

**Abatacept, adalimumab, etanercept and
tocilizumab for treating juvenile idiopathic
arthritis (including review of TA35) [ID783]**

Assessment Report

Commercial in Confidence stripped version for consultation

Produced by: Southampton Health Technology Assessment Centre (SHTAC)

CONFIDENTIAL UNTIL PUBLISHED

Technology Assessment Report commissioned by the NIHR HTA Programme on behalf of the National Institute for Health and Clinical Excellence

The clinical and cost-effectiveness of abatacept, adalimumab, etanercept and tocilizumab for treating juvenile idiopathic arthritis: a systematic review and economic evaluation

The Assessment group has presented the results of its economic modelling using the list prices for abatacept, adalimumab, etanercept and tocilizumab in this Assessment Report. A separate confidential appendix reporting the results incorporating the confidential patient access schemes for abatacept and tocilizumab has been prepared by the Assessment Group. The confidential appendix will not be released publically

Produced by Southampton Health Technology Assessments Centre (SHTAC)

Authors Jonathan Shepherd
Keith Cooper
Petra Harris
Joanna Picot
Micah Rose

Correspondence to Corresponding author:
Dr Jonathan Shepherd
Southampton Health Technology Assessments Centre (SHTAC)
University of Southampton
First Floor, Epsilon House
Enterprise Road, Southampton Science Park
Southampton, SO16 7NS, UK.
Tel: +44(0)23 8059 7055
Fax: +44(0)23 8059 5639
email: jps@soton.ac.uk
www.southampton.ac.uk/shtac

Date completed: 6th July 2015

Note: This document and any associated economic model are protected by intellectual property rights (IPR), which are owned by the University of Southampton. Anyone wishing to modify, adapt, translate, reverse engineer, decompile, dismantle or create derivative work based on the economic model must first seek the agreement of the property owners.

Word count: 63,707

Keywords: juvenile idiopathic arthritis, disease modifying anti-rheumatic drugs, anti-TNF therapy, systematic review, economic evaluation, health technology assessment

Source of funding: This report was commissioned by the NIHR HTA Programme as project number 14/64/01 and will be published in full in *Health Technology Assessment* (<http://www.journalslibrary.nihr.ac.uk/hta>).

Declared competing interests of authors

None declared.

All authors have completed the unified competing interest form at www.icmje.org/coi_disclosure.pdf (available on request from the corresponding author) and declare 1) no financial support for the submitted work from anyone other than their employer; 2) no financial relationships with commercial entities that might have an interest in the submitted work; 3) no spouses, partners, or children with relationships with commercial entities that might have an interest in the submitted work; and 4) no non-financial interests that may be relevant to the submitted work.

Acknowledgements

We would like to thank members of our advisory group who provided expert advice and comments on the protocol and a draft of this report: Professor Michael Beresford, Professor in Child Health, University of Liverpool; Professor Rachel Elliot, Lord Trent Professor of Medicines and Health, University of Nottingham; Dr Kimme Hyrich, Reader in Rheumatic Disease Epidemiology, University of Manchester; Dr Alice Leahy, Paediatric Rheumatology Consultant, Southampton General Hospital; Helen Strike, Nurse Specialist Paediatric and Adolescent Rheumatology, Bristol Royal Hospital for Children; Professor Lucy Wedderburn, Professor of Paediatric Rheumatology, Institute of Child Health, London.

Thanks to Karen Welch, Information Specialist, at SHTAC for conducting the literature searches; and thanks to Dr Jeremy Jones, Principal Research Fellow at SHTAC, for reviewing a draft of the report. Thanks to Maria Chorooglou (Senior Research Fellow at SHTAC) and Diana Mendes (independent health economist) for assisting with the systematic reviews of cost-effectiveness and health related quality of life.

Rider on responsibility for the report

The views and opinions expressed in this report are those of the authors and do not necessarily reflect those of the HTA programme, NIHR, NHS or the Department of Health. Any errors are the responsibility of the authors.

This report should be referenced as follows:

The clinical and cost-effectiveness of abatacept, adalimumab, etanercept and tocilizumab for treating juvenile idiopathic arthritis: a systematic review and economic evaluation. *Health Technology Assessment 2015*

Contribution of authors

Jonathan Shepherd (Principal Research Fellow) project managed the study, developed the research protocol, contributed to drafting the background section, assisted in the development of the search strategy, assessed studies for inclusion, performed data extraction and critical appraisal of included clinical-effectiveness studies, synthesised evidence, assessed the company submissions and drafted and edited the final report.

Keith Cooper (Senior Research Fellow) developed the research protocol, assessed cost-effectiveness and HRQoL studies for inclusion, synthesised evidence, led the development of the economic evaluation, assessed the company submissions and drafted the final report.

Petra Harris (Research Fellow) developed the research protocol, contributed to drafting the background section, assessed studies for inclusion, performed data extraction and critical appraisal of included clinical-effectiveness studies, synthesised evidence, assessed the company submissions and drafted the final report.

Joanna Picot (Senior Research Fellow) developed the research protocol, contributed to drafting the background section, assessed studies for inclusion, performed data extraction and critical appraisal of included clinical-effectiveness studies, synthesised evidence, assessed the company submissions and drafted the final report.

Micah Rose (Research Fellow) contributed to the development of the economic evaluation, assessed the company submissions and drafted the report.

Commercial in confidence information (CIC) is highlighted in blue. Academic in confidence information (AIC) is highlighted in yellow.

ABSTRACT

Background:

Juvenile idiopathic arthritis (JIA) is characterised by joint pain, swelling and limitation of movement caused by inflammation. Subsequent joint damage can lead to disability and growth restriction.

Treatment includes disease modifying anti-rheumatic drugs (DMARD) with methotrexate, the most commonly used DMARD in the UK. Clinical practice now favours newer drugs termed biologic DMARDs where indicated.

Objective: To assess the clinical and cost-effectiveness of four biologic DMARDs (etanercept, abatacept, adalimumab and tocilizumab - with or without methotrexate where indicated) for the treatment of JIA (systemic or oligoarticular JIA excluded).

Data sources: Electronic bibliographic databases including MEDLINE, EMBASE, The Cochrane Library and DARE were searched for published studies from inception to May 2015 for English language articles. Bibliographies of related papers, systematic reviews and company submissions were screened and experts were contacted to identify additional evidence.

Review methods: Systematic reviews of clinical-effectiveness, health-related quality of life and cost-effectiveness were undertaken according to the general principles recommended in the Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement. A cost-utility decision analytic model was developed to compare estimated cost-effectiveness of biologic DMARDs versus methotrexate for JIA. The base case time horizon was 30 years and the model took a National Health Service (NHS) perspective, with costs and benefits discounted at 3.5%.

Results: Four RCTs met the inclusion criteria of the clinical-effectiveness review (one RCT evaluating each biologic DMARD). Only one RCT included UK participants. All four RCTs were withdrawal trials with a placebo comparator. Participants had to achieve an American College of Rheumatology Pediatric (ACR Pedi) 30 response to open-label lead-in treatment in order to be randomised. An exploratory adjusted indirect comparison suggests that the four biologic DMARDs are similar with fewer disease flares and greater proportions with ACR Pedi 50 and 70 responses among participants randomised to continued biologic DMARD. However, confidence intervals were wide, the number of trials was low and there is clinical heterogeneity between the trials. Open-label extensions of the trials showed that generally ACR responses remained constant or even increased after the double-blind phase. The proportions of adverse events and serious adverse events were generally similar between treatment and placebo groups. The incremental cost-effectiveness ratio (ICER) for adalimumab, etanercept and tocilizumab versus methotrexate was £38,127, £32,526 and £38,656 per QALY, respectively. The ICER for abatacept versus methotrexate as a second line biologic was £39,536 per QALY.

Limitations: The model does not incorporate the natural history of JIA in terms of long-term disease progression, as the current evidence is limited. There are no head-to-head trials of biologic DMARDs and clinical evidence for specific JIA subtypes is limited.

Conclusions: Biologic DMARDs are superior to placebo treatment in RCTs enrolling children with (predominantly) polyarticular course JIA, an insufficient response to previous treatment. Randomised head-to-head comparisons of biologic DMARDs with long-term follow-up of safety and efficacy are needed to establish comparative effectiveness. RCTs for JIA subtypes where evidence is lacking are also required.

Word count: 483

TABLE OF CONTENTS

Scientific Summary	10
1 BACKGROUND	20
1.1 Description of underlying health problem	20
1.2 Measures of response to treatment and definition of remission	26
1.3 Current service provision	28
1.4 Description of technology under assessment	29
2 DEFINITION OF THE DECISION PROBLEM	35
2.1 Decision problem	35
2.2 Overall aims and objectives of assessment	36
3 METHODS	36
3.1 Identification of studies	36
3.2 Inclusion and exclusion criteria	37
3.3 Data extraction strategy	38
3.4 Critical appraisal strategy	39
3.5 Method of data synthesis	39
4 CLINICAL-EFFECTIVENESS	39
4.1 Results	39
4.1.1 Quantity and quality of research available	39
4.1.2 Assessment of clinical-effectiveness - biologic DMARDs vs placebo (with methotrexate where permitted)	49
4.1.3 Assessment of clinical-effectiveness - biologic DMARDs vs each other (with methotrexate where permitted)	66
4.1.4 Summary of the systematic review of clinical-effectiveness	71
4.2 Review of clinical-effectiveness in company submissions to NICE	74
4.3 Ongoing trials	79
4.4 Additional supporting evidence	85
4.4.1 Enthesitis-related arthritis (ERA) and psoriatic arthritis (PA)	85
4.4.2 JIA-associated uveitis	91
5 ECONOMIC ANALYSIS	93
5.1 Introduction	93
5.2 Systematic review of cost-effectiveness evidence	93
5.3 Systematic review of health related quality of life studies (HRQoL)	103
5.4 Review of cost-effectiveness in company submissions to NICE	107
5.5 Independent economic evaluation	117
5.6 Methods for independent economic analysis	118
5.6.1 Data sources	121
5.7 Results of the independent economic analysis	129
5.7.1 Sensitivity analysis	132
5.7.2 Scenario analysis	134
5.7.3 Sub-groups	140
5.8 Comparison of the economic models	142
5.9 Discussion	143
6 ASSESSMENT OF FACTORS RELEVANT TO THE NHS AND OTHER PARTIES	144
7 DISCUSSION	146
7.1 Statement of principal findings	146
7.1.1 Clinical-effectiveness	146
7.1.2 Cost-effectiveness	149
7.2 Strengths and limitations of the assessment	151
7.3 Uncertainties	153
8 CONCLUSIONS	154
8.1 Implications for service provision	154
8.2 Suggested research priorities	154
9 REFERENCES	155
10 APPENDICES	168

APPENDICES

Appendix 1 Search dates and example Medline search strategies for clinical-effectiveness, cost-effectiveness and HRQoL	168
Appendix 2 Screening Phase 1 – Titles and abstracts for systematic review of clinical-effectiveness	172
Appendix 3 Screening Phase 2 – Full papers for systematic review of clinical-effectiveness	173
Appendix 4 Table of excluded and unclear studies from systematic review of clinical-effectiveness	175
Appendix 5 Clinical-effectiveness data extraction tables	178
Appendix 6 Table of excluded studies for systematic review of cost-effectiveness.....	196
Appendix 7 Table of excluded studies for systematic review of health-related quality of life.....	197
Appendix 8 Cost-effectiveness studies – data extraction forms	199
Appendix 9 Health related quality of life systematic review– data extraction forms	205
Appendix 10 Cost-effectiveness data extraction forms for the company submissions.....	209
Appendix 11 Parameters used in the independent model PSA	227

TABLES

Table 1 JIA classification according to the revised ILAR Criteria	20
Table 2 Long term outcomes for different sub-types of JIA.....	23
Table 3 Proportions of different sub-types of JIA	24
Table 4 Overview of the drug treatment pathway for JIA	29
Table 5 Summary of licensed indications of the biologic DMARDs under consideration in this assessment.....	30
Table 6 Dosing regimen for abatacept	32
Table 7 Dosing regimen for adalimumab	33
Table 8 Dosing regimen for etanercept.....	34
Table 9 Dosing regimen for tocilizumab	35
Table 10 Summary characteristics of included studies	42
Table 11 Summary of outcomes measured.....	43
Table 12 Selected baseline characteristics of trial participants.....	48
Table 13 Summary of risk of bias assessment	49
Table 14 Disease flare during the randomised withdrawal phase.....	50
Table 15 ACR paediatric responses relative to baseline	52
Table 16 ACR Pedi core variables.....	53
Table 17 Joint-related outcomes (<i>other than ACR Pedi</i>)	54
Table 18 Pain	55
Table 19 Childhood Health Questionnaire (CHQ).....	56
Table 20 Adverse events (AEs)	58
Table 21 ACR Pedi response by background medication use at baseline at the end of the double-blind RCT phase.....	60
Table 22 ACR Pedi outcomes from trial open-label extension periods (OLE)	61
Table 23 OLE adverse events for abatacept.....	64
Table 24 OLE adverse events for etanercept	65
Table 25 OLE adverse events for tocilizumab.....	65
Table 26 Indirect comparisons of biologic DMARDs: disease flare	69
Table 27 Indirect comparisons of biologic DMARDs: ACR Pedi 50 response.....	69
Table 28 Indirect comparisons of biologic DMARDs: ACR Pedi 70 response.....	70
Table 29 Baseline characteristics.....	80
Table 30 Results week 12	80
Table 31 AEs week 12	81
Table 32 AEs >1 dose of adalimumab week 52.....	81
Table 33 Flares: 24-week double-blind phase	84
Table 34 Key baseline characteristics	86
Table 35 ACR Pedi response and inactive disease results at week 12 and week 96	88

Table 36 Mean change from baseline week effectiveness measures at week 12 (observed cases).....	89
Table 37 Adverse events at Week 96.....	90
Table 38 Characteristics of economic evaluations.....	94
Table 39 Critical appraisal checklist for economic evaluations.....	95
Table 40 Characteristics of included quality of life studies.....	104
Table 41 Characteristics of included HRQoL study by Hendry and colleagues.....	105
Table 42 Characteristics of patients included in Prince and colleagues.....	107
Table 43 Unit costs and dosages used in the BMS company model.....	109
Table 44 Results of the BMS model base case (CS Table 13).....	110
Table 45 ACR Pedi response rates from Roche submission (CS Table 21).....	113
Table 46 Roche base case results: combination therapy.....	114
Table 47 Roche base case results: biologic DMARD monotherapy.....	114
Table 48 Corrected Roche model results: combination therapy.....	115
Table 49 Risk of disease flare.....	122
Table 50 Discontinuations during the trials' lead-in time (1st cycle).....	123
Table 51 HRQoL utility values.....	124
Table 52 Drug acquisition costs and dosages (Source BNFC).....	125
Table 53 Concomitant biologic DMARD and methotrexate use.....	126
Table 54 Resource use and unit costs.....	127
Table 55 Summary of the input parameters used in the SHTAC economic model.....	128
Table 56 Summary of the total undiscounted QALYs in each health state for treatment with 1 st line biologic compared to methotrexate.....	130
Table 57 Summary of the total undiscounted costs in each health state for treatment with 1 st line biologic compared to methotrexate.....	130
Table 58 Cost-effectiveness of 1 st line biologic DMARDs versus methotrexate only.....	130
Table 59 Summary of the total undiscounted QALYs in each health state for treatment with 2 nd line biologics compared to methotrexate.....	131
Table 60 Summary of the total undiscounted costs in each health state for treatment with 2 nd line biologic DMARDs compared to methotrexate.....	131
Table 61 Cost-effectiveness of 2 nd line biologic DMARDs compared with methotrexate using list price.....	132
Table 62 Deterministic sensitivity analysis for adalimumab versus methotrexate only.....	132
Table 63 Deterministic sensitivity analysis for etanercept versus methotrexate only.....	133
Table 64 Deterministic sensitivity analysis for tocilizumab versus methotrexate only.....	133
Table 65 Deterministic sensitivity analysis for abatacept versus methotrexate only.....	134
Table 66 Cost-effectiveness for first line biologics versus methotrexate only with patients discontinuation of treatment for clinical remission.....	135
Table 67 Summary of the cost-effectiveness for adalimumab, etanercept and tocilizumab versus methotrexate only using health state costs from Prince and colleagues.....	135
Table 68 Cost-effectiveness for etanercept versus methotrexate using discount rate of 6% for costs and 1% for benefits.....	136
Table 69 Cost-effectiveness for first line biologics versus methotrexate only with inclusion of disutility for caregivers.....	137
Table 70 Cost-effectiveness for three lines of biologic therapy.....	137
Table 71 Cost-effectiveness of first line biologics with a starting age of 6 years.....	138
Table 72 Cost-effectiveness of 2 nd line biologics with a starting age of 6 years.....	138
Table 73 Summary of the probabilistic sensitivity results for first line biologics versus methotrexate only.....	139
Table 74 Summary of the probabilistic sensitivity results for 2 nd line biologics versus methotrexate only.....	140
Table 75 Comparison of the drug costs in the assessment report model with the Roche CS model (20 year discounted, no PAS).....	143
Table 76 Comparison of the drug costs in the assessment report model with the BMS CS model (20 year discounted, no PAS).....	143

FIGURES

Figure 1 Flow-chart for the identification of studies	41
Figure 2 Indirect comparison of biologic DMARDs	66
Figure 3 Summary forest plot of biologic DMARDs versus placebo: disease flare	68
Figure 4 Summary forest plot of biologic DMARDs versus placebo: ACR Pedi 50 response.....	69
Figure 5 Summary forest plot of biologic DMARDs versus placebo: ACR Pedi 70 response.....	70
Figure 6 Flow chart of identification of studies for inclusion in the review of cost-effectiveness	94
Figure 7 Flow chart of identification of studies for inclusion in the review of QoL studies	104
Figure 8 Schematic of the SHTAC JIA model structure.....	120
Figure 9 CEAC from the PSA for 1 st line biological treatments compared to methotrexate	139
Figure 10 CEAC from the PSA for 2 nd line biological treatments compared to methotrexate	140

Scientific Summary

Background

The term juvenile idiopathic arthritis (JIA) encompasses all forms of arthritis of unknown cause with onset prior to the age of 16 years with persisting symptoms for more than six weeks. Suggested incidence (1.6 to 23 per 100,000) and prevalence rates (3.8 to 400 per 100,000) vary widely. The disease is characterised by joint pain, swelling and limitation of movement caused by inflammation of the synovial membrane of the affected joints. Left untreated, this inflammation causes a progressive erosive arthritis, potentially leading to disability and growth restriction. However, severity of disease and long term outcome is variable both between different JIA subtypes and between different individuals with the same JIA subtype. At onset the particular sub-type of JIA will be diagnosed according to the presenting features as either oligoarthritis, polyarthritis, enthesitis-related JIA (ERA), psoriatic arthritis (PA), systemic-onset JIA, or undifferentiated arthritis. Polyarticular course JIA applies to patients who at a particular point in time six months or more after the onset of disease (JIA of any onset type) have five or more active joints. Polyarticular course JIA can typically include rheumatoid factor (RF) positive and RF negative polyarthritis, extended oligoarthritis, ERA, PA and systemic JIA (providing there have been no active systemic symptoms during the previous six months).

Treatment of JIA includes non-steroidal anti-inflammatory drugs, intra-articular corticosteroids and disease modifying anti-rheumatic drugs (DMARDs), with methotrexate the most common conventional (non-biologic) DMARD used in the UK. Clinical practice now favours earlier step-up treatment to biologic DMARDs, where indicated.

Objectives

The aim of this multiple technology appraisal (MTA) is to assess the clinical and cost-effectiveness of biologic DMARDs etanercept, abatacept, adalimumab, and tocilizumab in combination with methotrexate, where permitted, in the treatment of JIA. It updates and extends a previous NICE appraisal of etanercept conducted in 2002 (NICE TA35). The licensed indication for etanercept has broadened since 2002 and three newer biologic DMARDs have been licensed. This appraisal includes all sub-types of JIA with the exception of systemic JIA or persistent oligoarticular JIA.

Methods

Clinical-effectiveness systematic review

Electronic bibliographic resources including MEDLINE, EMBASE, The Cochrane Library, Embase, and DARE were searched for published studies from inception to May 2015 for English language

articles. Bibliographies of included articles and systematic reviews were also searched for additional studies, as were the company submissions to NICE. An expert advisory group was contacted to identify additional published and unpublished evidence.

Titles and abstracts were screened for eligibility by two reviewers independently using inclusion criteria that were defined *a priori*. Inclusion criteria were applied to full texts by one reviewer and checked by a second reviewer. Inclusion criteria were as follows:

- Population: patients with JIA including polyarthritis (both rheumatoid factor positive and negative, and extended oligoarthritis, both onset and course), ERA and PA.
- Intervention: biologic DMARDs abatacept, adalimumab, etanercept and tocilizumab (in combination with methotrexate where permitted), evaluated within their licensed indication. Studies of biologic DMARDs without concomitant methotrexate were permitted if patients were intolerant to it or if treatment with methotrexate was inappropriate.
- Comparators: DMARDs such as methotrexate (best supportive care if DMARDs are not tolerated), as well as abatacept, adalimumab, etanercept and tocilizumab compared with each other.
- Outcomes: disease activity, disease flares, physical function, joint damage, pain, corticosteroid reducing regimens, extra-articular manifestations (such as uveitis), body weight and height, mortality, adverse effects of treatment and health-related quality of life (HRQoL).
- Design: randomised controlled trials (RCTs). Non-randomised studies could be considered where RCT data were not available.

Data extraction and quality assessment were undertaken by one reviewer and checked by a second reviewer. Differences in opinion were resolved by discussion at each stage or in consultation with a third reviewer where necessary.

Data were synthesised through narrative reviews with tabulation of the results of included studies. An adjusted pairwise indirect comparison of the four biologic DMARDs was presented.

Economic evaluation

A systematic review of cost-effectiveness studies, and a systematic review of HRQoL studies was conducted to identify relevant evidence to inform the economic evaluation. Studies were included in the systematic review of cost-effectiveness if they were full economic evaluations (cost-effectiveness, cost-utility or cost benefit-analyses or cost-consequence).

A cost-utility decision analytic model was developed to compare the cost-effectiveness estimates of biologic DMARDs versus methotrexate. The model used a Markov approach to estimate the costs and

health benefits for patients with JIA. The model consisted of three health states: on treatment (with biologic DMARD), off treatment and death, with a further health state 'clinical remission off treatment' also included in a scenario analysis. The model cycles were three months in length to be consistent with timing between outpatient appointments in clinical practice. Patients discontinued treatment due to adverse events, inefficacy of the treatment or remission. The model also included the cost and disutility of disease flares. The perspective of the analysis was that of the NHS and Personal Social Services. The model used a time horizon of 30 years and discount rates of 3.5% for costs and health benefits. The outcome of the economic evaluation is reported as cost per quality-adjusted life year (QALY) gained.

Results

Clinical-effectiveness

From 2554 references screened on title and abstract, 56 full texts were retrieved. One further conference abstract was identified from a pharmaceutical company submission to NICE. From these, nine full papers and 12 conference abstracts met the inclusion criteria. The included papers and abstracts collectively described four multi-centre RCTs, with one RCT each evaluating abatacept, adalimumab, etanercept and tocilizumab. Only the tocilizumab study included UK participants. All four studies were described as being withdrawal trials starting with an open-label lead-in phase (12 to 16 weeks) in which participants had to achieve an American College of Rheumatology (ACR) Pedi 30 response to be eligible for entry to the randomised double-blind withdrawal phase of the study (16 to 32 weeks), followed by an open-label extension (OLE). All studies used a placebo as the comparator. With the exception of the etanercept trial, the majority of patients in the trials received methotrexate in addition to the biologic DMARD or placebo. The distribution of patients across the sub-types of JIA was only reported for two of the trials, with polyarthritis being the predominant sub-type. The other two trials appeared to include patients with polyarticular course JIA. Overall, the quality of the RCTs was reasonable with a low risk of bias for most domains, but some aspects were rated as unclear primarily due to insufficient reporting.

Significantly fewer patients who continued to receive biologic DMARDs during the randomised withdrawal phase of the studies had arthritis flares compared to those receiving placebo in all four trials. Time to disease flare for participants receiving biologic DMARDs was statistically significantly longer (reported for abatacept and etanercept only). A greater proportion of those treated with biologic DMARDs achieved ACR Pedi responses of ≥ 30 and had inactive disease (reported for abatacept and tocilizumab only). Generally, the individual ACR Pedi core variables (reported for abatacept, etanercept and tocilizumab) were improved by biologic DMARDs when compared to placebo, as were joint-related outcomes (reported for etanercept only) and pain in two out of three studies (etanercept and tocilizumab, not in abatacept). Not all studies reported a statistical comparison for

each of these outcomes. Three studies (adalimumab, etanercept and tocilizumab) reported mortality, with no treatment-related deaths. Differences between trial-arms in HRQoL reported in one study (abatacept) were not statistically significant. The proportions of adverse events (AE) and serious adverse events (SAEs) were generally similar between the treatment groups. One study (tocilizumab) reported sub-group data, albeit without statistical comparisons between treatment groups. None of the studies reported data for outcomes such as corticosteroid dose reduction, extra-articular manifestations (such as uveitis), height or weight for the randomised withdrawal phase of the trials.

An adjusted indirect comparison suggests that the four biologic DMARDs appear to be similar in terms of disease flare and ACR Pedi 50 and 70 responses, with wide confidence intervals and clinical heterogeneity between the trials.

There were differences across the trials in the eligibility criteria for the OLE phase, and in how the results are reported. In some studies it was not possible to differentiate between participants treated continuously with a biologic DMARD (i.e. from open-label lead-in and randomised withdrawal phase) and those who received placebo before being offered a biologic DMARD at entry to the OLE. Generally, patients' ACR responses remained constant over time or even increased after the double-blind phase. Limited data for adalimumab and tocilizumab reported in abstracts at week 104 appears to support the positive effect of these drugs on growth, but the use of different outcome measures prevents a comparison between the drugs.

In addition to the four RCTs, seven relevant ongoing trials were identified and summarised in this report (three investigating adalimumab, and four investigating etanercept).

There is limited evidence for the clinical-effectiveness of biologic DMARDs in specific JIA disease subtypes. An observational study (CLIPPER) assessing the safety and efficacy of etanercept in children and adolescents with extended oligoarticular JIA (EO), ERA and PA found variations in response to treatment between JIA disease sub-types

■. By week 96 similar ACR Pedi 90 (62% to 72%) and ACR Pedi 100 (51% to 60%) responses were achieved by participants with different JIA subtypes, and proportions with inactive disease varied between 29% (ERA and PA) and 37% (EO).

Evidence from observational studies suggests that biologic DMARDs can improve uveitis symptoms, such as intraocular inflammation, in children with JIA. Adalimumab appears to be more effective than etanercept in improving uveitis.

Four pharmaceutical companies made submissions in support of their drugs to NICE. Only one of these (Pfizer, etanercept) provided a systematic review of clinical-effectiveness. This was judged to be of good standard. None of the submissions included any relevant RCTs that were additional to those identified in this assessment report.

Cost-effectiveness

The systematic review of published economic evaluations identified 388 potentially relevant publications. Of these, four studies (described in five publications) met the inclusion criteria. The studies were conducted in UK, the Netherlands, Canada and Russia. There were two cost-utility-studies, one cost-effectiveness study and one cost-consequence study. The studies were assessed for quality and generalisability to the UK but all contained limitations in the methodological quality or generalisability to the UK NHS. The study conducted in the UK was the assessment report for the previous NICE appraisal for etanercept in children with JIA (NICE TA35). The systematic review of HRQoL identified two studies reporting health-state utility values for patients in JIA.

In terms of the company submissions to NICE, Roche (manufacturer of tocilizumab), constructed a Markov state-transition model that compared tocilizumab to adalimumab for children with JIA. The base case results conclude that tocilizumab is of similar effectiveness and is less expensive than adalimumab. Two companies, BMS (manufacturer of abatacept) and Pfizer (manufacturer of etanercept) assumed that the biologic DMARDs were equivalent in clinical-effectiveness. They submitted cost analyses to compare the biologic DMARDs. BMS concluded that abatacept was the least costly treatment option and tocilizumab was slightly cheaper than adalimumab. Pfizer concluded that for most ages, etanercept is the biologic treatment with the lowest acquisition cost compared to tocilizumab and adalimumab. AbbVie (manufacturer of adalimumab) did not submit an economic analysis and cited a number of methodological limitations to producing an economic model. Two companies, Roche (tocilizumab) and BMS (abatacept) submitted a confidential patient access scheme discount.

The independent model developed for this assessment report modelled one line of biologic treatment for the comparison of adalimumab, etanercept and tocilizumab versus methotrexate. From this model, the incremental cost-effectiveness ratio (ICER) for adalimumab, etanercept and tocilizumab versus methotrexate is estimated at £38,127, £32,526 and £38,656 per QALY gained, respectively, using the list price drug acquisition costs. Abatacept is licensed for second-line biologic therapy after discontinuation of an anti-TNF. Abatacept was compared with methotrexate as a second-line biologic treatment, following etanercept as the first-line biologic. In this analysis, abatacept had an ICER of £39,536 per QALY gained.

The model results are most sensitive to changes to the HRQoL utility values. The changes to the clinical-effectiveness parameters, such as treatment discontinuation and disease flare had minimal effect on the model results. The differences in cost-effectiveness of the biologic DMARDs are primarily the effect of the differences in the drug acquisition cost.

Discussion

Biologic DMARDs (plus methotrexate where indicated) are superior to placebo (plus methotrexate where indicated) treatment across a number of outcome measures in children with JIA and an insufficient response to previous treatment. Due to the withdrawal trial design results of the double-blind phase are only applicable to patients who have already achieved an initial (low) degree of benefit from a biologic DMARD. Long-term treatment effectiveness in terms of ACR Pedi response appears to be sustained for all four included RCTs and the occurrence of AEs generally similar between biologic DMARD and placebo-treated patients. SAEs seem to be uncommon and the long-term safety profile of the biologic DMARDs is relatively favourable. An incremental analysis and the costs and health benefits of the four biologic DMARDs was not presented, as the DMARDs were similar in effects and costs.

There was insufficient evidence available for all input parameters to permit a cost-effectiveness subgroup analysis for each of the respective types of JIA within the scope of the appraisal. The modelled patient population is people with JIA, though it is primarily relevant to those with polyarticular course JIA.

The strengths of this assessment include use of standard methods for evidence synthesis and economic modelling, and the transparent reporting of the scope and methods a priori in a published protocol. Limitations include lack of head-to-head trial comparisons of biologic DMARDs, necessitating an indirect comparison, and lack of available data to inform the economic evaluation, particularly HRQoL utility estimates (which were the most influential parameters of cost-effectiveness), long-term discontinuation rates, and the long-term impact of treatment on disease progression. Assumptions have been made where possible based on best available evidence and expert opinion.

Conclusions

Implications for service provision

Given that biologic DMARDs are currently used in the treatment of JIA any recommendation supporting their use is unlikely to have significant implications for service provision (e.g. in terms of changes to infrastructure, staff training).

Suggested research priorities

Randomised head-to-head comparisons of biologic DMARDs are necessary to establish comparative effectiveness. Trials should be sufficiently powered, with long-term follow-up of safety and efficacy, and should include an economic evaluation to assess cost-effectiveness.

Word count: 2395

Plain English Summary

Juvenile idiopathic arthritis (JIA) is a term for all forms of arthritis of unknown cause that start before the age of 16 years and persist for more than six weeks. Patients suffer from joint pain, swelling and limitation of movement caused by inflammation surrounding affected joints. The joint damage caused by the inflammation can lead to disability and growth restriction. Treatment includes disease modifying anti-rheumatic drugs, commonly abbreviated to DMARDs and methotrexate is the most used DMARD in the UK. The preferred treatment now includes the use of newer drugs termed biologic DMARDs. Using a systematic approach, we identified the most up-to-date evidence for four biologic DMARDs called abatacept, adalimumab, etanercept and tocilizumab, which are used to treat different forms of JIA (apart from systemic or oligoarthritis JIA). The evidence was assessed using recognised methods to evaluate whether treatment with a biologic DMARD (with or without methotrexate) benefits patients with JIA, taking into account treatment costs and health.

Four studies were identified, one for each drug. Each compared the biologic DMARD against a placebo treatment and, with the exception of the etanercept trial, the majority of patients also received methotrexate. To enter the study, patients had to have a positive response to the relevant biologic DMARD in a starting phase before they could be randomised to either continue biologic DMARD or a placebo treatment. Those who continued treatment with a biologic DMARD experienced significantly fewer disease flare ups than those who were switched to placebo treatment and the abatacept and etanercept trials found these occurred later. Continued DMARD treatment also led to a greater response level measured by the American College of Rheumatology Pediatric (ACR Pedi) criteria with the abatacept and tocilizumab trials reporting more participants with inactive disease. No trials directly compared the drugs against each other so a statistical method was used to compare them indirectly. It suggested that the four biologic DMARDs are similarly effective in terms of disease flare, and ACR Pedi response levels 50 and 70. These results must be treated with caution because of differences between the trials and the patients. Generally, patients' ACR responses remained constant or even increased after the randomised phase. The proportions of adverse events and serious adverse events were generally fairly similar between the biologic DMARD and placebo groups.

To assess cost-effectiveness, biologic DMARDs were compared to methotrexate treatment, as there was insufficient evidence to compare them to each other. Costs and health benefits appear to be similar for all the biologic DMARDs. Treatment of children and young people with biologic DMARDs may therefore be an effective therapy. However, due to the lack of evidence from direct comparisons between biologic DMARDs, a number of uncertainties remain.

Word count: 447 words

LIST OF ABBREVIATIONS

ABA	Abatacept
ACR Pedi 20, 30, 50, 70, 90 or 100	American College of Rheumatology Pediatric Response levels: 20, 30, 50, 70, 90 or 100
ADA	Adalimumab
AE	Adverse events
AIC	Academic in confidence
BMS	Bristol-Myers Squibb
BNF	British National Formulary
BNFC	British National Formulary for Children
BSA	Body surface area
BSPAR	British Society for Paediatric and Adolescent Rheumatology
BW	Body weight
CHAQ	Childhood Health Assessment Questionnaire
CHQ	Child Health Questionnaire
CI	Confidence Interval
CIC	Commercial in confidence
CRP	C reactive protein
CS	Company submission
DMARDs	Disease modifying anti-rheumatic drugs
EMA	European Medicines Agency
EO	Extended oligoarthritis
ERA	Enthesitis-related arthritis
ESR	Erythrocyte sedimentation rate
ETA	Etanercept
EOW	Every other week
GBP	Great British Pound
HAQ	Health Assessment Questionnaire
HRQoL	Health-related quality of life
HTA	Health Technology Assessment
HUI	Health Utilities Index
ICER	Incremental cost-effectiveness ratio
ILAR	International League of Associations for Rheumatology
ITT	Intention to treat
IQR	Interquartile range
JADAS	Juvenile Arthritis Disease Activity Score
JIA	Juvenile idiopathic arthritis
LOCF	Last Observation Carried Forward
LOM	Limitation of motion
MTA	Multiple Technology Assessment
MTX	Methotrexate
NICE	National Institute for Health and Care Excellence
NR	Not reported
NSAIDs	Non-steroidal anti-inflammatory drugs
OLE	Open-label extension
PBO	Placebo
PA	Psoriatic arthritis
PAS	Patient access scheme
PedsQL	Pediatric Quality of Life
PGA	Physician global assessment of disease activity
PSA	Probabilistic sensitivity analysis
PSSRU	Personal Social Services Research Unit

QoL	Quality of life
QALY	Quality-adjusted life year
RCT	Randomised controlled trial
RF-ve	Rheumatoid factor positive
RF+ve	Rheumatoid factor negative
RR	Relative risk
SAE	Serious adverse events
SDS	Standard deviation scores
SF36	Short Form (36) Health Survey
SHTAC	Southampton Health Technology Assessments Centre
SPC	Summary of Product Characteristics
TCZ	Tocilizumab
VAS	Visual analogue scale

1 BACKGROUND

1.1 Description of underlying health problem

Juvenile idiopathic arthritis (JIA) is an umbrella term that encompasses all forms of arthritis with onset before the age of 16 years and symptoms that persist for more than six weeks for which the cause is unknown.^{1,2} The role of infections (either bacterial or viral) in the development of JIA has been investigated but no unequivocal evidence to either support or rule out an association has been clearly demonstrated.³ The term JIA has been in use since 1995 and was proposed by the International League of Associations for Rheumatology (ILAR) committee to replace the older terms of ‘juvenile rheumatoid arthritis’ and ‘juvenile chronic arthritis’ which were the chief terms in use in the USA and in Europe respectively.⁴ JIA is characterised by joint pain, swelling and limitation of movement caused by inflammation of the synovial membrane of the affected joints. If untreated this inflammation causes a progressive erosive arthritis which can lead to disability and growth retardation.⁵ JIA is classified according to the Revised ILAR Criteria⁶ into seven subtypes: systemic arthritis, oligoarthritis (subcategories persistent and extended), polyarthritis - rheumatoid factor (RF) negative (RF-ve), polyarthritis - RF positive (RF+ve), psoriatic arthritis (PA), enthesitis-related arthritis (ERA), and undifferentiated arthritis (Table 1) and some forms of the disease are associated with extra-articular features such as uveitis (inflammation of the middle layer of the eye).

Table 1 JIA classification according to the revised ILAR Criteria

JIA classification ⁶ & features ⁷⁻¹²	Included in NICE appraisal scope?
<p>Oligoarthritis</p> <ul style="list-style-type: none"> • The most common type of JIA accounting for over 50% of JIA cases in the UK¹³ • Usually starts before six years of age and more common in girls than boys • Affects four or fewer joints in the first six months, most commonly one or both knees and/or ankles, which are swollen and may be painful • Regular checks for chronic anterior uveitis (painless eye inflammation) required <p>The ILAR classification recognises two subcategories</p> <ul style="list-style-type: none"> • Persistent oligoarthritis: affecting four or fewer joints throughout the disease course, accounts for about 48% of JIA cases in the UK¹³ • Extended oligoarthritis: affecting a total of more than four joints after the first six months of disease, accounts for about 	<p>No</p> <p>Yes</p>

6% of JIA cases in the UK ¹³	
<p>Polyarthritis - RF+ve: accounts for about 4% of cases in the UK¹³</p> <p>Polyarthritis - RF-ve: accounts for about 21% of cases in the UK¹³</p> <ul style="list-style-type: none"> • Polyarthritis is the second most common type of JIA affecting about one in four children with arthritis • Usually starts either before seven years of age or later in childhood • Causes painful swelling of five or more joints in multiple sites. The same joints on both sides of the body will often be affected. • RF negative is the most common form. RF positive subtype is more often seen in teenage girls. • Associated with chronic uveitis (painless eye inflammation) 	Yes (all forms)
<p>Enthesitis-related arthritis (ERA)</p> <ul style="list-style-type: none"> • Accounts for about 6% of JIA cases in the UK¹³ • Affects the entheses (sites where tendon attaches to bone) often of lower limb and pelvic joints as well as the joints themselves (spine or peripheral joints). • Can affect girls and boys although teenage onset disease mainly affects boys. • Associated with acute uveitis (red painful eye) 	Yes
<p>Psoriatic arthritis (PA)</p> <ul style="list-style-type: none"> • Accounts for about 7% of JIA cases in the UK¹³ • Joint pain associated with the skin condition psoriasis (although the typical rash of psoriasis may not occur until many years after the onset of arthritis) or with a family history of psoriasis. Typically affects finger and toe joints. • Usually starts around six years of age and about twice as common in girls as in boys. • Chronic anterior uveitis is fairly common. 	Yes
<p>Systemic arthritis</p> <ul style="list-style-type: none"> • Accounts for about 6% of JIA cases in the UK¹³ • Usually starts before five years of age and affects boys and girls about equally • General illness with fever, tiredness, rash, loss of appetite and weight loss as well as joint pain. May also have enlarged glands, spleen and liver. More rarely pericarditis (inflammation of sac surrounding the heart) 	Not active systemic onset JIA alone. Those who go on to have a form of JIA that is included (e.g. polyarthritis) do match the remit.
Undifferentiated arthritis	

- | | |
|--|--|
| <ul style="list-style-type: none"> • JIA that does not fit into any of the above categories or that has features of more than one. Accounts for about 4% of JIA in the UK.¹³ | |
|--|--|

At onset the particular sub-type of JIA will be diagnosed according to the presenting features corresponding to one of the seven ILAR categories. As JIA progresses more joints may become affected. For some, where JIA was classified at onset as oligoarthritis, problems with five or more joints develop after six months and the JIA type is then described as extended oligoarthritis. Similarly the term polyarthritis also applies to patients who at a particular point in time six months or more after the onset of disease (JIA of any onset type) have five or more active joints. In this case they are said to have polyarticular course JIA. The concept of polyarticular course JIA has been used for clinical trials and can typically include RF positive and RF negative polyarthritis, extended oligoarthritis, ERA, PA and undifferentiated arthritis. Systemic JIA may also be included in the definition of polyarticular course JIA providing there have been no active systemic symptoms during the previous six months.¹⁴

Severity of disease and long term outcome is variable both between different JIA subtypes and between different individuals with the same JIA subtype (Table 2). Analyses of historical cohorts of JIA patients (comprising a mix of JIA sub-types) have shown that more than 50% of patients continued to have active disease as long as 17 years after disease onset and such patients would require treatment into adulthood.^{15;16} However, it should be noted that in historical studies the patients, particularly at disease onset, were unlikely to have been treated with methotrexate or biologic disease modifying anti-rheumatic drugs (DMARDs), which were not available. Even when biologic DMARDs became available they may not have been widely used. Consequently for all types of JIA outcomes in general are likely to have improved due to more widespread use of the newer treatment strategies, particularly early in the disease course. Nevertheless a third or more of children will still require treatment for JIA in adult life. JIA which persists into adulthood is distinct from adulthood rheumatoid arthritis and should not be considered similar.

