sectional study by Lindgren  
33 et al 2008. The primary aim of the study was to assess the utility loss among  
34 stroke survivors at different time points following the stroke. The EQ-5D  
35 questionnaire was sent to 393 patients, divided into groups with three, six, nine  
36 and 12 months having passed since the stroke. The study patients had to be  
37 above the age of 18 and below the age of 76 years. This was done to avoid  
38 patients with a high degree of co-morbidities such as dementia. Furthermore, the  
39 sampling process aimed to identify at least 50 patients with ischemic stroke in  
40 each of the four groups listed above, and as many hemorrhagic strokes as were  
41 encountered. The study was conducted among stroke patients at six different  
42 centres that reported data to the Swedish national stroke register. The  
43 recruitment of patients was done consecutively at the study centres during a one  
44 month period. The questionnaire responses were converted to utility scores using  
45 the UK social tariff that were elicited with the time trade-off methodology. The  
46 utility scores for stroke were 0.65, 0.75, 0.63 and 0.67 for patients who have  
47 had stroke for 3, 6, 9 and 12 months respectively. The mean utility score for all

1 patients was 0.67 and mean age of study population was 64.4 years. The QALY  
2 gain due to stroke was estimated using a utility multiplier and duration of 4.7  
3 years. We estimated the utility multiplier, 0.85, as the ratio of the utility of 0.67,  
4 the mean utility score, and 0.79, the utility of a person aged 64.4 years old in  
5 the UK population. The starting age in the model is 79 years and we have used  
6 the utility multiplier to adjust the utility of an average person aged 79 years.  
7 The utility score for stroke that we used in the model was 0.62.

8

## 9 **16.5.4 Cost of Treatment Interventions**

10

### 11 **16.5.4.1 Haloperidol**

12 The costing of haloperidol is based on the oral dosage, 0.5 to 1mg every eight  
13 hours for up to five days. This is based on the dosage that was reported in the  
14 review of treatment interventions (chapter 14) for patients over 60 years. We  
15 have chosen this dosage as the starting age of our model is 79 years. The net  
16 price of 28-tab pack of haloperidol 500 micrograms is 91p (BNF 57,  
17 [<http://bnf.org/bnf/bnf/current/3225.htm#this>] accessed on 19/08/09]). Using  
18 an average of 0.75 mg thrice daily for five days will require 22.5 tablets. We  
19 have therefore used £0.73 as the cost of haloperidol in our model. We did not  
20 consider additional drug administration costs. In a sensitivity analysis we used the  
21 higher dosage of 2.5 to 5mg every eight hours for five days. This dosage was  
22 meant to be for patients less than 60 years old. The net price of 28-tab pack of  
23 haloperidol 5 mg is £3.87. Using 2.5 mg thrice daily for five days will cost  
24 £2.59 and we used this estimate in a sensitivity analysis.

25

### 26 **16.5.4.2 Olanzapine**

27 We have estimated the cost of olanzapine based on the oral dosage, 2.5 mg  
28 daily for up to five days. This dosage was reported for the treatment of patients  
29 over 60 years (chapter 14) and we have chosen this dosage in our base case  
30 analysis as the starting age of our model is 79 years. The net price of 28-tab  
31 pack of olanzapine 2.5 mg is £33.29 (BNF 57,  
32 [<http://bnf.org/bnf/bnf/current/56912.htm#this>], accessed on 19/08/09]).  
33 Using 2.5 mg daily for five days will require only five tablets and will cost  
34 £5.94. In a sensitivity analysis, we used the dosage of five mg daily for five  
35 days. This is the dosage for those less than 60 years old. This will require 10  
36 tablets and will cost £11.89.