A recent (2014) systematic review of the prevalence and incidence of JIA in Europe¹⁷ found that rates varied greatly among published studies. Incidence rates ranged from 1.6 to 23 per 100,000 (33 studies) and prevalence rates from 3.8 to 400 per 100,000 (29 studies). The estimated annual incidence of JIA in England 1989-1991 was 11 per 100,000.¹⁸ Prevalence in the UK has not been estimated since 1959 when a figure of 65 per 100,000 was reported.¹⁸ The Children's Chronic Arthritis Association website states that annual incidence is approximately one per 10,000 (i.e. 10 per 100,000) and prevalence is about one per 1,000 (i.e. 1000 per 1,000,000).¹⁹

Table 2 Long term outcomes for different sub-types of JIA

Long-term outcome ⁷⁻¹²
<p>Persistent oligoarthritis</p> <ul style="list-style-type: none"> • Often mild and may resolve with little or no lasting damage to joints, has the best outlook of all the types of JIA. • Approximately half of children will have symptoms for at least ten years, a third or more of children will have arthritis continuing into adulthood. • Chronic anterior uveitis may cause blindness or visual loss if not detected and treated early enough.
<p>Extended oligoarthritis</p> <ul style="list-style-type: none"> • Causes damage to joints so early treatment to minimise this is needed. • Can be destructive and disabling. • Approximately half of children will have symptoms for at least ten years, a third or more of children will have arthritis continuing into adulthood. • Chronic anterior uveitis may cause blindness or visual loss if not detected and treated early enough.
<p>Polyarthritis</p> <ul style="list-style-type: none"> • Approximately half of children will have symptoms for at least ten years and at least one third of children will have arthritis continuing into adulthood (most likely with the RF positive type, which is more severe and can require more aggressive treatment). • Joints may become damaged if inflammation is not controlled, leading to potential need for joint replacement or serious disability.
<p>Enthesitis-related arthritis (ERA)</p> <ul style="list-style-type: none"> • May evolve to ankylosing spondylitis in the adult years (especially those with teenage onset) and may require long term disease modifying or biologic agents.
<p>Psoriatic arthritis (PA)</p> <p>Although there is not much long term data, disease course may be similar to chronic arthritis (either oligoarthritis or polyarthritis) and is likely to continue into adulthood.</p>
<p>Systemic-onset JIA</p> <ul style="list-style-type: none"> • A third of children will have one or two episodes that settle with treatment, a third will have relapses and need intermittent treatment, a third require ongoing treatment into adulthood and are at risk of joint damage.
<p>Undifferentiated arthritis</p> <p>Although there is not much long term data clinical advisors indicate that the long-term outcome is likely to depend on the predominant features of the arthritis and whether persistent oligoarthritis or polyarticular course arthritis.</p>

The sources of these data are not given, however the same data are available in the Interim Clinical Commissioning Policy Statement for biologic therapies for the treatment of JIA.²⁰ Based on the mid-2013 population estimates for those aged 17 years and under in England (approximately 11.5 million) and Wales (approximately 630,000)²¹ these incidence and prevalence values equate to an estimated

incidence of 1150 cases a year in England and 63 cases a year in Wales with an estimated 11,500 and 630 children overall in England and Wales respectively with JIA.

A 2012 oral conference presentation¹³ presented data from a multi-centre long-term prospective inception cohort study of children with newly diagnosed inflammatory arthritis [The Childhood Arthritis Prospective Study (CAPS)]. This provides information on JIA subtypes classified using the ILAR criteria for 1014 newly diagnosed children [Median disease duration: 5.2 months, interquartile range (IQR) 2.5 to 10.9]. Amongst this cohort extended oligoarthritis and polyarticular course JIA may be under-represented because median disease duration is less than six months. Nevertheless, the proportions of each JIA subtype are similar to those reported by an older study (2002)²² for a smaller group of children (n=521) as shown in Table 3.

Table 3 Proportions of different sub-types of JIA

JIA classification ⁶	Newly diagnosed children (n=1014) ¹³	From 17 centres within the UK (n=521) ²²	
Oligoarthritis:			
Persistent oligoarthritis	48.2% (502/1041)	30.1% (157/521)	
Extended oligoarthritis	5.5% (57/1041)	15.2% (79/521)	
Polyarthritis - RF+ve	3.6% (37/1041)	7.1% (37/521)	
Polyarthritis - RF-ve	20.6% (214/1041)	19.6% (102/521)	
Enthesitis-related arthritis	5.6% (58/1041)	6.5% (34/521)	
Psoriatic arthritis	7.0% (73/1041)	7.1% (37/521)	
Systemic arthritis	6.0% (62/1041)	14.4% (75/521)	
Undifferentiated arthritis	3.7% (38/1041)	not reported	
Not recorded	N/A	N/A	

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

In addition to the immediate impacts of joint pain, swelling and limitation of movement that characterise JIA there are longer term problems and other issues that may arise over time. Progressive joint damage can lead to permanent disability and eventually a need for joint replacement. A retrospective review of 154 adolescents (aged 16-21 years) found that 14% had undergone a joint operation with 30 separate surgeries (e.g. synovectomies, reconstructive finger or toe joint operations) having been undertaken including one hip replacement.²⁴ Growth impairment affects about 10-20% of patients with JIA (mainly those with systemic or polyarticular JIA and who require high doses of glucocorticoids)²⁵ and decreased bone mass which can lead to the development of osteoporosis is also a recognised problem.²⁶

JIA is associated with a range of extra-articular manifestations, including uveitis, inflammatory bowel disease, and psoriasis. Uveitis commonly occurs in children with oligoarthritis and is less common in other subtypes of JIA. It is characterised by inflammation of the middle layer of the eye, the uveal tract. In severe cases which do not respond to treatment uveitis can be associated with complications such as cataract, glaucoma, and macular oedema, and can lead to sight impairment and blindness. Inflammatory bowel disease (e.g. Crohn's disease and ulcerative colitis) is typically associated with ERA, whilst psoriasis is associated with PA.

The incidence of childhood uveitis in North America and Europe is estimated to be 4.3 to 6 per 100,000 children and the prevalence at 30 per 100,000 children.²⁷ Between 20-25% of uveitis cases in children are associated with JIA. The prevalence of uveitis in JIA is between 8-30%, but in children with oligoarticular onset JIA it may be between 45-57%.²⁸ Uveitis in patients with JIA commonly occurs with the early onset of arthritis (mean age of onset 3-5 years). Presentation in younger children may be delayed due to their inability to articulate symptoms. Screening for uveitis has therefore been implemented for children with JIA in England.²⁸ Complications are present in between 30-50% of children with JIA with uveitis at diagnosis. Fifty to seventy per cent of children with severe uveitis will develop visual impairment.²⁹

A recent systematic review of qualitative studies that explored the experiences of children living with JIA highlighted the profound effect that JIA has on children's lives. In particular, pain was a constant reminder of their disease and limited their ability to participate in normal life including social events and schooling. Their physical limitations meant that they had to look for alternative activities and

potential career options which they would be able to pursue. Many children and adolescents felt misunderstood and some kept their illness a secret from their peers and others.³⁰

1.2 Measures of response to treatment and definition of remission

The aim of JIA treatment is to achieve clinical remission (complete absence of active disease). Aggressive early treatment aims to control inflammation and thus symptoms (e.g. joint pain); to decrease the number of actively affected joints in order to prevent joint damage, loss of function and disability; and to maintain or improve quality of life. Response to treatment is assessed in clinical trials by a validated core set of variables that were adopted by the American College of Rheumatology (ACR) in 1997. This defining of response is now known as the ACR Pediatric definition of improvement.³¹ The lowest level of improvement is known as ACR 30 (or ACR Pedi 30). The ACR Pediatric 30 core variables are:

- 1) physician global assessment of disease activity using a visual analogue scale (VAS) (range from best score 0 to worst score 100mm (although in some studies reported as 0 to 10cm)
- 2) patient or parent global assessment of overall well-being using a VAS (range 0-100mm, 0 is the best score)
- 3) functional ability as measured by the patient or parent using the Childhood Health Assessment Questionnaire (CHAQ, range 0-3, 0 is the best score)
- 4) number of joints with active arthritis
- 5) number of joints with limited range of motion
- 6) laboratory marker of inflammation (erythrocyte sedimentation rate (ESR) or C reactive protein (CRP) level)

Response to ACR Pedi 30 level is defined as an improvement in three of any six of the core variables by at least 30%, and no more than one of the remaining variables worsened by more than 30%. In addition to the ACR Pediatric 30, higher levels of response can also be defined - the ACR Pediatric 50, 70, 90 and 100 levels of response require at least 50%, 70%, 90% or 100% improvement respectively in at least three of any six of the core set variables, with no more than one of the remaining variables worsening by more than 30%.^{14;32} It should be noted that according to expert advice, ACR Pedi 30 is no longer accepted as a response but considered a non- or inadequate response, with response levels of at least ACR Pedi 50 or 70 looked for from a drug intervention.

More recently in 2009 the Juvenile Arthritis Disease Activity Score (JADAS) was proposed and validated.³³ The JADAS is a composite score that can be quickly calculated because it is the arithmetic sum of the scores from the following four individual component measures:

- 1) physician global assessment of disease activity, measured on a 10 cm VAS (range 0 = no activity and 10 = maximum activity)

- 2) parent/patient global assessment of well-being, measured on a 10 cm VAS (range 0 = very well and 10 = very poor)
- 3) count of joints with active disease
- 4) ESR

The component measures are also measures used in the ACR Pediatric definition of improvement.³¹

The count of joints with active disease in the JADAS is primarily based on a 27-reduced joint count (JADAS-27, total score range 0-57) although scores based on a full 71 joint count (JADAS-71, total score range 0-101) and a ten joint count (JADAS-10, total score range 0-40) have also been validated.³³ Further studies have shown that a 3-item JADAS that does not use ESR data is also a robust measure^{34,35} which is of particular benefit for children who do not need to provide a blood sample for routine medication monitoring. As the JADAS has become more widely used further proposals have been made that would define low, medium and high disease activity^{35,36} and define improvement.³⁷ With these definitions in place the future management goal would be to achieve minimal disease activity (MDA) for all children with JIA.³⁸

Preliminary criteria to define clinical remission in oligoarticular (persistent and extended), RF positive and RF negative polyarticular, and systemic JIA have also been developed.³⁹ Two levels of clinical remission have been proposed, clinical remission on medication and clinical medication off medication. The criteria for both types of clinical remission are based on achieving inactive disease, which is defined as:

- no joints with active arthritis
- no fever, rash, serositis, splenomegaly, or generalised lymphadenopathy attributable to JIA
- no active uveitis
- normal ESR or CRP (or both normal if both tested)
- physician's global assessment of disease activity indicates no disease activity

Clinical remission on medication is then proposed to have been achieved if all the criteria for inactive disease have been met for a minimum of six continuous months while the patient is on medication.

Clinical remission off medication is proposed to have been achieved if all the criteria for inactive disease have been met for a minimum of 12 continuous months while the patient is off all anti-arthritis and anti-uveitis medications.

Since the original publication of the preliminary criteria to define clinical remission,³⁹ validation of the criteria for defining clinical inactive disease in oligoarticular (persistent and extended),

polyarticular (RF positive and RF negative) and systemic JIA has been undertaken. This has led to three changes: the addition of a definition for no active uveitis [as defined by the Standardization of Uveitis Nomenclature (SUN) Working Group]; clarification that the ESR or CRP level should be within the normal limits in the laboratory where tested or, if elevated, not attributable to JIA; and one additional criterion (duration of morning stiffness of 15 minutes or less).⁴⁰

In addition to definitions of response to treatment and clinical remission some publications also report on the outcome of disease flare (periods when symptoms worsen). A preliminary definition based on the ACR Pediatric 30 core response variables was obtained from single small study (n=51).⁴¹ This preliminary definition was worsening in any 2/6 core response variables by 40% or more without concomitant improvement of more than one of the remaining core response variables by 30% or more. However, other studies have used different flare definitions e.g. a worsening of $\geq 30\%$ in three of six ACR Pediatric 30 variables.⁴²

1.3 Current service provision

There is currently no NICE clinical guideline on the treatment of JIA however there are two pieces of NICE guidance:

- Guidance on the use of etanercept for the treatment of juvenile idiopathic arthritis (NICE TA35) 2002⁴³ (this assessment report will inform an update of this guidance)
- Tocilizumab for the treatment of systemic juvenile idiopathic arthritis (NICE TA238) 2011⁴⁴ (active systemic JIA is not included within this assessment report)

There are currently two interim commissioning statements: Biologic Therapies for the treatment of Juvenile Idiopathic Arthritis (JIA) and the draft NHS Clinical Commissioning Policy for severe refractory uveitis in paediatric patients. The first interim clinical commissioning policy statement has been published (January 2015) by NHS England Clinical Reference Group for Paediatric Medicine²⁰ in the absence of NICE guidance for other biologic DMARDs and to cover more recent changes to the licenced indications to etanercept and is being consulted on. The purpose of the interim policy statement is to provide guidance for the use of biologic DMARDs in patients with JIA until the planned NICE guidance is published. The statement has a broader remit than the planned NICE guidance [Arthritis (juvenile idiopathic) - abatacept, adalimumab, etanercept and tocilizumab (including review of TA35)] as it includes all biologic DMARDs and all types of JIA (i.e. including persistent oligoarticular JIA and systemic JIA which are not included in the NICE scope for the planned guidance). A summary of the key features of the drug treatment pathway is provided in Table 4.

Clinical advice to the authors of this assessment report (hereafter referred to as the assessment group) suggested that the interim statement largely reflects current practice. However, it was acknowledged that there would still be some variability across the country due to differences in interpretation and limitation on access and prescribing.

Table 4 Overview of the drug treatment pathway for JIA

When/why & who	What	Notes
At diagnosis to induce disease remission, all patients	Corticosteroids: <i>Either</i> intra-articular to all affected joints	In patients with mild disease limited to <5 joints intra-articular steroids may induce remission of >6 months, particularly if the long-acting corticosteroid triamcinolone hexacetonide is used.
	<i>Or</i> systemic, preferably intravenous (due to the side effects (e.g. effect on growth or increased risk of osteoporosis) of oral corticosteroids)	Patients with more severe disease may need intravenous steroids to induce remission although intra-articular steroids are used in some patients as an alternative.
To maintain remission, patients with arthritis affecting ≥5 joints or arthritis severely affecting crucial joints (e.g. spine, ankles, hips, wrists)	MTX	This accounts for around half of all children who develop JIA. Effective in reducing the amount and severity of arthritis but only induces complete remission in 30-50% of patients.
When JIA remains active despite optimal MTX dosing <u>OR</u> when patient is intolerant of MTX	Biologic DMARD (many given in co-administration with MTX to optimise their effect)	Estimated that a third of all children who start treatment with MTX need to progress to a biologic DMARD.

MTX - methotrexate

According to the second interim clinical commissioning policy statement (The draft NHS Clinical Commissioning Policy for severe refractory uveitis in paediatric patients²⁹), patients with JIA-associated uveitis may be managed initially with topical corticosteroids, or systemic corticosteroids if required. In more severe cases a DMARD can be used, with methotrexate a standard treatment. If disease is not controlled with DMARDs the next line of treatment is use of a TNF inhibitor (tumour necrosis factor alpha is shown to be implicated in the pathogenesis of uveitis). TNF inhibitors include etanercept, adalimumab, infliximab, golimumab, and certolizumab, but the latter two may not be easily available in the UK, and only etanercept and adalimumab are licensed for the treatment of JIA in children in Europe. For severe refractory uveitis in paediatric patients the draft NHS Clinical Commissioning Policy states that etanercept is not suitable for use in JIA patients with uveitis, or uveitis not associated with JIA.²⁹ Adalimumab is recommended where methotrexate does not control symptoms, with infliximab used in patients in whom adalimumab is not tolerated, or not effective.²⁹

1.4 Description of technology under assessment

Four biologic DMARDs are within the scope of the NICE appraisal and are therefore included in this assessment report: abatacept, adalimumab, etanercept and tocilizumab. The licenced indication

differs across these interventions (e.g. in terms of the age range of children and young people eligible for treatment, the previous treatment that they should have received and the sub-type of JIA) as summarised in Table 5. The Interim Clinical Commissioning Policy Statement: Biologic Therapies for the treatment of JIA²⁰ provides a pragmatic estimate of 950 children with JIA in England who are currently receiving a biologic DMARD. This estimate is based on current data from the biologics databases in the UK which indicate that in England alone 890 children are receiving a biologic DMARD for JIA (most of which are NICE approved biologic DMARDs). Clinical advice to the assessment group suggested that this figure may be an underestimate. An alternative estimate of 1500 was suggested by one clinician.

Table 5 Summary of licensed indications of the biologic DMARDs under consideration in this assessment

Drug (chief mode of action)	Polyarthritis (polyarticular)			Enthesitis (ERA)	Psoriatic (PSA)	Systemic onset
	RF+ poly	RF- poly	Ext oligo			
Abatacept (prevents t-cell activation)	Yes	Yes	Yes	-	-	-
	With MTX. Patients 6 years & over with insufficient response to DMARDs including at least 1 TNF inhibitor					
Adalimumab (TNF-inhibitor)	Yes	Yes	Yes	Yes	-	-
	With MTX unless not tolerated/not appropriate. Patients 2 years & over, with inadequate response to 1 or more DMARDs.			Patients 6 years and over with inadequate response to or intolerant of conventional therapy		
Etanercept (TNF-inhibitor)	Yes	Yes	Yes	Yes	Yes	-
	Children and adolescents from age of 2 years with inadequate response to or intolerant of MTX			Adolescents from age of 12 years with inadequate response or intolerant of conventional therapy	Adolescents from age of 12 years with inadequate response or intolerant of MTX	
Tocilizumab (IL-6 inhibitor)	Yes	Yes	Yes	-	-	Yes
	With MTX unless not appropriate. Patients 2 years & over, have responded inadequately to previous treatment with MTX					With MTX unless not appropriate. Patients 2 years & over with inadequate response to NSAIDs and systemic corticosteroids

Note: Patients with active systemic onset JIA alone will not be addressed in this MTA. Patients with systemic onset JIA and a form of JIA that is included in the MTA (such as polyarthritis) will be addressed in this MTA.

Where systemic onset, enthesitis and psoriatic arthritis go on to have a polyarticular course they could be interpreted as falling within the marketing authorisations for all 4 of the drugs.

NSAIDS = Non-steroidal anti-inflammatory drugs

As noted earlier in section 1.3 the interim clinical commissioning policy indicates that the initial biologic DMARDs to be considered for use would be a TNF-inhibitor, which for the purposes of this assessment would be either adalimumab or etanercept (however etanercept is not suitable for use in JIA patients with uveitis). If a treatment switch was required the second-line biologic DMARD would initially be the alternative TNF-inhibitor (i.e. switch from adalimumab to etanercept or vice-versa). If a further switch was necessary the 3rd line biologic would either be abatacept or tocilizumab and the final switch possible would be to change abatacept to tocilizumab or vice-versa. However, in terms of the marketing authorisations the licence for abatacept indicates that there should have been a prior insufficient response to at least one TNF-inhibitor. There is no such indication in the licence for tocilizumab.

The Summary of Product Characteristics (SPCs) for each biologic DMARD should be consulted for the specific contraindications, special warnings and precautions for use however, there are some aspects that are common to all biologic DMARDs which are summarised here.⁴⁵⁻⁴⁸ These drugs block aspects of normal immune system signalling and consequently it is recommended that all patients receiving a biologic DMARD carry an alert card to indicate that they are at increased risk of developing a serious infection. Patients are not only at risk of typical bacterial and viral infections but also opportunistic infections including invasive fungal infections. Existing latent infections (e.g. latent hepatitis-B, latent tuberculosis) could potentially reactivate. Consequently, if patients have an existing infection, treatment with a biologic DMARD is not recommended until the infection is treated. Patients should be screened for latent infections and childhood vaccinations be brought up to date prior to beginning therapy with a biologic DMARD.

The SPCs for each of the four biologic DMARDs included in the review do not explicitly specify licenced upper age limits for treatment. Clinical advisors have indicated that if adolescents are responding to treatment then this should be continued into adulthood as required. Furthermore, some JIA patients may need to re-start a biologic DMARD in adulthood and some JIA patients may require a biologic DMARD for the first time in adulthood.

Abatacept

Abatacept (Orencia®, Bristol-Myers Squibb) in combination with methotrexate is indicated for the treatment of moderate to severe active polyarticular JIA in paediatric patients six years of age and older, who have had an insufficient response to other DMARDs including at least one TNF inhibitor.⁴⁵

Abatacept is a fusion protein produced by recombinant DNA technology in Chinese hamster ovary cells. It inhibits T-cell activation by specifically binding to CD80 and CD86 thereby selectively

inhibiting a costimulatory pathway that is required for full activation of T lymphocytes.^{45;49} Through this mechanism, abatacept modulates the downstream T lymphocyte-dependent antibody responses and inflammation that cause the symptoms of JIA.

Treatment should be initiated and supervised by specialist physicians experienced in the diagnosis and treatment of JIA at the appropriate dosage as indicated in Table 6. Abatacept is not recommended in combination with TNF-inhibitors.⁴⁵

Table 6 Dosing regimen for abatacept

Mode of administration & cost	Dose (child 6-17 years)		Notes
Intravenous infusion given during a period of 30 minutes Cost: powder for reconstitution, net price 250-mg vial = £302.40	Body-weight < 75 kg	10 mg/kg, repeated 2 weeks and 4 weeks after initial infusion, then every 4 weeks	Review treatment if no response within 6 months. Dosing for patients 75 kg and over follows the adult dosing regimen.
	Body-weight 75-100 kg	750 mg, repeated 2 weeks and 4 weeks after initial infusion, then every 4 weeks	
	Body-weight > 100 kg	1 g, repeated 2 weeks and 4 weeks after initial infusion, then every 4 weeks	

Adalimumab

Adalimumab (Humira®, AbbVie) in combination with methotrexate is indicated for the treatment of active polyarticular JIA in patients from the age of two years who have had an inadequate response to one or more DMARDs. Adalimumab can be given as monotherapy in the case of intolerance to methotrexate, or when continued methotrexate treatment is inappropriate. Adalimumab is also indicated for the treatment of active ERA in patients six years of age and older, who have had an inadequate response to, or who are intolerant of, conventional therapy.⁴⁶

Adalimumab is a fully human monoclonal antibody drug initially tested as a treatment for rheumatoid arthritis (hence the trade name Humira - **H**uman **M**onoclonal antibody **I**n **R**heumatoid **A**rthritis). It binds specifically to the inflammatory cytokine TNF thereby neutralising its biological function⁴⁶ and modifying the inflammatory disease process. The European Medicines Agency (EMA) therapeutic indication for adalimumab was extended to the treatment of JIA in July 2008.

Treatment should be initiated and supervised by specialist physicians experienced in the diagnosis and treatment of JIA at the appropriate dosage as indicated in Table 7.⁴⁶ The concomitant administration

of Adalimumab with other biologic DMARDs (e.g. anakinra and abatacept) or other TNF-antagonists is not recommended.⁴⁶

Table 7 Dosing regimen for adalimumab

Mode of administration & cost	Dose for polyarticular JIA		Notes
Subcutaneous injection given every other week (volume for injection is selected from a chart based on patient height and weight) Cost: net price 40-mg prefilled pen or prefilled syringe = £352.14; 40 mg/0.8-mL vial = £352.14.	Patients aged 2 to < 4 years	24mg/m ² body surface area up to a maximum single dose of 20mg	A clinical response is usually achieved within 12 weeks of treatment. Continued therapy should be carefully reconsidered in a patient not responding within this time period.
	Patients aged 4 to 12 years	24mg/m ² body surface area up to a maximum single dose of 40mg adalimumab	
	Patients 13 years and older	40mg administered every other week regardless of body surface area	
	Dose for ERA		Notes
	Patients 6 years and older	24mg/m ² body surface area up to a maximum single dose of 40mg	No indication for stopping treatment is provided.

Etanercept

Etanercept (Enbrel®, Pfizer) is a fully humanised soluble TNF receptor fusion protein produced by recombinant DNA technology in Chinese hamster ovary cells. It is a dimer with two copies of the extracellular domain of TNF receptor (p75) linked with the Fc component of human IgG1, binding to TNFa.⁵⁰ The mechanism of action of etanercept is thought to be its competitive inhibition of TNF binding to cell surface TNFR, preventing TNF-mediated cellular responses by rendering TNF biologically inactive. Etanercept may also modulate biologic responses controlled by additional downstream molecules (e.g. cytokines, adhesion molecules, or proteinases) that are induced or regulated by TNF.⁴⁷ The EMA therapeutic indication for etanercept in the treatment of JIA was extended in July 2012 to include:

- treatment of polyarthritis (RF+ve or RF-ve) and extended oligoarthritis in children and adolescents from aged ≥ 2 years who have had an inadequate response to, or who have proved intolerant of, methotrexate.
- treatment of PA in adolescents from aged ≥ 12 years who have had an inadequate response to, or who have proved intolerant of, methotrexate.

- treatment of ERA in adolescents from aged ≥ 12 years who have had an inadequate response to, or who have proved intolerant of, conventional therapy.

The age for treating polyarticular disease has been reduced from four to two years of age and the upper age limit of 17 years has been removed.

Treatment should be initiated and supervised by specialist physicians experienced in the diagnosis and treatment of JIA at the appropriate dosage as indicated in Table 8. The combined use of etanercept and anakinra or etanercept and abatacept is not recommended.

Table 8 Dosing regimen for etanercept

Mode of administration & cost	Dose for JIA	Notes
Subcutaneous injection Cost: net price 10-mg vial (with solvent) = £35.75; 25-mg vial (with solvent) = £89.38; 25-mg prefilled syringe = £89.38; 50-mg prefilled pen or prefilled syringe = £178.75.	0.4 mg/kg (up to a maximum of 25mg per dose) given twice weekly with an interval of 3-4 days between doses OR 0.8 mg/kg (up to a maximum of 50 mg per dose) given once weekly.	Consider discontinuation in patients who show no response after 4 months.

Tocilizumab

Tocilizumab (RoActemra®, Roche) in combination with methotrexate is indicated for the treatment of juvenile idiopathic polyarthritis (RF+ve or RF-ve and extended oligoarthritis) in patients two years of age and older who have responded inadequately to previous therapy with methotrexate. When the patient is intolerant to methotrexate or where continued treatment with methotrexate is inappropriate tocilizumab can be given as monotherapy.⁴⁸ Tocilizumab is also indicated for the treatment of active systemic JIA but this indication is not included within the current NICE appraisal.

Tocilizumab is a humanised, monoclonal, antihuman interleukin-6 receptor (IL-6R) antibody that binds to membrane and soluble IL-6R, inhibiting IL-6-mediated signalling - a key cytokine in rheumatoid arthritis pathogenesis.⁵¹ IL-6 is involved in causing inflammation and is found at high levels in patients with rheumatoid arthritis, systemic JIA and polyarticular JIA. By preventing IL-6 attaching to its receptors, tocilizumab reduces the inflammation and other symptoms of these diseases.⁴⁸ The EMA was granted a licence for tocilizumab in the treatment of JIA in May 2011.

Treatment should be initiated by healthcare professionals experienced in the diagnosis and treatment of JIA at the appropriate dosage as indicated in Table 9.

Table 9 Dosing regimen for tocilizumab

Mode of administration & cost	Dose for polyarticular JIA in patients above 2 years of age		Notes
Intravenous infusion over 1 hour Cost: net price 4 mL (80-mg vial) = £102.40, 10 mL (200-mg vial) = £256.00, 20 mL (400-mg vial) = £512.00.	Body-weight < 30 kg	10 mg/kg once every 4 weeks	Dose interruptions (including discontinuation) are recommended for liver enzyme abnormalities, low absolute neutrophil count and low platelet count according to the tables provided in the SPC. Clinical improvement is expected within 12 weeks of initiation of treatment. Continued therapy should be carefully reconsidered in a patient exhibiting no improvement within this timeframe.
	Body-weight ≥ 30 kg	8 mg/kg once every 4 weeks	

SPC, Summary of Product Characteristics.

2 DEFINITION OF THE DECISION PROBLEM

2.1 Decision problem

In line with the scope of the NICE appraisal, the clinical- and cost-effectiveness of abatacept, adalimumab, etanercept and tocilizumab for the treatment of JIA will be assessed.

The comparators for this assessment are: DMARDs (such as methotrexate), if DMARDs can be tolerated; best supportive care, if DMARDs are not tolerated; biologic DMARDs (etanercept, abatacept, adalimumab and tocilizumab) compared with each other within their licensed indications where appropriate.

The relevant population are children and young people with JIA diagnosed either at onset as polyarthritis (RF+ve and RF-ve) or those with extended oligoarthritis, and those with other forms of polyarticular course arthritis e.g. ERA, PA or undifferentiated arthritis. Children/young people with JIA and uveitis are also relevant. The age of the children/young people may vary by intervention because of differences in the licenced indications.

As specified in the NICE scope the following clinical-effectiveness outcome measures are relevant to the decision problem: disease activity; disease flares; physical function; joint damage; pain; reduced

use of corticosteroids; occurrence of extra-articular manifestations (such as uveitis); changes in body weight and height; mortality; adverse effects of treatment; and health-related quality of life.

2.2 Overall aims and objectives of assessment

The aim of this MTA is to assess the clinical and cost-effectiveness of abatacept, adalimumab, etanercept and tocilizumab for treating JIA.

The objectives are:

- To undertake systematic reviews of the clinical- and cost-effectiveness of abatacept, adalimumab, etanercept and tocilizumab for the treatment of JIA, and of the health-related quality of life (HRQoL) of people with JIA.
- To critique the companies' submissions (CS) to NICE from AbbVie (adalimumab), BMS (abatacept), Pfizer (etanercept) and Roche (tocilizumab), and to identify the strengths and weaknesses of the respective submissions.
- To conduct an economic evaluation establish the cost-effectiveness of abatacept, adalimumab, etanercept and tocilizumab for the treatment of JIA.

Patients with systemic onset JIA exhibiting typical systemic features such as spiking fever and rash are excluded from this MTA but if those features are no longer present (no active systemic symptoms during the previous six months) and the patients have gone on to have polyarticular course JIA they will be included. Similarly, patients with ERA and PA that has a polyarticular course will also be included.

3 METHODS

The *a priori* methods for systematically reviewing the evidence of clinical-effectiveness and cost-effectiveness are described in a research protocol published on the NICE website and registered with the PROSPERO international prospective register of systematic reviews database (registration number CRD42015016459). The protocol was sent to our expert advisory group (see Acknowledgements) for comment. Minor amendments were made as appropriate. None of the comments received identified specific problems with the methods of the review.

3.1 Identification of studies

Sensitive search strategies were developed and refined by an experienced information specialist. Separate searches were conducted to identify studies of clinical-effectiveness, cost-effectiveness and HRQoL.

The following databases were searched for published studies and ongoing research from inception to May 2015: The Cochrane Library including the Cochrane Database of Systematic Reviews (CDSR),

the Cochrane Central Register of Controlled Trials, CRD (University of York) Database of Abstracts of Reviews of Effectiveness (DARE), the NHS Economic Evaluation Database (NHS EED) and the Health Technology Assessment (HTA) database; Medline (Ovid); Embase (Ovid); Medline In-Process and Other Non-Indexed Citations (Ovid); Web of Science with Conference Proceedings: Science Citation Index Expanded (SCIE) and Conference Proceedings Citation Index - Science (CPCI) (ISI Web of Knowledge); Biosis Previews (ISI Web of Knowledge); Zetoc (Mimas); NIHR-Clinical Research Network Portfolio; Clinical Trials.gov, ISRCTN (International Standard Randomised Clinical Trial Number), UKCTG (UK Clinical Trials Gateway) and WHO ICTRP (International Clinical Trials Research Platform). In addition, Psychinfo (Ebsco) was searched for HRQoL studies. Searches were not limited to particular trial designs and although searches were not restricted by language, only full texts of English-language articles were retrieved during the study selection process. Cost-effectiveness and HRQoL searches were conducted from database inception to May 2015. References were downloaded into a Reference Manager database and de-duplicated where necessary.

Bibliographies of included articles and systematic reviews were also searched. The company submissions (CS) to NICE were searched for any additional studies that met the inclusion criteria (see section 4.2 and section 5.4). Members of our advisory group were asked to identify additional published and unpublished evidence. Further details including search dates for each database and an example search strategy can be found in Appendix 1.

3.2 Inclusion and exclusion criteria

The following inclusion/exclusion criteria were applied to the clinical-effectiveness review:

- Interventions: Etanercept, abatacept (with or without methotrexate), adalimumab (with or without methotrexate) and tocilizumab (with or without methotrexate). Each drug was evaluated within their licensed indication. Studies of treatment without methotrexate were permitted if patients were intolerant to methotrexate or for whom treatment with methotrexate is inappropriate.
- Comparators: DMARDs (such as methotrexate which is the most common conventional treatment in the UK) if DMARDs can be tolerated and best supportive care if DMARDs are not tolerated. Etanercept, abatacept, adalimumab and tocilizumab compared with each other.
- Population: Patients with JIA including
 - Polyarthritis (rheumatoid factor +ve, rheumatoid factor -ve and extended oligoarthritis, both onset and course)
 - ERA
 - PA

Studies of patients with systemic JIA were not included, as this was the subject of a separate NICE appraisal (NICE TA 238).⁴⁴

- Outcomes: Studies reporting one or more of the following outcomes were included:
 - Disease activity
 - Disease flares
 - Physical function
 - Joint damage
 - Pain
 - Corticosteroid reducing regimens
 - Extra-articular manifestations (such as uveitis)
 - Body weight and height
 - Mortality
 - Adverse effects of treatment
 - HRQoL
- Study design: Randomised controlled trials (RCTs). Any relevant systematic reviews identified in the systematic review of clinical-effectiveness were used as a source of references. Studies published as abstracts or conference presentations were only included if published from 2012 onwards and sufficient details were presented (or available elsewhere e.g. in a full paper reporting on the same RCT) to allow an appraisal of the methodology and the assessment of results to be undertaken.

The inclusion/exclusion criteria for the cost-effectiveness and HRQoL studies are presented in section 5.2 and section 5.3, respectively.

3.3 Data extraction strategy

Reference screening

All studies were selected for inclusion through a two-stage process. Titles and abstracts were screened independently by two reviewers for potential eligibility, using a standardised and piloted eligibility selection worksheet (Appendix 2 – clinical-effectiveness) containing the inclusion/exclusion criteria detailed above.

Full paper screening

Full texts for potentially relevant studies were obtained and screened using a standardised and piloted eligibility section worksheet (Appendix 3) by one reviewer, checked by a second and a final decision regarding inclusion was agreed. At each stage any disagreements were resolved by discussion or with the involvement of a third reviewer when necessary.

3.4 Critical appraisal strategy

Clinical-effectiveness studies were appraised using the Cochrane Risk of Bias criteria (e.g. selection bias, detection bias, performance bias, attrition bias, and selective reporting bias).⁵² Aspects of study quality including statistical procedures, outcome measurement and generalisability were also assessed.

Critical appraisal of the included clinical-effectiveness and cost-effectiveness studies (section 5.2) was conducted by one reviewer and checked by a second reviewer. Any disagreements were resolved by consensus or in consultation with a third reviewer where necessary.

3.5 Method of data synthesis

Details of the trial outcomes in the clinical-effectiveness review were synthesised through narrative review with tabulation of the results of included studies. Quantitative pooling of outcomes across clinical-effectiveness studies in a meta-analysis was not possible as the identified evidence included only one trial per biologic DMARD, all using placebo as the comparator. It was not considered appropriate to meta-analyse the four biologic DMARDs together due to clinical heterogeneity.

An adjusted indirect comparison of the four biologic DMARDs was performed using the method described by Bucher and colleagues (1997).⁵³ An indirect comparison refers to the synthesis of data from trials in which the technologies of interest have not been compared in head-to-head trials, but have been compared indirectly using data from a network of trials that compare the technologies with other interventions. A distinction is often made between adjusted and naïve (unadjusted) indirect comparisons. In the adjusted indirect comparison, the comparison of the interventions of interest is adjusted by preserving the strength of randomisation. Unadjusted indirect comparisons are considered to be observational evidence and therefore not recommended.^{54;55}

4 CLINICAL-EFFECTIVENESS

4.1 Results

4.1.1 Quantity and quality of research available

Titles and, where available, abstracts of a total of 2651 references identified by searches (after de-duplication) were screened and full copies of 60 references were retrieved. Of these 29 were excluded after inspection of the full article as shown in Figure 1 and these are listed in Appendix 4. The most common reasons for exclusion of a reference was an irrelevant study design (e.g. systematic reviews, (which were used as a source of references), commentaries). One full text⁵⁶ was of unclear relevance to the review because the type of JIA was not stated and it was not clear whether participants met the

licenced indication for etanercept therapy in respect of having inadequate response or intolerance to methotrexate. One full paper and eight conference abstracts relating to four ongoing studies that appeared relevant were tagged for inclusion in Section 4.3 'Ongoing studies' (note that a further three ongoing studies were identified from a separate search specifically undertaken for ongoing studies which is not represented in Figure 1 hence a total of seven ongoing studies is summarised in Section 4.3).

Nine full texts and 12 conference abstracts described four RCTs (each described by at least one full paper) that met the inclusion criteria of the review (Figure 1). As the full texts provided the most complete data, these were the primary source of information for this review.

One of the RCTs evaluated abatacept⁵⁷⁻⁶⁰ (the AWAKEN trial), one RCT evaluated adalimumab (Lovell et al. 2008⁶¹⁻⁶⁴), one RCT evaluated etanercept (Lovell et al. 2000^{42;65-67}) and one RCT evaluated tocilizumab⁶⁸⁻⁷⁶ (the CHERISH trial). For the sake of brevity, generally only the key reference of each RCT will be cited in the report. All four RCTs used placebo as the comparator, however, with the exception of the etanercept trial, the majority of the patients in the trials received methotrexate in addition to the biologic DMARD or placebo. The key characteristics of the trials are presented in **Error! Reference source not found.** with the primary and secondary outcomes measured in trials summarised in Table 11. All studies were multi-centre RCTs with the number of centres ranging from nine in the etanercept study⁴² to 58 in the tocilizumab study.⁶⁸ Locations of the studies included the USA (all four studies), Canada (one study⁴²), Europe (three studies^{57;61;68} with only the tocilizumab study⁶⁸ including UK centres), Latin America (two studies^{57;68}), Australia (one study⁶⁸ and Russia (one study⁶⁸). In each study, participants were initially treated in an open-label phase with the biologic DMARD under investigation and had to achieve at least an ACR Pedi 30 response to the biologic DMARD to be eligible for entry to the randomised double-blind withdrawal phase, with the number of participants randomised ranging from 51 in the etanercept study to 166 in the tocilizumab study. As each study investigated a different biologic DMARD study specific details are provided below by study drug.

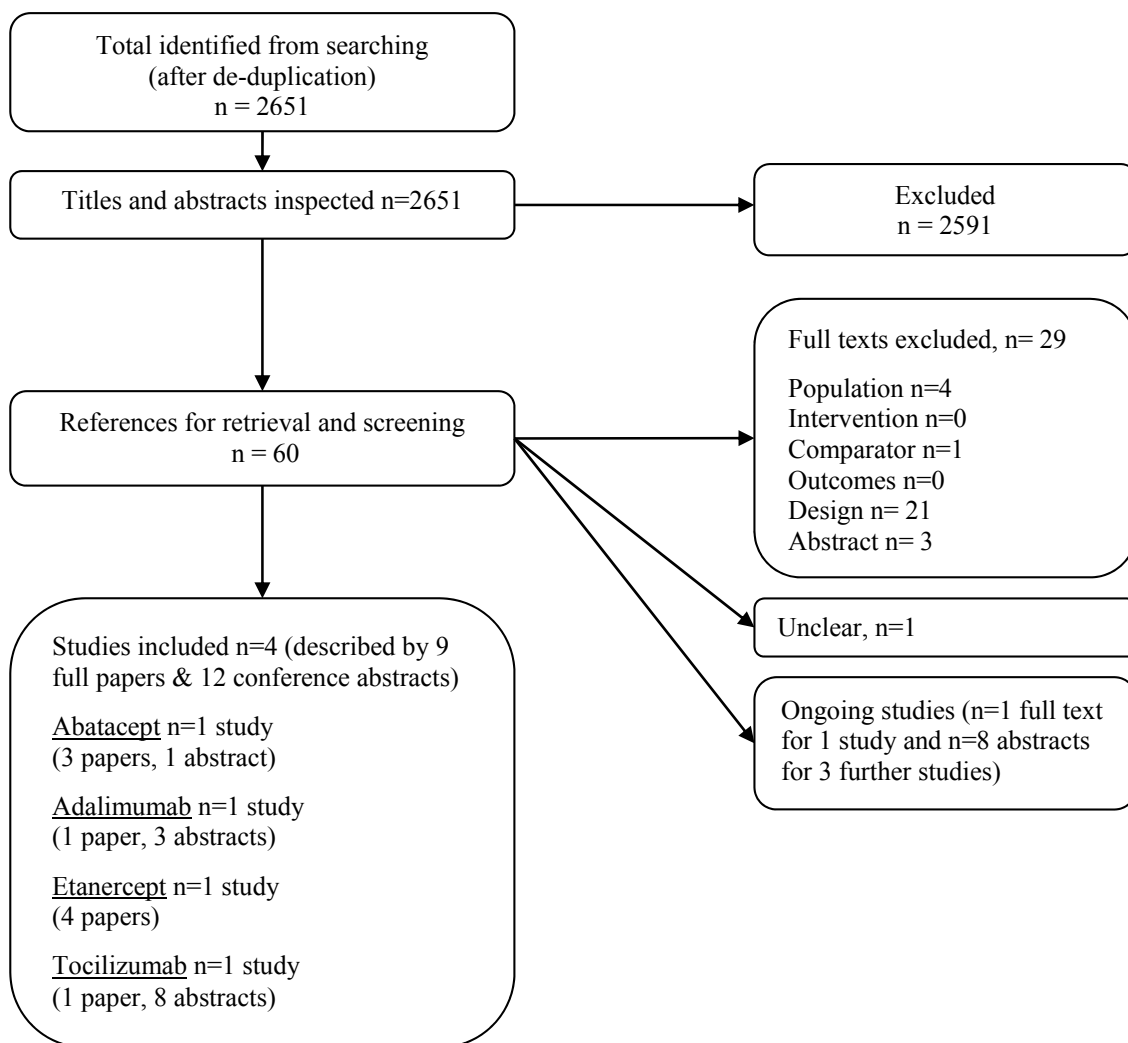


Figure 1 Flow-chart for the identification of studies

Abatacept

The abatacept RCT⁵⁷ was funded by Bristol-Myers Squibb (BMS) and consisted of three phases: a four-month open-label lead-in phase (days 1-113); a six-month double-blind randomised withdrawal phase (days 114-283), and an open-label extension phase [up to day 1681 (5.5 years) for efficacy and up to 7 years for safety]. Enrolled participants all received abatacept intravenously (10 mg/kg to a maximum of 1000 mg) and were permitted to continue to take stable methotrexate during the four months lead-in phase. Those achieving an ACR Pedi 30 response were then eligible to be randomised in a 1:1 ratio to continued abatacept (n=60) or placebo (n=62). In the six-month randomised withdrawal phase, abatacept was given at randomisation and about 28 day intervals (**Error! Reference source not found.**).

Patients were eligible for the trial if they were aged six to 17 years and had extended oligoarticular, polyarticular (RF+ve or RF-ve) or systemic JIA without systemic manifestations.

Table 10 Summary characteristics of included studies

	Abatacept	Adalimumab	Etanercept	Tocilizumab
Study details	AWAKEN, Ruperto et al. ⁵⁷⁻⁶⁰ Multi-centre withdrawal RCT at 45 centres in Europe (not UK), Latin America & USA	Lovell et al. 2008 ⁶¹⁻⁶⁴ Multi-centre withdrawal RCT at 31 centres in Europe (not UK) & USA	Lovell et al. 2000 ^{42,65-67} Multi-centre withdrawal RCT at 9 centres in Canada & the USA	CHERISH, Brunner et al. ⁶⁸⁻⁷⁶ Multi-centre withdrawal RCT at 58 centres in Australia, Europe (inc. UK), Latin America, Russia & USA
Study phases¹	16-week open-label lead-in 24-week randomised double-blind withdrawal Open-label extension	16-week randomised open-label 32-week randomised double-blind withdrawal Open-label extension	12-week open-label 16-week randomised double-blind withdrawal Open-label extension	16-week open-label lead-in 24-week randomised double-blind withdrawal Open-label extension
Intervention²	Abatacept: n=60 Abatacept 10 mg/kg at about 28-day intervals for 24 weeks or until disease flare	Adalimumab /Methotrexate: n=38 Adalimumab 24 mg/m ² BSA (to max. 40 mg) every other week for 32 weeks + MTX ≥10 mg/m ² BSA/week	Etanercept: n=25 Etanercept 0.4 mg/kg twice weekly until disease flare or for 16 weeks	Tocilizumab: n=82 10 mg/kg <30kg BW, n= 16; 8 mg/kg <30 kg BW n=11; 8 mg/kg ≥30kg BW n=55
Comparator²	Placebo n = 62	Placebo/Methotrexate: n= 37 Placebo + Methotrexate ≥10 mg/m ² body surface area/week	Placebo: n = 26	Placebo: n=84 10 mg/kg <30kg BW, n=15; 8 mg/kg <30kg BW n=13; 8 mg/kg ≥30kg BW n=56
Key inclusion criteria	Age 6-17 years Active ³ JIA (extended oligoarticular, polyarticular, RF+ve or RF-ve, systemic without systemic manifestations) Inadequate response or intolerance to ≥1 DMARD including biological agents ACR Pedi 30 for entry to randomised double-blind phase	Age 4-17 years Active ³ polyarticular-course JIA (any onset type) Inadequate response to NSAIDs ACR Pedi30 at week 16 for entry to double-blind withdrawal phase	Age 4-17 years Active ³ JIA Inadequate response to NSAIDs and methotrexate at doses of ≤10 mg/m ² body surface area/week	Age 2-17 years Active ³ polyarticular course or extended oligoarticular JIA (RF+ve or RF-ve) for ≥6months. Inadequate responses to or intolerant of methotrexate. Either never treated with biologics or had discontinued for a specified minimum period

DB, double-blind. LOM, limitation of motion. NSAIDs, non-steroidal anti-inflammatory drugs. RF, rheumatoid factor. +ve, positive. -ve, negative.

¹ the key phase of interest for efficacy outcomes is in bold text with lengths of phases reported in weeks for all studies for ease of comparison.

² during randomised double-blind withdrawal phase.

³ Inclusion criteria for active disease were very similar for the adalimumab, etanercept and tocilizumab studies (key aspects were at least five swollen joints and at least three joints with LOM). The abatacept study required at least five active joints (with swelling or LOM accompanied by pain or tenderness) and active disease (at least two active joints and 2 joints with LOM)

Table 11 Summary of outcomes measured

Parameter	Abatacept	Adalimumab	Etanercept	Tocilizumab
<i>Primary outcome</i>	Time to disease flare	Proportion of participants not receiving methotrexate with disease flares (week 16 to 48)	Number of patients with disease flare	Proportion of patients in whom a JIA-flare occurred during part 2 (up to and including week 40) compared with week 16
<i>Secondary outcomes:</i>	Proportion of patients at end of 6 months double-blind phase who had disease flare	Adverse events	Not specifically stated (ACR core variables, mortality and adverse events amongst others reported)	JIA-ACR 30/50/70/90 responses (week 40)
	Changes from baseline in ACR core variables			Change from baseline in ACR core response variables (week 40)
	Pain			Clinically inactive disease (week 40)
	Assessment of safety and tolerability			
	HRQoL			

Participants were required to have at least five active joints (defined as swelling or, in the absence of swelling, limited range of motion, accompanied by either pain or tenderness), active disease (defined as at least two active joints and two joints with a limited range of motion) and an inadequate response to, or intolerance to, at least one DMARD which could include biologic agents (e.g. etanercept, infliximab and adalimumab). Exclusion criteria included active uveitis, any major concurrent medical conditions and pregnancy or lactation.

The primary outcome measure was time to disease flare during the double-blind period. Disease flare was defined in three ways depending on the measure used: worsening of 30% or more in at least three of the six ACR core-response variables for JIA, and at least 30% improvement in no more than one variable during the double-blind period; a worsening of 20 mm or more on the 100 mm VAS if a global assessment by either physician or parent was used; worsening in two or more joints if the number of active joints or joints with limited range of motion was used. Clinical assessments preceded drug administration at each visit. Secondary outcomes included the proportion of patients at the end of six-month double-blind phase who had disease flare, changes from baseline in each of the six ACR core variables, pain, assessment of safety and tolerability and health related quality of life (HRQoL).

Adalimumab

The Lovell and colleagues (2008) RCT⁶¹ was funded by a research grant from Abbott Laboratories and consisted of three phases: a 16-week randomised open-label phase, a 32-week randomised

double-blind withdrawal phase and an open-label extension phase. Enrolled participants all received adalimumab subcutaneously (24 mg per square metre of body surface area, to a maximum of 40 mg) every other week and methotrexate (at least 10 mg per square meter of body surface area per week) during the four-month lead-in phase. Those achieving an ACR Pedi 30 response were then eligible to be randomised in a 1:1 ratio to continued adalimumab plus methotrexate (n=38) or placebo plus methotrexate (n=37) (**Error! Reference source not found.**). The trial included two further study arms (adalimumab only and placebo only), but because the majority of participants in these arms had never received methotrexate, they do not meet the licenced indication and are not included in this report.

Patients were eligible for the trial if they were aged four to 17 years and had polyarticular-course JIA of any onset type. If systemic onset then patients had to be free of any systemic JIA manifestations for at least three months prior to study qualification.⁷⁷ Participants were required to have active disease (defined as five or more swollen joints and three or more joints with limited range of motion), had an inadequate response to NSAIDs, and had either not previously been treated with methotrexate or if previously treated with methotrexate, had had adverse events or an inadequate response. Exclusion criteria included clinically significant deviations in haematologic, hepatic or renal indicators; ongoing infection or a recent major infection that had required hospitalisation or intravenous antibiotics; recent receipt of live or attenuated vaccines. Patients who had previously been treated with other biologic agents at any time or who had received recent treatment with intravenous immune globulin, cytotoxic agents, investigational agents, DMARDs (other than methotrexate) or corticosteroids were also excluded from participation.

The primary outcome for the study (percentage of participants not receiving methotrexate who had a disease flare during the double-blind period) related to the two study arms that, as noted above, do not meet the licenced indication and are therefore not included in this report. Disease flare was reported for the two study arms relevant to this assessment and it was defined in different ways depending on the measure used: worsening of 30% or more in at least 3 of the 6 core criteria for JIA, and at least 30% improvement in no more than one of the criteria during the double-blind period; an increase of more than 30% on the 0-100 VAS if a global assessment was used; an increase in the number of active joints to at least two when the patient had none or only one if the number of active joints was used, with the same approach used for defining flare using joints with loss of motion. Outcomes were assessed every 12 weeks. The occurrence of adverse events was a secondary outcome.

Etanercept

The Lovell and colleagues (2000) RCT^{42,65-67} was funded by the Immunex Corporation and consisted of three phases: an open-label lead-in phase of up to three months; a four-month double-blind

randomised withdrawal phase, and an open-label extension phase. All enrolled participants received etanercept subcutaneously (0.4 mg/kg twice weekly) during the four-month lead-in phase. Those who improved and achieved an ACR Pedi 30 response were then eligible to be randomised to continued etanercept (n=25) or placebo (n=26) during the withdrawal phase (**Error! Reference source not found.**).

Patients were eligible for the trial if they were aged four to 17 years and had active polyarticular JIA despite treatment with NSAIDs and methotrexate doses of at least 10 mg per square metre of body-surface area per week. Active disease was defined as at least five swollen joints and at least three joints with limited motion with pain, tenderness or both. Exclusion criteria included any major concurrent medical conditions and pregnancy or lactation.

The primary outcome measure was number of patients with disease flare during the double-blind withdrawal period. Disease flare was defined depending on the measure used: worsening of 30% or more in at least three of the six ACR core-response variables for JIA, at least 30% improvement in no more than one variable and a minimum of two active joints; a change of at least two units on a scale from 0 to 10 if a global assessment was used. Clinical assessments during the withdrawal phase took place on day one, day 15 and at the end of each month. Secondary outcomes were not specifically listed.

Tocilizumab

The tocilizumab RCT⁶⁸ consisted of three phases: a 16-week open-label lead-in phase; a double-blind randomised withdrawal phase (week 16 to week 40), and an open-label extension phase (64 weeks). Some funding for manuscript preparation was provided by H. Hoffmann-La Roche Ltd. Enrolled participants were permitted to receive methotrexate and all received tocilizumab intravenously (three groups, those with body weight < 30kg randomised to either 10 mg/kg or 8 mg/kg every four weeks. Those with body weight 30kg or more received 8 mg/kg every 4 weeks) during the 16-week lead-in phase. Those achieving an ACR Pedi 30 response were then eligible to be randomised in a 1:1 ratio to continued tocilizumab (n=82) or placebo (n=84) given every four weeks until week 40 unless they experienced disease flare (**Error! Reference source not found.**).

Patients were eligible for the trial if they were aged two to 17 years and had polyarticular-course or extended oligoarticular JIA that was either RF+ve or RF-ve for six months or more. Systemic JIA or any other categories of JIA were excluded from the trial.⁷⁸ Participants were required to have at least five active joints with a limited range of motion in at least three active joints and have an inadequate response to, or intolerance to methotrexate. If participants were taking methotrexate (10-20 mg/m²) or low dose oral glucocorticoids (≤ 0.2 mg/kg/day, daily maximum 10 mg) the dose had to have been

stable for eight or more weeks (methotrexate) or four or more weeks (oral glucocorticoids). Patients had to be treatment-naïve for biologics or had discontinued for a specified minimum period. No other exclusion criteria were specified.

The primary outcome measure was proportion of participants with disease flare during the double-blind period (up to and including week 40 compared to week 16). Disease flare was defined as worsening of 30% or more in at least three of the six ACR core-response variables for JIA, and at least 30% improvement in no more than one variable during the double-blind period. Outcomes were assessed every four weeks. Secondary outcomes included the ACR Pedi 30/50/70/90 responses, the change from baseline in JIA core response variables and clinically inactive disease (physician global assessment indicating no disease activity plus the absence of all the following: joints with active arthritis, uveitis and erythrocyte sedimentation rate greater than 20 mm/hour).

Overview of the participants in the withdrawal phases of the included studies

For three of the four trials (abatacept,⁵⁷ adalimumab,⁶¹ and etanercept⁴²) baseline characteristics are provided for the participants who had achieved an ACR Pedi 30 and who were randomised to the double-blind withdrawal phase of each trial. The tocilizumab trial publication,⁶⁸ however, presented participant baseline characteristics for participants as randomised to the initial open-label lead-in phase, where three groups of participants all received the study drug (if body weight <30 kg then randomised to either 10 mg/kg or 8 mg/kg every 4 weeks; if body weight ≥30 kg then received 8 mg/kg every 4 weeks). Selected baseline characteristics are presented in Table 12, with the full set of characteristics available in the data extraction forms (Appendix 5). The mean age of trial participants reflected the differing entry criteria for the trials. Participants in the abatacept trial⁵⁷ (ages 6-17 years eligible) had the highest mean age (12-13 years) whereas those in the adalimumab⁶¹ and etanercept trials⁴² (ages 4-17 years eligible) had a slightly lower mean age (approximately 9-12 years) which was similar to those enrolled in the open-label phase of the tocilizumab study⁶⁸ (ages 2-17 eligible, mean age approximately 11 years). The majority of participants in all four studies were female (ranging from 67% in the etanercept study⁴² to 80% in the adalimumab study⁶¹) and of white ethnicity (73% in the etanercept study⁴² to 96% in the adalimumab study⁶¹). The proportion of patients across the sub-types of JIA were only reported for two of the trials (abatacept⁵⁷ and etanercept⁴²). In these two trials polyarthritis was the predominant sub-type. In the abatacept trial just under 20% of patients had systemic JIA (without systemic manifestations),⁵⁷ whilst in the etanercept trial around a third had systemic JIA (with apparent systemic manifestations: spiking fever and rheumatoid rash).⁴² None of the trials included patients with PA or ERA (based on the eligibility criteria given). The proportion of participants who were RF+ve ranged from 22% in the adalimumab study⁶¹ to 29% in the tocilizumab study⁶⁸ and the duration of JIA from just under four years in the abatacept study⁵⁷ to approximately six years in the etanercept study.⁴²

The treatment groups in the abatacept study⁵⁷ appear similar on most variables although the placebo group had a smaller proportion of RF+ve patients than the abatacept group (19% versus 32%). The adalimumab study report⁶¹ indicated that there were no significant differences in baseline characteristics between the placebo and adalimumab groups. Groups were described as well balanced in the etanercept study⁴² with the exceptions of age group (4-8 years old: 52% etanercept vs 19% placebo, $p<0.02$), race (white race: 56% etanercept vs 88% placebo, $p<0.02$) and corticosteroid use (corticosteroid use at wash out: etanercept 24% vs 50% placebo, $p=0.05$). The tocilizumab study did not report baseline characteristics for those participants who entered the double-blind wash out phase of this study.

Assessment of the risk of bias of included studies

The Cochrane risk of bias criteria⁵² focus on various aspects of study design, conduct and reporting which may help to gauge the internal validity (whether the study answered the research question in a manner that was free from bias) of the individual studies. The risk of bias in the included trials is summarised in Table 13 and further details are presented in the data extraction tables (Appendix 5).

Only the abatacept trial⁵⁷ reported sufficient details on the methods for generating the random sequence (computer generated) and allocation concealment (interactive voice-randomisation system) to establish that there was a low risk of selection bias in this trial. In the three other trials (adalimumab,⁶¹ etanercept⁴² and tocilizumab⁶⁸) the risk of selection bias associated with randomisation and allocation were unclear because either no details were reported or there was insufficient information to make a judgement. The randomised withdrawal phases of all four trials were described as double-blind with three of the trials providing some information to support this statement (e.g. placebo identical in appearance,⁵⁷ indication of who was unaware of treatment assignment^{42;61}). The risk of performance bias and detection bias was judged to be low for all four trials. Attrition bias (systematic differences in withdrawals between trial arms) was judged to be low for all outcomes in three trials^{42;61;68} (either because attrition was similar between groups or because incomplete data were addressed). In the abatacept trial,⁵⁷ however, a larger proportion of patients dropped out of the placebo group in the double-blind phase (placebo 50%, abatacept 18%), with the chief reason being due to a lack of efficacy.

Table 12 Selected baseline characteristics of trial participants

Baseline characteristics	Abatacept ⁵⁷		Adalimumab ⁶¹		Etanercept ⁴²		Tocilizumab ⁶⁸		
	Abatacept (n=60)	Placebo (n=62)	Adalimumab (n=38)	Placebo (n=37)	Etanercept (n=25)	Placebo (n=26)	<i>TCZ 8 mg/kg <30kg (n=34)¹</i>	<i>TCZ 10 mg/kg <30kg (n=35)¹</i>	<i>TCZ 10 mg/kg ≥30kg (n=119)¹</i>
Age, years; mean (SD)	12.6 (3)	12.0 (3)	11.7 (3.3)	10.8 (3.4)	8.9	12.2	<i>7.6 (2.71)</i>	<i>6.9 (3.02)</i>	<i>13.1 (2.78)</i>
Sex female, n (%)	43 (72)	45 (73)	30 (79)	30 (81)	19 (76)	15 (58)	<i>24 (71)</i>	<i>30 (86)</i>	<i>90 (76)</i>
Ethnicity white, n (%)	46 (77)	49 (79)	36 (95)	36 (97)	14 (56)	23 (88)	NR	NR	NR
Black	5 (8)	4 (7)	0	0	3 (12)	1 (4)			
Hispanic	NR	NR	NR	NR	6 (24)	2 (8)			
Other	9 (15)	9 (15)	2 (5)	1 (3)	2 (8)	0			
Type of JIA									
Pauciarticular ²					2 (8)	1 (4)	Eligible patients had RF +ve or RF -ve polyarticular-course JIA or extended oligoarticular JIA but no further detail provided.		
Persistent oligoarthritis	0	2 (3)	Described as 'polyarticular course' no further detail (This older nomenclature could have included patients who would now be defined as having ERA or PA)						
Extended oligoarthritis	9 (15)	7 (11)							
Polyarthritis (RF +ve)	14 (23)	12 (19)			14 (56)	17 (65)			
Polyarthritis (RF -ve)	26 (43)	28 (45)							
Systemic	11 (18) ³	12 (19) ³			9 (36)	8 (31)			
RF +ve, n (%)	19 (32)	12 (19)	10/37 (27) ⁴	6/36 (17) ⁴	4 (16)	8 (31)	<i>2 (6)</i>	<i>4 (11)</i>	<i>48 (40)</i>
Duration of JIA, years; mean (SD)	3.8 (3.7)	3.9 (3.5)	4.3 (4.1)	4.0 (3.5)	5.3	6.4	<i>3.5 (2.57)</i>	<i>3.4 (2.39)</i>	<i>4.7 (4.16)</i>

NR, not reported. RF+ve, rheumatoid factor positive. RF -ve, rheumatoid factor negative. SD, standard deviation.

¹ Baseline data in italics for the tocilizumab study were presented only for all patients randomised to the initial open lead-in phase of the study. Of these participants, 15/188 (7.9%) did not achieve ACR Pedi 30 response and were not randomised to the double-blind withdrawal phase of the study.

² Pauciarticular arthritis would now be called oligoarticular arthritis. It is not clear from the paper whether these participants had persistent oligoarthritis or extended oligoarthritis.

³ Systemic without systemic manifestations

⁴ Calculated by reviewer

Although this was addressed for some outcomes (e.g. analysis of ACR variables), it was not addressed for HRQoL where the analysis was based on available data at each time point, hence the risk of attrition bias is high for this outcome. Selective reporting bias was judged low for all the trials as all outcomes were reported on. The only other uncertainty surrounding study biases was the risk of bias due to inter-centre variability in the adalimumab⁶¹ etanercept⁴² and tocilizumab⁶⁸ trials where inter-centre variability was not discussed. In contrast the abatacept study⁵⁷ reported that training was in place for joint assessors from each centre who had specific and standardised joint assessment training.

Table 13 Summary of risk of bias assessment

Criteria	Abatacept ⁵⁷⁻⁵⁹	Adalimumab ⁶¹	Etanercept ⁴²	Tocilizumab ⁶⁸
Selection bias				
Random sequence generation	Yes	Unclear	Unclear	Unclear
Allocation concealment	Yes	Unclear	Unclear	Unclear
Performance bias				
Blinding of participants and personnel	Yes	Yes	Yes	Yes
Detection bias				
Blinding of outcome assessment	Yes	Yes	Yes	Yes
Attrition bias				
Incomplete outcome data addressed				
Non-HRQoL outcomes	Yes	Yes	Yes	Yes
HRQoL outcome	No	N/A	N/A	N/A
Reporting bias				
Selective reporting	Yes	Yes	Yes	Yes
Other bias				
Other sources	Unclear	Unclear	Unclear	Unclear

Yes (low risk of bias). N/A, not applicable.. No (high risk of bias). Unclear (uncertain risk of bias).

4.1.2 Assessment of clinical-effectiveness - biologic DMARDs vs placebo (with methotrexate where permitted)

Disease Flare

The primary outcome for all four trials was disease flare, albeit with some differences in the way this outcome was reported. Data on disease flare from the trials contribute to the economic model in this assessment report (section 5.6.1). The definitions for disease flare were broadly consistent between the studies (a worsening of at least 30% in three or more of the six core criteria for JIA, and an improvement of 30% or more in no more than one of the criteria), with some studies also including

flare definitions based on global assessments and number of active joints. In all four studies, there were statistically significantly fewer arthritis flares in patients being treated with biologic DMARDs compared to those receiving placebo and in the two studies that reported time to disease flare this was statistically significantly longer in patients being treated with biologic DMARDs compared to those receiving placebo (Table 14).

Table 14 Disease flare during the randomised withdrawal phase

Study (Length: OL, RCT¹), Outcome	Intervention	Comparator	
Abatacept⁵⁷ (4mo OL, 6mo RCT)	ABA (n=60)	PBO (n=62)	p value
Time to flare, median months	Not reached	6	0.0002
Disease flares, n (%)	12 (20)	33 (53)	0.0003
Disease flares, hazard ratio	0.31 (95% CI 0.16 to 0.59)		NR
Adalimumab⁶¹ (4mo OL, 8mo RCT)	ADA (n=38)	PBO (n=37)	p value
Disease flares, n/N (%)	14/ 38 (37)	24/37 (65)	0.02
Etanercept⁴² (3mo OL, 8mo RCT)	ETA (n=25)	PBO (n=26)	p value
Disease flare, n (%)	7 (28)	21 (81)	0.0031 ²
Corticosteroid use at baseline ³			0.05
Yes	3/6 (50)	12/13 (92)	
No	4/19 (21)	9/13 (69)	
Time to flare, median days	>116	28	p<0.001
Tocilizumab⁶⁸ (4mo OL, 6mo RCTs)	TCZ (n=82)	PBO (n=81)⁴	Difference⁵ TCZ vs PBO (95% CI); p value
Proportion with JIA flare, n (%)	21 (25.6)	39 (48.1)	-0.21 (-0.35, 0.08); 0.0024

ABA, abatacept. ADA, adalimumab. ETA, etanercept. mo, months. NR, not reported. OL, open-label. PBO, placebo. TCZ, tocilizumab.

¹ For ease of comparison lengths of open-label and RCT phases are presented as time in months (where originally presented in weeks value has been divided by 4, where originally presented in days value has been divided by 28)

² p<0.001 after adjustment for baseline characteristics in logistic regression model.

³ Authors state that with the exception of corticosteroid use at base line (p=0.05), none of the baseline characteristics were significant predictors of flare rates (p>0.15).

⁴ Of the 84 participants who achieved at least ACR Pedi 30 and were then randomised to placebo, three discontinued (1 insufficient therapeutic response, 2 due to adverse events). These three did not receive any study drug in the randomised part of the study and so were excluded from the analyses.

⁵ Adjusted for baseline stratification factors (background use of MTX and oral glucocorticoids).

In the abatacept study,⁵⁷ by the end of the RCT period disease flare had occurred in 20% of patients receiving the study drug compared with 53% of patients receiving placebo (p=0.0003). Median time to disease flare was six months for the placebo group and statistically significantly greater compared to the abatacept group (p=0.0002), but authors state that insufficient events occurred in the abatacept group for this to be assessed. The risk of disease flare in patients randomised to continued abatacept during the RCT phase was just under a third of that for those receiving placebo (hazard ratio 0.31, 95% CI 0.16, 0.95, no p value reported).

Disease flare occurred in 37% of patients receiving adalimumab⁶¹ compared with 65% of those receiving placebo (p=0.02).

In the etanercept study, disease flare occurred in 28% of patients receiving etanercept⁴² compared with 81% receiving placebo (p=0.003). The authors of the etanercept study state that after adjustment for the effects of baseline characteristics, the rates of flare remained significantly lower in the etanercept group (p<0.001), with only corticosteroid use at baseline being a significant predictor of flare rates (p=0.05). The median time to disease flare with etanercept was greater than 116 days compared with 28 days for the placebo group, with 13/25 patients still receiving etanercept at the end of the study (day 116) (p<0.001).

In those receiving tocilizumab, disease flare occurred in 26% of patients compared with 48% receiving placebo (adjusted difference in flare rate: -0.21; 95% CI -0.35 to -0.08; p=0.0024), with authors stating that flares in the placebo group were evident as early as 28 days after randomisation.

American College of Rheumatology (ACR) Pediatric (Pedi) Responses

ACR Pedi 30, 50 and 70 responses were reported by all four studies, with all but the etanercept study⁴² also reporting ACR Pedi 90 responses. The abatacept and tocilizumab studies^{57;68} additionally report inactive disease, which was defined similarly in the two studies [no joints with active arthritis, normal erythrocyte sedimentation rate (ESR) of 20 mm per hour or less, physician's global assessment (PGA) <10 on a 100mm on a VAS (VAS)⁵⁷ or PGA also <10 on a 100mm on a VAS, indicating no disease activity with the tocilizumab study⁶⁸ also including an absence of uveitis (patients with uveitis were excluded from the abatacept study)]. In all groups that continued to receive biologic DMARDs during the randomised withdrawal phase of the study the proportion of participants with ACR Pedi responses of 30 or more were greater than in the placebo group and when a p-value was reported the differences were statistically significant in all but two instances (Table 15).

While more patients receiving abatacept (82%) achieved an ACR Pedi 30 response compared to patients receiving placebo (69%), the difference was not statistically significant (p=0.1712).⁵⁷ However, a statistically significantly greater proportion of patients in the abatacept treatment group achieved an ACR Pedi 50, 70 or 90 response compared with those receiving placebo (p=0.0071, p=0.0185 and p=0.0062, respectively). In addition statistically significantly more patients treated with abatacept (30%) compared with those receiving placebo (11%; p=0.0195) were classified as having inactive disease.

A statistically significantly higher percentage of patients being treated with adalimumab achieved ACR Pedi 30, 50 and 70 responses compared with those receiving placebo (p=0.03, p=0.03 and p=0.002, respectively).⁶¹ The percentage of patients with ACR Pedi 90 response rates was also greater for adalimumab-treated patients compared to placebo (42 vs 27 placebo), however this difference was not statistically significant (p=0.17).

In the etanercept study ACR 30, 50 and 70 responses were achieved by a greater proportion of patients being treated with etanercept during the randomised withdrawal phase than those receiving placebo. However, a statistical comparison showing that this difference was statistically significant was only reported for ACR Pedi 30 (p<0.01).

A statistically significantly higher proportion of tocilizumab-treated patients achieved ACR Pedi 30, 50 and 70 responses compared to those receiving placebo during the randomised withdrawal phase of the study (p=0.0084, p=0.0050 and p=0.0032, respectively). While ACR Pedi 90 response was also higher in the tocilizumab group, no p value was provided. The proportion of patients with inactive disease was 36.6% for the tocilizumab group compared with 17.3% for the placebo group (no p value provided).

Table 15 ACR paediatric responses relative to baseline

Study (Length: OL, RCT), Outcome	Intervention	Comparator	
Abatacept⁵⁷ (4mo OL, 6mo RCT)			
ACR Pedi, n (%)¹	ABA (n=60)	PBO (n=62)	p value
30	49 (82)	43 (69)	0.1712
50	46 (77)	32 (52)	0.0071
70	32 (53)	19 (31)	0.0185
90	24 (40)	10 (16)	0.0062
Inactive disease ²	18 (30)	7 (11)	0.0195
Adalimumab⁶¹ (4mo OL, 8mo RCT)			
ACR Pedi, %	ADA (n=38)	PBO (n=37)	p value
30	63	38	0.03
50	63	38	0.03
70	63	27	0.002
90	42	27	0.17
Etanercept⁴² (3mo OL, 4mo RCT)			
ACR Pedi, n (%)³	ETA (n=25)	PBO (n=26)	p value
30	20 (80)	9 (35)	p<0.01
50	18 (72)	6 (23)	NR
70	11 (44)	5 (19)	NR
Tocilizumab⁶⁸ (4mo OL, 6mo RCT)			
ACR Pedi, n (%)	TCZ (n=82)	PBO (n=81)	Difference⁴ TCZ vs PBO (95% CI); p value
30	61 (74.4)	44 (54.3)	0.09 (0.05, 0.33); 0.0084
50	60 (73.2)	42 (51.9)	0.20 (0.06, 0.34) ; 0.0050
70	53 (64.6)	34 (42.0)	0.22 (0.07, 0.37); 0.0032
90	37 (45.1)	19 (23.5)	0.21 (0.07, 0.35); NR
Inactive disease	30 (36.6)	14 (17.3)	0.18 (0.05, 0.32); NR

ABA, abatacept. ADA, adalimumab. D-B, double-blind. ETA, etanercept. mo, months. NR, not reported. OL, open-label. PBO, placebo. TCZ, tocilizumab.

¹ Assessed after the 6-month RCT phase or at the time of flare for patients who did not complete this period

² Defined as no of joints with active arthritis, a physician's assessment of ≤10 on a 100mm VAS and a normal ESR rate.

³ If a patient had a flare they were classified as having no response (ACR Pedi <30) from that point on, regardless of their ACR Pedi response at that time. Missing values were also imputed as non-responses.

⁴ Adjusted for baseline stratification factors (background use of methotrexate and oral glucocorticoids).

ACR Pedi core variables

The adalimumab study⁶¹ did not report outcomes for the ACR Pedi core variables. The three remaining studies however reported data as mean (abatacept⁵⁷), median (etanercept⁴²) or an adjusted mean for change from baseline (tocilizumab⁶⁸). In addition, the etanercept study reports additional joint and pain outcomes, while the abatacept and tocilizumab studies also report additional pain outcomes (see following sections *Joint Damage* and *Pain*).

Generally, the core-response variable outcomes were in favour of treatment with the biologic DMARDs compared to placebo. However, as can be seen in Table 16, there were some exceptions. For abatacept,⁵⁷ differences in the adjusted mean percentage change (adjustment based on an ANCOVA model with treatment as factor, and baseline value as covariate) over the double-blind period (from day 113 to day 282) for the parent's global assessment and ESR rates between the treatment groups were not significantly different (p=0.6992 and p=0.9562, respectively). The mean scores for the CHAQ disability index at the end of the double-blind withdrawal trial period (day 282) are the same for both groups (0.8). However, when the difference in the adjusted mean percentage change values for the CHAQ disability index from the start to the end of the double-blind period (day 113 to day 282) are compared a statistically significant p value is reported in favour of the abatacept group (p=0.0388).

For etanercept,⁴² all core variable outcomes appear to be in favour of the etanercept group when compared to placebo, however no statistical comparisons between treatment groups were reported. For the tocilizumab study⁶⁸ differences in adjusted mean changes from baseline between treatment groups for physician's global assessment of disease and number of active joints are reported to be statistically significant in favour of tocilizumab (p values of 0.0031 and, 0.0435 respectively), no p values for the remaining outcomes were reported.

Table 16 ACR Pedi core variables

Study (Length: OL, RCT), Outcome	Intervention	Comparator	p value¹
Abatacept⁵⁷ (4mo OL, 6mo RCT)	ABA (n=60)	PBO (n=62)	p value¹
Core-response variables, mean (SD) ²			
Physician's global assessment (VAS: 100mm)	14.7 (18.9)	23.2 (21.8)	0.0004
Parent's global assessment (VAS: 100mm)	17.9 (22.2)	23.9 (21.6)	0.6992
Physical function (CHAQ disability index: 0-3, best-worst)	0.8 (0.9)	0.8 (0.7)	0.0388
No. of active joints (number assessed not stated)	4.4 (7.0)	6.0 (5.8)	0.0245
No. of joints with LOM (number assessed not stated)	8.8 (12.8)	8.6 (12.0)	0.0128
ESR (mm per hour) ³	25.1 (26.4)	30.7 (30.1)	0.9562
Etanercept⁴² (3mo OL, 4mo RCT)	ETA (n=25)	PBO (n=26)	p value
JIA core set criteria, median ²			

Physician's global assessment of disease severity (0-10, best-worst)	2	5	NR
Patient/parent global assessment of overall well-being (0-10, best-worst)	3	5	NR
Scores of CHAQ (0-3, best-worst)	0.8	1.2	NR
Total number of active joints (out of 73 joints)	7.0	13.0	NR
No. of joints with LOM and with pain, tenderness, or both (out of 71 joints; 0-10, best-worst)	1.0	4.5	NR
ESR (normal ranges 1-30mm per hour for females, 1-13mm per hour for males) ³	18	30	NR
Tocilizumab⁶⁸ (4mo OL, 6mo RCT)	TCZ (n=82)	PBO (n=81)	Difference⁴ TCZ vs PBO (95% CI); p value
JIA- core response variables, change from baseline - adjusted mean ²			
Physician's global assessment of disease severity (0-100, 0= inactive disease)	-45.2	-35.2	-9.9 (-16.5, -3.4); 0.0031
Patient global assessment of well-being (0-100, 0 = very poor)	-32.1	-24.7	-7.4 (-14.8, 0.0); NR
CHAQ - Disability index score (0-3, 0 = no disability)	-0.8	-0.6	-0.2 (-0.4, 0.0); NR
No. of active joints (range 0-71)	-14.3	-11.4	-2.9 (-5.7, -0.1); 0.0435
No. of joints with LOM (range 0-67)	-9.5	-7.7	-1.8 (-4.1, 0.5); 0.1229
ESR (mm/h)	-26.3	-12.0	-14.3 (-19.6, -9.0), NR

ABA, abatacept. CHAQ, Childhood Health Assessment Questionnaire. ESR, Erythrocyte sedimentation rate. ETA, etanercept. LOCF, last-observation carried forward. LOM, limitation of motion. Mo, months. PBO, placebo. NR, not reported. RCT, randomised controlled trial. SD, standard deviation. TCZ, tocilizumab. VAS, visual analogue scale.

¹ Abatacept study - p-values are based on the difference in the adjusted mean percentage change from day 113 to day 282 (start and end of the double-blind period).

² Missing values were imputed with LOCF.

³ C-reactive protein values were also reported by these studies: mean (SD) ABA 0.16 (0.25) vs PO 0.29 (0.54); p=0.0255.⁵⁷ Median ETA: 0.4 vs PO 3.0 (normal range 0-0.79 mg per decilitre).⁴²

⁴ adjusted for baseline stratification factors (background use of methotrexate and oral glucocorticoids).

Joint-related outcomes

None of the trials reported any radiographic outcomes. However, in addition to the ACR Pedi core variable outcomes that capture number of active joints and joints with limited range of motion, the etanercept study⁴² presented data for some additional joint related outcomes (Table 17).

No statistical comparisons for these outcomes were reported between the etanercept and the placebo group, however the median number of swollen joints (4.0 vs 11.0 placebo), number of joints with LOM (9 vs 22 placebo), articular severity score (38 vs 66 placebo) and duration of stiffness (5 vs 38 placebo) all favour the etanercept treatment group (Table 17).

Table 17 Joint-related outcomes (other than ACR Pedi)

Study (Length: OL, RCT), Outcome	Intervention	Comparator	
Etanercept⁴² (3mo OL, 4mo RCT)	ETA (n=25)	PBO (n=26)	p value
No. of swollen joints (out of 66), median	4.0	11.0	NR
No. of joints with LOM (out of 71), median ¹	9	22	NR
Articular severity score (0-962, best-worst), median	38	66	NR

Duration of morning stiffness (min), median	5	38	NR
---	---	----	----

ETA, etanercept. LOM, limitation of motion. mo, months. No, number. NR, not reported. OL, open-label. PBO, placebo. RCT, randomised controlled trial.

¹ While 'number of joints with LOM' are part of the ACR Pedi core variables, authors reported results under 'other' rather than under the core variables in the publication.

Pain

The adalimumab study⁶¹ did not report a pain outcome. All the other studies (abatacept,⁵⁷ etanercept⁴² and tocilizumab⁶⁸) report pain assessed on a VAS, albeit reporting the data differently (mean, median and mean change from baseline, respectively). The difference between the abatacept (mean pain 15 mm) and the placebo (mean pain 21 mm) treatment groups was not statistically significant (p=0.105) and reported mean pain scores were lower for patients being treated with abatacept. The etanercept study did not report a statistical comparison between treatment groups, however median pain scores for patients being treated with etanercept (VAS 1.5 cm) were less than half of those for patients receiving placebo (VAS 3.5 cm). The tocilizumab study⁶⁸ reported the adjusted mean change from baseline, which statistically compared to the placebo group, was in favour of the tocilizumab treatment group (p = 0.0076).

Table 18 Pain

Study (Length: OL, RCT), Outcome	Intervention	Comparator	
Abatacept⁵⁷ (4mo OL, 6mo RCT)	ABA (n=60)	PBO (n=62)	p value
Pain (parent global assessment of pain, CHAQ VAS:100mm), mean	15 ¹	21 ¹	0.105
Etanercept⁴² (3mo OL, 4mo RCT)	ETA (n=25)	PBO (n=26)	p value
Pain (VAS: 0-10cm, best-worst), median	1.5	3.5	NR
Tocilizumab⁶⁸ (24 weeks)	TCZ (n=82)	PBO (n=81)	Difference² TCZ vs PBO (95% CI); p value
Pain (VAS: no details reported), adjusted mean change from baseline	-32.4	-22.3	-10.2 (-17.6, -2.7); 0.0076

ABA, abatacept. CHAQ, Childhood Health Assessment Questionnaire. ETA, etanercept. mo, months. NR, not reported. OL, open-label. PBO, placebo. RCT, randomised controlled trial. TCZ, tocilizumab.

¹ Read off from graph by reviewer. Analysis based on available data but number of patients at this time-point contributing data to this outcome unclear, 49/60 in the abatacept group and 31/62 in the placebo group completed the 6-month double-blind period.

² adjusted for baseline stratification factors (background use of methotrexate and oral glucocorticoids).

Corticosteroid reducing regimens

None of the included RCTs reported the effectiveness of biologic DMARDs on reducing the need for corticosteroids.

Extra-articular manifestations (such as uveitis)

None of the included RCTs reported outcomes for extra-articular manifestations. Of note, one of the trials (abatacept) excluded patients with active uveitis.⁵⁷

Height and body weight

None of the studies reported differences in height or body weight between the treatment groups for the double-blind, randomised controlled withdrawal phase of the trial.

Mortality

No deaths occurred in the adalimumab,⁶¹ etanercept⁴² and tocilizumab⁶⁸ studies, while this outcome was not reported in the abatacept study.⁵⁷

Quality of life – Child Health Questionnaire (CHQ)

The outcome measures for QoL in the abatacept study were summary physical scores, summary psychosocial scores (both measured on a 100-mm VAS) and 15 CHQ health concepts.⁵⁸ Differences between the abatacept and placebo treatment groups were not statistically significant for either the reported summary scores (p=0.666 and p=0.056, respectively), although there appears to be a positive trend for the latter (Table 19). Abatacept-treated patients (n=52) had improved scores for 14 of the 15 subscales and placebo-treated patients (n=34) for 6 of the 15 CHQ subscales (p >0.05 for abatacept versus placebo for all subscales; details not data extracted).

Table 19 Childhood Health Questionnaire (CHQ)

Study (Length: OL, RCT), Outcome	Intervention	Comparator	p value
Abatacept⁵⁷ (4mo OL, 6mo RCT)	ABA (n=52)⁵⁸	PBO (n=34)⁵⁸	p value
CHQ - Physical summary score	43.6	41 ²	p=0.666
CHQ - Psychosocial summary score	51.7	47 ²	p=0.056

ABA, abatacept. CHAQ, Childhood Health Assessment Questionnaire mo, months. OL, open-label. PBO, placebo. RCT, randomised controlled trial.

¹ Original group sizes were ABA n=60 & placebo n=62 but not all participants contributed data to these analyses.

² Estimated from graph by reviewer. Number of patients in the trial arms not clear.

Adverse events (AE)

A summary of AE reporting during the double-blind withdrawal trial phases is provided here with complete details for the AEs reported by each of the studies available in the data extraction forms (Appendix 5). Adverse events reported during trial OLEs are presented in a following section (*Adverse events OLE*).

Abatacept⁵⁷

During the six-month double-blind withdrawal period there were no statistically significant differences in AEs between the abatacept and placebo treatment groups. The total number of AEs (occurring in ≥5% of patients in the open-label and double-blind phase) was 62% for the abatacept and 55% for the placebo group, with two serious AEs occurring in the placebo group but none in the abatacept group (Table 20). The most common class of AEs in both treatment groups were infections

and infestations (44 - 45%). Adverse events were also reported under the headings of gastrointestinal disorders, general disorders and administration site conditions, nervous system disorders and respiratory, thoracic and mediastinal disorders (Appendix 5).

Adalimumab⁶¹

There were 155 AEs in the adalimumab group (10.3 per patient-year) and 234 in the placebo group (12.8 per patient-year) during the eight-month double-blind period. No statistical comparisons in AEs between treatment groups were reported. Only one serious AE possibly related to study drug was reported and this was gastroduodenitis which occurred in one patient in the adalimumab group. The most common AEs were related to injection-site reactions (ADA: 57 events in 3.8 patient-years; PBO: 73 events in 4.0 patient-years). Other reported AEs were contusion, nasopharyngitis, upper respiratory tract infection, viral infection, vomiting and excoriation (Appendix 5). No AE lead to the discontinuation of the treatment drug (Table 20). Sixteen percent of patients (27/171) had at least one positive test for anti-adalimumab antibody during the open-label and double-blind phases (methotrexate: 5/85 -6%, No methotrexate: 22/86 - 26%), but this did not lead to a greater rate of discontinuation of the study drug, nor did it increase the incidence of serious AEs. The study authors state that there was no occurrence of opportunistic infections, malignant conditions, demyelinating diseases or lupus-like reactions.

Etanercept⁴²

Two patients who received etanercept needed hospitalisation for serious AEs (one for depression and a personality disorder, and the other for gastroenteritis-flu syndrome). It is not clear at what point in the trial these events occurred. One patient withdrew after the first dose of etanercept (presumably at the start of the open-label period) because of urticaria. There were only two reported injection-site reactions during the double-blind phase of the trial, one in each treatment group (Table 20). All other AEs were reported to be of mild-to-moderate intensity, with no significant difference in the frequency of AEs between the treatment groups during the double-blind phase. There were no laboratory abnormalities requiring urgent treatment in the etanercept group. No patient had persistent elevations in autoantibodies or had signs or symptoms of another autoimmune disease. Two patients tested positive for non-neutralising antibody to etanercept.