37

38 A summary of the input parameter estimates used in the model is in table 16.8  
39 below.

40

1 Table 16.8: other inputs used in base case analysis in the economic model

Model input	Point Estimate (95% CI)	Source
<b>Baseline risk</b>		
Complete recovery	17.2%	Hu et al 2006
Stroke	1.5%	Wooltorton 2002
<b>Absolute risk</b>		
NH admission or death in patients with complete recovery	67.7%	McAvay et al 2006
NH admission or death in patients with delirium at discharge	85.9%	
Proportion of people with death only, nursing home admission only, and "nursing home admission and death"		
NH admission only in patients with complete recovery	61.9%	
Death only in patients with complete recovery	33.3%	
NH admission and death in patients with complete recovery	4.8%	
NH admission only in patients with delirium at discharge	55.0%	
Death only in patients with delirium at discharge	5.0%	
NH admission and death in patients with delirium at discharge	40.0%	
<b>Unit cost</b>		
Stay in long-term care (per week)	£656	PSSRU 2007, Netten et al 1998
Stroke (first year)	£8486	Grieve et al 2000
Stroke (subsequent years)	£4827	NICE guideline on stroke (2008). Assumed that 38% of strokes cases are dependent and 62%, independent
<b>Utility</b>		
Stay in long-term care	0.18	Ekman et al, 2007 (reported 0.25 for moderate dementia, GDG suggested it should be used to estimate utility for this outcome)
Stroke	0.62	Lindgren et al 2008 (reported 0.67 as mean utility score)
<b>Duration</b>		
Stay in long-term care (months)	18.9	Netten et al 2001
Life expectancy for stroke (years)	4.7*	NICE, 2008
<b>Intervention Efficacy</b>		
Haloperidol	3.95 (1.75, 8.9)	Hu et al 2006
Olanzapine	3.68 (1.63, 8.33)	
<b>Intervention Cost</b>		
Haloperidol	£0.73	BNF 57 (dosage for people over 60 years as stated in treatment review)
Olanzapine	£5.94	BNF 57 (dosage for people over 60 years as stated in treatment review)
<b>Relative risk of stroke as a side effect of using antipsychotic agents</b>		
All antipsychotic agents	1.73 (1.60, 1.87)	Douglas and Smeeth 2008
Haloperidol	1.69 (1.55, 1.84)	
Olanzapine	2.32 (1.73, 3.11)	

\*Life expectancy for a patient without any event is 5.3 years

### 16.5.5 Sensitivity Analyses

As described previously for the prevention model, we have used a DSA to explore the importance of the various model assumptions and probabilistic sensitivity analysis to explore the impact of parameter uncertainty associated with the various model inputs. In the first DSA we included the impact of stroke in the model as this was not done in the base case analysis. We used the relative risk of 1.73 for both haloperidol and olanzapine in the first sensitivity analysis. In the second analysis, we used drug specific relative risk estimates (haloperidol = 1.69, olanzapine = 2.32).

One of the adverse consequences included in the model was nursing home admission and death. In the base case, we assumed that death will occur after the patient has spent six months in long-term care. In another DSA we assumed the patient will spend 12 months in long-term care. Further analysis was carried out by assuming that only 70% of the cost of long-term care will be publicly financed. The model parameters, the type of distribution and distribution parameters used in PSA are listed in the table below (table 16.9)

Table 16.9: input parameters, type of distribution and distribution parameters used in PSA

Parameter	Type of distribution	Point estimate	Distribution parameters	Source
<b>Baseline Risk</b>				
Complete recovery	Beta	17.2%	$\alpha = 5, \beta = 24$	Hu et al 2006
<b>Absolute Risk</b>				
NH admission or death in patients with complete recovery	Beta	67.7%	$\alpha = 21, \beta = 10$	
NH admission or death in patients with delirium at discharge	Beta	85.9%	$\alpha = 9, \beta = 1$	
NH admission only in patients with complete recovery	Dirichlet	61.9%	$\alpha = 13$	McAvay et al 2006
Death only in patients with complete recovery		33.3%	$\alpha = 7$	
NH admission and death in patients with complete recovery		4.8%	$\alpha = 1$	
NH admission only in patients with delirium at discharge	Dirichlet	55.0%	$\alpha = 11$	
Death only in patients with delirium at discharge		5.0%	$\alpha = 1$	
NH admission and death in		40.0%	$\alpha = 8$	

patients with delirium at discharge				
Post-discharge survival				
Difference in mortality between delirious and non-delirious patients	Lognormal	HR = 1.71	Log (mean) = 0.54, se = 0.26	Rockwood et al 1999
Cost				
Stay in long-term care	Gamma	£656	Mean = £656, se = £84	PSSRU 2007
Haloperidol	Gamma	£0.73	Mean = £0.73, se = £0.09	BNF 57
Olanzapine	Gamma	£5.94	Mean = £5.94, se = £0.76	BNF 57
Utility				
Stay in institution	Beta	0.25	$\alpha = 293, \beta = 880$	Ekman et al 2007
Population utility	Multinomial	Linear relationship with age	Age-Utility intercept: 1.06; Age-Utility gradient: -0.00	Based on a re-analysis of data from Kind et al 1998 in Ward et al 2007
Efficacy of treatment interventions				
Haloperidol	Lognormal	3.95	Log (mean) = 1.37, se = 0.41	Hu et al 2006
Olanzapine	Lognormal	3.68	Log (mean) = 1.30, se = 0.42	