Tocilizumab⁶⁸

The safety population consisted of all patients who received ≥ 1 dose of study medication. During the double-blind period, the total number of patients with at least one AE was 58 in the tocilizumab group (454.7 AEs per 100 patient-years) and 60 in the placebo group (514.4 AEs per 100 patient years). The most frequently reported AE in both treatment groups was nasopharyngitis (TCZ: 17%, PO 11%). Other reported AEs occurring were headache, upper respiratory infection, cough, pharyngitis, nausea,

diarrhoea, rhinitis, vomiting, abdominal pain, oropharyngeal pain, and rash (Appendix 5). Two AEs led to drug discontinuation, one in each treatment group (TCZ: increased blood bilirubin level; PO: gastroenteritis) and 3.7% of patients in each treatment group had ≥ 1 serious AE. One patient in the tocilizumab group suffered with ≥ 1 infectious serious AE. Rates of serious AEs per 100 patient-years were similar between groups (TCZ: 9.3; PO: 10.9), while the rate of infections serious AEs per 100 patient-years was 3.1 for the tocilizumab group. Other reported serious AEs, included pneumonia, upper limb fracture, uveitis, psychosomatic disease, enterocolitis and complicated migraine, with one case in each category and varying by treatment group (Appendix 5).

Table 20 Adverse events (AEs)

Study, Outcome	Intervention	Comparator	p value
Abatacept⁵⁷ (during 6-month DB period)	ABA (n=60)	PBO (n=62)	
Total serious AEs, n (%)	0	2 (3)	0.50
Total AEs, ¹ n (%)	37 (62)	34 (55)	0.47
Adalimumab⁶¹ (during 8-month DB period)	ADA (+MTX) (n=37; 15 Pt-yrs)	PBO (+MTX) (n=38; 18.3 Pt-yrs)	
Any AE, n of events (n of events per patient-year)	155 (10.3)	234 (12.8)	
Serious AEs, possibly related to study drug, ² n of events (n of events per patient-year)	1 (0.1)	0	
AEs leading to the discontinuation of the drug, n	0	0	
Etanercept⁴² (time period unclear unless stated below)	ETA (n=25)	PBO (n=26)	
Hospitalisation for serious AEs, n	2	0	
Injection-site reactions during the 4-month double-blind period, n	1	1	
Most common AEs - injection-site reaction, no. of events (no. of events per patient-year)	57 (3.8)	73 (4.0)	
Tocilizumab⁶⁸ (during 6-month DB period)	TCZ⁴ (n=82)	PBO⁴ (n=81)	
Serious AEs and AEs occurring $\geq 5\%$ of patients, n (%)			
Duration in study (years)	32.33	27.41	
Patients with ≥ 1 AE	58 (70.7)	60 (74.1)	
Total no. of AEs ⁵	147	141	
Rate of AEs per 100 patient-years	454.7	514.4	
Serious AEs			
Patients with ≥ 1 serious AE	3 (3.7)	3 (3.7)	
Rate of serious AEs per 100 patient-years	9.3	10.9	
Patients with ≥ 1 infectious serious AE	1 (1.2)	0	
Rates of infectious serious AEs per 100 pt-years	3.1	0	
AEs leading to study drug discontinuation	1 (1.2) ⁶	1 (1.2) ⁷	

ABA, abatacept. ADA, adalimumab. AE, adverse event. DB, double-blind. ETA, etanercept. MTX, methotrexate. no, number. PBO, placebo. Pt-yrs, patient years. NR, not reported. TCZ, tocilizumab.

¹ AEs that occurred in $\geq 5\%$ of patients in the open-label and double-blind phases.

² Serious AEs were death or any event that was life-threatening; required hospitalisation or prolongation of existing hospitalisation; resulted in persistent or significant disability, congenital anomaly, or spontaneous or elective abortion; or required medical or surgical intervention to prevent another serious outcome.

³ After 1st dose of etanercept (responded to oral antihistamines).

⁴ AE data on open-label TCZ escape therapy were excluded

⁵ Multiple occurrences of the same AE in one individual were counted.

⁶ Increased blood bilirubin level, highest total bilirubin reading, 50 μ mol/L (normal range, 3–24 μ mol/L); 2 consecutive readings >51 μ mol/L mandated withdrawal per protocol. The event resolved without sequelae.

⁷ Gastroenteritis occurred 46 days after the last of five doses of placebo.

Sub-group analyses

Only the tocilizumab study⁶⁸ reported sub-group analyses, which reported on ACR Pedi 70 and ACR Pedi 90 response at week 40 in three sub-groups: patients with background treatment of methotrexate, background treatment of glucocorticoid and previous biologic agent use at baseline. It is unclear if these analyses were pre-planned or post-hoc. The trial authors also stated that no differences were observed in response to tocilizumab between patients who were RF+ve and those who were not, but no data in support of this statement were presented. No statistical comparisons between treatment groups were reported and it is therefore unclear if differences between the sub-groups were statistically significant.

Background methotrexate

Patients receiving background methotrexate in both the tocilizumab and the placebo groups had higher ACR Pedi 70 and 90 response rates at the end of the double-blind RCT withdrawal phase compared to those who were not in receipt of background methotrexate (Table 21). However, patients receiving tocilizumab with or without background methotrexate had better response rates than patients in the corresponding placebo groups.

Background glucocorticoid

At the end of the double-blind RCT phase (week 40) in the tocilizumab group, a slightly higher proportion of participants receiving background glucocorticoid achieved a ACR Pedi 70 and 90 response compared to those who were not in receipt of background glucocorticoid (Table 21). However, among participants in the placebo group the opposite pattern was observed, with a lower proportion of those who were in receipt of background glucocorticoid achieving ACR Pedi 70 and 90 (Table 21). Response rates for both ACR Pedi 70 and 90 were higher in sub-groups of patients receiving tocilizumab compared to placebo sub-groups regardless of whether or not patients received background glucocorticoid.

Previous biologic agent

Patients in either the tocilizumab or placebo groups who had received previous treatment with a biologic agent (primarily comprising anti-TNF agents) had lower ACR Pedi 70 responses at the end of the double-blind RCT phase (week 40) compared to patients who had not previously been treated with a biologic agent (Table 21). Patients receiving placebo who had not received previous treatment with a biologic agent had better ACR Pedi 70 and 90 response rates compared to patients on tocilizumab who had previous biologic agent experience (Table 21).

Table 21 ACR Pedi response by background medication use at baseline at the end of the double-blind RCT phase

Tocilizumab⁶⁸ (4months OL, 6 months RCT)					
Proportion of patients in the ITT population with ACR Pedi 70 and ACR Pedi 90 response at the end of the double-blind phase (week 40) by background methotrexate, glucocorticoid and previous biologic agent use at baseline ¹					
Concomitant therapies and previous exposure to biologic agent, n/N (%)	Response level	TCZ (n=82)		PBO (n=81)	
		Yes	No	Yes	No
Background methotrexate	ACR Pedi 70	45/67 (67.2)	8/15 (53.3)	30/64 (46.9)	4/17 (23.5)
	ACR Pedi 90	32/67 (47.8)	5/15 (33.3)	18/64 (28.1)	1/17 (5.9)
Background glucocorticoid	ACR Pedi 70	23/33 (69.7)	30/49 (61.2)	4/38 (36.8)	20/43 (46.5)
	ACR Pedi 90	16/33 (48.5)	21/49 (42.9)	5/38 (13.2)	14/43 (32.6)
Previous biologic agent	ACR Pedi 70	13/27 (48.1)	40/55 (72.7)	2/23 (8.7)	32/58 (55.2)
	ACR Pedi 90	5/27 (18.5)	32/55 (58.2)	2/23 (8.7)	17/58 (29.3)

ITT, intention to treat. OL, open-label. PBO, placebo. TCZ, tocilizumab.

¹ Patients who withdrew or escaped to open-label TCZ or for whom the end point could not be determined were classified as non-responders.

Results - Open-label extensions (OLE)

All four studies included OLEs with some differences in which participants were eligible to enter, and how data were presented. ACR Pedi results are presented below and with additional outcomes presented either in the study data extraction forms (Appendix 5) or published papers (Adalimumab: minimal disease activity; Abatacept: ACR Pedi component items, analysis according to prior exposure to biologic agents, ACR Pedi data for those in the OLE who had not taken part in the double-blind phase and information on anti-abatacept and anti-CTLA-4 antibody production; Etanercept: ACR Pedi component items, minimal disease activity; Tocilizumab: ACR Pedi component items, minimal disease activity).

Abatacept⁵⁹

The abatacept study reported ACR Pedi data separately for those who had been treated with abatacept continuously (lead-in, double-blind & OLE phases) and those whose abatacept had been interrupted by placebo during the double-blind-RCT phase. The OLE included 85% of the abatacept group and 76% of the placebo group from the double-blind phase. For those receiving continuous abatacept therapy treatment length ranged from 31 to 52 months (participants who had entered the study earliest had been treated longest). Those who received placebo during the double-blind phase usually received abatacept for a shorter period (length not stated), but the ACR Pedi scores achieved were similar for ACR Pedi 30, 50 and 70 to those whose abatacept treatment had been continuous (Table 22). The proportions of participants who had received placebo during the double-blind phase achieving ACR Pedi 90 and ACR Pedi 100, and having inactive disease are lower than those whose abatacept treatment had been continuous.

Table 22 ACR Pedi outcomes from trial open-label extension periods (OLE)

Study (follow-up), Outcome	Intervention	Comparator (during RCT phase)
Abatacept⁵⁹ (OLE day 589)		
ACR Pedi, n/N (%)	ABA (n= 51)	PBO (n= 47)
30	46/51 (90%)	41/47 (87%)
50	45/51 (88%)	39/47 (83%)
70	38/51 (75%)	35/47 (75%)
90	29/51 (57%)	19/47 (40%)
100	20/51 (39%)	9/47 (19%)
Inactive disease	22/51 (43%)	11/47 (23%)
Adalimumab⁶¹ (OLE week 104)		
ACR Pedi, %	ADA (n=128^a)	PBO group from RCT phase not separately reported
30	89%	
50	86%	
70	77%	
90	59%	
100	40%	
Etanercept⁶⁷ (OLE up to 8 years)		
ACR Pedi response, 8 years (LOCF^b), n/N (%)	ETA (n=58^c)	PBO group from RCT phase not separately reported
ACR Pedi 30	40/48 (83%)	
ACR Pedi 50	36/47 (77%)	
ACR Pedi 70	28/46 (61%)	
ACR Pedi 90	19/46 (41%)	
ACR Pedi 100	8/45 (18%)	
Tocilizumab^{69:71:74} (104 weeks)		
ACR Pedi, proportion of patients with improvement relative to baseline, n (%)⁶⁹	TCZ (n= 82)	PBO (n= 73^d)
ACR Pedi 70 ^e	71/82 (86.6%)	NR
ACR Pedi 90 ^e	58/82 (70.7%)	NR
Proportion with inactive disease ^f	52/82 (63.4%)	NR

ABA, abatacept. ADA, adalimumab. ETA, etanercept. LOCF, last observation carried forward. NR - not reported. OLE, open-label extension. PBO, placebo. TCZ, tocilizumab.

^a Only 71/128 (58%) of this group received methotrexate during the open-label and double-blind phases of the study and meet the licenced indication for adalimumab

^b A LOCF analysis was necessary because data were not available for all participants who entered the 8th year of follow-up (n=26) and because the remaining 32/58 (55%) of participants had discontinued the OLE already.

^c Total number of participants who entered the OLE. As this is greater than the total number of participants who took part in the double-blind phase of the study (n=51) it is presumed that some of these participants entered the OLE directly from the initial open-label treatment phase of the study.

^d n calculated by reviewer (155 completed 104 weeks - 82 TCZ group completed 104 weeks)

^e Two abstracts^{69:74} contain a table with a footnote to indicate patients who withdrew were excluded, however in the third abstract⁷¹ the table footnote states that patients who withdrew due to non-safety reasons are non-responders whereas patients who withdrew due to safety are included using LOCF.

^f no active joints, no active uveitis, ESR <20 mm/h, and physician global assessment VAS ≤10.

Adalimumab⁶¹

Results were reported for those who entered the OLE phase as a single group of participants (n=128). This group included 35 of 38 (92%) participants who received adalimumab and methotrexate and 36 of 37 (97%) participants who received placebo and methotrexate in the double-blind phase of the study. However, also within this group of 128 are 57 (45%) participants from two further study arms (adalimumab only and placebo only) which are not included in this report because the majority of

participants in these arms had never received methotrexate and therefore do not meet the licenced indication. Through the first 104 weeks of the OLE phase there was no diminution of the ACR Pedi responses, such that after 104 weeks of open-label treatment in the extension phase, 40% of participants had an ACR Pedi 100 response (Table 22).

Etanercept⁴²

All 69 participants who began the open-label lead-in phase of the study (51 of whom took part in the double-blind randomised withdrawal phase) were eligible to enter the OLE phase but only 58 did so. Of the 58 who took part in the OLE, 26 entered the eight year of follow-up; therefore a last observation carried forward (LOCF) analysis was used to calculate the ACR Pedi responses reported in Table 22. These responses appear to have remained constant over the OLE. While LOCF analyses are commonly used in drug trials, this method can be prone to bias when used in progressive diseases such as JIA and results should be interpreted with caution.

Tocilizumab⁶⁸

Results from the OLE of the tocilizumab study⁶⁸ are reported in conference abstracts.^{69;71;74} Only participants who achieved at least ACR Pedi 30 during the open-label-phase and who then continued into the double-blind RCT phase of the trial were eligible to enter the OLE, either after a JIA flare or when they completed the double-blind RCT phase. One hundred and sixty (96%) of the 166 participants eligible to enter the OLE did so and 155 (97%) completed 104 weeks of follow-up (16 week open-label + 24 weeks double-blind RCT + 64 weeks OLE). ACR Pedi 70, ACR Pedi 90 and proportion with inactive disease are presented (Table 22) only for the 82 participants who received continuous tocilizumab throughout the study and the proportion achieving each of these measures increased since the end of the double-blind phase (Table 22).

Growth

Adalimumab⁶¹

Two abstracts^{62;63} report limited data for growth from a post-hoc analysis of JIA patients who had taken part in any arm of the double-blind phase of the RCT and entered the OLE (this includes n=58 who were not receiving methotrexate and who were therefore not receiving adalimumab treatment according to the licenced indication). All patients who received ≥ 1 dose of adalimumab +/- methotrexate were included in the analysis (n=133). Patients were assigned by baseline weight into 2 groups: $\leq 33^{\text{rd}}$ percentile (41%, n=55) and $>33^{\text{rd}}$ percentile (59%, n=59) based on the US Centers for Disease Control and Prevention (CDC) growth charts. Missing data were analysed using LOCF. Those in the $\leq 33^{\text{rd}}$ percentile baseline weight group had a higher mean percentile change from baseline in height at week 104 than those in the $>33^{\text{rd}}$ percentile group (values for mean height percentile change from baseline estimated from graph by reviewer 5.5 and 3.3 respectively).⁶² Similar

patterns were stated to have been observed for weight and body mass index (BMI). At week 104, there were no statistically significant differences between the methotrexate and the non-methotrexate groups in mean changes from baseline in height, weight, or body mass index ($p>0.26$). Long-term adalimumab treatment appears to show improvement in growth for JIA patients who were in the $\leq 33^{\text{rd}}$ percentile weight group at baseline receiving adalimumab with or without the addition of methotrexate. However, caution in the interpretation of the results is recommended due to the limitations in the data presented and the absence of an appropriate control group.

Tocilizumab⁷⁶

The Roche CS included data for growth and glucocorticoid treatment at week 104 based on a conference abstract⁷⁶ from the CHERISH trial.⁶⁸ Most growth data came from a subset of patients ($n=123$) with the highest growth potential, represented by patients with Tanner stage <4 at baseline. The Tanner stages are based on a scale of physical development in children, adolescents and adults (boys - development of external genitalia; girls - breast development; boys and girls - pubic hair), with stage 5 being the final adult/mature stage. Growth measures included height standard deviation scores (SDS) and height velocity. The mean height SDS of patients with polyarticular JIA and Tanner stage <4 was below normal at baseline (-0.68 SD 1.23) and rose to -0.19 (SD 1.14) at week 104 ($n=103$) with the difference being statistically significant ($p<0.001$ vs baseline). Of these patients, 71.8% had an increased height SDS. The CS states (CS page 16) that there was no observed difference in patients who received placebo during the randomised phase of the trial (based on 154 patients of the growth population with height SDS data at both time points) however fewer than half the patients received placebo through the entire 24 weeks of the randomised phase of the trial, as most escaped to tocilizumab before week 40. For the entire growth population ($n=187$, i.e. not restricted to those with Tanner stage <4), the reported mean change in height SDS from baseline to week 104 was 0.25 (SD 0.54) (no p value for comparison with baseline reported). The mean daily oral glucocorticoid dose decreased from baseline (0.05 (SD 0.08) mg/kg) to week 104 ($n=103$) (0.02 (SD 0.05) mg/kg). A multiple linear regression analysis for the same 103 patients indicated that height velocity at week 52 was related to baseline age ($p<0.001$) and oral glucocorticoid use at the end of week 52 ($p=0.0002$). No data for week 104 was reported. Caution in the interpretation of the growth results is recommended due to the limitations in the data presented and the absence of an appropriate control group.

Adverse events

This is a summary of OLE AEs presented in the published papers.

Abatacept - OLE to day 589 and year 7

In the abatacept study,⁵⁷ common AEs (occurring in 10% or more of the total group, not data extracted) and common serious AEs (occurring in 1% or more of the total group) were reported separately for those who had been in the abatacept group and those who had been in the placebo group during the double-blind period of the trial, and those who had not entered the double-blind phase because they did not achieve an ACR Pedi 30 response during initial open-label treatment. Serious AEs by day 589 (approximately 20-21 months) occurring in 23/153 patients (Table 23), the most common were arthritis flares (n=6), arthralgia (n=2), foot deformity (n=2), pyrexia (n=2), and vomiting (n=2). The proportions of serious adverse events at day 589 were similar in the three groups. At 7 year follow-up (reported in an abstract⁶⁰), 30/153 (19.6%) patients had serious AEs. Most were unrelated and were primarily musculoskeletal or infectious events. The incidence rate (per 100 patient-years) of serious AEs in the OLE at 7 years (5.6/100 patient years) did not increase versus the 6-month double-blind rate (6.8/100 patient years).

Table 23 OLE adverse events for abatacept

Abatacept⁵⁷ (OLE: day 589,⁵⁹ 7 years⁶⁰)			
Serious AEs, n (%)	DB ABA (n=58)¹	DB PBO (n=59)¹	Patients with less than an ACR-Pedi 30 response initially (n=36)
Total serious AEs, n/N (%)	8/58 (14)	8/59 (14)	7/36 (19)
Most common serious AEs			
Arthritis flares ²	3 (5.2)		3 (8.3)
Arthralgia ²	1 (1.7)	1 (1.7)	1 (2.8)
Foot deformity ²	1 (1.7)	1 (1.7)	
Pyrexia	1 (1.7)	1 (1.7)	
Vomiting		1 (1.7)	
Serious AEs Year 7, n/N (%)	30/153 (19.6)		

ABA, abatacept. AEs, adverse events. DB, double blind OLE, open-label extension. PBO, placebo.

¹ patients who had been in the randomised double-blind phase.

² all related to underlying disease.

Adalimumab - OLE ongoing

The OLE was ongoing at the time the key trial publication was published and the time period for which events were reported is not clear.⁶¹ Serious AEs considered possibly related to study drug occurred in seven patients during the OLE (a table in the published paper⁶¹ suggests none were receiving methotrexate, in which case they were not receiving adalimumab treatment according to the licenced indication). Three patients discontinued treatment due to AEs during the OLE.

Etanercept – year 8

In the etanercept study,⁴² OLE the safety analyses captured serious AEs, medically important infections (MII) and mortality as well as some ‘Events of interest’ (these included opportunistic infections, tuberculosis, lupus, demyelinating disorders, malignancies, and lymphomas). Non-serious AEs were not recorded.⁶⁷ There were a total of 39 serious AEs based on 318 patient-years of etanercept exposure (n=69), with 26 patients entering their eighth year of etanercept treatment,

equating to 0.12 events per patient year (Table 24). There were nine MII resulting in the need for intravenous antibiotic therapy or hospitalisation, equating to 0.03 events per patient years, with only one reported MII since four years (pyelonephritis). The most common new serious AEs reported beyond four years of drug exposure were flares or worsening of disease, occurring in 6/9 patients (67%).

Table 24 OLE adverse events for etanercept

Etanercept⁴² (OLE: up to 8 years⁶⁷)				
Year of etanercept treatment from RCT (excluding gaps between RCT and OLE)	Serious AE¹		MII²	
	No. of events	No. of events/patient year	No. of events	No. of events/patient year
1 (n=69; 57 patient-years of drug exposure)	5	0.09	2	0.04
9 (n=14; 4 patient-years of drug exposure)	0	0	0	0
Total for all years (n=69; 318 patient-years of drug exposure)	39	0.12	9	0.03

AE, adverse events. MII, medically important infections. No, number. OPE, open-label extension.

¹ Serious AEs occurring during the study or within 30 days of the last dose of etanercept. Defined as events that were fatal or life-threatening, required hospitalisation or prolonged an existing hospitalisation, resulted in a persistent or significant disability or incapacity, or resulted in a congenital anomaly or birth defect.

² Defined as medically important infections resulting in the need for intravenous antibiotic therapy or hospitalisation.

Tocilizumab - 104 weeks

Long-term AEs rates based on a safety population of 188 patients (307 patient-years) were 406.5 per 100 patient-years over 104 weeks (approximately 2 years) in patients receiving tocilizumab, based on an abstract only.⁶⁹ The equivalent serious AEs rate was 11.1 per 100 patient-years (Table 25).

Infections categorised into the most common AEs and serious AEs were 151.4 and 5.2 per 100 patient years respectively. The study also reports AE safety population data for elevations of alanine aminotransferase and aspartate aminotransferase, grade 2/3/4 thrombocytopenia, grade 3 lowest neutrophil count and low-density lipoprotein cholesterol (see data extractions in Appendix 5).

Table 25 OLE adverse events for tocilizumab

Tocilizumab⁶⁸(OLE: 104 weeks⁶⁹)	
AEs and SAEs	Safety population=188 patients with 307 patient-years of tocilizumab exposure
AEs, rates/100 patient-years	406.5
Serious AEs, rates/100 patient-years	11.1
Most common AE - infections	151.4
Infections – serious AE	5.2

AE, adverse events. SAE, serious adverse events, OLE, open-label extension.

4.1.3 Assessment of clinical-effectiveness - biologic DMARDs vs each other (with methotrexate where permitted)

Background

None of the RCTs included in the systematic review of clinical-effectiveness directly compared any of the biologic DMARDs with each other. It was therefore necessary to undertake an indirect comparison of the drugs to inform the assessment of comparative clinical-effectiveness. One published indirect comparison was identified through literature searching, by Otten and colleagues (2012).⁷⁹ This was a systematic review of RCTs that constructed two separate evidence networks: polyarticular course JIA, and systemic JIA. For each network a series of pairwise indirect comparisons was conducted, with placebo as a common comparator, using the method described by Bucher and colleagues (1997).⁵³

Three RCTs were included in Otten and colleagues'⁷⁹ polyarticular course JIA network,^{42;57;61} all of which have been included in this assessment report. However, this network did not include tocilizumab, as at that time no RCT evidence for that drug in polyarticular course JIA was published. The network therefore only included comparisons of three of the four biologic DMARDs of relevance to the scope of this assessment (abatacept, adalimumab and etanercept). We have conducted a similar adjusted indirect comparison to Otten and colleagues'⁷⁹ including the recently published tocilizumab RCT by Brunner and colleagues (the CHERISH trial).⁶⁸ Figure 2 illustrates the design of the analysis, representing what is termed a star network.⁸⁰

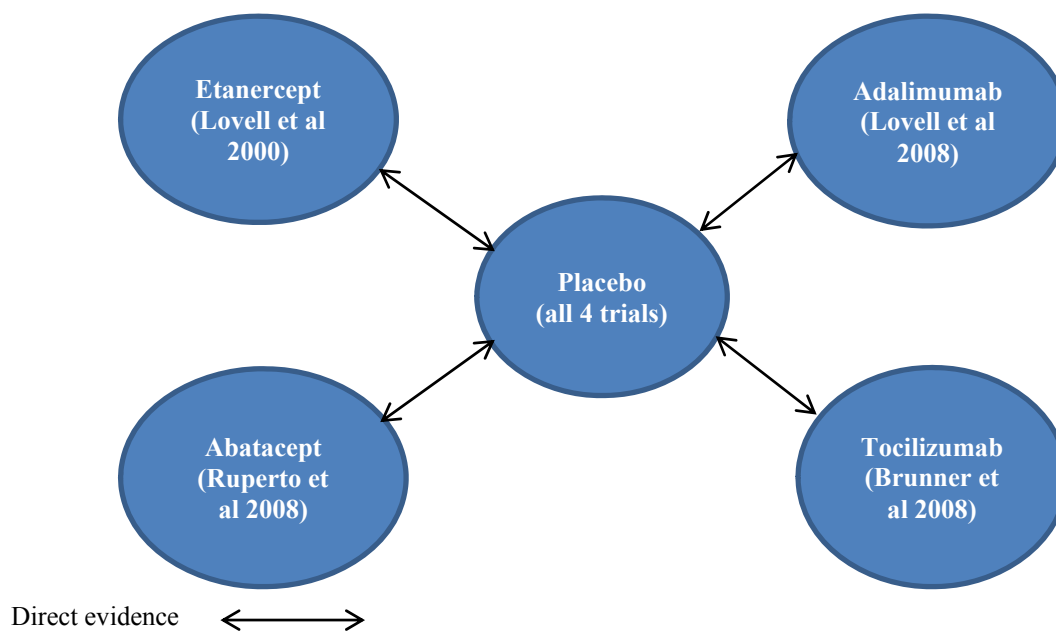


Figure 2 Indirect comparison of biologic DMARDs

Otten and colleagues⁷⁹ indirectly compared the drugs in relative risk (RR) of disease flare. We have similarly included disease flare as an outcome, and in addition have chosen ACR Pedi 50 and 70 response as an outcome. ACR Pedi 50 and 70 was chosen as opposed to ACR 30 as it was considered that a higher level would be a more clinically relevant level of treatment response. Furthermore, due to the design of the RCTs all patients who were randomised had achieved an ACR Pedi 30 response at the end of the open-label lead-in phase.

The adjusted indirect comparison should be considered to be exploratory rather than definitive due to limitations in the evidence base, and heterogeneity between the included trials. Specifically:

- There is only one trial available for each drug. Although the trials were considered to be of generally good methodological quality and low risk of bias, evidence networks are considered to be weaker if informed by small numbers of studies and small numbers of participants.⁸¹ The number of patients in the trials was also relatively low (ranging from 51 to 163).
- There is some variation in the proportion of sub-types of JIA in the included trials. Although the network is considered to be most applicable to polyarticular course JIA, in one trial (etanercept⁴²) around a third of patients were classified as having systemic JIA with apparent systemic manifestations (which is outside the scope of the current appraisal). There was insufficient evidence from RCTs to construct a network for PA, or ERA because outcome data for these subtypes of JIA were not reported separately in trials even though some cases with these subtypes may have been included.
- The duration of JIA ranged from just under four years⁵⁷ to approximately six years across the trials.⁴²

██
██

██ Disease duration has also been found to be a predictor of response to etanercept⁸³ and to methotrexate⁸⁴ among patients from the German BIKER registry. Differences between the trials in disease duration may therefore potentially confound results.

- Three of the four trials permitted patients to take methotrexate in addition to the biologic DMARD (in proportions of patients varying from 74% to 100%), whilst the fourth (etanercept) did not permit use of methotrexate.
- Previous therapy with biologic DMARDs had been received by approximately a third of participants who entered the initial open label run-in of two trials (abatacept⁵⁷ and tocilizumab⁶⁸). Prior therapy with another biologic DMARD was an exclusion criterion for the adalimumab trial⁶¹ and was not mentioned for the earliest trial (etanercept⁴²) presumably because no other biological

therapies were available at the time. Currently it is unclear whether prior biologic DMARD treatment influences the effectiveness of subsequent biologic treatment.

- The mean age of patients across the trials varied from around 7.5 years to 13 years. Part of this variation may reflect the age ranges specified in the inclusion criteria of the trials and potentially the mix of JIA subtypes in the trials which have a different mean age of onset. Age could be an effect modifier given the progressive nature of JIA.
- The duration of the double-blind randomised treatment phase of the trials varied, from four months⁴², to eight months⁶¹. Treatment duration may affect outcomes which are time dependent, such as disease flare.

Results

Figure 3 illustrates the results of the four included RCTs comparing the biologic DMARDs to placebo (with background methotrexate where permitted) on the outcome of disease flare (NB. The adalimumab trial⁶¹ stratified results according to whether or not patients received methotrexate background therapy, and we have only included data for patients who did receive methotrexate, in accordance with the licensed indication – this applies to disease flare and to ACR Pedi 50 / 70). Treatment with each of the four DMARDs resulted in a statistically significant reduction in the RR of a disease flare, ranging from 0.38 to 0.57. (NB. we have not presented a pooled RR given differences between the DMARDs and also heterogeneity between the trials).

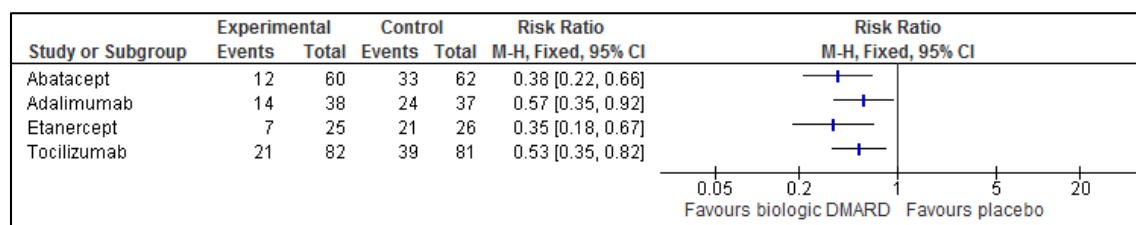


Figure 3 Summary forest plot of biologic DMARDs versus placebo: disease flare

Table 26 reports adjusted pairwise indirect comparisons for the four biologic DMARDs for the outcome of disease flare. The point estimate for risk of flare was lower for etanercept than the other three comparators. Abatacept had a lower risk of flare compared to adalimumab and to tocilizumab. Tocilizumab had a lower risk of flare compared to adalimumab only. Adalimumab was associated with a higher risk of disease flare than the other three comparators. The ranking of treatments in terms of risk of flare was therefore etanercept, abatacept, tocilizumab, and adalimumab. However, none of the comparisons demonstrated a statistically significant difference between the treatments being compared, with confidence intervals crossing one in every case. The results of our analysis match

those of Otten and colleagues,⁷⁹ with the exception of the comparison with tocilizumab which was not included in their polyarticular course JIA trial network (as discussed above).

Table 26 Indirect comparisons of biologic DMARDs: disease flare

Comparison	Relative risk
Etanercept vs adalimumab	0.61 (95% CI 0.27 to 1.38)
Etanercept vs abatacept	0.92 (95% CI 0.39 to 2.18)
Etanercept vs tocilizumab	0.65 (95% CI 0.30 to 1.43)
Adalimumab vs abatacept	1.51 (95% CI 0.72 to 3.15)
Adalimumab vs tocilizumab	1.07 (95% CI 0.56 to 2.04)
Abatacept vs tocilizumab	0.71 (95% CI 0.35 to 1.43)

CI, confidence interval.

Figure 4 illustrates the results of the four included RCTs comparing the biologic DMARDs to placebo (with background methotrexate where permitted) on the outcome of ACR Pedi 50 response.

Treatment with each of the four DMARDs led to a statistically significant greater proportion of participants with ACR Pedi 50 response, with RR ranging from 1.41 to 3.12.

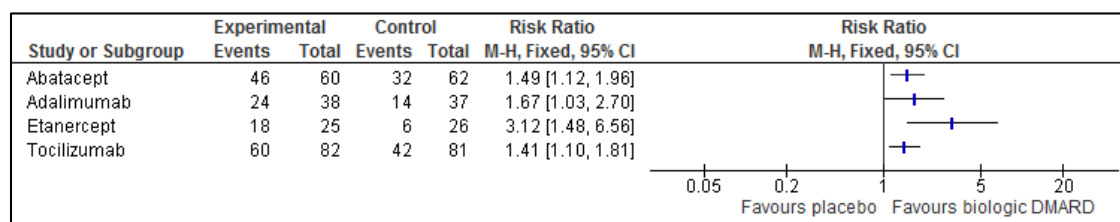


Figure 4 Summary forest plot of biologic DMARDs versus placebo: ACR Pedi 50 response

Table 27 reports adjusted pairwise indirect comparisons for the four biologic DMARDs for the outcome of ACR Pedi 50 response. Etanercept had a higher RR for treatment response than the other three comparators. Adalimumab had a higher RR for treatment response than abatacept and tocilizumab. Adalimumab had a higher RR for treatment response than tocilizumab. The ranking of treatments in terms of treatment response was therefore etanercept, adalimumab, abatacept, and tocilizumab. With the exception of etanercept compared with tocilizumab, none of the comparisons indicated a statistically significant difference between the treatments being compared, with confidence intervals crossing one.

Table 27 Indirect comparisons of biologic DMARDs: ACR Pedi 50 response

Comparison	Relative Risk
Etanercept vs adalimumab	1.87 (95% CI 0.77 to 4.53)
Etanercept vs abatacept	2.10 (95% CI 0.95 to 4.64)

Etanercept vs tocilizumab	2.21 (95% CI 1.01 to 4.84)
Adalimumab vs abatacept	1.12 (95% CI 0.65 to 1.96)
Adalimumab vs tocilizumab	1.18 (95% CI 0.69 to 2.02)
Abatacept vs tocilizumab	1.05 (95% CI 0.72 to 1.53)

CI, confidence interval.

Figure 5 illustrates the results of the four included RCTs comparing the biologic DMARDs to placebo (with background methotrexate where permitted) on the outcome of ACR Pedi 70 response.

Treatment with each of the four DMARDs led to a statistically significant greater proportion of participants with ACR Pedi 70 response, with RR ranging from 1.54 to 2.34.

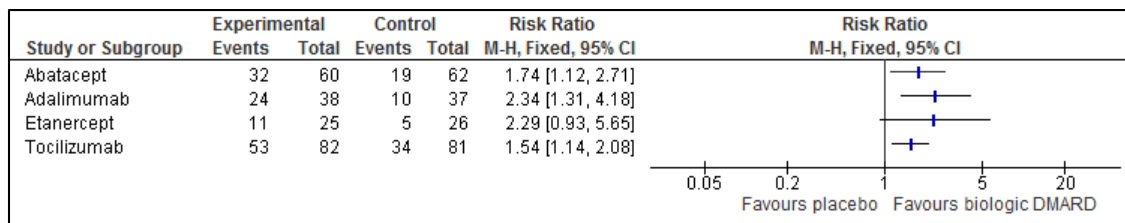


Figure 5 Summary forest plot of biologic DMARDs versus placebo: ACR Pedi 70 response

Table 28 reports adjusted pairwise indirect comparisons for the four biologic DMARDs for the outcome of ACR Pedi 70 response. Etanercept had a higher RR for treatment response than abatacept and tocilizumab but a slightly lower RR for treatment response compared to adalimumab.

Adalimumab had a higher RR for treatment response than abatacept and tocilizumab. Abatacept had a higher RR for treatment response than tocilizumab. The ranking of treatments in terms of treatment response for ACR Pedi 70 (adalimumab, etanercept, abatacept, and tocilizumab) was therefore different than for the ACR Pedi 50. None of the comparisons indicated a statistically significant difference between the treatments being compared, with confidence intervals crossing one.

Table 28 Indirect comparisons of biologic DMARDs: ACR Pedi 70 response

Comparison	Relative Risk
Etanercept vs adalimumab	0.98 (95% CI 0.33 to 2.87)
Etanercept vs abatacept	1.31 (95% CI 0.48 to 3.60)
Etanercept vs tocilizumab	1.49 (95% CI 0.57 to 3.85)
Adalimumab vs abatacept	1.34 (95% CI 0.65 to 2.79)
Adalimumab vs tocilizumab	1.52 (95% CI 0.79 to 2.92)
Abatacept vs tocilizumab	1.13 (95% CI 0.66 to 1.93)

CI, confidence interval.

The results of this exploratory analysis based on the limited evidence available currently (only one trial for each of the four biologic DMARDs) supports etanercept being more effective than the other three biologic DMARDs in terms of preventing disease flares, and achieving a response to treatment based on a composite index (ACR Pedi 50) whereas the ACR Pedi 70 exploratory analysis shows

adalimumab with a slight advantage over etanercept (though see comment below about confidence intervals). Abatacept appeared to be superior to tocilizumab for all outcome measures. Adalimumab appeared to be less effective than abatacept and tocilizumab in terms of preventing disease flare, but appeared to be more effective than these two comparators in terms of ACR Pedi 50 response. Therefore, there was no consistent ranking of treatment comparisons across these outcome measures. The indirect comparisons were generally not statistically significant, and confidence intervals were wide so caution is advised in the interpretation of these results. Furthermore, the etanercept trial⁴² appears to have some differences from the other trials which may confound the results, namely the absence of methotrexate background therapy and the longer duration of JIA disease. There was also a noticeably higher rate of flares in the placebo arm of that trial compared to the other three trials (81% compared to 48% – 65%) which may account for the bigger treatment effect seen. Taking the above limitations into account an overall interpretation of the results of the indirect comparison is that, due to the absence of statistically significant differences between the biologic DMARDs, currently they appear to be similar in treatment effectiveness. This accords with the conclusion reached by Otten and colleagues who suggested that the short-term efficacy of the biologic DMARDs in polyarticular course JIA seem similar.⁷⁹ Furthermore, the clinical advisors to the assessment group felt that these data generally reflect clinical experience in that when used for the same indication in the same population effectiveness was likely to be similar. However there was also a recognition that for individual patients and potentially for particular sub-groups of JIA patients differential effects of each biologic DMARD might be apparent but these differential effects have not yet been captured by current trial data.

4.1.4 Summary of the systematic review of clinical-effectiveness

- Four multi-centre RCTs, one each evaluating abatacept, adalimumab, etanercept and tocilizumab met the inclusion criteria of this review. Only the tocilizumab RCT included UK patients. Seven additional RCTs (3 for adalimumab and 4 for etanercept) are described as ongoing (details summarised in section 4.3 ‘Ongoing trials’ below).
- Each RCT had three phases, an open-label lead-in period, a randomised withdrawal period, and an open-label extension. The lengths of the lead-in and randomised phases varied between studies (open-label lead-in: 12 to 16 weeks; randomised double-blind withdrawal phase 16 to 32 weeks). In each study patients had to achieve an ACR Pedi 30 response during the initial open-label phase in order to be eligible for entry to the randomised double-blind withdrawal phase, Hence results are only applicable to patients who have already achieved an initial (low) degree of benefit from a biologic DMARD.
- The quality of the included RCTs was reasonable overall, with a low risk of bias judged for most items, although some aspects were rated as unclear, mainly due to a lack of reporting.

- Disease flare: this was the primary outcome in all four RCTs, with definitions broadly consistent between the studies. Patients who continued to receive biologic DMARDs during the randomised withdrawal phase of the studies had statistically significantly fewer arthritis flares compared to those receiving placebo in all four studies, while time to disease flare reported in two studies (abatacept and etanercept) was statistically significantly longer in those treated with biologic DMARDs.
- ACR Pedi: a greater proportion of patients receiving biologic DMARDs during the randomised withdrawal phase of the studies achieved ACR Pedi responses of ≥ 30 compared with placebo-treated patients, with differences statistically significant in all but two instances where p-values were reported. The proportion of biologic DMARD-treated patients with inactive disease was more than twice that of placebo-treated patients in the two studies (abatacept and tocilizumab) reporting this outcome.
- ACR Pedi core variables: in the three studies reporting this outcome (abatacept, etanercept and tocilizumab) results were generally in favour of treatment with biologic DMARDs when compared to placebo.
- Joint related outcomes: one study (etanercept) reported additional joint outcomes without statistical comparisons. All outcomes favoured etanercept when compared to placebo.
- Pain: three studies reported pain (abatacept, etanercept and tocilizumab). A statistically significant difference in the mean change from baseline favoured the tocilizumab group. While pain scores were lower for those receiving biologic DMARDs in the remaining two studies, differences between treatment groups were not statistically significant in the abatacept study and no statistical comparison reported in the etanercept study.
- Mortality: no treatment-related deaths were reported in the three studies reporting this outcome (adalimumab, etanercept and tocilizumab).
- Outcomes not reported by the included RCTs for the randomised withdrawal phase of the trials were corticosteroid reducing regimens, extra-articular manifestations (such as uveitis), height and weight.
- HRQoL: reported by the abatacept study (abatacept) only, with differences between treatment groups for the physical and psychosocial summary scores not statistically significant. Those treated with abatacept had improved scores for 14 of the 15 CHQ subscales compared to 6 out of 15 for placebo-treated patients.
- AEs: during the randomised withdrawal phase of the trials the proportions of AEs and serious AEs were generally fairly similar between the biologic DMARDs and the placebo groups.
- Sub-group analyses: the tocilizumab study reported data for sub-groups but no statistical comparisons between treatment groups were reported.

- OLE: all four studies included an OLE phase. There were differences in eligibility criteria between studies and in how data were presented. Only results for ACR Pedi, AEs and growth are included in this report.
- OLE ACR Pedi:
 - Abatacept: the proportion of patients achieving ACR Pedi 30, 50 and 70 scores were similar for those with continuous abatacept therapy and those who received placebo during the double-blind phase, but was greater in achieving ACR Pedi 100 and inactive disease in abatacept-treated patients (ACR Pedi 100 abatacept 39%, placebo 19%; inactive disease abatacept 43%, placebo 23%).
 - Adalimumab: There was no diminution of ACR Pedi responses and 40% of patients had an ACR Pedi 100 response after open-label treatment in the extension phase, but results included patients not meeting the licensed indication.
 - Etanercept: 26/58 (45%) patients who took part in the OLE entered the 8th year of follow-up. ACR Pedi responses appear to have remained constant over the OLE.
 - Tocilizumab: limited results based on conference abstracts for 82 patients who received continuous tocilizumab throughout the study. The proportion of patients achieving ACR Pedi 70 and 90 increased since the end of the double-blind phase, with 63% having inactive disease.
- OLE AEs:
 - Abatacept: at seven year follow-up, 19.6% of patients had serious AEs, with similar incidence rates between the OLE phase (5.6 per 100 patient-years) and the six-month double-blind phase (6.8 per 100 patient years).
 - Adalimumab: serious AEs considered possibly related to study drug occurred in seven patients during the OLE, but would appear to be in patients not in line with licensed indication (OLE phase ongoing, time period unclear). Three patients discontinued treatment due to AEs.
 - Etanercept: there were a total of 39 serious AEs based on 318 patient-years of etanercept exposure, with 26/69 patients entering their 8th year of etanercept treatment (0.12 events per patient year). Nine medically important infections resulted in the need for intravenous antibiotic therapy or hospitalisation (0.03 events per patient year).
 - Tocilizumab: AEs rates were 406.5 per 100 patient-years and the serious AEs rate 11.1 per 100 patient-years over around two years, with the most common AEs and serious AEs related to infections (151.4 and 5.2 per 100 patient years respectively).
- Growth: limited data reported in abstracts at week 104 for adalimumab and tocilizumab appears to support the positive effect of these drugs on growth, but the use of different outcome measures prevents a comparison between the drugs.

- An exploratory adjusted indirect comparison found that there was a lack of statistically significant differences between the four biologic DMARDs in terms of disease flare and ACR Pedi 50/ 70 response, with wide confidence intervals and clinical heterogeneity between the trials.

4.2 Review of clinical-effectiveness in company submissions to NICE

Four companies made submissions in support of their drugs to NICE: BMS for abatacept, AbbVie for adalimumab, Pfizer Ltd for etanercept and Roche for tocilizumab. A review of the information presented about the economic evaluation of biologic DMARDs for treatment of JIA in the CSs can be found in Section 5.4 of this report.

Review of Bristol-Myers Squibb (BMS) company evidence submission for abatacept

The company did not report a systematic review of clinical-effectiveness of abatacept.⁸⁵ There is no indication that any databases were searched and no search strategies were supplied. Furthermore, there is no search or report for any ongoing studies. The majority of the clinical-effectiveness information in the CS comes from published papers with a few details that are CIC which come from the clinical study reports.

The CS includes one phase III double-blind randomised withdrawal study, the AWAKEN trial. Although not clearly summarised the CS draws on two published papers,^{57;58} one conference presentation⁸⁶ and the trial clinical study reports. The published papers^{57;58} had met the inclusion criteria of this assessment report. One published paper⁵⁹ relating to the AWAKEN trial that was identified in this assessment report was not cited by the CS but the data in this appear to have been superseded by the more recent conference presentation.⁸⁶ However, the more recent efficacy data are not presented according to the randomised groups in the double-blind period whereas the safety summary data are.⁸⁶ Furthermore there is limited detail regarding the length of follow-up which is stated to be ≥ 56 months and up to seven years of total follow-up. No critical appraisal is reported for any of the studies cited in the CS.

A summary of the AWAKEN trial is provided in the CS which is broadly similar to the information presented in the published papers.^{57;58;86} Information from the OLE phase drawn from the conference presentation⁸⁶ is more recent than the data from the published paper,⁵⁹ which is included in the assessment report. Furthermore an analysis was conducted (using Fisher's exact test) to compare serious AEs during the double-blind phases of trials of abatacept, adalimumab etanercept and tocilizumab (CS section 3.4.6). The CS highlights the lack of statistical power due to low numbers of patients and event rates, which should be taken into account in the interpretation of their finding that the incidence of serious AEs was likely to be similar between the biologic DMARDs.

The CS focuses on abatacept with very little information provided regarding the other biologic DMARDs included in this MTA. However information is presented in the CS on indirect pairwise comparisons for the four biologic DMARDs for the outcome of disease flare. The comparisons for abatacept, adalimumab and etanercept are taken from a published paper by Otten and colleagues⁷⁹ and this is supplemented by new indirect pairwise comparisons with tocilizumab taken data from an RCT⁶⁸ that has been published since the Otten and colleagues study (the CHERISH trial)⁶⁸. The results of the indirect comparisons reported in the CS (Tables 4 and 5) match those reported in the indirect comparisons conducted for this assessment report (see Section 4.1.3).

In summary, the CS has not conducted a systematic review of clinical-effectiveness but has summarised data from the AWAKEN trial,⁵⁷ presented indirect pairwise comparisons of the four biologic DMARDs included in this appraisal and conducted an analysis to compare serious AEs during the double-blind phases of trials of the four biologic DMARDs. No additional RCTs were included in the CS that would have met the inclusion criteria of this assessment report.

Review of AbbVie company evidence submission for adalimumab

AbbVie submitted a report to NICE on adalimumab as a treatment for JIA.⁷⁷ The clinical-effectiveness evidence has been briefly appraised.

The company did not conduct a formal systematic review of the clinical-effectiveness evidence, but provided what they describe as ‘an iterative literature review’ (CS page 15). The company asserts that all RCTs of adalimumab in the treatment of JIA have been identified (CS page 15). It would appear that RCTs were identified chronologically from an adalimumab trial programme and there is no mention that any databases were searched and no search strategies were provided. There is no search for or report of any ongoing studies, but the CS does contain information about a trial in progress, the SYCAMORE RCT.⁸⁷ This trial is evaluating the clinical-effectiveness, safety and cost-effectiveness of adalimumab in combination with methotrexate for the treatment of JIA-associated uveitis (further information on this trial is given in section 4.3 ‘Ongoing trials’ of this assessment report). Data from abstracts/conference proceedings are also presented in the CS.

The submission contains narrative summaries for the pivotal RCT by Lovell and colleagues (2008),⁶¹ which formed the basis of the original marketing authorisation in 2008; an ongoing RCT of adalimumab treatment in patients with ERA by Burgos-Vargas and colleagues⁸⁸⁻⁹³ (see section 4.3 ‘On going trials’ of this assessment report for details of this study); two open-label single-arm studies^{94;95} and supporting data from an ongoing registry (STRIVE) funded by AbbVie. The multi-national STRIVE registry is assessing the long-term safety and effectiveness of adalimumab in

patients with moderate to severe polyarticular JIA. Some data from this registry are given in the CS, for efficacy outcomes up to one year, and safety outcomes for longer (mean duration of drug exposure 643 days for methotrexate patients and 653 days for adalimumab and methotrexate patients). The registry does not appear to include patients from the UK. Other evidence such as case series, open-label trials, a systematic review⁹⁶ and data from an Italian registry were included to provide evidence for the effectiveness of adalimumab in JIA-associated uveitis (see section 4.4.2 ‘JIA-associated uveitis’ of this assessment report for details of these).

Based on the Lovell and colleagues RCT,⁶¹ the key outcomes of disease flares and ACR Pedi responses are the same in the CS and the assessment report.

The CS notes several methodological concerns which prevented the presentation of a network meta-analysis comparing the four biologic DMARDs. An indirect comparison was therefore not presented.

In summary, the CS has not conducted a systematic review of clinical-effectiveness but has summarised data separately for RCTs and other non-randomised studies, as well as data from a registry. No indirect comparison of the biologic DMARDs was conducted by the company. No additional RCTs were included in the CS that met the inclusion criteria for systematic review in this assessment report, however, some of the non-randomised study evidence in the CS is presented in this assessment report for patient sub-groups where randomised evidence is lacking (i.e. ERA, and JIA-associated uveitis) (section 4.4).

Review of Pfizer Ltd company evidence submission for etanercept

The company report a systematic review of clinical-effectiveness of etanercept (in addition they report a systematic review of observational evidence on etanercept-associated innovation, caregiver burden and treatment adherence, plus a systematic review of HRQoL associated with etanercept).⁹⁷ Details of the literature search strategy are provided and the search appears to be comprehensive, up to date and reproducible. A search for ongoing studies was also conducted. A systematic process was followed to screen studies for inclusion, with titles and abstracts and full texts screened independently by two reviewers. The inclusion criteria are in-keeping with the scope of the appraisal, with the exception that only studies of etanercept were included, not the other biologic DMARDs. A broad range of study designs were eligible, with the exception of case reports. The majority of the data are in the public domain although some information is either AIC or CIC.

The review included 11 publications relating to five primary interventional studies and three extension studies (see CS Table 7, CS page 44). It also included 41 observational studies (including registry studies) plus 2 unpublished studies (see CS Table 18, CS page 81). Of the included five primary

interventional studies only one meets the inclusion criteria for the systematic review in this current report – Lovell and colleagues (2000).⁴² Of the remaining four studies, three were not relevant as they were single arm studies, and a fourth was an RCT reported in a conference abstract⁹⁸ with only limited detail available. However, one of the single-arm studies - the CLIPPER study⁹⁹ - is noteworthy as it focuses specifically on the JIA sub-types that were absent from the pivotal Lovell and colleagues trial,⁴² namely extended oligoarticular JIA, ERA and psoriatic arthritis. Details are presented in Section 4.4 of this assessment report.

Of the three extension studies, only one was relevant to the inclusion criteria of this assessment report – the long-term follow-up publications⁶⁵⁻⁶⁷ of the Lovell and colleagues RCT,⁴² all of which have been included in the data extraction for this study (see Appendix 5). The other two extension studies included a Japanese open-label single arm multi-centre study followed by a double-blind, randomised dose-down extension study (2 doses of etanercept - no comparator), and an open-label multi-centre phase 3b long-term safety and efficacy study of the CLIPPER study (reported in Section 4.4 of this assessment report).⁹⁹

A critical appraisal of the interventional studies is provided in the CS section 4.7. The company's appraisal of the Lovell and colleagues RCT⁴² is provided in CS Table 13 (CS page 62). Our critical appraisal differs slightly from the company's (section 4.1.1). Specifically, we did not consider that adequate details had been provided of the study's randomisation method or concealment of allocation. We also note that there was a large imbalance in drop-outs between the randomised groups (see Table 13 in this report, and also Appendix 5).

A narrative synthesis of the interventional and observational studies is provided in the CS, with detailed tabulation of study characteristics and results. A meta-analysis was not considered feasible or appropriate by the company and an indirect comparison was not conducted as it was not considered feasible to conduct one due to differences in respective marketing authorisations across biologic treatments, paucity of data and heterogeneity.

Of note, some of the observational studies of etanercept included in the CS reported (limited) data for outcomes relevant to the scope of the appraisal that were not included in the RCT by Lovell and colleagues,⁴² namely corticosteroid reduction, growth and disease activity according to the JADAS (see CS section 4.12.5).

In summary, the systematic review of clinical-effectiveness reported in the CS appears to be of good standard and no additional RCTs were included in the CS that met the inclusion criteria for the systematic review in this assessment report.

Review of Roche company evidence submission for tocilizumab

The company did not conduct a formal systematic review of the clinical-effectiveness evidence, but provided ‘most relevant literature’ on the use of tocilizumab in patients with polyarticular JIA and extended oligoarthritis (CS page 7). There is no evidence that searches were conducted and no search strategies were reported. The CS does state that a systematic literature review was completed for the indirect comparison presented in the submission, though provides no further detail. The CS did not report searching conference proceedings or details of any ongoing trials, but data from abstracts/conference proceedings are included in the submission. CIC data is limited to the economic model.

The submission contains narrative summaries of two studies. One of the studies is an RCT comparing tocilizumab to placebo (CHERISH)⁶⁸ linked to six additional conference publications/abstracts.^{69;71-73;76;100} The CHERISH RCT⁶⁸ met the inclusion criteria of the this assessment report and was reported earlier in Section 4.1. Of the six conference publications/abstracts linked to the CHERISH RCT, only two were related to the randomised phase of the trial^{72;73} and the remaining four were related to the OLE phase. One of these four conference abstracts was not identified by searches for this assessment report, but if it had been, it would not have met the inclusion criteria as none of the outcomes reported were relevant.¹⁰⁰

The other study was a single-arm open-label study of efficacy, pharmacokinetics, and safety of adalimumab in Japanese patients with polyarticular JIA.⁹⁵

The CS presents all the evidence separately for each study in the form of a narrative summary. Individual tables of baseline patient characteristics, as well as details of methods and design are reported for both the CHERISH trial⁶⁸ and the open-label Japanese study.⁹⁵ No quality assessment of the studies is presented. The CS reports growth data from the open-label extension phase of the CHERISH trial at week 104, which has been included in this assessment report. The assessment report contains additional data for ACR Pedi 90 responses relative to baseline at week 40 and inactive disease from the CHERISH trial, both of which are not reported in the CS.

The CS includes a hierarchical Bayesian indirect treatment comparison of adalimumab and tocilizumab, conducted in WinBUGS software and using methods described by Dias and colleagues 2013.¹⁰¹ Limited detail is provided on the specific methods used to conduct the indirect comparison (e.g. which adalimumab trial was used to compare against tocilizumab – likely to be Lovell and colleagues⁶¹ but not explicitly stated). An indirect comparison with abatacept was not considered

possible due to the difference in trial design, the fact that it is not approved (appraised) by NICE, and also because of slight differences in licences (i.e. lower age for which treatment is indicated). We note the heterogeneity between the RCTs that increases uncertainties in any indirect comparison, though the fact that abatacept has not been appraised by NICE is not an adequate justification for not performing the comparison. The CS provides an additional analysis which assumes a class effect across anti-TNF drugs (based on the indirect comparison with adalimumab which showed ‘overlapping ACR response rates’), permitting a comparison between tocilizumab with etanercept (CS section 5.17). The exploratory pairwise indirect comparisons of all four biologic DMARDs presented in this assessment report showed no statistically significant differences between the drugs (section 4.1.3).

Most of the adverse events and safety data are presented for the CHERISH RCT.⁶⁸ The adverse event data reported at week 40 (end of the randomised phase of the RCT) in the CS does not include any data for the placebo group, which is presented in this assessment report.

In summary, the CS has not conducted a systematic review of clinical-effectiveness but has summarised data from the CHERISH trial and an open-label Japanese study, presented an indirect pairwise comparisons of two of the biologic DMARDs included in this appraisal (tocilizumab and adalimumab), as well as exploratory analysis comparing tocilizumab with etanercept. No additional RCTs were included in the CS that met the inclusion criteria of the assessment report.

4.3 Ongoing trials

As stated in section 4.1.1 above, citations relating to four ongoing RCTs were identified from the electronic bibliographic database literature search (Figure 1) and a separate search specifically for ongoing studies identified a further three ongoing RCTs. Three trials are investigating adalimumab and four etanercept. Each trial is described in turn below, with preliminary results presented where possible. It should be noted that on-line clinical trial registers generally provide less information (and no outcome data) in comparison with published conference abstracts.

Adalimumab ongoing trial 1

This is a phase 3, multi-centre, randomised, double-blind study (NCT01166282) described by six conference abstracts (Burgos-Vargas and colleagues⁸⁸⁻⁹³) in children aged ≥ 6 to < 18 years with ERA based on ILAR criteria, with active disease not responsive to ≥ 1 NSAID and ≥ 1 DMARD. No full paper appears to have been published so far and the six abstracts provide limited information, hence preventing a full assessment of the methodology and trial quality and risk of bias. In addition, baseline characteristics were only reported for the overall trial population and not separately for each randomised group. The estimated study completion date is December 2015.

Forty-six patients were randomised in a 2:1 ratio (adalimumab n=31; placebo n=15) to receive blinded adalimumab (24 mg/m² body surface area up to 40 mg every other every other week [EOW]) or placebo for 12 weeks followed by open-label adalimumab EOW up to 144 weeks. It is unclear if patients also received methotrexate. A table in one of the abstracts⁹³ shows that 11/15 placebo patients and 21/31 adalimumab patients received DMARDs at baseline.

The primary endpoint of this study was percent change from baseline in the number of active joints with arthritis (AJC) at week 12 and secondary variables assessed included enthesitis count (EC), tender and swollen joint counts, and ACR Pedi 30/50/70 responses. Active disease was defined as ≥ 3 active joints (swelling or loss of motion and pain/tenderness) and enthesitis in ≥ 1 location (past or present). Safety was assessed in terms of adverse events (AE), laboratory values and vital sign measurements. Some interim data were reported for 52 weeks including discontinuation of concomitant medication (at the discretion of the treating physician).

Authors state that no children discontinued the double-blind period, while at the same time reporting that seven children ‘escaped early’ to open-label adalimumab.

Results

At baseline, children had a mean age of 12.9 years, with 2.6 mean years of ERA symptoms, a mean enthesitis count and active joint count of 8.1 and 7.8 respectively (Table 29). It is unclear if baseline characteristics between the treatment groups were balanced.

Table 29 Baseline characteristics

Parameter, mean (SD) ¹	All children
Age, years	12.9 (2.9)
ERA symptoms, years	2.6 (2.3)
AJC	7.8 (6.6)
EC	8.1 (8.4)

AJC, active joints count. EC, enthesitis count. ERA enthesitis related arthritis. SD, standard deviation.

¹ Not specially stated, but presumed to be standard deviation.

Only the primary outcome percent change from baseline in the number of active joints with arthritis showed a statistically significantly greater improvement (p=0.039) in the adalimumab treatment group (-62.6) compared with placebo (-11.6). Secondary outcomes were reported to be mostly numerically greater in the adalimumab group, but none of the improvements were statistically significant.

Table 30 Results week 12

Primary outcome ¹	ADA (n=31)	PBO (n=15)	p value
AJC, % change from baseline at week 12	-62.6 (59.5)	-11.6 (100.5)	p=0.039

	(median percent change -88.9%)	(median percent change -50.0%)	
Secondary outcomes, change from baseline,² mean (SD)			
Number of enthesitis sites (0-35)	-4.4 (6.2)	-2.7 (5.0)	NS
Tender joint count (0-72)	-7.9 (8.3)	-4.5 (9.0)	NS
Swollen joint count (0-8)	-3.5 (5.6)	-2.4 (4.7)	NS
ACR Pedi Response,³ (n, %)			
ACR Pedi 30 responder	21 (67.7)	10 (66.7)	NS
ACR Pedi 50 responder	20 (64.5)	7 (46.7)	NS
ACR Pedi 70 responder	16 (51.6)	4 (26.7)	NS

ADA, adalimumab. AJC, active joints count. PBO, placebo. SD, standard deviation.

¹ Presumed mean and SD, but not specifically stated.

² Last observation carried forward.

³ Analysed with non-responder imputation.

Adverse Events (AEs)

Only one patient (in the adalimumab group) experienced a serious AE (abdominal pain and headache). Around two thirds of the children in the adalimumab treatment group and just over half in the placebo group experienced any AE, while nearly a third of children in the adalimumab treatment group but only a fifth in the placebo group experienced infectious AEs (Table 31).

Table 31 AEs week 12

AEs, %	ADA (n=31)	PBO (n=15)
Any AEs	67.7	53.3
Serious AEs	3.21 (1 patient)	0
Infectious AEs	29.0	20.0

AD, adalimumab. AE, adverse events. PBO, placebo.

Open-label week 52 results

The authors state that treatment response was maintained with continued adalimumab therapy up to 52 weeks (% change from baseline at week 52 in AJC: -88.7, SD 26.1). In those receiving at least one dose of adalimumab through to week 52, over 91% of children experienced an AE and over 76% experienced infectious AEs (Table 32). Serious AEs were reported in approximately 11% of children, with no reported deaths, tuberculosis or malignancies. Eight (19%) of the 43 participants who remained in the study at week 52 had completely discontinued concomitant ERA medication.

Table 32 AEs >1 dose of adalimumab week 52

AEs, %	ADA¹
Any AEs	91.3
Serious AEs	10.9
Infectious AEs	76.1

ADA, adalimumab. AE, adverse events.

¹ Number of patients receiving ADA treatment during open-label not reported.

Adalimumab ongoing trial 2

The second adalimumab RCT (SYCAMORE, ISRCTN10065623)⁸⁷ is funded by the NIHR HTA programme and Arthritis Research UK. The study is assessing adalimumab combined with methotrexate compared to placebo combined with methotrexate for JIA associated uveitis in participants aged between 2 and 18 years. All participants will receive 18 months of treatment with a three-year follow-up. The study will also include an assessment of cost-effectiveness. Originally expected to report findings in 2020, it has recently been announced that the trial has closed for recruitment early following interim analysis showing a favourable effect for adalimumab. Analysis of the primary outcome is underway and key findings will therefore be available earlier than expected. Collection and analysis of health economic data will continue as planned.

Adalimumab ongoing trial 3

This third adalimumab RCT (Effect of Adalimumab for the Treatment of Uveitis in Juvenile Idiopathic Arthritis (ADJUVITE); NCT01385826) is also currently in progress and is not expected to report findings until June 2016. The study is set in France (seven hospital ophthalmology departments) and assesses the efficacy of two-month adalimumab treatment versus placebo treatment on reduction of ocular inflammation quantified by laser flare photometry in patients aged ≥ 4 years, with JIA associated uveitis resistant to steroid therapy. The investigators plan to include 40 patients, follow-up appears to be 12 months and the final data collection date for the primary outcome measure is November 2015. The primary endpoint of this study is improvement of uveitis.

Etanercept ongoing trial 1

This is an RCT evaluating the efficacy of etanercept in 124 Chinese JIA patients (no clinical trial registration number has been found for this study).⁹⁸ No full paper appears to have been published so far and only one conference abstract was identified, which includes very limited information. The abstract states that a “randomised principle was applied” to divide the JIA patients into a control and a treatment group. While no baseline characteristics were reported, the authors of the abstract state that there were no significant differences of clinical classification and basic treatment between the groups.

Sixty-two patients in the treatment group (oligoarticular JIA n=17, polyarticular JIA n=15 and systemic JIA n=30) received 0.8 mg/kg per week of subcutaneous etanercept for six months. No details for the control group are reported.

ACR Pedi 30, 50 and 70 responses were used to assess the clinical efficacy (primary outcome not stated) and adverse reactions were recorded.

Results

Authors state that the remission rates ‘of different cases’ (this is presumed to mean different types of JIA) in the treatment group differed at each time point (three- and six-month time points are mentioned), with no obvious difference in ACR Pedi 30, 50 and 70 remissions for patients with oligoarticular and polyarticular JIA. Eighty percent of these patients had ACR Pedi 50 remission after six-month treatment and more than 50% had ACR Pedi 70 remission. The remission rate of systemic JIA cases was lower compared with the two other types (data not extracted). While the differences between the randomised groups are said to be significant, no data were reported.

Adverse events (AE)

There were no reported AEs for patients with oligoarticular or polyarticular JIA. Details of AEs for the systemic JIA sub-group are reported (not extracted).

Etanercept ongoing trial 2

The second placebo-controlled etanercept RCT [Remission Induction by Etanercept in Enthesitis related Arthritis JIA-Patients (REMINDER) Study, EudraCT Number: 2010-020423-51) was identified from the search of ongoing clinical trials registers. The trial is set in Germany and has a start date of February 2016. This study has a withdrawal RCT design, with a 12-week open-label treatment phase prior to the controlled randomised double-blind phase. The study will assess the safety and effectiveness of etanercept in patients diagnosed with ERA-JIA age ≥ 6 years and < 18 years having met all criteria for eligibility for treatment with etanercept according to SPC and local guidelines, with expectation of the requirement of a minimum of five affected joints. The on-line record does not provide the treatment time for the double-blind phase or report any follow-up period. The primary endpoint of the study will be inactive disease of ERA-JIA.

Etanercept ongoing trial 3

The third multi-centre etanercept RCT was identified from a conference abstract¹⁰² which does not provide a clinical trial registration number. This trial appears to be set in Germany, has enrolled patients with ERA and has a withdrawal design, with a 24-week open-label etanercept treatment phase prior to the 24-week placebo-controlled double-blind withdrawal phase. Patients had to achieve at least an ACR Pedi 30 response in order to be randomised to the double-blind phase (terminated in case of a disease flare or at week 48, whichever occurs earlier). No details of the study’s inclusion or exclusion criteria are reported in the abstract.

Results

Forty-one patients entered the open-label phase, of which two patients discontinued prematurely (one due to intolerance and one to protocol deviation). Thirty-eight patients (93%) achieved at least an ACR Pedi 30 response and were randomised to the double-blind phase. As can be seen in Table 33, during the double-blind phase the majority of flares occurred in the placebo group, with 10 (56%) placebo-treated patients compared to 18 (90%) etanercept-treated patients reaching week 48 without a flare (OR 7.2 PBO vs. ETA (1.3 to 40.7 - although not stated this is presumed to be a 95% CI), p= 0.016).

Table 33 Flares: 24-week double-blind phase

Time	ETA (n=20)	PBO (n=18)
Week 28	2	2
Week 32	0	4
Week 48	0	2

The authors state that in patients continuously treated with etanercept at week 48 (n=20), JADAS 10 decreased to a mean of 3.4 (17.4 to 1.9 at week 24), with 12 (60%) patients reaching JADAS-minimal disease activity and 11 (55%) JADAS remission. The equivalent data for the placebo group are not reported.

Adverse events (AEs)

There were 166 AEs in 39 patients and three serious AEs. All were said to be considered unrelated and resolved without sequelae. It is unclear if these data are for the combined open-label and double-blind phases, but considering the number of patients reported (n=39) this may indeed be the case.

Etanercept ongoing trial 4

The fourth multi-centre etanercept RCT (set in the Netherlands) assessed when and in whom to stop etanercept after successful treatment of JIA (NCT01287715) and was identified from the search of ongoing clinical trials registers. Estimated enrolment was 50, the study completed in September 2013 and final data collection for the primary outcome measure is reported as September 2012 in the on-line record. No publication of data has been identified. Patients aged 4 to 17 years and in remission were selected from the ABC-register, an observational study including all Dutch JIA patients on etanercept therapy. The inclusion criteria states no or low dose of methotrexate and it is therefore unclear if patients were intolerant or had a previous inadequate response to methotrexate. All JIA subtypes were included in the study. Patients were randomised to a stop-arm (discontinuation of etanercept - half of the dose for 3 months and discontinuation thereafter) or a control-arm (etanercept

continued for another 9 months and, if still meeting the eligibility criteria, discontinued thereafter). The primary outcome of the study was flare-rate.

4.4 Additional supporting evidence

This section includes additional non-randomised study evidence relating to aspects of JIA where adequate RCT data in the systematic review of clinical-effectiveness were lacking. This evidence has been identified from the systematic review search itself, and from relevant studies included in the company submissions to NICE (see Section 4.2). Evidence relating to two aspects is presented: JIA sub-types ERA and PA, and JIA-associated uveitis.

4.4.1 Enthesitis-related arthritis (ERA) and psoriatic arthritis (PA)

The most informative study available for these sub-types is the CLIPPER study.⁹⁹

This is a single arm, phase 3b open-label, multi-centre interventional study funded by Wyeth (subsequently acquired by Pfizer Inc.). The study was designed to assess the safety and efficacy of etanercept in children and adolescents with three JIA subtypes as classified using the ILAR criteria: extended oligoarticular JIA (EO), ERA and PA. There are two parts to the study. Part 1 (which has been published⁹⁹) has investigated the efficacy and safety of etanercept in the three JIA subtypes over an initial 12-week period with a primary endpoint of the percentage of patients achieving ACR Pedi 30 criteria at week 12. Part 2 of the study is a 96-week open-label extension, assessing the long-term safety and efficacy of etanercept in JIA subtypes, which is currently published in poster-format only.¹⁰³ This study formed part of the evidence base supporting the licence extension for etanercept across the JIA subtypes in 2012. The assessment group has extracted 12-week data from the published paper⁹⁹ on ACR Pedi response rates and, inactive disease, change in CHAQ score, PGA of pain, number of active joints, number of joints with LOM and JIA category-specific assessments (data at week 12 compared with historical placebo data, historical active control data and data from a meta-analysis have not been data extracted). Data from the conference poster¹⁰³ on ACR Pedi response rates and inactive disease have also been extracted and this summary is supplemented with some data presented in the Pfizer CS.

The study included 127 patients with the JIA subtypes of ERA (n=38, age 12-17 years), EO (n=60, age 2-17 years) and PA (n=29, age 12-17 years) who received 0.8 mg/kg of etanercept once weekly (maximum dose 50 mg/week). Key inclusion criteria were ≥ 2 active joints (swollen or LOM accompanied by either pain or tenderness); history of intolerance or unsatisfactory response to at least a three-month course of ≥ 1 DMARD or, only for ERA, unsatisfactory response to at least a one-month course of ≥ 1 NSAID (i.e. for ERA prior methotrexate treatment was not required). A stable dose of concomitant medication (only one DMARD, one oral corticosteroid and one NSAID) was permitted. The inclusion criteria extended below the threshold number of active joints for the

classification of polyarticular disease. Key exclusion criteria included other rheumatic diseases, active uveitis within 6 months of baseline, and any prior receipt of biologic DMARDs. A total of five patients failed to complete Part 1 of the study (completed Part 1: EO 97%, ERA 95% and PA 97%) and 13 patients Part 2 (completed Part 2: EO 90%, ERA 79% and PA 86%).

Key baseline characteristics can be seen in Table 34 (additional baseline characteristics are available in the published paper⁹⁹).

Table 34 Key baseline characteristics

Parameter [mean (SD) unless stated otherwise]	All patients (n=127)	EO (n=60)	ERA (n=38)	PA (n=29)
Age, years	11.7 (4.5)	8.6 (4.6)	14.5 (1.6)	14.5 (2.0)
Female, %	56.7	68.3	21.1	79.3
JIA duration	26.8 (26.4)	31.6 (31.7)	23.0 (19.8)	21.8 (20.2)
Age at onset, months	9.5 (4.8)	6.1 (4.5)	12.5 (2.1)	12.6 (2.7)
Concomitant medication use, n (%)				
Any DMARD	109 (85.8)	54 (90.0)	32 (84.2)	23 (79.3)
Oral corticosteroid	16 (12.6)	7 (11.7)	8 (21.1)	1 (3.5)
Oral NSAID	74 (58.3)	32 (53.3)	26 (68.4)	16 (55.2)
No. active joints	6.7 (4.6)	7.6 (5.1)	5.2 (3.6)	7.0 (4.3)
No. joints with LOM	5.7 (4.2)	6.3 (4.4)	4.8 (4.0)	5.6 (4.1)
No. of painful joints	6.4 (5.2)	5.5 (4.1)	6.7 (4.9)	7.8 (7.0)
No. of swollen joints	5.5 (4.2)	6.5 (4.8)	3.8 (2.8)	5.6 (3.7)
CHAQ score	0.8 (0.6)	0.9 (0.7)	0.7 (0.5)	0.7 (0.6)
PGA pain (VAS)	5.1 (2.5)	4.8 (2.6)	5.8 (2.5)	4.6 (2.3)
JIA category-specific characteristics				
Tender enthesal score			5.9 (9.4)	
Overall back pain VAS, mm			25.9 (28.0)	
Nocturnal back pain VAS, mm			16.4 (27.8)	
Modified Schober's test, cm			15.0 (1.9)	
Psoriasis BSA, %				10.4 (13.4)
PGA of psoriasis				1.8 (1.4)

CHAQ, Childhood health assessment questionnaire (0 - 3 scale, no disability-severe disability).EO, Extended oligoarthritis. JIA, ERA, Enthesitis-related arthritis. LOM, Limitation of motion. N/A, Not applicable. PGA, Parent global assessment. PA, Psoriatic arthritis. VAS, Visual analogue scale (0 - 10).

Patients with EO and PA had a higher number of active joints and number of joints with limitation of motion (EO: mean 7.6 (SD 5.1) and 6.3 (SD 4.4) respectively; PA: mean 7.0 (SD 4.3) and 5.6 (SD 4.1) respectively) at baseline compared with ERA patients (mean 5.2 (SD 3.6) and 4.8 (SD 4.0)

respectively). The number of painful joints was highest in PA patients [mean 7.8 (SD 7.0)] compared with the other two sub-groups, while the number of swollen joints was the lowest in ERA patients [mean 3.8 (SD 2.8)] (see Table 34). Mean CHAQ sub-group scores ranged between 0.7 and 0.9, while the parent global assessment of pain VAS ranged between 4.6 and 5.8. Also reported are JIA category-specific assessments (ERA: tender enthesal score, back pain (VAS) and modified Schober's test; PA: body surface area and PGA of psoriasis) at baseline. Limitations of the study noted by the authors was the difference in concomitant medication use at baseline which may have affected efficacy responses and the lower age limit of 12 years set for inclusion of EO patients (the licensed indication is from 2 years of age).

Results

At week 12, the overall ACR Pedi 30 response rate for patients was almost 89%, with response for the separate JIA disease type sub-groups varying from around 83% to 93% (see Table 35). The overall ACR Pedi 90 response rate for patients was just under 30% and 12% of patients had inactive disease. JIA disease sub-groups varied in ACR Pedi 90 response rates between

[REDACTED]

[REDACTED] At week 12, 12%, 17% and 7% of EO, ERA and PA patients respectively had inactive disease as can be seen in Table 35.

By week 96, around 99% of all patients achieved a ACR Pedi 30 response and over 65% a ACR Pedi 90 response. Thirty four percent of all patients had inactive disease. Overall, patients in the ERA sub-group appeared to have received the greatest benefit from etanercept therapy at 12 weeks, but by 96 weeks the sub-groups achieved similar levels of ACR Pedi 90 (62% to 72%) and ACR Pedi 100 (51% to 60%). Inactive disease at 96 weeks varied between 29% (ERA and PA) and 37% (EO).

[REDACTED]

[REDACTED]

[REDACTED]

Table 35 ACR Pedi response and inactive disease results at week 12 and week 96

Parameter, %	All patients (n=127)		EO (n=60)		ERA (n=38)		PA (n=29)		
	Week 12	N	% (95% CI)	n	% (95% CI)	n	% (95% CI)	n	% (95% CI)
ACR Pedi 30			88.6 (81.6, 93.6)		89.7 (78.8, 96.1)		83.3 (67.2, 93.6)		93.1 (77.2, 99.2)
ACR Pedi 50			81.1 (73.1, 87.7)						
ACR Pedi 70			61.5 (52.2, 70.1)						
ACR Pedi 90			29.8 (21.8, 38.7)						
Inactive disease	12.1 (6.9, 19.2)		11.9 (4.9, 22.9)		16.7 (6.4, 32.8)		6.9 (0.8, 22.8)		
Week 96¹	All patients (n=108)		EO² (n=53)		ERA² (n=30)		PA² (n=25)		
ACR Pedi 30	99.1% (95% CI 94.9, 100)		99%		100%		98%		
ACR Pedi 50	98.1% (95% CI 93.5, 99.8)		99%		97%		98%		
ACR Pedi 70	92% ²		94%		87%		98%		
ACR Pedi 90	65.4% (95% CI 55.6, 74.4) (n=107)		62%		67%		72% (n=24)		
ACR Pedi 100	54% ² (n=107)		54%		51%		60% (n=24)		
Inactive disease	34 (25.0, 43.8) (n=106)		37%		29% (n=28)		29%		

¹ Efficacy analyses were based on observed data. ² Data estimated by reviewer from poster graphs using Engauge digitizer 4.1 software¹⁰³ CIC data are unpublished, taken from the CS.

At week 12 mean change from baseline for CHAQ scores were similar for the sub-groups (improvement of 51% to 58%), but there were differences in the parent global assessment of pain VAS. The lowest mean change in pain VAS occurred in patients with ERA and PA (decreases of 45% and 47% respectively) compared to patients with EO (59% decrease). The mean decreases from baseline for the number of active joints ranged between 70% and 78% and between 64% and 72% for the number of joints with limitation of motion (see Table 36). It is unclear why details about changes from baseline for the number of painful and the number of swollen joints are not reported.

The greatest improvements in JIA category-specific assessments was a 58% improvement from baseline in tender enthesal score at 12 weeks in patients with ERA, and for patients with PA a 48% improvement in body surface area of psoriasis and a 40% improvement in PGA of psoriasis (see Table 36).

Table 36 Mean change from baseline week effectiveness measures at week 12 (observed cases)

Parameter mean (95% CI) [% change]	Overall (n=123)	EO (n=58)	ERA (n=36)	PA (n=29)
CHAQ	-0.5 (-0.6 to -0.4) [-53.6%]	-0.5 (-0.7 to -0.4) [-52.2%]	-0.5 (-0.7 to -0.3) [-57.8%]	-0.4 (-0.6 to -0.2) [-51.3%]
Parent GA of child's pain VAS	-3.0 (-3.5 to -2.6) [-51.9%]	-3.2 (-3.8 to -2.5) [-58.9%]	-3.2 (-4.2 to -2.2) [-44.9%]	-2.6 (-3.4 to -1.8) [-46.6%]
No. of active joints	-5.1 (-5.8 to -4.3) [-73.0%]	-5.5 (-6.7 to -4.2) [-69.8%]	-4.3 (-5.4 to -3.1) [-77.7%]	-5.2 (-6.8 to -3.6) [-73.8%]
No. of joints with LOM	-4.1 (-4.8 to -3.4) [-66.9%]	-4.5 (-5.6 to -3.3) [-64.1%]	-3.4 (-4.1 to -2.6) [-67.4%]	-4.3 (-5.7 to -2.9) [-71.7%]
JIA category-specific assessments				
Tender enthesal score			-4.4 (-6.3 to -2.4) [-57.8%]	
Overall back pain VAS, mm			-12.5 (-21.3 to -3.7) [-21.2%]	
Nocturnal back pain VAS, mm			-8.9 (-16.7 to -1.2) [-6.8%]	
Modified Schober's test, cm			0.35 ¹ (-0.02 to 0.72) [9.7%] (n=35)	
Psoriasis BSA, %				-6.7 (-10.6 to -2.9) [-48.2%]
PGA of psoriasis				-1.0 (-1.4 to -0.6) [-39.6%] (n=28)

BSA, body surface area. GA, global assessment. VAS, visual analogue scale.

¹ Change from baseline calculated after subtracting 10 from the baseline and week 12 scores

Adverse Events

The highest number of treatment emergent AEs per patient year of etanercept exposure (EXP) occurred in the ERA sub-group (1.827; EO: 1.313; PA: 1.036), which also had the lowest number of treatment emergent infections per patient year of etanercept treatment (0.979; EO sub-group: 2.114;

PA: 1.514). As Table 37 illustrates, patients with ERA appear to experience more treatment-related injection site reactions than patients with either EO or PAA.

Treatment emergent AEs leading to patient withdrawal only occurred in the ERA sub-group (events 3, EXP 7.9), and there were two events of treatment emergent infections causing withdrawal (one each in the EO and PA groups). The rate of serious treatment emergent AEs and serious treatment emergent infections appears to be low in all three sub-groups (see Table 37), as does the rate of treatment emergent autoimmune disorder events. Of the two cases of uveitis (EO n=1, PA n=1), one was reported in a patient with EO after 7.8 months of etanercept plus methotrexate. This resolved and the patient completed the 96-week study. There were a total of three cases of Crohn's disease in patients with ERA, of which two cases were considered to be unrelated to etanercept therapy.

Table 37 Adverse events at Week 96

All values are reported as no. of events (EPPY of EXP to ETN) unless otherwise stated	Overall (n=127) EXP=215.086	EO (n=60) EXP=103.603	ERA (n=38) EXP=61.298	PA (n=29) EXP=50.185
Treatment emergent AEs ¹	300 (1.395)	136 (1.313)	112 (1.827)	52 (1.036)
Treatment emergent infections	355 (1.651)	219 (2.114)	60 (0.979)	76 (1.514)
Treatment emergent ISRs	63 (0.293)	22 (0.212)	29 (0.473)	12 (0.239)
Treatment emergent AEs causing withdrawal, n (%) ¹	3 (2.4)	0	3 (7.9)	0
Treatment emergent infections causing withdrawal, n (%)	2 (1.6)	1 (1.7)	0	1 (3.4)
Serious treatment emergent AEs ¹	16 (0.074)	2 (0.019)	11 (0.179)	3 (0.060)
Serious treatment emergent infections	10 (0.046)	4 (0.039)	3 (0.049)	3 (0.060)
Opportunistic infections ²	1 (0.005)	0	1 (0.016)	0
Infections considered preventable by vaccination in patients not previously vaccinated	7 (0.033)	5 (0.048)	1 (0.016)	1 (0.020)
Infections considered preventable by vaccination in patients previously vaccinated	1 (0.005)	1 (0.010) ³	0	0
Treatment emergent autoimmune disorders ⁴	4 (0.019)	1 (0.010)	2 (0.033)	1 (0.020)

AE, Adverse events. EPPY, Events per patient year. EXP, Exposure. ISR, Injection site reaction.

¹Excluding infections and ISRs.

²1 case of herpes zoster affecting 2 dermatomes was considered to be an opportunistic infection and 1 case of latent tuberculosis (purified protein derivative conversion) was not considered to be an opportunistic infection.

³1 case of rubella.

⁴ 2 cases of uveitis (EO and PA subtypes), 1 case of iridocyclitis (a subtype of uveitis; ERA subtype) and 1 case of Crohn's disease (ERA subtype) were treatment emergent. One case of Crohn's disease (ERA subtype) was not considered treatment emergent based on missing last-dose data.

4.4.2 JIA-associated uveitis

As stated earlier, the effects of biologic DMARD on extra-articular manifestations such as uveitis were not assessed by the included RCTs. However, evidence from non-randomised studies is available, as summarised by systematic reviews.

A recently published systematic review by Simonini and colleagues⁹⁶ assessed the effectiveness of anti-TNF drugs for childhood uveitis. To be included studies had to include patients with autoimmune uveitis refractory to topical and/or systemic steroids and at least one immunosuppressive therapy (e.g. methotrexate). The anti-TNF alpha drugs of relevance to the review were etanercept, infliximab and adalimumab (NB. infliximab is not within the scope of this NICE appraisal). The primary outcome was improvement in intraocular inflammation, with additional outcomes including tapering/stopping systemic steroid administration, improvement in visual acuity, and treatment discontinuation amongst others. A number of bibliographic databases were searched from January 2000 to October 2012.

The review included 23 studies, mainly retrospective chart reviews with very small patient numbers. Of these 23 studies only seven were conducted exclusively in JIA uveitis patients (1 RCT of etanercept; 2 retrospective studies of etanercept; 2 retrospective studies of infliximab; and 2 retrospective studies of adalimumab). Eleven studies comprised mixed study populations with uveitis associated with a range of conditions, including JIA. The remaining five studies included populations that did not include any children with JIA. It was not possible to analyse results separately by uveitis-associated condition. However, of the 229 children included across all the studies, 152 had chronic uveitis associated with JIA. The results can therefore be interpreted as being generally relevant to JIA uveitis.

A pooled analysis of the observational studies found that adalimumab and infliximab were more efficacious at improving intraocular inflammation than etanercept. The proportion of children with improved intraocular inflammation (responders) was 87% (95% CI 75 to 98%) for adalimumab, 72% (95% CI 64 to 79%) for infliximab, and 33% (95% CI 19 to 47%) for etanercept. There was no statistically significant difference in the proportion of responders between adalimumab and infliximab ($p=0.08$), but there was a significant difference for both compared to etanercept ($p=0.001$ for both comparisons).