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3 **16.5.6 Results**

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5 The costs, QALYs and cost-effectiveness estimates of the treatment model are  
6 presented in the table xxx below. In the deterministic base case analysis  
7 haloperidol and olanzapine were both cost-effective when compared to usual  
8 care. Haloperidol and olanzapine were estimated to have INMB of £10,340  
9 and £9,390 respectively. In the PSA, the mean total cost of the three treatment  
10 strategies, usual care, haloperidol and olanzapine were £31,120, £25,630, and  
11 £26,090 respectively. The mean total QALYs were 0.615, 1.035 and 1.004  
12 respectively. The use of haloperidol or olanzapine reduced cost and increased  
13 QALYs when compared to usual care. The ICERs for the two drugs were -  
14 £13,040 and -£12,920 respectively and the INMB were £13,900 and £12,820  
15 respectively. Haloperidol dominates olanzapine because it saves more costs and  
16 generates more QALYs.

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18 Table 16.10: costs, QALYs and cost-effectiveness of haloperidol and olanzapine  
19 treatment intervention compared to usual care

		Usual Care	Haloperidol	Olanzapine
Deterministic	Incr NMB	N/A	£10,340	£9,390
Probabilistic	Mean cost	£31,120	£25,630	£26,090
	Mean QALYs	0.615	1.035	1.004

	Incr Cost		-£5,490	-£5,030
	Incr QALYs	N/A	0.420	0.390
	Incr Cost / QALY		-£13,040	-£12,920
	Incr NMB		£13,900	£12,820
	% of simulations where strategy was most cost-effective	0%	54%	45%

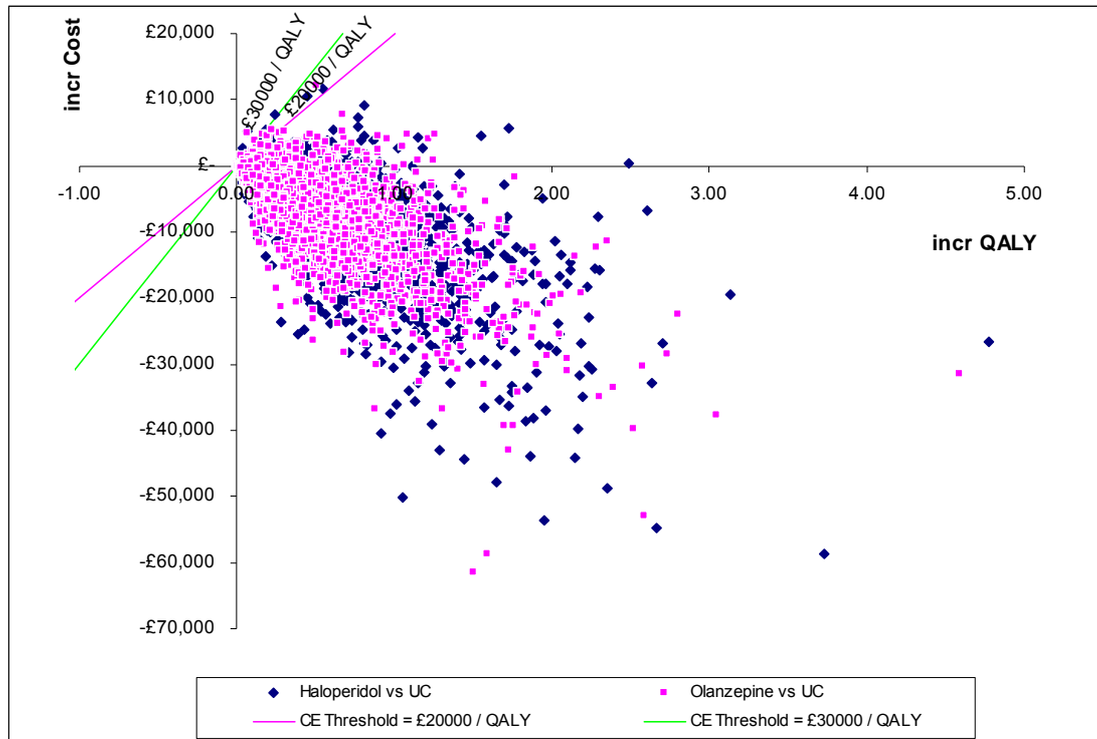
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2 At a cost-effectiveness threshold of £20,000 per QALY, the use of haloperidol  
 3 was the most cost-effective in 54.4% of the simulations that were run in the PSA  
 4 (figure 16.4). The use of Olanzapine was most cost-effective in 45.4% of the  
 5 simulations. Usual care was the most cost-effective strategy in only 0.3% of all  
 6 simulations. Haloperidol increased cost and reduced QALYs in 0.00% of the  
 7 simulations while olanzapine increased cost and reduced QALYs in 0.02% of the  
 8 simulations. When compared to usual care and at a threshold of £20,000 per  
 9 QALY, haloperidol was cost-effective 99.74% of all the 5000 simulations. For  
 10 olanzapine, it was 99.72%. At a cost-effectiveness threshold of £30,000 per  
 11 QALY, it was 99.92% and 99.90% for haloperidol and olanzapine respectively.