Simonini and colleagues⁹⁶ did not pool the results of the single RCT identified in the systematic review with the observational studies.⁵⁶ This was a small RCT (n=12 children) of treatment with etanercept. The authors state that this study did not report substantial benefits for the biologic treatment. (NB. Due to limitations in reporting, this RCT was judged to be unclear for inclusion in our systematic review of clinical-effectiveness, as it was not clear whether the etanercept was given with in its licensed indication). Caution is advised in the interpretation of the findings of the Simonini and colleagues⁹⁶ systematic review, given the weaknesses of the study designs included.

The assessment group are aware of only one other recent systematic review of biologic DMARD treatment of children with uveitis, by Cordero-Coma and colleagues.¹⁰⁴ The most recent search date for literature was October 2011. This review had a broader inclusion criteria than that of Simonini and colleagues⁹⁶ and a total of 61 studies were included. Again, much of the included evidence was from observational studies. A total of 14 studies assessed adalimumab, 11 assessed etanercept, and 50 studies assessed infliximab (studies assessing certolizumab and golimumab were also included). Of the 1093 patients included across the studies, 316 (30%) were classed as having JIA uveitis. The review does not provide any formal synthesis and quantification of the effectiveness of treatment, but provides a narrative conclusion for each biologic DMARD and a level of evidence. Adalimumab and infliximab were considered by the authors to be effective in autoimmune uveitis, both based on level 2b evidence (individual cohort study, or low-quality RCT). Etanercept was judged ineffective, based on level 1b evidence (individual RCTs with narrow confidence intervals).

The Abbvie company submission to NICE⁷⁷ provides narrative summaries of five selected studies published since the Simonini and colleagues⁹⁶ systematic review. All of them were observational in design (3 case series¹⁰⁵⁻¹⁰⁷; 1 Italian registry-based study¹⁰⁸; 1 comparative cohort study¹⁰⁹), and all assessed treatment with adalimumab (with 1 also assessing infliximab¹⁰⁹ in uveitis patient populations with varying proportions of JIA uveitis. We have not performed an independent critical appraisal of these studies in this assessment report. From the summaries given it appears that adalimumab is associated with improvements in intraocular inflammation and visual acuity, and a decrease in use of corticosteroids. Adverse events appeared to be minor. The other company submissions to NICE did not present much detail of studies of treatment of JIA uveitis with other biological DMARDs.

In summary, the evidence from observational studies suggests that biologic DMARDs can improve uveitis symptoms in children with JIA, such as intraocular inflammation. Adalimumab and infliximab appear to be more effective than etanercept in improving uveitis. The effects of the treatments in terms of arthritis outcomes in JIA uveitis patients have not been reported. As noted in Section 4.3 ('Ongoing trials') the UK-based SYCAMORE RCT⁸⁷ has investigated adalimumab in the treatment of JIA uveitis patients, and the results of the trial (which will be available sooner than expected, though

an exact date has not been specified) will provide more rigorous evidence for effectiveness than that currently available.

5 ECONOMIC ANALYSIS

5.1 Introduction

The aim of the economic evaluation is to assess the cost-effectiveness of abatacept, adalimumab, etanercept and tocilizumab for people with JIA, compared to alternative treatments. The economic evaluation comprises:

- a systematic review of the cost-effectiveness of biologic DMARDs for people with JIA (Section 5.2);
- a systematic review of studies of the HRQoL of people with JIA (Section 5.3);
- a critical appraisal of the submissions from the relevant drug companies received as part of the NICE appraisal process (Section 5.4); and,
- a *de novo* economic model and cost-effectiveness evaluation developed by SHTAC to inform the NICE appraisal (Section 5.5).

5.2 Systematic review of cost-effectiveness evidence

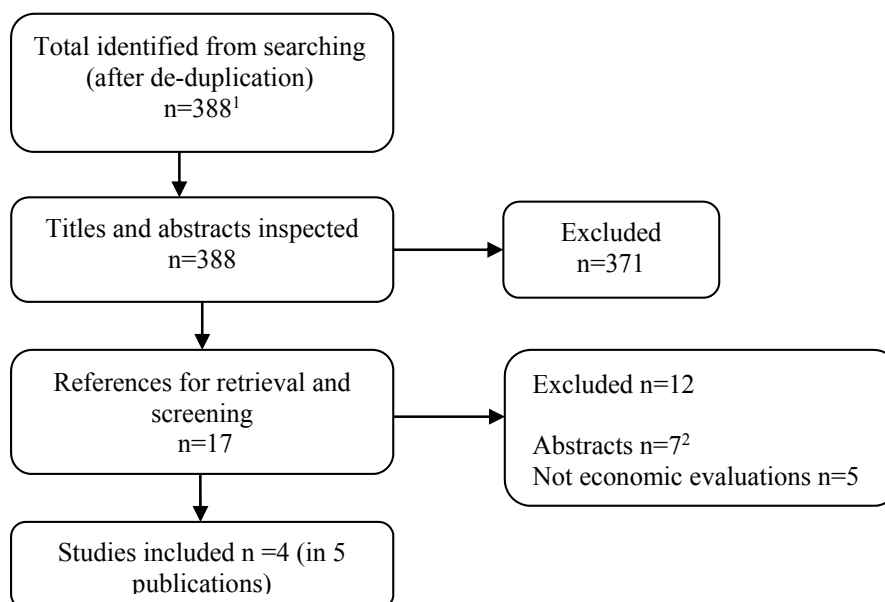
Methods for the systematic review

A systematic literature search was undertaken to identify economic evaluations of the biologic DMARDs, within the NICE scope for this appraisal. Studies were included if they were full economic evaluations (cost-effectiveness, cost-utility, cost-consequence, or cost benefit-analyses) conducted in children and young people with JIA that compared one or more biologics with a DMARD, such as methotrexate. Studies that were not reported in the English language or did not provide sufficient information on the model structure, data and results were excluded. This systematic review aimed to summarise the currently available evidence and inform the construction of a *de novo* model.

Results of the systematic review

Searches for economic evaluations identified 387 potentially relevant references and a further study was identified through *ad hoc* searching. The full texts for 17 papers were retrieved for further screening. A summary of the selection process and the reasons for exclusion are presented in Figure 6 and a list of excluded studies in Appendix 6. Although seven studies reported as abstracts appeared to meet the *a priori* inclusion criteria, they did not contain sufficient information on the methods used and the results to permit formal data extraction or critical appraisal.¹¹⁰⁻¹¹⁶ Five studies were found not to be economic evaluations.¹¹⁷⁻¹²¹ Four studies were included, described in a total of five

publications.¹²²⁻¹²⁶ The characteristics of the four included economic evaluations are shown in Table 38. Data extraction forms for the studies are in Appendix 8.



¹ Including 1 study found through hand searching

² The abstracts provided insufficient details of methods and results to allow inclusion in the systematic review

Figure 6 Flow chart of identification of studies for inclusion in the review of cost-effectiveness

Table 38 Characteristics of economic evaluations

Author	Cummins¹²²	Prince¹²³	Simpson¹²⁴	Ungar^{125;126}
Publication Year	2002	2011	2012	2011
Country	UK	Netherlands	Russia	Canada
Funding source	UK HTA Programme	Dutch Board of Health Insurance and Wyeth International	Not stated	Ontario Ministry of Health and Long-term Care Drug Innovation Fund
Analysis type	Cost-utility analysis	Cost-consequence analysis	Cost-utility analysis	Cost-effectiveness analysis
Perspective	Health care system	Health care system	Health care system and societal	Societal
Study population	Children with polyarticular juvenile rheumatoid arthritis	Dutch JIA patients younger than 18 years eligible for treatment with etanercept; various types of JIA	Patients from adalimumab trial ⁶¹ : children aged 4 to 17 years with JIA	Patients had JIA with a prior inadequate response or intolerance to DMARDs.
Intervention(s)	Etanercept	Etanercept	Adalimumab	Etanercept, adalimumab, abatacept and infliximab vs. methotrexate.

Intervention effect	Effect size measured in terms of CHAQ and mortality. Cost per HAQ point.	Six response variables measured, including overall assessment of well-being, CHAQ score and number of active joints. HUI3 also measured.	CHAQ scores and active joint counts	Proportion of patients who had a reduction in symptoms at 1 year according to the ACR Pedi 30 criteria.
Currency base	UK pounds (GBP, £)	Euros (EUR €)	Russian Roubles (RUB)	Canadian dollars (CAD, \$)
Model type, health states	Not clear	None	Markov model	Decision analysis model
Time horizon	Life course	27 months	7 years / Lifetime	1 year
Base case results	Incremental cost £28,022; incremental effectiveness in terms of QALY 1.7; ICER £16,082. Sensitivity analysis ICER varied from £3,900 to £34,000.	HUI3 score increases from 0.53 to 0.78 after 28 months; Total direct medical costs were €12,478 per patient year after start of etanercept compared to €3720 before start.	For a lifetime horizon, the incremental cost-utility ratio for adalimumab vs. conventional non-biologic therapy was 1,571,500 roubles / QALY.	Cost per ACR Pedi 30 responder were \$26,061, \$46,711, \$16,204 and \$31,209 for etanercept, adalimumab, abatacept and infliximab respectively compared to methotrexate.

CHAQ, Childhood Health Assessment Questionnaire. DMARDs, disease modifying anti-rheumatic drugs. HAQ, Health Assessment Questionnaire. ICER, Incremental Cost-Effectiveness Ratio. QALY, Quality Adjusted Life Year.

Table 39 Critical appraisal checklist for economic evaluations

Item	Cummins¹²²	Prince¹²³	Simpson¹²⁴	Ungar^{125;126}
1. Is the decision problem (including interventions compared and patient group) relevant to the UK?	Yes	Yes	Yes	Yes
2. Is the setting comparable to the UK?	Yes	Unclear	Unclear	No
3. Is the analytical and modelling methodology appropriate?	No	Yes ¹	Yes	Yes
4. Are all the relevant costs and consequences for each alternative identified?	No	Yes	Yes	Yes
5. Are the data inputs for the model described and justified?	Yes	N/A	No	Yes
6. Are health outcomes measured in QALYs?	Yes	No	Yes	No
7. Is the time horizon considered appropriate?	Yes	No	Yes	No
8. Are costs and outcomes discounted?	Yes	No	Yes	No
9. Is an incremental analysis performed?	Yes	Unclear ¹	Yes	Yes
10. Is uncertainty assessed?	Yes	No ¹	Yes	Yes

QALYs, quality-adjusted life years.

¹ The methodology is appropriate for a cohort-based evaluation; however, a full incremental cost-utility analysis has not been performed.

Critical appraisal of the studies

The cost-effectiveness studies were assessed using a critical appraisal checklist (Table 39). The checklist assessed study quality and generalisability to the UK. The checklist was adapted by the review authors from checklists by Philips and colleagues,¹²⁷ Drummond and colleagues¹²⁸ and methodological requirements stated in the NICE reference case.¹²⁹

Cummins and colleagues was conducted in the UK,¹²² whilst the generalisability of the others to the NHS is unclear. The other studies were conducted in the Netherlands,¹²³ Russia¹²⁴ and Canada,¹²⁶ used appropriate modelling methodology; and included relevant costs.

In terms of the analytical and modelling methodology used, the studies were generally considered appropriate, except for the model reported in Cummins and colleagues¹²² which was based upon a number of questionable assumptions due to limitations in the data available at that time (2002).

The data inputs for the model were clearly described and justified by two studies,^{122;126} but the description of some of the data inputs are missing from Simpson and colleagues.¹²⁴ Prince and colleagues¹²³ conducted a cost-consequence analysis based on a prospective observational study that collected cost and utility data. The study did not measure quality-adjusted life years (QALYs).

Two of the studies, Cummins and colleagues¹²² and Simpson and colleagues,¹²⁴ used appropriate time horizons, measured the health outcomes in QALYs and discounted costs and outcomes. The model by Ungar and colleagues¹²⁶ did not use QALYs in the model and used a one year time horizon, eliminating the need for discounting.

All three modelling studies analysed results incrementally and assessed uncertainty through sensitivity analyses.^{122;124;126}

In summary, the cost-effectiveness studies have certain limitations with regard to methodology, reporting of results or generalisability to the UK NHS (Table 39).

Cummins and colleagues

Approach

Cummins and colleagues¹²² consisted of a HTA conducted as part of the NICE appraisal of etanercept for JIA (NICE TA35).¹³⁰ The HTA includes a systematic review of clinical-effectiveness and a critical appraisal of a CS to NICE from Wyeth Laboratories (manufacturer of etanercept). The HTA does not provide an independent economic model due to considerable uncertainties in the available evidence

for JIA at that time. The CS contained a cost-utility analysis of etanercept in patients with JIA, compared with other treatment options. The cost-utility model was based upon a model developed for rheumatoid arthritis in adults. The model used the results from the etanercept RCT by Lovell and colleagues.⁴² The model assumed a positive linear relationship between the Health Assessment Questionnaire (HAQ) score and costs, modelling responders, non-responders, and deaths at each time point.

The model used EQ-5D values derived from mapping HAQ values in adult rheumatoid arthritis patients. Mortality was related to HAQ values, with a 38% increase in mortality per unit change in HAQ. The model assumed a relative risk of mortality in JIA.

Estimation of effectiveness

The HAQ progression rate was 0 for responders for 0 to 4 years, 0.034 for responders after four years and 0.0669 for non-responders. No definition was given for response.

Cummins and colleagues¹²² reported the evidence limitations due to limited or non-existent long-term data on efficacy and lifelong impacts of the disease and treatment. The Juvenile Rheumatoid Arthritis 30 efficacy measure (JRA30) was assumed equivalent to ACR Pedi 20, HAQ and CHAQ were assumed equivalent, and utility and mortality were derived from an adult rheumatic arthritis trial. Due to limited evidence on potential adverse effects, disease progression and long-term prognosis for treatment-resistant JIA insufficiently supported assumptions were made in the economic evaluation. The authors of the review expressed concerns about the validity of the economic model and the assumptions made to extrapolate beyond the limited evidence base.

Estimation of QALYs

Utility values were derived from EQ-5D estimates for adults with rheumatoid arthritis, as there was limited evidence on HRQoL in JIA. The model assumed the HAQ was equivalent to CHAQ and that adult values were therefore appropriate for children. The HAQ score for the placebo arm was 1.3 at baseline and 1.2 after seven months, and the HAQ score for the etanercept arm was 1.6 at baseline and 0.8 at seven months (lower scores indicate better health). In the base-case, the model reported a 1.7 incremental QALY gain in favour of etanercept. However, this result was questioned by Cummins and colleagues¹²² due to the limitations in the evidence for HRQoL.

Estimation of costs

Resource use was considered similar to that for the adult rheumatoid arthritis population. Information regarding resource use was not available from the JIA etanercept trial.⁴² Costs were discounted at 6% per annum and benefits at 1% per annum. The cost offset per HAQ point was £860.

Results

The incremental QALYs were 1.74 for the patients on etanercept as compared to placebo. The incremental cost-effectiveness ratio (ICER) was £16,082 per QALY gained in the base-case analysis and in the sensitivity analyses ICERs ranged between £3,900 (cost-offsets assumption changed to exclude nursing home and home help costs but to include indirect costs) and £34,000 (SF-36 regression used). Probabilistic results were not reported.

Key issues

- There were concerns about the validity of the results due to lack of suitable evidence for model input parameters, particularly with regard to HRQoL.

Prince and colleagues

Approach

Prince and colleagues¹²³ reported a cost-consequence analysis of etanercept therapy in patients with JIA in the Netherlands, who had an insufficient response to the maximum dose of methotrexate. Forty-nine JIA patients were evaluated at start of treatment and after 3, 15 and 27 months of therapy from the National Arthritis and Biologicals in Children (ABC) register. For all included patients, data were collected on the use of etanercept, disease activity and HRQoL. Most of the patients had polyarticular JIA (45%), followed by 22% for both extended oligoarticular and systemic JIA. The remainder had ERA (4%) or juvenile arthritis psoriatica (6%). The median age of patients at the start of etanercept treatment was 11.6 years and median disease duration was 3.6 years.

Estimation of effectiveness

The outcome measure used to assess disease activity consisted of six response variables: (i) overall assessment of disease activity by the physician through the visual analogue scale (VAS); (ii) CHAQ by the patient or parent; (iii) overall assessment of well-being by the patient or parent through the VAS; (iv) number of active joints (joints with swelling and/or limited motion with pain or tenderness); (v) number of joints with limited motion; and (vi) a laboratory marker of inflammation, ESR.

After three months use of etanercept, the mean number of active joints decreased from 16.7 to 3.99 per person and the CHAQ score decreased from 1.70 to 1.00. These outcomes further improved by 27 months to 2.45 active joints per person and a CHAQ score of 0.50 (lower score indicates better outcome).

Estimation of QALYs

HRQoL data were collected for patients in the study using the Health Utility Index-3 (HUI). The questionnaire consists of eight health domains: vision, hearing, speech, ambulation, dexterity, emotion, cognition and pain, each with five or six levels representing the range of functioning. HRQoL was collected by proxy by the parents of the study participants since children were considered unable to value health states. The HRQoL data are reported in more detail in the systematic review of HRQoL studies (section 5.3 of this assessment report).

Estimation of costs

Costs were collected for direct medical costs (i.e. medication, diagnostic and hospitalisation costs). The base year was 2008 for all costs with costs retrieved from other years converted to 2008 Euros using the general Dutch price index rate. Unit costs for medication were retrieved from the Pharmacotherapeutic Compass provided by the Dutch Board of Health Insurances, and treatment costs were calculated with the exact dose of medication and administration period as reported in the patients' files. Prices for all hospital-related costs were based on real prices from the coordinating centre (Erasmus MC Sophia Children's Hospital). The etanercept unit cost was estimated at €10,478 per year.

Results

Mean total direct medical costs after the start of etanercept were on average €12,478 per patient-year compared with €3720 before the start of etanercept treatment. The utility for patients was 0.53 before start of etanercept treatment and increased to 0.78 over 27 months of etanercept treatment.

Key issues

- The study does not report cost-effectiveness.
- The study was based in the Netherlands, so unclear how generalisable results are to the NHS.

Simpson and colleagues

Approach

Simpson and colleagues¹²⁴ reported results from a Markov model developed to assess the cost-effectiveness of adalimumab relative to methotrexate for the treatment of JIA. Cost-effectiveness analyses were performed from the perspective of the Russian health care system (base model) and society as a whole (secondary model). The base case model reported outcomes for a cohort of 100 children, mean age 11 years old. Sensitivity analyses assessed variation of the age of children at treatment initiation. The model has two parts. The first part followed children from age 11 to 18 years with four-month long cycles. This part of the model used data derived from the adalimumab RCT (Lovell and colleagues (2008)⁶¹). Additional analyses followed patients aged seven years at treatment

initiation for a period of 11 years. The second part of the model was derived from the literature on adult rheumatoid arthritis and modelled the remaining lifetime of the patients (age >18 years). The adalimumab RCT⁶¹ compared adalimumab plus methotrexate to placebo plus methotrexate for the treatment of JIA in children aged 4 to 17 years (further detail of this trial can be found in Section 4 of this assessment report).

Disease activity was defined as mild, moderate, or severe using the CHAQ scores and active joint counts from the adalimumab RCT.⁶¹ Health states from the childhood model were used to capture the effects of joint damage and need for hip replacement for the treatment of adulthood JIA (>18 years). The base model included five health states (remission no disease, remission disease, activity mild, activity moderate and activity severe). The remission disease group was introduced to capture the effect of joint damage and the need for joint replacement.

Estimation of effectiveness

Effectiveness estimates were based on observed changes from the adalimumab trial⁶¹ assessed using the CHAQ. These effects within the model were translated to HUI2 utility values, using a mapping algorithm developed by the authors.¹³¹

Estimation of QALYs

For the first part of the model (<18 years) CHAQ items were transformed to HUI2 utility values, using the mapping algorithm. QALY estimates were based on these utility values. Utility values for the second part of the model were derived from the literature and were based on adult patients with rheumatoid arthritis. Mean predicted utility values varied from 0.56 to 0.98 (range 0.18 to 1.00).

Estimation of costs

Health care costs were derived from a study by Yagudina and colleagues¹³² reporting the cost of JIA during a 15-month period in Russia. The costs were for the year 2011. Given that this study reported the cost of one month inpatient and 14 months outpatient treatments, the cost attributable to each health state was adjusted using this as starting point. Base case costs were discounted at a rate of 3%, while additional sensitivity analysis used a discounting rate of 5%.

Results

Relative to conventional non-biologic therapy, adalimumab was assessed to be cost-effective when used to treat JIA patients whose disease severity is comparable to that of participants in the adalimumab RCT.⁶¹ Adalimumab plus methotrexate was reported to be more effective and more costly than methotrexate with an incremental cost per QALY ratio of approximately 1,437,480

roubles (£16,974 at current exchange rate) for the base case (7 years) and 119,496 roubles (£1,411) adopting a lifetime horizon.

Key issues

- There was uncertainty related to the predicted utility values used to estimate QALYs; a recent study indicated how using different algorithms to convert HAQ to utility values affects the cost-effectiveness and health technology assessment results.¹³³
- The lifelong model uses utility estimates derived from adult patients with rheumatoid arthritis. JIA which persists into adulthood has a different disease process to rheumatoid arthritis, and therefore assumptions of similarity between the two conditions are not valid.
- Cost estimates may not be applicable to the UK.
- Mortality rates are assumed to be equal to published rates for the Russian Federation; it is not clear if this refers to the general population or to JIA and age-specific mortality rates.

Ungar and colleagues

Approach

Ungar and colleagues¹²⁶ developed a decision analysis model for etanercept, infliximab, adalimumab and abatacept for polyarticular-course JIA patients in Canada, with an inadequate response or intolerance to DMARDs. The model had a one year time horizon and consisted of two consecutive six month cycles, with no discounting. The model incorporated the probabilities that patients would, based on their response at six months, either continue with the same treatment or switch to an alternative treatment. Patients switched due to lack of response, intolerance to therapy or adverse events (AE). Where data on switching biologic DMARDs were not available in paediatric studies, the relative risk of switching from biologic DMARDs due to non-response or AEs was extrapolated from studies of rheumatoid arthritis in adults. Patients who switched from methotrexate were assumed to receive a biologic DMARD for the next six months, where the cost of the biologic was represented by the average cost of all the biologic DMARDs.

Estimation of effectiveness

The model compared each of the biologic DMARDs against methotrexate, but did not compare them with each other since head-to-head trials were not available and the study populations differed by JIA onset type. The effectiveness measure was the proportion of patients who had a reduction in symptoms at one year according to the ACR Pedi 30 criteria. To derive six-month response rates for each biologic DMARD, data from the key RCTs (Lovell and colleagues, 2000);⁴² Lovell and colleagues, 2008;⁶¹ Ruperto and colleagues, 2007);¹³⁴ Ruperto and colleagues, 2008⁵⁷) were combined with data from registry and observational studies in a meta-analysis. For the base case analysis the

proportion of patients achieving ACR Pedi 30 at six months varied between 79% and 82% for the biologic DMARDs, with the assumption that 30% of patients treated with methotrexate would achieve ACR Pedi 30. Probabilities for switching due to non-response and AEs were estimated from the RCTs and observational studies.

Estimation of QALYs

HRQoL was not included in the analysis.

Estimation of costs

The model included the costs of medication, monitoring costs and costs associated with treating serious infections. Costs were in Canadian dollars and the price year was 2008. In the base case analysis a 40 kg patient was assumed, based upon the mean weight of patients in the 2 paediatric trials that reported weight. The direct medical costs included drug acquisition costs for biologic DMARDs and methotrexate, concomitant drug costs, drug administration materials, nursing time, dispensing fees, physician assessments and laboratory tests. Unit prices of health resources were obtained from public sources, including Quebec and Ontario provincial drug plan formularies for medications, and Ontario Ministry of Health and Long-Term Care fee schedules (laboratory tests and physician fees).

Results

The model reports results as additional cost per additional ACR Pedi 30 responder at one year of \$26,061, \$46,711, \$16,204 and \$31,209 for etanercept, adalimumab, abatacept and infliximab vs. methotrexate, respectively. Probabilistic sensitivity analyses were conducted for each treatment versus methotrexate and cost-effectiveness acceptability curves were calculated. If a decision maker was willing to pay no more than \$30,000 per additional responder, then the probability that etanercept would demonstrate a net economic benefit would be 95%. The willingness to pay points at which the biologic DMARDs had a 50% probability of cost-effectiveness were \$45,000, \$17,000 and \$27,500 for adalimumab, abatacept and infliximab, respectively.

Key issues

- The time horizon was inadequate to model treatment of a long-term condition (only 1 year).
- This was not a cost-utility study as no HRQoL data were included.

Summary of published economic evaluations

- A systematic review of economic evaluations of biologic treatments included four studies. Two of these studies were cost-utility analyses, one a cost-effectiveness study and one was a cost-consequence study.
- The evaluations were published between 2002 and 2012 in the UK,¹²² the Netherlands,¹²³ Russia¹²⁴ and Canada.¹²⁶

- The studies varied in design and structure. The time horizons varied between one year¹²⁶ and lifetime.¹²²
- The comparators differed between studies. One study compared etanercept with methotrexate,¹²² one adalimumab with methotrexate,¹²⁴ one etanercept, adalimumab, abatacept and infliximab with methotrexate¹²⁶ and the remaining study compared a cohort before and after receiving etanercept.¹²³
- There were limitations in the methodological quality in all the studies identified, including limited reporting of model parameters and assumptions. The UK study¹²² is now considered out of date, and it is unclear how generalisable the results from the other studies are given the methodological limitations.

5.3 Systematic review of health related quality of life studies (HRQoL)

Methods for the systematic review

A systematic literature review was undertaken to assess the HRQoL of people with JIA treated with biologic DMARDs. The aim of the review was to provide data to populate the *de novo* economic model in this report with health state utility values to calculate QALYs. The description of the search strategy is shown in Appendix 1. The inclusion criteria were to include primary studies that investigated HRQoL in people with JIA. To be eligible, the study should report health utility values using any generic preference based HRQoL measure (e.g. EQ-5D, SF-6D) or choice-based valuation methods (e.g. time trade off, standard gamble). Studies that were not reported in the English language or did not provide sufficient information were excluded. The methodology used for searching and data extraction is outlined in section 3 of this assessment report.

Results of the systematic review

The database searches identified 2249 references, with one further study retrieved by hand searching, making the total references identified 2250. Full text papers for 28 references were retrieved, meeting the *a priori* inclusion criteria. Figure 7 presents a flow chart of the selection process and the excluded studies with reasons for exclusion are listed in Appendix 7. Six references were considered to have insufficient information on the study methods, population and results, nine included an inappropriate population and ten did not report a relevant outcome measure. Two studies, described in three publications, met the inclusion criteria and the characteristics of these studies are presented in Table 40. Data extraction forms for the included studies are in Appendix 9.

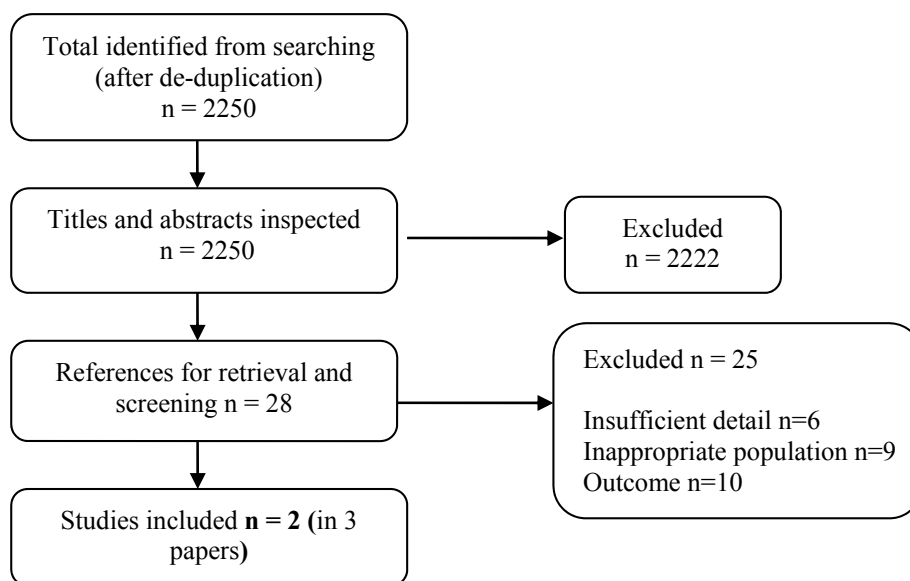


Figure 7 Flow chart of identification of studies for inclusion in the review of QoL studies

Table 40 Characteristics of included quality of life studies

Author	Hendry et al ¹³⁵	Prince et al ^{123;136}
Publication Year	2013	2010, 2011
Country	UK	Netherlands
Study type	RCT	Prospective observational study
Study population	Children / adolescents with JIA and inflammatory joint disease affecting the foot /ankle (n=44)	Children and adolescents with refractory JIA from the national Arthritis and Biologicals in Children register (n=49)
Study population age (mean)	10 years old	11.6 years old
Intervention(s)	Multidisciplinary foot care intervention informed by musculoskeletal ultrasound	Etanercept therapy
Comparator population	Standard care	No treatment
QoL instrument used	EQ-5D	HUI3
Time period where HRQoL instruments administered	At baseline and 12 months	Baseline, 3 months, 15 months and 27 months
Methodology of collecting HRQoL data	EQ-5D was completed by patient (using EQ-5D-Y) and by proxy (using EQ-5D-3L).	The parents of the JIA patients completed the HUI3
Results	EQ-5D was 0.57 and 0.69 for the intervention group at baseline for the self-reported and proxy groups. Results were similar at 12 months and in the control group.	Utility was 0.53 at baseline and increased to 0.78 after 27 months.

HRQoL, Health-related quality of life. JIA, juvenile idiopathic arthritis. QoL, quality of life. RCT, randomised controlled trial.

The two HRQoL studies are each now described in more detail.

Hendry and colleagues

Hendry and colleagues¹³⁵ conducted an exploratory RCT to assess the effectiveness of a multidisciplinary foot care programme in children with JIA, and to investigate the methodological considerations of such a trial.

Children and adolescents with a definitive diagnosis of JIA and inflammatory joint disease affecting the foot/ankle were recruited at a single hospital - the Royal Hospital for Sick Children, Glasgow, UK. Participants were included if they satisfied at least one of the following criteria: (i) previously documented foot arthritis including small joints derived from medical case notes, (ii) previously documented foot arthritis in one or more large joints derived from medical case notes, or (iii) current widespread polyarthritis involving large and small foot joints derived from clinical examination by a consultant paediatric rheumatologist. Patients with an unconfirmed diagnosis of JIA, and/or only upper limb, jaw, or neck involvement were excluded. Hence, a sub-group of the JIA patient population of relevant to this assessment report (i.e. those whose disease had not affected the foot/ankle) was excluded from the study (Table 41).

Table 41 Characteristics of included HRQoL study by Hendry and colleagues

	Multidisciplinary foot care	Standard care
Participants, n	21	23
Age (years), mean (SD)	10.1 (4.22)	10.0 (3.39)
Sex, n	M 7, F 14	M 6, F 17
Disease subtypes, n (%)		
- Persistent oligoarthritis	7 (33)	4 (17)
- Extended oligoarthritis	4 (19)	5 (22)
- Polyarthritis rheumatoid factor negative	6 (29)	10 (43)
- Polyarthritis rheumatoid factor positive	0 (0)	2 (9)
- PA	2 (10)	1 (4)
- ERA	2 (10)	0 (0)
- Undifferentiated	0 (0)	1 (4)
Pharmacological management, n (%)		
- Analgesics	2 (9)	3 (13)
- NSAIDs	2 (9)	3 (13)
- Methotrexate	18 (86)	16 (70)
- Etanercept	7 (33)	5 (22)
- Methotrexate & etanercept	5 (24)	5 (22)
- Sulphasalazine	1 (5)	0 (0)
- Rituximab	0 (0)	1 (4)
EQ-5D utility index at baseline		
- Self, mean (SD)	0.57 (0.31)	0.58 (0.35)
- Self, median (IQR)	0.62 (0.52 to 0.76)	0.66 (0.52 to 0.75)
- Proxy, mean (SD)	0.69 (0.29)	0.60 (0.33)
- Proxy, median (IQR)	0.69 (0.58 to 1)	0.62 (0.55 to 0.82)
Change in EQ-5D utility index at 12 months, median (IQR)		
- Self	0 (-0.1 to 0.01)	0 (-0.04 to 0.04)
- Proxy	0 (0 to 0.11)	0 (0 to 0.1)

IQR, interquartile range. NSAIDs, non-steroidal anti-inflammatory drugs. SD, standard deviation.

Enrolled participants (n=44) were randomly allocated to the intervention group receiving multidisciplinary foot care (individualised care packages including foot orthoses and targeted home exercise programs) or to the control group treated with standard care (normal outpatient medical care from their consultant paediatric rheumatologists). Treatment groups were similar in terms of pharmacological treatment and both had a proportion of patients receiving etanercept. There were small differences in proportions of JIA disease subtypes, but there were no statistically significant differences in baseline characteristics (Table 41).

Patients' HRQoL was collected at baseline and 12 months using the EQ-5D-Y (patients) and EQ-5D-3L (parents/guardians) questionnaires. There were no significant differences in HRQoL between treatment groups at 12 months and both self- and proxy-reported outcomes were similar (Table 41).

Prince and colleagues

Prince and colleagues^{123;136} evaluated changes in HRQoL in patients with refractory JIA who were being treated with etanercept, following an insufficient response to the maximum tolerated dose of methotrexate (NB. This study was also included in section 5.2 'systematic review of existing cost-effectiveness evidence' of this assessment report). Data was collected from Dutch patients registered at the national Arthritis and Biologicals in Children (ABC) register, supplemented by prospectively collected additional data from patients who started etanercept treatment from 2003 until 2006. Three HRQoL questionnaires were used, one of which was the HUI3 preference-based HRQoL instrument. HRQoL questionnaires were completed at the start and after 3, 15 and 27 months of treatment.

Prince and colleagues^{123;136} report the results in two publications and the results differ slightly between the publications. In the publication including costs,¹²³ four fewer patients are included as these patients did not continue treatment with etanercept for at least 27 months, whilst the publication reporting only QoL reports the results for all 53 patients. For the purposes of this assessment report, the smaller dataset is of more relevance,¹²³ but results from both are shown in

Table 42.

The results from the study indicated a statistically significant improvement in the HUI3 utility score from baseline of 0.53 to 0.78 at 27 months follow-up. Mean utility values were 0.69 at three months and 0.74 at 15 months follow-up. For the cohort with more patients there was a mean utility improvement of 0.25 during the 27 months of treatment.¹³⁶ The baseline mean utility value was 0.51 and significant changes were observed in the domains of pain, ambulatory and dexterity.

Table 42 Characteristics of patients included in Prince and colleagues

Characteristic	Prince et al (2010) ¹³⁶	Prince et al (2011) ¹²³
Participants, n	53	49
Age, median (IQR)	11.9 (8.1-14.9)	11.6 (7.9-14.6)
Sex, %	Male 38%, Female 62%	Male 31%, Female 59%
Proportion of sample with systemic JIA	26% (14/53)	22% (11/49)
Proportion of sample receiving methotrexate	80%	79%
HUI utility value (SD) baseline	0.51 (0.04)	0.53 (0.04)
HUI utility value (SD) follow-up (27 months)	0.77 (0.08)	0.78 (0.07)

IQR, Interquartile range. SD, standard deviation.

Summary and conclusions of the HRQoL review

The included studies assessed the HRQoL of children and adolescents with JIA, applying EQ-5D and HUI3 preference-based utility measures. While both studies reported utility values, they are not directly comparable. The study by Hendry and colleagues¹³⁵ assessed the effectiveness of a foot care programme in an RCT, while the Prince and colleagues^{123;136} conducted an observational study reporting HRQoL and costs from patients in the Dutch ABC registry to assess the effect of treating patients with etanercept. The mean utility values reported for baseline by Hendry and colleagues¹³⁵ for the intervention and control group (0.57 and 0.58 respectively) are relatively similar to the baseline values reported by Prince and colleagues study (0.51 and 0.53).^{123;136} The HRQoL values may be higher for Hendry and colleagues¹³⁵ due to 23% of patients receiving etanercept, while no patients received etanercept at baseline in the Prince and colleagues study.^{123;136} The sample size of both cohorts is considered relatively small, but reasonable given the population group. The cohort used in the Prince and colleagues study^{123;136} included patients with systemic JIA which are out of the scope of the current review; however, the proportion of patients within the group with systemic JIA is relatively small (<30%). Neither of the studies can be considered fully informative for the *de novo* economic evaluation in this assessment report. However, estimates provided by Prince and colleagues^{123;136} are considered reasonably appropriate for use in the economic evaluation, despite not being considered directly generalisable to the UK population.

5.4 Review of cost-effectiveness in company submissions to NICE

All four pharmaceutical companies submitted evidence to be considered for the NICE appraisal. Two of these submissions (BMS (abatacept) and Roche (tocilizumab))^{78;85} consisted of a written report and an electronic economic model, and the other two submissions (AbbVie (adalimumab) and Pfizer (etanercept))^{77;97} just comprised a written report.

A structured data extraction form was used by the assessment group to assess the company submissions (CS) (Appendix 10). A description and critique of each of the submissions in turn is provided in the following sub-sections. Greater description is provided of the Roche and the BMS submissions as these conducted economic models. (NB a description and critique of the companies' clinical-effectiveness evidence is given in section 4.2 of this assessment report).

Review of BMS submission to NICE (abatacept)

The company submitted a *de novo* economic model that included all comparators specified in the NICE scope except for methotrexate monotherapy: i.e. abatacept, etanercept, adalimumab, and tocilizumab.⁸⁵ The company states that methotrexate monotherapy was not included due to inconsistency with the clinical-effectiveness data, i.e. all patients in the RCTs either did not have sufficient response with methotrexate or were refractory to methotrexate. The scope reflects the licensed indication of abatacept, polyarticular JIA patients aged six and above who have received at least one tumour necrosis factor α (TNF α) inhibitor (etanercept or adalimumab). All included drugs were assumed to be administered with subcutaneous methotrexate.

Modelling approach

The model presented in the CS is a cohort-based cost-minimisation model, in which all drugs were assumed to have identical efficacy. The base case model presents a cohort of 12-year old polyarticular JIA patients and follows them until age 18 years in four week cycles. The model is essentially a one state model. Patients gain weight and height as they age, but their disease does not change; only the costs associated with treating the disease increase due to weight and body surface area based dosing among the drugs. The drug acquisition cost values within the model were appropriately derived from MIMS data.¹³⁷

Assumptions

The model contained a number of assumptions that appear reasonable: a 52-week year, dosing regimen for methotrexate consistent between under 16 year olds and over 16 year olds, constant weight after age 20, methotrexate dosing based on an algorithm for lowest cost when more than 30 mg methotrexate is necessary, normal distribution for height and weight, truncation of model starting age in the probabilistic sensitivity analysis (PSA) to represent only between age 6 and 16 years as starting ages, assumptions for standard errors where they were unavailable, and no vial sharing of drugs.

Critical appraisal of model

While the comparators and population within the model were generally consistent with the NICE scope, the model does not adequately represent all the available evidence for the treatment pathway or natural history of the disease.

The model time horizon and structure are inadequate to capture long-term treatment effects or the treatment pathway of the disease as many JIA patients continue to receive treatment into adulthood. The model does not allow for drug discontinuation or treatment switching, which is known to happen in clinical practice. The CS indicates that the model was validated by an internal reviewer but full details were not reported of this validation.

Estimation of effectiveness

The model did not include clinical-effectiveness data to represent clinical outcomes or to represent events that incur costs such as disease flare, vision loss or joint surgery. The effectiveness of the biologic DMARDs was assumed to be equivalent as a justification for the use of cost-minimisation methods. The CS cites the systematic review and indirect comparison by Otten and colleagues (2013) for evidence of equivalent effectiveness (as discussed earlier in this assessment report, Section 4).⁷⁹

Estimation of QALYs

HRQoL was not assessed in the model and the CS indicates this was due to uncertainty in the QoL values for JIA in the literature.

Estimation of costs

Intervention dosages and prices were derived from MIMS,¹³⁷ while costs for subcutaneous injection and infusion drug delivery methods were derived from a previously published HTA of biologic DMARDs in rheumatoid arthritis.¹³⁸ A confidential patient access scheme (PAS) was incorporated for abatacept (██████████). Sensitivity analyses were run for the price of tocilizumab using various assumed percentage price discounts as a confidential PAS has been agreed for tocilizumab.

The costs were derived from appropriate sources and are clearly reported, but it is assumed that the drugs had identical AE costs, discontinuation rates, and clinical-effectiveness. The details for drug costs and dosages used in the BMS company model are shown in Table 43.

Table 43 Unit costs and dosages used in the BMS company model

Drug	Cost, £	Dose
Abatacept	302.40	250mg
Etanercept	35.75	10mg
	89.38	25mg
	178.75	50mg

Adalimumab	352.14	40mg
Tocilizumab	102.40	80mg
	256.00	200mg
	512.00	400mg
Methotrexate	14.85	7.5mg
	15.29	10mg
	16.50	12.5mg
	16.57	15mg
	17.50	17.5mg
	17.84	20mg
Administration Method		
Infusion	154.00	
Subcutaneous Injection	3.05	

Results

The model was a cost-minimisation and only analysed costs, assuming equivalent clinical-effectiveness for all biological DMARDs. The results of the base case analysis are shown in Table 44 and Table 44. The model results presented by the manufacturer include a PAS discount of ■■■ for abatacept.

Table 44 Results of the BMS model base case (CS Table 13)

Results for the base-case model (12 year olds, 6 year time horizon, from Excel model)				
	Abatacept	Adalimumab	Etanercept	Tocilizumab
Drug costs	■■■■■	■■■■■	■■■■■	■■■■■
Administration costs	£11,797	£871	£871	£11,646
Total costs	■■■■■	■■■■■	■■■■■	■■■■■
Cost savings with abatacept		■■■■■	■■■■■	■■■■■

In the base case, the company found that abatacept was the least costly biologic DMARD. The CS states that abatacept has similar efficacy and safety to the other biologic DMARDs. Deterministic sensitivity analyses and PSA were conducted for a variety of scenarios. The findings were found to be robust to a wide range of scenarios.

The company undertook a number of deterministic and scenario sensitivity analyses:

- A sensitivity analysis was undertaken wherein the infusion costs for tocilizumab were increased due to the longer infusion time.
- The starting age of patients in the model was varied between 6 and 16 years. In the biologic DMARD trials the mean age was 11 years at baseline, but the drug licenses were for much younger ages.
- The time horizon of the model was varied between 6 months and 20 years. Longer time horizons were meant to represent that a third of children with JIA will have it continue into adulthood.

- PAS discount for tocilizumab was tested for a range of percentage of list price reductions and calculated to show the tocilizumab discount with identical drug costs to abatacept.
- Methotrexate was excluded from the etanercept arm

There were no analyses that varied more than one parameter at a time. None of the one-way sensitivity analyses were accompanied by a probabilistic analysis that reassessed the probability that abatacept was the least costly biologic for second line biologic DMARD therapy of polyarticular JIA. Given that the model was simple and the number of simulations for the PSA was only 1000, these analyses would have been simple and quick to do. It would have been especially informative to do this for the tocilizumab cost sensitivity analyses. No analyses looked at sub-groups of patients, and there was no discussion of potential sub-groups.

The PSA results using 1000 simulations were within 5% of the base case analysis results. Abatacept was the least costly option in 67% of simulations, while etanercept was the least costly option in the remaining 33%. Not all distributions used in the sensitivity analysis were appropriate. For the infusion costs and subcutaneous administration costs, a normal distribution was used for the cost data although the use of a normal distribution could lead to simulations with negative cost values; a gamma or lognormal distribution would be more appropriate. The drug cost data does not appear to have been subjected to uncertainty. Given that there is a PAS for tocilizumab, using a lower average cost value for tocilizumab and subjecting it to PSA could have been an alternative approach to only conducting deterministic analysis for tocilizumab. This would give more realistic estimates of the cost uncertainty between treatments.

Critique of the company's submission

The company constructed a cost-minimisation model, assuming that there were no differences between the biologic DMARDs in clinical-effectiveness, AEs and discontinuation rates. Patients do not discontinue or switch treatments, which does not reflect current clinical practice or the available evidence.

Overall, the model can be considered limited for decision making due to factors such as inadequate time horizon and structural limitations. However, the methods for integrating variable dosing over time used within the model may be useful for building a more comprehensive model.

Review of the Roche submission to NICE (tocilizumab)

The CS includes an economic model and reports the total costs, the QALYs gained and cost-effectiveness of tocilizumab in the treatment of polyarticular JIA.⁷⁸ The model evaluates the lifetime

costs and benefits for tocilizumab compared to adalimumab. The perspective of the analysis is that of the NHS and Personal Social Services (PSS).

Modelling approach

A *de novo* Markov state-transition decision model was developed in Microsoft Excel with three health states (uncontrolled disease / off treatment, on treatment and dead). The model has six-month cycles and a time horizon of 25 years. Costs and benefits are discounted at 3.5%. Patients entering the model have active JIA and have previously experienced an inadequate response to, or were intolerant of methotrexate. Patients in the model have a mean age of 11 years and are based on those in the CHERISH RCT.⁶⁸

Patients start with uncontrolled disease at cycle 0 then move to first line biologic treatment. Once all lines of treatment are exhausted, patients move into uncontrolled disease health state. Mortality is included in the model and assumes a 1% six-month mortality rate across all years. The model includes the occurrence of serious AEs. The mortality rate used in the model is about 100 times higher than the annual mortality rate for the general paediatric population of 0.02%. We consider that a lower mortality rate should be used in the model.

Assumptions

The CS states that due to differences in terms of trial design, patients, methods of imputation and quality, only adalimumab and tocilizumab could be compared. The CS states that an indirect comparison of safety was not possible and so the risk of serious AEs was assumed to be the same for both biologic DMARDs. The model assumes that patients discontinue at a rate proportional to their ACR response, i.e. no response (ACR Pedi <30), moderate response (ACR Pedi 30 – ACR Pedi 70) and good response (ACR Pedi ≥ 70).

Critical appraisal of model

The submission meets all of the requirements for methodological quality and generalisability, except that it did not fully explore uncertainty or provide any evidence that the economic model had been validated.

The evaluation provided a clear statement of the decision problem to be addressed, including the population, which appeared to follow the scope for the appraisal issued by NICE. The comparators included (adalimumab and tocilizumab) were appropriate as these are being routinely used or considered for use within the NHS in England and Wales. The model also included etanercept in an exploratory analysis, but did not include abatacept and the CS states that this was not possible due to differences in trial design, patients, methods of imputation and quality. The 25-year time horizon,

reflects the chronic nature of the disease and allows for all relevant costs and benefits to be included. The model structure was clearly presented with a description and justification of the key assumptions and data inputs used. Benefits for the model are measured in QALYs using the HUI3 for measuring utility. All benefits and costs are discounted at 3.5% as required by NICE.¹²⁹ The CS does not assess uncertainty in sensitivity analysis. It was unclear if the model had been fully validated as no details were provided.

Estimation of effectiveness

The company reported a systematic review of biologic DMARDs in the treatment of JIA. The CS states that it was only possible to compare adalimumab and tocilizumab by indirect comparison as these two treatments had greater similarities in trial design, patients, methods of imputation and quality. Results from each study were combined using a hierarchical Bayesian indirect treatment comparison, using an ordered probit model in WinBUGS software to estimate the relative treatment effects and achieving different levels of ACR response. The ACR response rates were estimated for the biologic DMARDs with and without methotrexate and are shown in Table 45. The response is generally similar for adalimumab and tocilizumab (both with methotrexate).

Table 45 ACR Pedi response rates from Roche submission (CS Table 21)

		ACR Pedi 30	ACR Pedi 50	ACR Pedi 70	ACR Pedi 90
Without methotrexate	Placebo	31%	28%	25%	12%
	Tocilizumab	62%	59%	54%	35%
	Adalimumab	52%	49%	44%	26%
With methotrexate	Methotrexate	52%	51%	41%	25%
	Tocilizumab	72%	70%	61%	44%
	Adalimumab	76%	75%	66%	49%

The discontinuation rate used in the model was derived according to ACR response from the Dutch Arthritis and Biological in Children (ABC) register.¹²³ An exponential distribution was fitted to the data for no response (ACR Pedi response < 30), moderate response (ACR Pedi response > 30 and <70) and good response (ACR Pedi response ≥ 70). The six-month discontinuation rate was 0.126 for no response, 0.09 for moderate response, and 0.042 for good response.

Estimation of QALYs

The company conducted a literature review that identified one study reporting utility values suitable for use in the model (Prince and colleagues¹²³). This study reported utility scores obtained using the HUI3 questionnaire to JIA patients starting treatment with etanercept in the Dutch ABC register. Based on these data, the company used for values at time 0 for the patients who are off treatment (utility of 0.53), and used values at time one year for patients on treatment (utility of 0.73).

The assessment group identified a couple of errors in the Roche model with regard to estimation of QALYs. Firstly, utility values for patients have been applied as if patients were on treatment for some time after finishing the first-line biologic treatment, when these patients should have been assigned the off treatment utility. Secondly, utility values have been incorrectly calculated as the utility value for one year has been assigned to each cycle of six months. Corrections to these errors can be found in Table 48 of this assessment report.

Estimation of costs

The costs associated with each health state was obtained from Prince and colleagues (2011),¹²³ who report costs data from the Dutch ABC register for the year before and after starting etanercept. The total six-month health state cost for patients on treatment is £912.33 and off treatment is £1591.43. Treatment unit costs and doses were reported. Tocilizumab was provided with a confidential PAS discount [REDACTED]. The model included costs for both IV infusion and for subcutaneous injection, as required by the treatment. The administration cost of an infusion (for tocilizumab) was £152.24, using inflated costs from Barton and colleagues (2004).¹³⁹ The cost of an administration of a subcutaneous injection was £6.10 for children and £3.05 for a young person, assuming that a proportion of these patients would require nurse assistance.

Cost-effectiveness results

Table 46 and Table 47 show the cost-effectiveness results from the CS for tocilizumab compared to adalimumab when used in combination with methotrexate or as a monotherapy. The CS states that the results indicate that both treatments are of similar clinical- and cost-effectiveness whether used in combination with methotrexate or as a monotherapy. The company urges caution in interpretation of the QALY estimates, but conclude that tocilizumab is less expensive and therefore represents better value to the NHS.

Table 46 Roche base case results: combination therapy

	Adalimumab + methotrexate	Tocilizumab + methotrexate	Incremental Difference	ICER (£ per QALY)
Total QALYs	18.76	18.72	-0.0303	
Total Cost	£81,827	£70,707	-£11,120	South West Quadrant ¹

ICER, Incremental Cost-Effectiveness Ratio. QALY, Quality Adjusted Life Year.

¹ Adalimumab vs. tocilizumab has an ICER of £367,551

Table 47 Roche base case results: biologic DMARD monotherapy

	Adalimumab	Tocilizumab	Incremental Difference	ICER (£ per QALY)
Total QALYs	18.65	18.7	0.0455	
Total Cost	£74,576	£68,560	-£6,015	Dominant

ICER, Incremental Cost-Effectiveness Ratio. QALY, Quality Adjusted Life Year.

The CS does not include any sensitivity analyses. It includes an exploratory analysis with etanercept. This analysis assumed a class effect across anti-TNFs in polyarticular JIA.

Critique of the company’s submission

There are some concerns over the reliability of the model results in the Roche submission due to errors found by the assessment group in the calculation of QALYs.

The assessment group has corrected the errors in the Roche model by applying the off treatment utility values when patients finished the first-line biologic treatment and assigning the 6-month utility value to each cycle. In addition the mortality rate has been reduced to 0.03% per cycle to reflect that of the general population. The results for this analysis are shown in Table 48. The corrected results show reduced QALYs and increased costs for adalimumab and tocilizumab in combination with methotrexate compared to the base case results. However the incremental QALYs and costs between the tocilizumab and adalimumab are similar in the corrected results to the base case results.

Table 48 Corrected Roche model results: combination therapy

	Adalimumab + methotrexate	Tocilizumab + methotrexate	Incremental Difference	ICER (£ per QALY)
Total QALYs	10.10	10.05	-0.05	
Total Cost	£95,761	£83,593	-£12,168	South West Quadrant ¹

ICER, Incremental Cost-Effectiveness Ratio. QALY, Quality Adjusted Life Year.

¹ Adalimumab vs. tocilizumab has an ICER of £251,208

Review of AbbVie submission to NICE (adalimumab)

The company did not provide a systematic review of cost-effectiveness studies or an economic evaluation.⁷⁷ They discussed the interventions in the NICE scope: adalimumab, etanercept, abatacept, tocilizumab, and methotrexate. In sections 5.8 to 5.10 of the CS, the company provides justifications for not conducting an economic evaluation. Other sections of the CS provide details on what the company would consider important in conducting an economic evaluation in JIA, including an evaluation of the costs associated with surgeries and vision loss.

The company states that an economic evaluation was not conducted due to a lack of the appropriate utility data for HRQoL, heterogeneity in study methods and populations between the interventions that complicated indirect comparisons, and a lack of long-term effectiveness data. The company identified one HRQoL study (Prince and colleagues, 2011¹²³), which collected HUI3 utilities in addition to other JIA clinical variables such as the CHAQ score. The data collected from Prince and colleagues (2011)¹²³ was deemed unsuitable to map the CHAQ to HUI3 due to insufficient sample

size, but was considered the most suitable source of utility data by the company. The CS discusses the use of an algorithm by Khan and colleagues¹⁴⁰ that mapped the PedsQL instrument (Pediatric Quality of Life Inventory) to EQ-5D in secondary school pupils. The company notes the potential limitations in the use of this method in JIA, as PedsQL was not collected in any of the JIA biologic DMARD RCTs. The company considers that the biologic DMARD trial populations and study methods were not sufficiently similar to allow indirect comparison through network meta-analysis. The CS concluded that using current data and methods would lead to “untenable” uncertainty (CS page 90).

Review of the Pfizer submission to NICE (etanercept)

Pfizer did not submit any cost-effectiveness evidence.⁹⁷ The CS notes the limitations raised in the previous submissions for NICE of etanercept TA35⁴³ and tocilizumab TA238.⁴⁴ These relate to the limitations in the HRQoL data and the limited evidence on the long-term outcomes and the effectiveness of the treatments. The CS states that any cost-effectiveness evidence would be associated with considerable and unresolvable uncertainty. The company submitted a cost-analysis that compared the annual costs for the first year of treatment based on etanercept against adalimumab and tocilizumab in patients with polyarticular JIA. The CS states that the cost-analysis showed that etanercept is the biologic DMARD with the lowest acquisition cost compared to list prices for tocilizumab and adalimumab.

Comparison of economic models in companies’ submissions

The companies’ submissions differ in the approach to providing economic evidence for biologic DMARDs for JIA. Only one company (Roche) constructed an economic model for tocilizumab that included both costs and outcomes. Two companies (BMS for abatacept and Pfizer for etanercept) submitted cost-analyses and assumed that the biologic DMARDs were equivalent, whilst the remaining company (AbbVie for adalimumab) considered that there were too many limitations with any potential analysis and therefore did not submit an economic analysis.

While AbbVie has raised valid concerns about uncertainty in the data available for conducting an economic evaluation, we consider that concerns about uncertainty are an insufficient justification for not building an economic model. A model provides a representation of current knowledge in a subject and uncertainty is part of that current knowledge. A model, even an uncertain one with limitations noted by the company, gives a more transparent description of available knowledge and allows more informed decision making than simply presenting clinical trial data from trials that only represent a highly selected sub-group of the drug licenses. Modelling also allows the exposure of the most valuable areas for future research enquiries.

BMS and Pfizer consider that all of the treatments are equivalent. It is noted that the available evidence base consists of small trials that lack the statistical power to justify this assumption and Otten and colleagues (2013)⁷⁹ do not conclude that there is equivalent efficacy between the treatments, but that the treatments are similar.

Briggs and O'Brien (2001)¹⁴¹ argue that cost-minimisation analysis should only be conducted when equivalence of comparators has been statistically demonstrated. Dakin and Wordsworth (2013)¹⁴² argue that the limitations of cost-minimisation analysis do not allow appropriate assessment of uncertainty or value of future research and may lead to biased conclusions. It is also the case that equivalence of one clinical outcome does not mean equivalence of all clinical outcomes. Patients may have the same QoL on treatment, but have different adherence and discontinuation, or different AEs for example. For these reasons, cost-minimisation analysis is generally foregone in favour of cost-effectiveness analysis and/or cost-utility analysis.

Roche have provided a cost-utility analysis that compared two of the biologic DMARDs (adalimumab and tocilizumab). The model appears to be a reasonable attempt at modelling JIA, albeit only in two of the biologic DMARDs. However, we have noted errors in the calculation of QALYs in the model, which limit the credibility of the results.

5.5 Independent economic evaluation

The models described in our systematic review of economic evaluations (Section 5.2) had certain methodological limitations and were not wholly generalisable to the NHS. Furthermore, the economic evaluation used to inform the NICE appraisal of tocilizumab for systemic JIA (NICE TA238) was subject to a number of concerns from the Appraisal Committee, especially with regard to the estimation of HRQoL.⁴⁴ Given the limitations of existing available models we therefore constructed a *de novo* economic model to inform this current appraisal.

The model estimates the costs, benefits and cost-effectiveness of the four biologic DMARDs in patients with JIA and inadequate responses to, or intolerance of, methotrexate. The model compares the biologic DMARDs (in combination with methotrexate, where permitted) with a DMARD (e.g. methotrexate), as specified in the NICE scope. The model does not compare the biologic DMARDs with best supportive care (e.g. NSAIDs; corticosteroids) for patients who cannot tolerate a DMARD as there are limited data available to make this comparison. Furthermore, patients who are intolerant to a DMARD such as methotrexate would be offered a biologic DMARD rather than best supportive care, particularly to avoid potential adverse effects of long-term corticosteroid use.¹⁴³

The evidence used in the model was taken from data sources for such as the RCTs of biologic DMARDs (in which a number of JIA sub-types were represented, with polyarticular course JIA being the predominant sub-type), and data sources such as registry studies comprising mixed JIA populations (primarily comprising polyarticular and oligoarthritis JIA patients, but also small proportions of patients with ERA, PA and systemic JIA). However, there was insufficient evidence available for all input parameters to permit a cost-effectiveness sub-group analysis for each of the respective types of JIA within the scope of the appraisal. Therefore the modelled patient population is people with JIA, with the results of particular relevance to people with polyarticular course JIA (extended oligoarthritis, and RF +ve and RF -ve polyarthritis). The biologic DMARDs are assessed in this report within their licenced indications (e.g. the cost-effectiveness estimates for some of the biologic DMARDs cannot be applied to JIA sub-types for which they are not licensed, such as abatacept and tocilizumab for the treatment of ERA and PA).

The model was populated with clinical-effectiveness data from the included RCTs in our systematic review of clinical-effectiveness (Section 4), HRQoL data from our systematic review of HRQoL studies (Section 5.3) and cost data derived from published studies (where available), as well as national and local NHS unit costs.

The economic evaluation was from the perspective of the NHS and PSS, with only these direct costs included. The model estimates the long term costs and benefits from each of the treatments. The costs and benefits were discounted at 3.5%, as recommended by NICE.¹²⁹ The base price year for the costs was 2014. The intervention effect, in terms of reducing disease flare, was derived from the systematic review of clinical-effectiveness reported in Section 4. The outcome of the economic evaluation is reported as incremental cost per QALY gained.

5.6 Methods for independent economic analysis

A Markov model was developed in Microsoft Excel to assess the cost-effectiveness of the biologic DMARDs. The model contains health states for 'on treatment', 'off treatment', 'remission off treatment' and 'death'. A diagram of the model is shown in Figure 8. The model uses three-month cycles to be consistent with the usual time between outpatient appointments for JIA patients. A time horizon of 30 years was modelled as the base case, with shorter and longer horizons tested in sensitivity analyses. This time horizon was considered sufficiently long to capture the costs and effects of biologic DMARDs for paediatric patients, given the uncertainty around the long term clinical outcomes for adults with JIA. The model structure is based upon the clinical pathway of patients who participated in the withdrawal RCTs, described in section 4, the natural history of JIA described in registry data and discussions with clinical experts. The starting age of patients in the

model is 11 years old, based upon the mean age of patients in the RCTs. Patients' height and weight are assumed to be similar to those in the general population.

Patients treated with a biologic DMARD continue on treatment unless they die or withdraw from treatment due to adverse effects, loss of efficacy or clinical remission off treatment. Patients with clinical remission who have their treatment discontinued may relapse and resume treatment with a biologic DMARD. Based on clinical advice, we assumed that clinicians would be reluctant to stop treatment with a biologic DMARD due to remission because of the risk of relapse and so for the base case analysis we assume that no patients discontinue treatment due to remission. We investigate discontinuation due to remission in more detail in a scenario analyses. Therefore, for the base case analysis no patients enter the 'clinical remission off treatment' health state and this is indicated in the diagram by dotted lines for this health state.

In the base case analysis, patients treated with adalimumab, etanercept and tocilizumab receive only one line of biologic DMARD treatment. Patients treated with abatacept receive two lines of biologic DMARD treatment, as abatacept is licensed for use only after a preceding anti-TNF. Following withdrawal from these biologic DMARDs, patients continue on a standard treatment regimen that does not contain a biologic DMARD.

In a scenario analysis (scenario v), we investigate multiple lines of treatment with biologic DMARDs, which reflects the range of strategies used in clinical practice. Patients continue on the subsequent treatments until they die or withdraw from treatment, due to adverse events, loss of efficacy or clinical remission. Following withdrawal from the final biologic DMARD, patients continue on a standard treatment regimen that does not contain a biologic DMARD.

Patients treated with methotrexate only (i.e. not receiving a biologic DMARD) are assumed to continue on treatment unless they die or withdraw from treatment due to adverse events, loss of efficacy.

The model incorporates disease flares to estimate the clinical-effectiveness of treatment. This was the primary outcome measure in the RCTs of the biologic DMARDs. Patients who have a disease flare continue in their current health state in the model but are allocated a HRQoL disutility and an additional health care cost during that cycle.

The costs in the model comprise drug treatment, consultation, and monitoring costs and costs for treating AEs. Costs used in the model are described in more detail in Section 5.6.1. HRQoL is estimated according to each health state.

In each cycle the total costs and QALYs are calculated by multiplying the individual costs and HRQoL by the number of people in the cohort still alive for each of the treatments. The total long term costs and QALYs are calculated by aggregating the costs and QALYs for all cycles. The total discounted QALY gain and cost of treatments are calculated and compared to give the cost-effectiveness of the treatments.

Assumptions are applied to all treatment options unless explicitly stated otherwise. These assumptions have been made due to an absence of data and have been informed by discussion with our clinical advisors. The model includes the following main assumptions:

- There are few studies for biologics, other than etanercept, that report long term discontinuation due to adverse events and inefficacy. The discontinuation rate is assumed to be similar for each of the biologic DMARDs.
- Our clinical-effectiveness review concluded that there is no evidence of a difference in efficacy between biologics and therefore we assumed that the quality of life utility values are the same for all biologic DMARDs ('On treatment').
- It is currently unclear whether the effectiveness of subsequent lines of biologic DMARDs would be reduced or remain the same. The effectiveness of the biologic DMARDs is assumed to be similar taken as a first or subsequent line biologic treatment. This applies to the abatacept as second line biologic DMARD in the base case, and to the scenario analysis that models three lines of biologic DMARD treatment (scenario v).

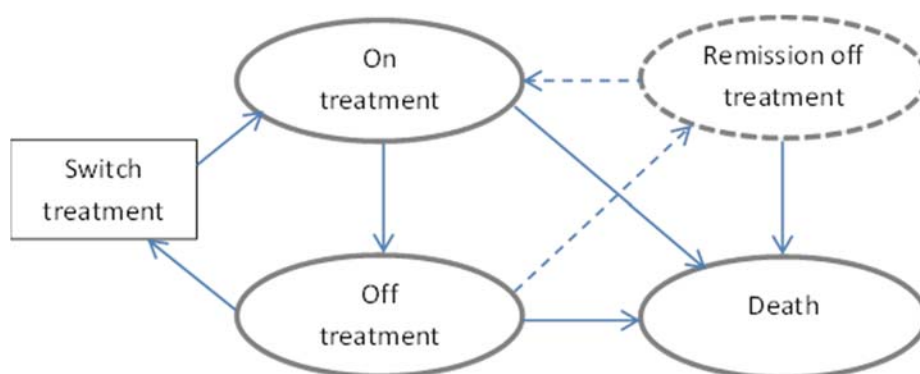


Figure 8 Schematic of the SHTAC JIA model structure

Evaluation of uncertainty

The evaluation of the cost-effectiveness of treatment for JIA is based on uncertain information about variables such as the clinical effect, HRQoL and resource use. This uncertainty was evaluated using

deterministic and PSA. One-way deterministic sensitivity analyses were conducted to evaluate the influence of individual parameters on the model results and test the robustness of the cost-effectiveness results to variations in the structural assumptions and parameter inputs (Section 5.7). Where possible, the parameters were varied according to the ranges of the confidence intervals of these parameters based on the published estimates. Where these data were not available an alternative range was chosen.

Multi-parameter uncertainty in the model was addressed using PSA (Section 5.7).¹⁴⁴ In the PSA, probability distributions are assigned to the parameter point estimates used in the base case analysis. The model is run for 1000 iterations, with a different set of parameter values for each iteration by sampling parameter values at random from their probability distributions.

The uncertainty surrounding the cost-effectiveness of the treatment is represented on a cost-effectiveness acceptability curve (CEAC) according to the probability that the intervention will be cost-effective at a particular willingness to pay threshold. Appendix 11 reports the parameters included in the PSA, the form of distribution used for sampling each parameter, and the upper and lower limits assumed for each variable.

Model validation

The model was validated by checking the model structure, calculations and data inputs for technical correctness by another researcher. The structure was reviewed by clinical experts from the advisory group for its appropriateness for the disease and its treatment. A senior health economist from the advisory group reviewed the methods and assumptions of the economic evaluation. The robustness of the model to changes in input values was tested using sensitivity analyses to ensure that any changes to the input values produced changes to the results of the expected direction and magnitude.

5.6.1 Data sources

Effectiveness data

Disease flare

The risk of disease flare was included in the model as a relative risk compared to methotrexate (Table 49), as derived from our systematic review of clinical-effectiveness (Section 4.1.2). The baseline risk of flare for methotrexate was a weighted average of the risk of flare estimates from the placebo arms of the abatacept, adalimumab and tocilizumab trials, converted to a three month cycle risk. For each biologic DMARD, the risk of flare was derived using the relative risk for that treatment, compared to methotrexate from the relevant RCT, multiplied by the baseline risk.

Table 49 Risk of disease flare

Disease Flare	Risk of flare per cycle	Source
Methotrexate	0.25	Ruperto et al. ⁵⁷ , Lovell et al. 2008 ⁶¹ Brunner et al. ⁶⁸
Abatacept	0.09	Ruperto et al. ⁵⁷
Adalimumab	0.14	Lovell et al. 2008 ⁶¹
Etanercept	0.09	Lovell et al. 2000 ⁴²
Tocilizumab	0.14	Brunner et al. ⁶⁸

Treatment discontinuation

Treatment discontinuation was assumed to be due to AEs, lack of efficacy or clinical remission. Estimates for treatment discontinuation were identified through a literature search of trial and registry data. The first model cycle has certain different treatment discontinuation parameters because it was designed to represent the open label lead-in phase of the RCTs.

The estimates for the first model cycle were taken from the RCTs of the biologic DMARDs^{42;57;61;68} for the open-label lead-in period (Table 50). For the period after the treatment lead-in, the clinical trials of the biologic DMARDs also include other categories of withdrawal (such as patient/guardian consent or physician decision), but it is unclear whether these categories would also apply to clinical practice or were particular to the trials. As a result, the discontinuation rate after the first model cycle was not taken from these clinical trials but from Tynjala and colleagues (2009)¹⁴⁵ who conducted a retrospective observational study on JIA patients in Finland taking etanercept or infliximab with a four-year follow-up.

There was little long-term data on treatment discontinuation identified except for etanercept and hence we assumed that the discontinuation rate would remain constant over time and would be the same for all biologic DMARDs (based on data for from Tynjala and colleagues¹⁴⁵). The discontinuation rate was 7% for AEs and 28% for inefficacy over four years. The discontinuation rate for inefficacy for methotrexate was taken from a retrospective analysis of the German Methotrexate Registry,¹⁴⁶ which collected data on the efficacy and safety of methotrexate treatment since 2005. The discontinuation rate for inefficacy used in the model was 0.4% per cycle. The STRIVE registry (mentioned in section 5.4 of this report) reported a methotrexate discontinuation rate for adverse events of 2.3% per year and this was converted to a three-month rate and used in the model.⁷⁷

Table 50 Discontinuations during the trials' lead-in time (1st cycle)

	Abatacept⁵⁷	Adalimumab⁶¹	Etanercept⁴²	Tocilizumab⁶⁸
Adverse events	0.5%	1.8%	1.4%	1.6%
Loss of efficacy	9.5%	3.5%	2.9%	8.0%
Total discontinuation	10.0%	5.3%	4.3%	9.6%

Mortality

Patients are assumed to have the same mortality rate as for the general population. Mortality was taken from age-related statistics from the Office of National Statistics.²¹

Health related quality of life (HRQoL)

Our systematic review of HRQoL studies (see Section 5.3) identified two potentially relevant studies that reported generic preference-based HRQoL studies of people with JIA who received a biologic DMARD.^{123;135} Furthermore, none of the clinical trials of the biologic DMARDs under review collected HRQoL data that could be used as health state utility values. We investigated methods for mapping HRQoL to treatment response, for example from CHAQ or ACR Pedi to HUI3, but concluded that the data available were insufficient to provide a reliable fit for modelling. Therefore the utility values used in the model were taken directly from the Dutch ABC registry by Prince and colleagues,¹²³ as this study was considered to be of most relevance from the available literature. This study is described in more detail in section 5.3 of this report. It consists of patients who have a polyarticular course JIA and in whom the response to maximum dose of methotrexate is not sufficient. The utility value for patients who had not yet received etanercept in that study is assumed to be representative of those patients with uncontrolled disease not currently receiving a biologic DMARD, i.e. those patients in the methotrexate only arm and those patients who discontinue biologic DMARD therapy. In the study by Prince and colleagues,¹²³ most of the patients were still receiving methotrexate. In the absence of any other utility data, we assumed that all biologic DMARDs would have the same utility values as each other, and this would increase over time as seen in the Prince and colleagues study.¹²³ For simplicity, for the scenarios with additional lines of biologic treatments (scenario analysis v), we assumed that treatment with the biologic DMARD would have a constant utility value of 0.74 (i.e. the value after 15 months of treatment for second and third line biological DMARD treatment). The annual health state utility values used are shown in Table 51.

Patients with a disease flare are assumed to have an associated disutility. We assumed that patients with a disease flare would have a similar HRQoL to patients with uncontrolled disease, but with appropriate treatment would recover from their disease flare within three months (one model cycle). Assuming patients recovered HRQoL at a constant rate over the model cycle, then the average

HRQoL for these patients during that cycle would be 0.655, and converting this to an annual disutility would be equivalent to 0.03 per flare.

Table 51 HRQoL utility values

HRQoL utility values	Per year	Source
No treatment	0.53	Prince <i>et al.</i> ¹²³
Treatment with first line biologic, 0-3 months	0.53	Prince <i>et al.</i> ¹²³
Treatment with first line biologic, 3-15 months	0.69	Prince <i>et al.</i> ¹²³
Treatment with first line biologic, 15-27 months	0.74	Prince <i>et al.</i> ¹²³
Treatment with first line biologic, 27+ months	0.78	Prince <i>et al.</i> ¹²³
Treatment with second and third line biologic	0.74	Prince <i>et al.</i> ¹²³
Disutility for disease flare	0.03	Assumption

Caregiver disutility

We conducted a literature search of studies reporting the quality of life impact on caregivers of patients with JIA and did not identify any studies that reported HRQoL as utility values. The precise quality of life impact on primary caregivers of patients with JIA, in terms of change in HRQoL utility values, is unclear. A study by Bruns and colleagues¹⁴⁷ examined the HRQoL and disease burden of primary caregivers of 70 patients with JIA. They used the CHAQ, SF-36, and the psychiatric screening questionnaire (SRQ-20). The burden of disease on the caregivers was measured by the caregiver burden scale (CB Scale). They concluded that there was a high prevalence of psychoemotional disturbance in JIA caregivers and the burden of disease on the caregivers was primarily related to patients' emotional status (rather than their physical status).

In the absence of suitable HRQoL data for caregiver disutility, we assumed in the base case analysis there was no utility benefit for parents of children and young people and varied this assumption in a scenario analysis (scenario iv).

Estimation of costs

Drug costs

Drug unit costs and doses were based on the British National Formulary for Children (BNFC) 2015.¹⁴⁸ A summary of the dose and unit cost of treatment for each of the comparators is given in Table 52. The manufacturers of abatacept and tocilizumab have provided a confidential PAS. Cost-effectiveness results for these treatments presented in section 5.7 of this report are based on the drug list price, whilst a commercial in confidence separate appendix to this report presents results with the confidential PAS discount applied. Patient height, weight and body surface area were taken from the British National Formulary (BNF) and reflect the increase in child's height and weight as they grow

older.¹⁴⁹ The administration costs for IV infusion was £154 based on an HTA monograph of disease modifying drugs in the treatment of rheumatoid arthritis (Stevenson and colleagues¹³⁸). We assumed that for patients taking methotrexate, half would receive oral and half subcutaneous administration, based upon clinical advice.

Table 52 Drug acquisition costs and dosages (Source BNFC)

Parameter	Methotrexate	Abatacept	Adalimumab	Etanercept	Tocilizumab
Drug dose	10 - 15 mg/m ²	10mg/kg	24 mg/m ² (max. 40 mg) aged 4-13 years; 40 mg (aged 13-18 years)	0.4 mg/kg (up to a maximum of 25 mg per dose)	10mg/kg for patients less than 30 kg; 8mg/kg for patients over 30 kg
Method	Subcutaneous injection / Oral	IV infusion	Subcutaneous injection	Subcutaneous injection	IV infusion
Dosing schedule	Once weekly	Infusions given at week 0, 2,4, 8,12, 16	Every other week	Twice weekly	Every 4 weeks
Unit cost	Oral: 2.5 mg, 24 tab pack £2.22, 28 tab pack £2.60; Subcutaneous: Metoject pre-filled syringe. 50 mg/mL, 0.15 mL=£14.85, 0.2 mL = £15.29, 0.3 mL=£16.57, 0.4 mL = £17.84, 0.5 mL = £18.48, 0.6 mL = £18.95	250-mg vial = £302.40	40-mg prefilled pen or prefilled syringe = £352.14	10-mg vial = £35.75; 25-mg vial = £89.38, 25-mg prefilled syringe = £89.38	3 mL (80-mg vial) = £102.40, 10 mL (200-mg vial) = £256, 20 mL (400-mg vial) = £512.00
Administration cost	0	£154	0	0	£154

Patients taking biologic DMARDs also receive concomitant methotrexate treatment as shown in Table 53. These values have been taken from the RCTs or registries for these treatments. Patients receiving etanercept in the model do not also receive methotrexate, according to etanercept's marketing authorisation. It was assumed that 20% of patients in the methotrexate comparator arm would be intolerant to methotrexate and therefore would not receive it.¹²³

Table 53 Concomitant biologic DMARD and methotrexate use

	Methotrexate only	Abatacept	Adalimumab	Etanercept	Tocilizumab
Methotrexate use, %	80%	80%	69%	0%	82%

Resource use

We conducted a literature search for costing studies in patients with JIA and identified two relevant studies. Thornton and colleagues¹³⁸ examined the resources used and associated patient-based costs during the first year after diagnosis for JIA patients in the UK. Prince and colleagues¹²³ analysed the costs of treatment for patients in the Dutch ABC register before and after receiving etanercept. There are limitations to both studies: the patients in the Thornton and colleagues¹³⁸ study are likely to have different resources and costs in the first year after diagnosis than the patients included in this assessment report; for example they may have had less severe disease. The resources used by patients in the Prince and colleagues study¹²³ are not reported and it is unclear how different Dutch health care costs would be to the NHS. Our clinical experts commented that the resources for monitoring patients costs were not substantially different between the patients treated with methotrexate only or with a biologic DMARD and were broadly similar to those in the Thornton and colleagues¹³⁸ study. We therefore used the resources described by Thornton and colleagues in the base case and explored the costs used by Prince and colleagues¹²³ in a scenario analysis (scenario ii). The assumed resources used by patients are shown in Table 54. Blood tests consisted of the combined cost of full blood count, C-reactive protein, urea and electrolytes, and a liver function test. Clinical imaging consisted of the combined cost of MRI scan, Dual-energy X-ray absorptiometry (DEXA) scan, ultrasound and X-ray. The total health care cost for patients on biological treatment and off biologic treatment using the resources shown in

Patients who experienced a disease flare received one or more injections of intra-articular steroids, and were treated as a paediatric rheumatology inpatient case at a cost of £429.97.¹⁵⁰

Adverse events (AE)

The database of studies from our systematic review was searched for studies reporting any AEs or discontinuation. In addition, the company submissions were consulted for any relevant data. Whilst the types and frequencies of AEs were reported, no cost data were identified in any of the studies that reported serious adverse events (SAE) or discontinuation rates, or in observational studies reported in the company submissions. In order to identify data, previous NICE technology appraisals were

searched. Neither of the JIA technology appraisals, TA35 and TA238, contained data on the cost of SAEs.^{43;44}

Table 54 was £724 per cycle.

Patients who experienced a disease flare received one or more injections of intra-articular steroids, and were treated as a paediatric rheumatology inpatient case at a cost of £429.97.¹⁵⁰

Adverse events (AE)

The database of studies from our systematic review was searched for studies reporting any AEs or discontinuation. In addition, the company submissions were consulted for any relevant data. Whilst the types and frequencies of AEs were reported, no cost data were identified in any of the studies that reported serious adverse events (SAE) or discontinuation rates, or in observational studies reported in the company submissions. In order to identify data, previous NICE technology appraisals were searched. Neither of the JIA technology appraisals, TA35 and TA238, contained data on the cost of SAEs.^{43;44}

Table 54 Resource use and unit costs

Resource per year	Resource use per year		Unit cost	Reference
	Off biologic treatment	On biologic treatment		
GP visit	10	10	£46.00	PSSRU 2014 ¹⁵¹
Hospital appointments				
Rheumatology paediatric consultant	5.58	5.58	£234.86	National reference costs 2013/14 ¹⁵⁰
Ophthalmologist	2.69	2.69	£114.73	National reference costs 2013/14 ¹⁵⁰
Specialist nurse	7.00	7.00	£40.00	PSSRU 2014 ¹⁵¹
Physiotherapist	4.00	4.00	£16.50	PSSRU 2014 ¹⁵¹
Occupational therapist	0.65	0.65	£16.50	PSSRU 2014 ¹⁵¹
Podiatry	0.61	0.61	£43.59	National reference costs 2013/14 ¹⁵⁰
Hospital tests				
Blood tests	1	1	£46.27	Thornton et al. 2008 ¹⁵² , updated to 2013/14 values using PSSRU HCHS Index
Clinical imaging	1	1	£386.42	
Disease flare				
Inpatient treatment per disease flare			£429.97	National reference costs 2013/14 ¹⁵⁰

PSSRU, Personal Social Services Research Unit.

Due to the paucity of data in JIA, technology appraisals of RA were also assessed. Of the six technology appraisal publications available on the NICE website,¹⁵³⁻¹⁵⁸ only one contained data for the cost of an adverse event, TA195: ‘Adalimumab, etanercept, infliximab, rituximab and abatacept for the treatment of rheumatoid arthritis after the failure of a TNF inhibitor’.¹⁵⁷ A Pfizer CS provided the only relevant cost data in TA195. Pfizer assumed an SAE involved two GP visits, seven days of hospitalisation, and a utility decrement of 0.05, with a total cost of £1181. No specific adverse events were identified by Pfizer. Further details on the types of SAE experienced by JIA patients are given in Section 4.1.2 and in Appendix 2 of this assessment report. The most common SAEs were serious infections and infestations, but SAEs also included autoimmune diseases and malignancies. All independent analyses in RA conducted by assessment groups to this point have been based on the Birmingham Rheumatoid Arthritis Model (BRAM),¹³⁹ which assigns a cost to increases in HAQ scores. In JIA, it is not possible to model SAE costs in this way due to lack of HAQ in the RCTs.

In order to model the cost of SAEs, healthcare resource group codes for intermediate and severe paediatric infections were consulted. Additionally, a study in etanercept patients by Otten and colleagues indicated that the median length of hospitalisation for SAEs was nine days (IQR 2-12).¹⁵⁹ Given this, we estimated inpatient costs by averaging all spells for intermediate and major paediatric infections (£1,532.87).

A summary of the input parameters used in the model are shown in Table 55.

Table 55 Summary of the input parameters used in the SHTAC economic model

Parameter	Mean	Higher value	Lower value	Source
Starting age, years	11	15	6	Assumption, based on RCTs
Time horizon, years	30	10	70	Assumption
Discount rate, costs	3.5%	6%	1.5%	NICE reference case ¹²⁹
Discount rate, benefits	3.5%	6%	1.5%	NICE reference case ¹²⁹
Utility values, per cycle				
No treatment	0.13	0.15	0.11	Prince <i>et al.</i> ¹²³
Treatment after 3 months	0.17	0.20	0.15	Prince <i>et al.</i> ¹²³
Treatment after 15 months	0.19	0.21	0.16	Prince <i>et al.</i> ¹²³
Treatment after 27 months	0.20	0.23	0.16	Prince <i>et al.</i> ¹²³
Disease flare disutility	0.03	0.04	0.02	Assumption
Disease flare, per cycle				
Placebo	0.25	0.34	0.16	Ruperto <i>et al.</i> ⁵⁷ , Lovell <i>et al.</i> 2008 ⁶¹ Brunner <i>et al.</i> ⁶⁸
Abatacept	0.09	0.16	0.05	Ruperto <i>et al.</i> ⁵⁷
Adalimumab	0.14	0.23	0.09	Lovell <i>et al.</i> 2008 ⁶¹

Etanercept	0.09	0.17	0.04	Lovell et al. 2000 ⁴²
Tocilizumab	0.14	0.20	0.09	Brunner et al. ⁶⁸
Adverse events 1st cycle				
Abatacept	0.53%	1.51%	0.00%	Ruperto et al. ⁵⁷
Adalimumab	1.75%	3.71%	0.00%	Lovell et al. 2008 ⁶¹
Etanercept	1.45%	4.19%	0.00%	Lovell et al. 2000 ⁴²
Tocilizumab	1.60%	3.36%	0.00%	Brunner et al. ⁶⁸
Loss of efficacy				
Abatacept	9.47%	13.59%	5.36%	Ruperto et al. ⁵⁷
Adalimumab	3.51%	6.25%	0.76%	Lovell et al. 2008 ⁶¹
Etanercept	2.90%	6.82%	0.00%	Lovell et al. 2000 ⁴²
tocilizumab	7.98%	11.90%	4.06%	Brunner et al. ⁶⁸
Further line treatment				
AEs, biologic DMARD	0.43%	0.82%	0.04%	Tynjala et al. ¹⁴⁵
Loss of efficacy biologic DMARD	2.00%	2.59%	1.41%	Tynjala et al. ¹⁴⁵
AEs, methotrexate	0.58%	0.82%	0.34%	STRIVE ⁷⁷
Loss of efficacy methotrexate	0.42%	0.79%	0.05%	Klein et al. ¹⁴⁶
Costs				
On biologic DMARD cost	£724	£940.92	£506.65	National Reference costs 2013/14 ¹⁵⁰
Off biologic DMARD cost	£724	£940.92	£506.65	PSSRU 2014 ¹⁵¹
SAE cost	£1,533	£1,993	£1,073	National Reference costs 2013/14 ¹⁵⁰
Disease flare cost	£430	£301	£559	National Reference costs 2013/14 ¹⁵⁰

5.7 Results of the independent economic analysis

This section reports the cost-effectiveness results for a person with JIA who received treatment with a biologic DMARD in combination with methotrexate (where permitted) compared to those who received methotrexate only. Results for costs and QALYs are presented for each treatment, with costs and benefits discounted at 3.5%. The results are presented for biologic DMARDs licensed for use as a first-line biologic treatment, i.e. adalimumab, etanercept and tocilizumab and then presented for abatacept as a second-line biologic treatment following previous treatment with an anti-TNF. The results shown in this section are for the drug list price and the results with the confidential PAS discount for abatacept and tocilizumab are presented in a commercial in confidence separate appendix to this report.

Licensed first line biologics: adalimumab, etanercept and tocilizumab

The undiscounted summary results of the analyses for adalimumab, etanercept and tocilizumab compared to methotrexate for the treatment effects are shown in Table 56 - Table 58. In the base case, total undiscounted QALYs vary between 14.98 for methotrexate to 17.99 for tocilizumab (Table 56).