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14 Figure 16.4: cost-effectiveness plane for haloperidol and olanzapine treatment  
 15 interventions compared to usual care



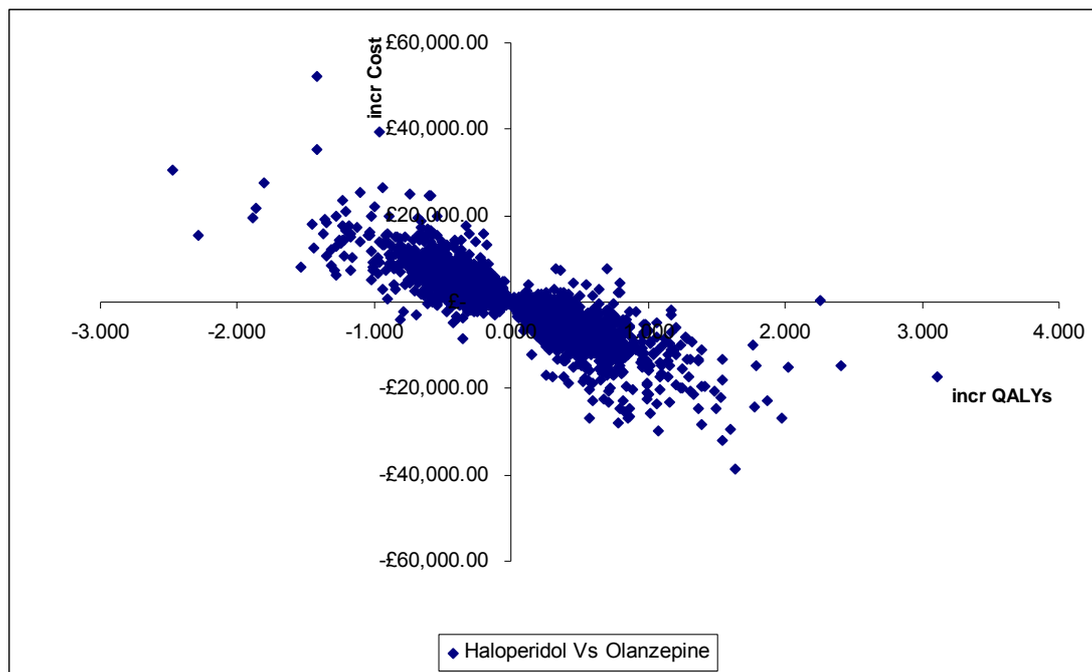
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1 When compared with olanzapine, haloperidol was associated with a mean cost  
 2 reduction of -£460 and a mean incremental QALY of 0.031. The ICER and INMB  
 3 were -£14,560 and £1,080 respectively. However, there is wide uncertainty  
 4 around the incremental cost-effectiveness of haloperidol compared to  
 5 olanzapine as shown in figure 16.5. Haloperidol was more cost-effective in  
 6 54.5% of the 5000 simulations and olanzapine was more cost-effective in the  
 7 rest (45.5%) of the simulations.

8  
 9 Figure 16.5: cost-effectiveness plane for haloperidol treatment interventions  
 10 compared to olanzapine



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 13 The two treatment intervention strategies in the model remained cost-effective in  
 14 all the univariate DSA that we conducted. When compared with usual care, the  
 15 use of the drugs resulted in higher INMB and became even more cost-effective  
 16 when the time a person stays in long-term care before death was increased to  
 17 12 months. They became less cost effective when the impact of stroke side effect  
 18 is included in the model. When compared to olanzapine, haloperidol was  
 19 estimated to have the higher INMB for all the analyses conducted.

20  
 21  
 22 Table 16.11: other deterministic sensitivity analyses on the cost-effectiveness of  
 23 haloperidol and olanzapine treatment interventions compared to usual care

	Incr NMB (Haloperidol)	Incr NMB (Olanzapine)
All model parameters excluding the side effect stroke (Base case)	£10,340	£9,390
All model parameters including the side effect stroke (RR for both	£9,950	£9,000

atypical antipsychotic = 1.73)		
Drug specific stroke relative risk (Hal=1.69, Ola=2.32)	£9,970	£8,680
Duration of stay in long-term care before death=12 months	£12,750	£11,580
Accounted for only 70% of cost of stay in long-term care	£9,100	£8,260
Increased cost of haloperidol due to increased dosage	£10,340	N/A
Increased cost of olanzapine due to increased dosage	N/A	£9,384

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## 4 16.6 SUMMARY OF THE RESULTS OF THE COST-EFFECTIVENESS

### 5 MODELS

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We estimated the cost-effectiveness of prevention and treatment interventions using an original economic evaluation model. The use of multi-component targeted interventions was found to be cost-effective in the prevention of delirium in the population groups considered in the model (elderly patients at risk of delirium who were admitted to the general medicine service and patients undergoing surgical repair of hip fracture). The use of haloperidol and olanzapine in the treatment of delirium was also cost-effective. On average, haloperidol was associated with a higher net monetary benefit but there is wide uncertainty around the incremental cost-effectiveness.

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There are a number of limitations with the model findings and the GDG considered these when interpreting the results of the analyses. In the prevention model we have assumed that the adverse outcomes on the branches of the decision tree are mutually exclusive. It is possible that a patient with delirium who experiences dementia will also be admitted to a nursing home and the total cost and QALY gain for that patient might be different from the modelled estimate as the two outcomes are occurring in the same patient rather than in separate individuals. We tried to test the impact of this assumption by considering that each of the six adverse outcomes was the only outcome to be associated with delirium therefore removing the risk of double counting. The results of the model were robust in that multi-component interventions remain cost-effective.

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In the prevention and treatment model, the baseline risk estimates we used for delirium in hospital, dementia, new admission to institution, complete recovery after delirium incidence and stroke were taken from studies in other countries. The baseline risk of complete recovery and efficacy of treatment interventions were taken from a Chinese study (Hu et al 2006). The absolute risk used in the treatment model for nursing home admission, death or nursing home admission and death were taken from a US study (McAvay et al 2006). We could not identify suitable UK studies for these outcomes and the ones chosen were the

1 best available in terms of study quality and applicability. We assumed that the  
2 relative risk of falls and pressure ulcer are the same. No other better studies  
3 could be identified for these outcomes. The GDG discussed the applicability of  
4 the studies that were used and considered them in the interpretation of the  
5 results.

6

7 The cost estimate used in the base case analysis for pressure ulcer in the  
8 prevention model was based on the assumption that it would be a grade 1  
9 pressure ulcer that would heal normally. We made an alternative assumption  
10 that it would be a grade 4 ulcer. We assumed in the base case analysis for the  
11 prevention and treatment models that all the cost of long-term care will be paid  
12 by the NHS and PSS. We made an alternative assumption that only 70% of this  
13 cost will be paid by the public. The cost of dementia in the prevention model  
14 included the cost of stay in long-term care. It could be argued that the cost of  
15 long-term care has been accounted for as a different model outcome and that  
16 we have double counted cost. We made an alternative assumption and removed  
17 the proportion of cost of dementia attributable to long-term care. In all the  
18 alternative assumptions the model results suggest that the prevention and  
19 treatment interventions considered above remained cost-effective. In the  
20 treatment model we have assumed, in base case analysis, that patients who  
21 experience nursing home admission and death will spend only six months in long-  
22 term care before death. The cost-effectiveness estimate from this assumption was  
23 conservative as an increase in the duration to 12 months showed that the  
24 treatment interventions were even more cost-effective.

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26 The point estimates used in the model were associated with some uncertainties  
27 which are normally captured in confidence intervals and ranges. We have tried  
28 to explore the effect of such uncertainties using probabilistic sensitivity analysis.  
29 The results of which did not change the findings that the use of multi-component  
30 treatment interventions was found to be cost effective in elderly patients that  
31 had surgery for hip fracture repair, or elderly patients at intermediate or high  
32 risk of delirium who were admitted in the general medicine services. The use of  
33 haloperidol and olanzapine were also found to be cost-effective in the treatment  
34 of delirium.

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31 **Appendices A–K are in separate files**