Patients on methotrexate only have higher QALYs in the off biologic DMARD health state than the patients on biologics as they spend more time in this health state. The summary results of the undiscounted drug costs are shown in Table 57. The total undiscounted drug acquisition cost of the biologic DMARDs varied between £103,497 and £128,071 for treatment first with etanercept and tocilizumab respectively compared to a total undiscounted cost of £7029 for patients treated with methotrexate only. The total patient costs varied between £107,299 and £225,797 for methotrexate only and tocilizumab respectively. As noted earlier, patients taking etanercept do not receive methotrexate, which partially explains the lower costs for the etanercept regimen.

The base case discounted cost-effectiveness results are shown in Table 58. Each of the biologic DMARDs is more expensive than methotrexate only, with the incremental cost ranging from £77,513 to £82,995 for etanercept and tocilizumab respectively. The incremental cost-effectiveness versus methotrexate only for adalimumab, etanercept and tocilizumab is £38,127, £32,256 and £38,656 per QALY gained respectively. The results are not presented as an incremental analysis of the biologic DMARDs as the costs and QALYs for each biologic DMARD are generally similar and it has been previously discussed in (Section 4.1.2) that the biologic DMARDs may be regarded as similar in effectiveness.

Table 56 Summary of the total undiscounted QALYs in each health state for treatment with 1st line biologic compared to methotrexate

Treatment	Health state QALYs			
	On biologic DMARD	Off biologic DMARD	Disease flare	Total
Methotrexate only	N/A	15.9	-0.9	14.98
Adalimumab	8.6	9.9	-0.8	17.77
Etanercept	8.5	10.0	-0.7	17.81
Tocilizumab	9.2	9.5	-0.8	17.99

Table 57 Summary of the total undiscounted costs in each health state for treatment with 1st line biologic compared to methotrexate

	Medical	Drug	AEs	Flare	Total
Methotrexate only	£86,938	£7,029	£498	£12,834	£107,299
Adalimumab	£86,938	£114,701	£248	£10,805	£212,693
Etanercept	£86,938	£103,497	£254	£9,766	£200,454
Tocilizumab	£86,938	£128,071	£269	£10,519	£225,797

Table 58 Cost-effectiveness of 1st line biologic DMARDs versus methotrexate only

	Costs, £	QALYs	Incremental costs	Incremental QALYs	ICER (£ per QALY gained) versus methotrexate ^a
Methotrexate only	£67,426	9.35			
Adalimumab	£145,047	11.40	£77,513	2.0	£38,127
Etanercept	£134,868	11.44	£67,334	2.1	£32,526
Tocilizumab	£150,530	11.52	£82,995	2.1	£38,656

^a Results presented compared to methotrexate; no incremental analysis presented.
 Note: Abatacept was not included in this analysis as the marketing authorisation is not for first line biologic DMARD

Licensed second line biologic: abatacept

Abatacept is licensed for use after at least one previous anti-TNF biologic DMARD. The results are shown for abatacept, adalimumab, etanercept and tocilizumab compared to methotrexate. For each biologic comparator, patients are assumed to have been treated initially with etanercept as the first line biologic. The summary results of the non-discounted treatment effects are shown in Table 59. In the base case, total undiscounted QALYs vary between 14.98 for methotrexate and 20.07 for abatacept. The summary results of the undiscounted costs are shown in Table 60. The total undiscounted drug acquisition cost of the DMARDs varied between £7,029 for methotrexate only to £222,533 for abatacept respectively. The total patient costs varied between £107,299 and £317,097 for methotrexate only and abatacept respectively.

Table 59 Summary of the total undiscounted QALYs in each health state for treatment with 2nd line biologics compared to methotrexate

Treatment	QALYs			
	On biologic DMARD	Off biologic DMARD	Disease flare	Total
Methotrexate only	N/A	15.9	-0.9	14.98
Abatacept	15.8	4.8	-0.5	20.07
Adalimumab	15.1	5.3	-0.6	19.80
Etanercept	15.0	5.4	-0.5	19.82
Tocilizumab	15.7	4.8	-0.6	20.00

Table 60 Summary of the total undiscounted costs in each health state for treatment with 2nd line biologic DMARDs compared to methotrexate

Treatment	Costs, £
-----------	----------

	Medical	Drug	AEs	Flare	Total
Methotrexate only	£86,938	£7,029	£498	£12,834	£107,299
Abatacept	£86,938	£222,533	£502	£7,124	£317,097
Adalimumab	£86,938	£184,594	£433	£8,118	£280,082
Etanercept	£86,938	£179,686	£440	£7,311	£274,374
Tocilizumab	£86,938	£205,174	£457	£7,840	£300,409

The base case discounted cost-effectiveness results are shown in Table 61. The costs and QALYs are different to those for the first line biologic cost-effectiveness analysis (Table 58) because this analysis includes the costs and QALYs of two lines of biologics. The cost-effectiveness of the abatacept compared to methotrexate is £39,536 per QALY. The results are not presented as an incremental analysis as the costs and QALYs for the biologic DMARDs are similar.

Table 61 Cost-effectiveness of 2nd line biologic DMARDs compared with methotrexate using list price

	Costs, £	QALYs	Incremental costs	Incremental QALYs	ICER (£ per QALY gained) versus methotrexate^a
Methotrexate only	£67,534	9.37			
Abatacept	£203,276	12.80	£135,742	3.4	£39,536
Adalimumab	£183,387	12.65	£115,853	3.3	£35,284
Etanercept	£179,580	12.67	£112,045	3.3	£33,948
Tocilizumab	£194,263	12.76	£126,728	3.4	£37,363

^a Results presented compared to methotrexate; no incremental analysis presented.

5.7.1 Sensitivity analysis

Deterministic sensitivity analysis

Table 62 to Table 64 show the results of the deterministic sensitivity analyses for each of the biologic DMARDs versus methotrexate for the most influential parameters. Other parameters, such as time horizon, cost and frequency of disease flare, complete response rate and utility values were varied in the sensitivity analyses but were found to only have a negligible effect on the results. For each of the treatments, the models are most sensitive to the utility values chosen whilst on biologic DMARD treatment. They are also sensitive to the discount rate and the health state costs.

The deterministic sensitivity results for adalimumab versus methotrexate are shown in Table 62 and varied between £26,571 and £67,470 per QALY gained.

Table 62 Deterministic sensitivity analysis for adalimumab versus methotrexate only

Adalimumab vs methotrexate	High CI	Low CI	Range
Base case ICER: £38,127			
Utility treatment, long-term ¹	£26,571	£67,470	£40,898
Utility no treatment	£59,814	£27,982	£31,832
Discount rate benefits	£45,936	£32,123	£13,813
Discount rate costs	£31,919	£45,016	£13,097
On biologic DMARD cost	£41,630	£34,624	£7,006
Off biologic DMARD cost	£34,624	£41,630	£7,006
Disease flare methotrexate	£35,871	£40,598	£4,727
AE adalimumab	£37,983	£33,308	£4,675

¹ After treatment for more than 27 months with biological DMARD

The deterministic results for etanercept versus methotrexate only varied between £22,886 and £56,196 per QALY gained (Table 63).

Table 63 Deterministic sensitivity analysis for etanercept versus methotrexate only

Etanercept vs methotrexate	High CI	Low CI	Range
Base case ICER: £32,526			
Utility, treatment, long-term ¹	£22,886	£56,196	£33,310
Utility no treatment	£50,511	£23,986	£26,525
Discount rate costs	£26,909	£38,783	£11,874
Discount rate benefits	£39,075	£27,478	£11,598
Start age	£35,045	£26,173	£8,873
Off biologic DMARD cost	£29,118	£35,934	£6,817
On biologic DMARD cost	£35,934	£29,118	£6,817
Disease flare Methotrexate	£30,566	£34,668	£4,102

¹ After treatment for more than 27 months with biological DMARD

The deterministic sensitivity results for tocilizumab versus methotrexate only varied between £26,835 and £69,092 per QALY gained (Table 64).

Table 64 Deterministic sensitivity analysis for tocilizumab versus methotrexate only

Tocilizumab vs methotrexate	High CI	Low CI	Range
Base case ICER: £38,656			
Utility, treatment, long-term ¹	£26,835	£69,092	£42,257

Utility no treatment	£58,865	£28,777	£30,088
Discount rate costs	£31,360	£46,904	£15,545
Discount rate benefits	£47,140	£32,196	£14,943
Start age	£42,589	£32,993	£9,596
On biologic DMARD cost	£42,130	£35,182	£6,948
Off biologic DMARD cost	£35,182	£42,130	£6,948
Disease flare methotrexate	£36,395	£41,130	£4,735

¹ After treatment for more than 27 months with biological DMARD

The deterministic sensitivity analysis results for abatacept versus methotrexate only varied between £31,259 and £52,995 per QALY gained (Table 65).

Table 65 Deterministic sensitivity analysis for abatacept versus methotrexate only

Tocilizumab vs methotrexate	High CI	Low CI	Range
Base case ICER: £39,536			
Utility, treatment, long-term	£31,529	£52,995	£21,467
Discount rate costs	£30,512	£50,137	£19,625
Utility treatment 15-27 months	£32,110	£51,430	£19,319
Discount rate benefits	£49,908	£31,906	£18,002
Utility no treatment	£50,345	£32,549	£17,796
Start age	£42,187	£33,234	£8,952
On biologic DMARD cost	£43,094	£35,978	£7,117
Off biologic DMARD cost	£35,978	£43,094	£7,117

¹ After treatment for more than 27 months with biological DMARD

5.7.2 Scenario analysis

We conducted several scenario analyses to investigate uncertainty for specific aspects of the modelling. The results of these analyses are presented for the first line biologics.

i) Discontinuation of treatment due to clinical remission

Patients with clinical remission off medication are at high risk of relapse. Baszis and colleagues¹⁶⁰ conducted a retrospective chart review in a cohort of 171 patients with JIA (of a range of sub-types but predominantly polyarticular course) in the United States treated with TNF α antagonists. They found that 12 months after stopping treatment only 33% still had clinical remission. Similarly, a retrospective chart review of 437 JIA patients from centres in the United States and Italy by Wallace

and colleagues¹⁶¹ estimated that 6% of patients, that had discontinued methotrexate therapy with clinical remission, had persistent remission after five years off treatment.

The rate of discontinuation of biologic treatment varies between studies. In a retrospective observational study Tynjala and colleagues, patients receiving etanercept were followed up for four years and 10% of patients had discontinued treatment due to inactive disease. In the study by Baszis and colleagues¹⁶⁰ 80% of patients discontinued TNF α antagonist treatment due to inactive disease. We varied the discontinuation rate between that seen by Tynjala and colleagues¹⁴⁵ (used in the base case) and Baszis and colleagues¹⁶⁰.

We assumed a relapse rate from Baszis and colleagues¹⁶⁰ of 67% for that analysis and 40% relapse rate as seen in Wallace and colleagues¹⁶¹ for the ‘Tynjala and colleagues’ analysis. We assumed no patients on the methotrexate only arm would discontinue as fewer patients on methotrexate would be in remission.

The results for the scenario with patients discontinuing treatment for clinical remission is shown in

Table 66 for first line biologics compared to methotrexate only. In the scenario with the highest discontinuation rate, the cost-effectiveness of the biologics improves from the base case by about £4000 per QALY.

Table 66 Cost-effectiveness for first line biologics versus methotrexate only with patients discontinuation of treatment for clinical remission

	Remission off treatment (per cycle)	Relapse rate	ICER (£/QALY) versus methotrexate		
			Adalimumab	Etanercept	Tocilizumab
Base case	0%		£38,127	£32,526	£38,656
Baszis et al. ¹⁶⁰	7.8%	67%	£33,744	£28,580	£34,214
Tynjala et al. ¹⁴⁵	0.66%	40%	£37,512	£31,970	£38,028

ii) Health state costs from Prince and colleagues

The base case analysis uses health state costs estimated by a UK study by Thornton and colleagues¹³⁸ of patients during the first year after diagnosis. However, as stated earlier, this may not necessarily reflect the patient group in this economic evaluation as patients in that study were newly diagnosed.

The Roche CS cost-effectiveness analysis uses health state costs based on the Prince and colleagues study.¹²³ Assuming hospital admissions would be for disease flare only, the health state costs per cycle are £589.51 and £408.91 for the off treatment and on treatment health states, respectively (compared to £724 in the base case). In this analysis the biologic DMARDs are slightly more cost-effective and the ICER decreases by about £2900 per QALY compared to the base case analysis, e.g. the ICER for adalimumab decreases to £35,214 per QALY (Table 67).

Table 67 Summary of the cost-effectiveness for adalimumab, etanercept and tocilizumab versus methotrexate only using health state costs from Prince and colleagues

	Costs, £	QALYs	Incremental costs	Incremental QALYs	ICER (£/QALY)
Methotrexate only	£57,306	9.37			
Adalimumab	£128,894	11.40	£71,589	2.0	£35,214
Etanercept	£118,771	11.44	£61,465	2.1	£29,691
Tocilizumab	£134,097	11.52	£76,792	2.1	£35,767

iii) Discount rates used in NICE appraisal of etanercept

The previous NICE appraisal of etanercept (NICE TA35⁴³) used a discount rate of 6% for costs and 1% for benefits¹²² (which were the recommended rates at the time). We ran the analysis for etanercept using that discount rate. Table 68 shows ICERs that are much reduced compared to the base case in the current assessment report; £21,718 per QALY. Using this discount rate etanercept would be cost-effective at a willingness-to-pay threshold of £20,000 - £30,000 per QALY.

Table 68 Cost-effectiveness for etanercept versus methotrexate using discount rate of 6% for costs and 1% for benefits

	Costs, £	QALYs	Incremental costs	Incremental QALYs	ICER (£/QALY)
Methotrexate only	£51,494	12.96			
Etanercept	£107,200	15.53	£55,707	2.6	£21,718

iv) Caregiver benefit

We were unable to find HRQoL utility values associated with caring for a child or young person with JIA and assumed in the base case analysis no disutility benefit for parents and caregivers. A study by Kuhlthau and colleagues.¹⁶² compared the well-being of parents of children with and without activity limitations. This list of conditions includes medical conditions that would commonly be considered disabling (e.g. paraplegia and blindness) as well as typically less disabling but chronic conditions (e.g.

The cost-effectiveness of the two scenarios varied between £36,982 and £38,152 per QALY (Table 70). The cost-effectiveness of three line biologic therapy is similar to that seen in the base case analysis for one line of biologic therapy (Table 58).

Table 70 Cost-effectiveness for three lines of biologic therapy

	Costs, £	QALYs	Incremental costs	Incremental QALYs	ICER (£/QALY)
Methotrexate only	£67,534	9.37			
Etanercept, adalimumab, tocilizumab	£207,565	13.16	£140,031	3.8	£36,982
Etanercept, adalimumab, abatacept	£212,562	13.17	£145,028	3.8	£38,152

vi) Younger biologic DMARD starting age

Adalimumab, etanercept, tocilizumab have a licensed indication from aged two years for patients with polyarticular arthritis and abatacept has a licensed indication from aged six years and over. In this scenario we investigated the cost-effectiveness of the biologics with a starting age of six years old (in the base case the starting age is 11 years). The results of the analysis for first line biologics are shown in Table 71. These indicate that there is minimal difference in the cost-effectiveness for adalimumab but a decrease in about £6000 in the cost-effectiveness of etanercept and tocilizumab.

Table 71 Cost-effectiveness of first line biologics with a starting age of 6 years

	Costs, £	QALYs	Incremental costs	Incremental QALYs	ICER (£/QALY)
Methotrexate only	£67,492	9.39			
Adalimumab	£145,089	11.42	£77,597	2.0	£38,124
Etanercept	£121,737	11.46	£54,245	2.1	£26,173
Tocilizumab	£138,421	11.54	£70,929	2.1	£32,993

The results of the analysis for second line biologics are shown in Table 72. These indicate a similar improvement in the cost-effectiveness and there is a reduction in the cost-effectiveness of abatacept of £6,302 per QALY.

Table 72 Cost-effectiveness of 2nd line biologics with a starting age of 6 years

	Costs, £	QALYs	Incremental costs	Incremental QALYs	ICER (£/QALY)
Methotrexate only	£67,492	9.39			
Abatacept	£181,776	12.83	£114,285	3.4	£33,234

Adalimumab	£170,364	12.68	£102,872	3.3	£31,283
Etanercept	£163,006	12.69	£95,514	3.3	£28,895
Tocilizumab	£176,066	12.78	£108,575	3.4	£31,961

Probabilistic sensitivity analyses (PSA)

In the PSA, all parameters were sampled probabilistically from an appropriate distribution using similar ranges as used in the deterministic sensitivity analyses. The parameters sampled were: treatment effectiveness, discontinuation rate, health state costs, disease flare parameters, and HRQoL. The distribution assigned to each variable included in the PSA and the parameters of the distributions are reported in Appendix 11.

First line biologics

One thousand simulations were run. The PSA results are presented in Table 73 for first line biologics and show similar results to the deterministic analyses (Table 58). The cost-effectiveness for biologics versus methotrexate only varied between £32,554 and £38,744 per QALY for tocilizumab.

Table 73 Summary of the probabilistic sensitivity results for first line biologics versus methotrexate only

	Costs, £	QALYs	Incremental Costs, £	Incremental QALYs	ICER (£/QALY)
Methotrexate only	£67,531	9.38			
Adalimumab	£145,933	11.43	£78,402	2.05	£38,181
Etanercept	£135,803	11.48	£68,272	2.10	£32,554
Tocilizumab	£151,800	11.55	£84,269	2.18	£38,744

The CEAC is shown in Figure 9 and indicates that at the £20,000 and £30,000 willingness-to-pay thresholds methotrexate has the highest probability of being cost-effective, of 0.98 and 0.62 respectively.

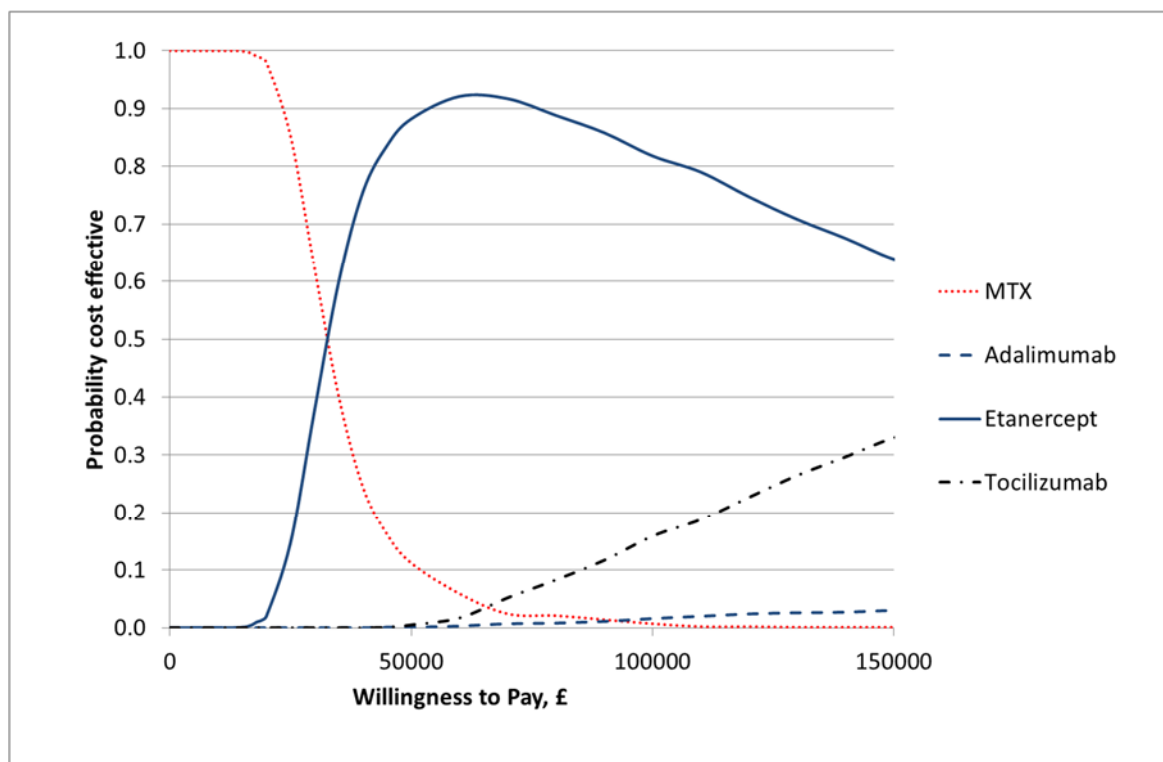


Figure 9 CEAC from the PSA for 1st line biological treatments compared to methotrexate

Second line biologics

The PSA results are presented in Table 73 for second line biologics and show similar results to the deterministic analyses (Table 61). The cost-effectiveness of abatacept in the PSA is £39,608 per QALY.

Table 74 Summary of the probabilistic sensitivity results for 2nd line biologics versus methotrexate only

	Costs, £	QALYs	Incremental Costs, £	Incremental QALYs	ICER (£/QALY)
Methotrexate only	£67,168	9.35			
Abatacept	£203,396	12.81	£136,041	3.43	£39,608
Adalimumab	£183,563	12.66	£116,208	3.29	£35,366
Etanercept	£179,807	12.67	£112,452	3.30	£34,053
Tocilizumab	£194,464	12.77	£127,109	3.39	£37,443

The CEAC is shown in Figure 10 and indicates that at the £20,000 and £30,000 willingness-to-pay thresholds methotrexate has the highest probability of being cost-effective of 0.99 and 0.71 respectively.

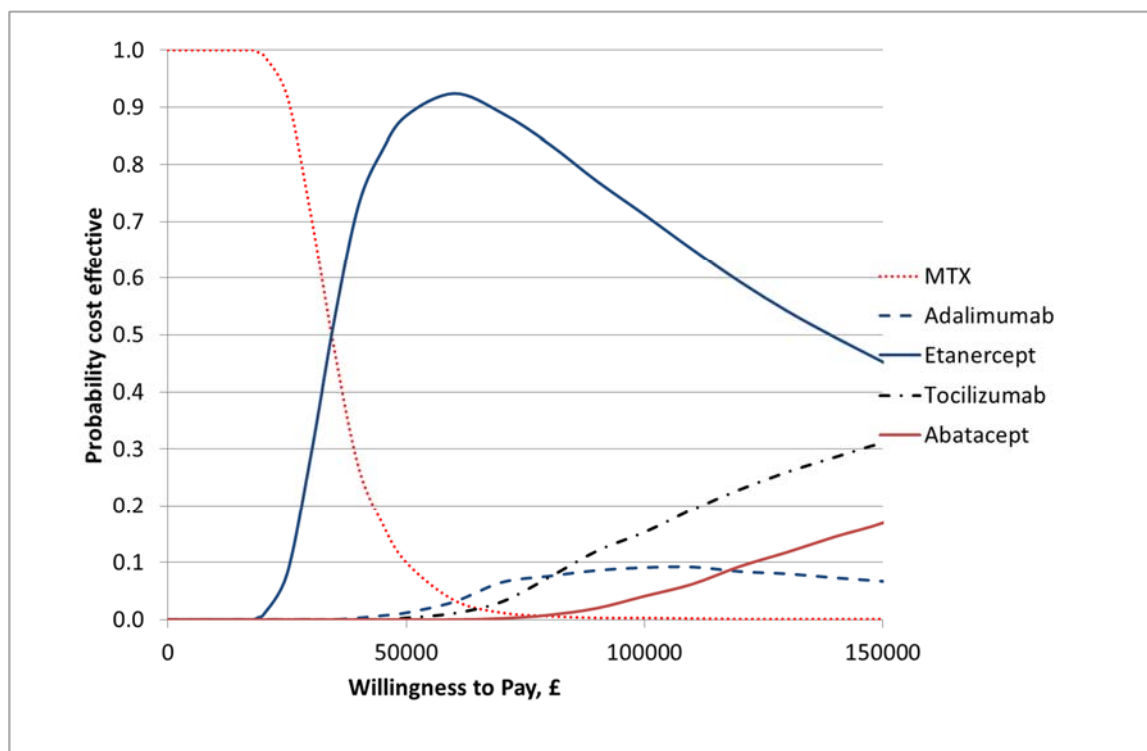


Figure 10 CEAC from the PSA for 2nd line biological treatments compared to methotrexate

5.7.3 Sub-groups

There are a number of potential sub-groups that were within the NICE scope including the sub-types of JIA (extended oligoarthritis, polyarticular arthritis, ERA, and PA) and patients with extra-articular manifestations such as uveitis. As stated earlier, sub-group analyses by sub-type of JIA was not possible due to insufficient evidence for input parameters to support modelling. The modelled patient population is therefore people with JIA, with the results of particular relevance to those with polyarticular course JIA (extended oligoarthritis, and RF + ve and RF – ve polyarthritis).

In considering the potential for modelling the clinical-effectiveness and cost-effectiveness of biologic DMARDs in patients with JIA-associated uveitis, a draft NHS clinical commissioning policy on the use of anti-TNF alpha agents in paediatric patients with severe refractory uveitis was consulted.²⁹ The policy discusses cost-effectiveness of treatment and the elements of an economic evaluation are given, though full reporting of the results from such an economic evaluation have not been reported.

The report states that infliximab and adalimumab in combination with methotrexate are widely used worldwide for the treatment of refractory uveitis, and that etanercept is not recommended for use in this patient group. The report also cites evidence from a systematic review by Simonini and colleagues⁹⁶ (as described in section 4.4.2 of this assessment report), which shows, based on a pooled

analysis of observational studies, the proportion of children with improved intraocular inflammation (responders) was 87% for adalimumab, 72% for infliximab, and 33% for etanercept. Potential modelling of the clinical-effectiveness and cost-effectiveness in JIA-associated uveitis in this report would therefore apply only to adalimumab as this is the only one of the four biologic DMARDs within the scope of the appraisal recommended for treating this patient sub-group.

With regards to quality of life, the clinical commissioning policy assumes that loss of vision causes detrimental effects on utility based on the results of a study of age-related macular degeneration by Reeves and colleagues.¹⁶⁴ This study measured HRQoL changes associated with loss of vision using data from the SF-6D and best corrected visual acuity. This population is quite different from JIA-associated uveitis, and the data do not capture aspects of JIA related to arthritic joints.

In our model, we have used utility data derived using the HUI3 generic preference instrument, from the study by Prince and colleagues.¹²³ This instrument is appropriate for conditions that involve vision impairment, as it includes a domain for vision. HUI3 is not compatible with the SF-6D, and the instruments will produce different quality of life estimates. Attempting to combine data from Reeves and colleagues (2009) and Prince and colleagues¹²³ would be inappropriate due to the differences in the populations of the studies, the incompatibility of SF-6D and HUI3. Moreover, if it is assumed that adding vision loss to the other quality of life decrements due to advancing JIA even partially decreases patient quality of life, then it follows that adalimumab will be more cost-effective in JIA patients with uveitis and joint inflammation, than it is in JIA patients without uveitis.

Likewise, if most of the costs related to uveitis relate to the management of vision loss, as stated in the clinical commissioning policy,²⁹ then any reduction of these costs due to improving vision would increase cost-effectiveness in the sub-group of JIA patients with uveitis. Additional analysis of cost-effectiveness in a JIA uveitis population that is refractory to methotrexate, as is indicated in the licensing for adalimumab, is therefore likely to have predictable results.

As discussed in section 4.3 of this report, the SYCAMORE trial of adalimumab and methotrexate in JIA-associated uveitis patients, has recently closed early following interim analysis showing a favourable effect for treatment.⁸⁷ The trial also includes a cost-effectiveness analysis, the results of which would likely concur with the logical implications discussed above.

5.8 Comparison of the economic models

The cost-effectiveness of biologic DMARDs estimated in this report varies between £30,000 and £40,000 per QALY gained compared to methotrexate only. This is higher than estimated by the

previous NICE appraisal for etanercept in patients with JIA, which estimated an ICER of £16,082 per QALY gained.¹²² The NICE Appraisal Committee accepted that the ICER for etanercept was likely to be in the region of £15–30,000 per QALY.⁴³ The model used in that NICE appraisal is not fully described and so it difficult to compare with the current model developed for this report. However, the discount rate for that appraisal was 6% for costs and 1% for benefits. Using these discount rates in the independent model in this assessment report gives cost-effectiveness estimates of between £20,000 - £30,000 per QALY gained.

A cost consequence analysis conducted by Prince and colleagues in the Netherlands did not estimate the cost-effectiveness of etanercept.¹²³ We have estimated the cost-effectiveness of etanercept compared to methotrexate from that study, by aggregating the costs and QALYs in each time period reported, to be £32,590 (43,300 Euros).

Comparing the results from the independent model in this assessment report with those submitted by the companies was complicated by differences in structure between the models. Roche, who manufacture tocilizumab, was the only company that submitted a full economic analysis to NICE, including costs and QALYs and with a 25 year time horizon. BMS, who manufacturer abatacept, submitted a model with a 20 year time horizon with only drug and administration costs. In addition, in one company model patients receive oral methotrexate (Roche model), and the other company model patients receive subcutaneous methotrexate (BMS Model). It was therefore only possible to compare drug costs between the three models with a 20 year time horizon and with discounting applied, to allow a level comparison between the models independent of structural assumptions. Table 75 shows the comparison with the Roche model with patients using oral methotrexate, whilst Table 76 shows the comparison with the BMS model with patients using subcutaneous methotrexate.

It should be noted that the Roche analysis has not compared the biologic DMARDs against methotrexate in their submission but has compared adalimumab with tocilizumab; however this analysis was present in their economic model.

Table 75 Comparison of the drug costs in the assessment report model with the Roche CS model (20 year discounted, no PAS)

	Assessment report model (using oral methotrexate)		Roche Model	
	Drug Costs, £	Total Costs, £	Drug Costs, £	Total Costs, £
Methotrexate	£393	£49,178	████	████
Adalimumab	£71,992	£119,269	████	████

Etanercept	£65,396	£111,941	██████	██████
Tocilizumab	£74,578	£121,725	██████	██████

Table 76 Comparison of the drug costs in the assessment report model with the BMS CS model (20 year discounted, no PAS)

	Assessment report model (using oral methotrexate)		BMS Model	
	Drug Costs, £	Total Costs, £	Drug Costs, £	Total Costs, £
Methotrexate	£8,012	£56,798	██████	██████
Adalimumab	£81,804	£129,081	██████	██████
Etanercept	£70,368	£116,914	██████	██████
Tocilizumab	£85,312	£132,459	██████	██████

As can be seen there was variation in costs between the models. The Roche model has lower drug costs and total costs than the assessment report model. This is due to their model using a higher discontinuation rate so patients remain on the biologic for a shorter duration, and with lower health state costs. The BMS model does not include discontinuation for any cause, which explains why it has the highest drug costs of the above models. Overall, the differences between the model results may be explained by differences in model structures and choices with regard to discontinuation, adverse events, and other costs.

5.9 Discussion

- A systematic search of the literature found four relevant economic evaluations of biologic DMARDs for patients with JIA. Two of the studies were presented as cost-utility studies, one was a cost-effectiveness study and the other was cost-consequence study. The evaluations were published between 2002 and 2012 in the UK, the Netherlands, Russia and Canada. One of the studies was the previous NICE appraisal of etanercept.¹²² The studies varied in design and structure, time horizons and the comparators included. The limitations in the methodological quality in all the studies identified include limited reporting of model parameters and assumptions.
- A systematic search of the literature found two HRQoL studies in children and adolescents with JIA. One study assessed the effectiveness of a foot care programme in a RCT setting, while the other evaluated the quality of life in a cohort of patients from the Dutch ABC registry before and after treatment with etanercept.

- Four pharmaceutical companies submitted evidence to NICE for consideration in this appraisal. Only one company (Roche) constructed a cost-utility analysis that included both costs and outcomes. Two companies (BMS and Pfizer) submitted cost-analyses and assume that the biologic DMARDs were equivalent in effectiveness, whilst AbbVie did not submit an economic analysis due to limitations identified with any potential analysis. Roche submitted a Markov state-transition model with health states for uncontrolled / off treatment, on treatment and dead. The model compared treatment with adalimumab to tocilizumab. The base case results from the submission conclude that tocilizumab is of similar effectiveness and is less expensive than adalimumab.
- We developed an independent cost-utility model comparing the biologic DMARDs to methotrexate alone. From this model, the incremental cost-effectiveness versus methotrexate only for adalimumab, etanercept and tocilizumab is estimated at £38,127, £32,526 and £38,656 per QALY gained respectively. An analysis comparing second line biologics with methotrexate only, estimated a cost-effectiveness ratio of £39,536 per QALY gained. The model results are most sensitive to changes to the HRQoL utility values.

6 ASSESSMENT OF FACTORS RELEVANT TO THE NHS AND OTHER PARTIES

Etanercept was recommended by NICE in 2002,¹³⁰ and it is known that adalimumab, abatacept, and tocilizumab are commonly used in practice (as well as infliximab – though not in the scope of this NICE appraisal).¹⁴³ It is unlikely that any positive NICE recommendations for the use of these biologic DMARDs will significantly increase the number of patients requesting treatment, and thus affecting budget impact.

Given that biologic DMARDs are currently used in the management of patients with JIA in the NHS it is unlikely that substantial modifications will be needed to services, such as infrastructure development or increased staff training. However, a survey of services for children, young people and families living with JIA in the UK by the National Rheumatoid Arthritis Society (NRAS) found that, among the 13 specialist (tertiary) centres surveyed, there was a shortfall of staff to adequately cover the services required.¹⁶⁵ These included paediatric rheumatology consultants and clinical nurse specialists, clinical psychologists, occupational therapists and physiotherapists. Further recruitment and training of professionals to make up the multi-disciplinary teams needed to provide effective treatment and care of JIA patients would appear necessary.

A long-term condition such as JIA can have a significant impact on children and young people's education. They may need to miss lessons to attend health care appointments, and may be absent for longer periods of time whilst experiencing symptoms (including disease flares) or if joint or other surgery is required. This can have a negative impact on educational attainment and, in turn, on their ability to gain employment in adulthood. It may also affect their social and psychological health, through reduced ability to participate in social and leisure activities and sport, and the general burden of a serious health condition during the sensitive period of adolescence. The effect of this may, therefore, widen socio-economic and health inequalities in this group. Only one of the RCTs included in the systematic review of clinical-effectiveness reported the impact of treatment (abatacept)⁵⁸ on missed school days. This outcome was not formally included in our review, but it was found that treated patients experienced a statistically significantly higher increase in school days (1.9 days) than placebo treated patients (0.9 days). This indicates the potential for biologic DMARDs to improve education as well as health outcomes, though further evidence is required, particularly in a UK context.

Schools and health services are required to liaise to ensure appropriate care for children and young people with JIA. The NRAS survey of 13 specialist centres found that all centres liaise with schools by letter or telephone, but less than half were unable to visit schools or only provided a limited service.¹⁶⁵ However, there were some examples of greater involvement, such as in one centre the clinical nurse specialist will visit schools and give talks if required. Effective liaison between health services and schools is important to ensure that the needs of children with JIA receiving biologic DMARDs are adequately met.

The impact of JIA on parents and caregivers can also be significant. For example, they may have to pay for child care, take time away from work, or even cease employment altogether to provide their own care. This will negatively affect their income and may increase dependency on welfare benefits (where available). Again, this is likely to increase socio-economic inequalities. The inability of parents and caregivers to work may have a negative impact on society and the economy, through reduced productivity, less income tax collection, and in some professions a shortage of skilled workforce capacity. The impact of treating JIA on parents and caregivers was generally not assessed by the RCTs in the systematic review of clinical-effectiveness. However, one of the RCTs (abatacept)⁵⁸ reported improvements in the number of days of normal activity per month missed by parents, including work and non-work activities, compared to placebo. The number of days when paid care was required remained stable in both trial arms (following an initial decline in the open-label lead-in phase with abatacept treatment). Further evidence on the impact of biologic DMARD treatment on parents and carers would be useful to gauge the full potential benefits of treatment beyond the patients themselves.

7 DISCUSSION

7.1 Statement of principal findings

7.1.1 Clinical-effectiveness

The systematic review of clinical-effectiveness conducted for this report found that biologic DMARDs are superior to placebo (with methotrexate where permitted) across a number of outcome measures in children with JIA (predominantly polyarticular course) and who had an insufficient response to previous DMARD treatment. With the exception of the etanercept trial the majority of patients in the trials received methotrexate in addition to the biologic DMARD/placebo. Biologic DMARD-treated patients had fewer arthritis flares, longer time to disease flare (applicable to abatacept and etanercept), were more likely to achieve a treatment response as defined by the ACR Pedi criteria, and to have inactive disease (only measured in the abatacept and tocilizumab trials). The latter outcome can be considered to be most clinically significant as absence of disease activity (e.g. no joints with active arthritis, physician's global assessment) is a key treatment goal. Treatment was associated with reduced pain scores, though this was only reported as statistically significant in one study (tocilizumab). HRQoL as measured by the CHAQ appeared to be higher for treated patients, though this was not always statistically significant.

The percentage of patients achieving ACR Pedi 30 in the open-label lead-in phases of the RCTs ranged from 65% to 94% across the trials. It should be acknowledged that due to the withdrawal design of the RCTs, in which only patients achieving an ACR Pedi 30 response during the open-label lead-in phase are eligible for randomisation, the results of the double-blind randomised phase of the trials are therefore only applicable to patients who have achieved an initial degree of treatment benefit. The effects seen during the double-blind period in the placebo group may not necessarily be the same for a placebo group who had not received a biologic DMARD prior to randomisation. However, expert clinical opinion suggests that ACR Pedi 30 can be considered an inadequate or partial response threshold, and higher rates, such as ACR Pedi 70 or above are considered more clinically significant. In this respect the patients responding to ACR Pedi 30 in the open-label lead-in phase (and eligible to be randomised) may not necessarily be considered atypical of patients eligible for treatment in clinical practice, as both would have active disease.

The clinical significance of the ACR Pedi 30 results of the randomised phases of the trials may also be questioned. ACR Pedi 30 response rates varied from 63% to 80% across the trials and declined with increasing response thresholds. Nonetheless, at ACR Pedi 70 (the highest threshold for which data were available across all four RCTs) the response rate varied from 44% to 65% and remained

higher in biologic DMARD-treated patients than placebo patients in all trials. Research is underway to further develop the JADAS tool as a clinically useful measurement tool³⁵⁻³⁷, though clinical trials are continuing to use the ACR Pedi criteria, albeit with effectiveness judged at thresholds higher than ACR Pedi 30.

In the longer-term, treatment effectiveness, in terms of ACR Pedi response, appears to be sustained as reported in the observational OLE studies for all four included RCTs. The longest follow-up available is for etanercept where ACR Pedi responses were maintained up to eight years of treatment.

The occurrence of AEs was generally similar between biologic DMARD and placebo-treated patients, based on non-statistically significant differences were reported. A range of AEs were reported, including viral and upper respiratory tract infections, injection-site reactions and nasopharyngitis. Serious AEs were uncommon. Discontinuations due to AEs were also uncommon (<3% patients). In the lead-in phase of the RCTs discontinuations due to AEs were low, ranging from 0.5% to 1.8%. The incidence of AEs and serious AEs during open-label long-term follow-up did not appear to be excessive. The safety profile of the biologic DMARDs therefore appears to be relatively favourable.

Sub-group analyses were reported in only one of the included RCTs (tocilizumab).⁶⁸ Patients receiving methotrexate background therapy had higher ACR Pedi response rates than those without, as did patients receiving background glucocorticoids. Patients who had received previous treatment with a biologic agent had lower ACR Pedi responses than those who were naïve to biologic DMARDs. It is not clear whether these sub-group analyses were pre-planned or post-hoc, so caution is advised in their interpretation.

Two recent published systematic reviews of the effectiveness of biologic DMARDs were identified during the production of this report.^{166;167} Both of these included a range of biologic DMARDs including the four relevant to the scope of this assessment. However, none of these reviews identified any additional RCT evidence to this assessment report. The only other relevant published systematic review of biologic DMARDs that we are aware of is by Otten and colleagues⁷⁹, (most recent search date January 2012). As discussed earlier in this report (section 4.1.3) Otten and colleagues⁷⁹ conducted an adjusted indirect comparison of adalimumab, abatacept and etanercept, using the same RCTs as included in this assessment report. We replicated the indirect comparison, extending it to include the tocilizumab RCT⁶⁸ which was not published during the timescale of the Otten and colleagues⁷⁹ review. Our results and conclusions match those of Otten and colleagues⁷⁹, that the biologic DMARDs appear similar in effectiveness in polyarticular course JIA, in terms of ACR Pedi response and preventing disease flares. Otten and colleagues⁷⁹ also share some of the caveats made in this assessment report about the limitations of the data included in the indirect comparison. Namely,

the small number of trials (and patient numbers), and differences between the trials in key patient characteristics and in treatment duration.

The conclusion that biologic DMARDs may be similar in clinical-effectiveness was supported by the expert advisers to this assessment report. In their experience there is similarity in effects between the drugs at a population level. However, it is noted that inter-patient variation in effects may occur, and comparative effectiveness of the biologic DMARDs may potentially vary between JIA sub-types. Currently there is a lack of clinical trial data to confirm this. Experts suggested that future trials of biologic DMARDs should stratify by disease phenotype to assess the differential effects of each treatment.

As noted earlier in this report, the RCTs of the biologic DMARDs included a mixture of JIA sub-types, broadly under the classification of polyarticular course JIA (including extended oligoarthritis). The trials did not appear to include patients with ERA or PA, thus we reviewed available evidence from trials in progress (see section 4.3) and from non-randomised studies (section 4.4) to gauge the effectiveness of biologic DMARD treatment in these groups. Much of the evidence is for etanercept (licensed for ERA and PA) with some available for adalimumab (licensed for PA). A broad comparison of the results of these studies with those of the RCTs included in this assessment report suggests that effectiveness is generally similar between these JIA sub-types. For example, ACR Pedi 70 response rates for biologic DMARDs were in the range of 44% to 65% across the RCTs (Table 15), compared to around 45% to 71% across the JIA sub-types in the CLIPPER study of etanercept⁹⁹ (Table 36) (notwithstanding differences in study variables such as length of follow-up). Evidence from trials in progress will provide greater clarity regarding efficacy and safety of biologic DMARDs in these JIA sub-types. At present there do not appear to be any studies of the comparative effectiveness of biologic DMARDs in these sub-types (e.g. adalimumab versus etanercept).

All of the RCTs were multi-national, with only one specifying including patients from the UK. The distribution of JIA sub-types within the trials, as far as reported (Table 12), appear reasonably similar to those seen in UK registry studies (Table 3), though this comparison may be limited by different reporting classifications used between studies. In addition, clinical practice in the RCTs (the oldest one was published in 2000⁴²) may not necessarily reflect current NHS care. The generalisability of the RCTs to the NHS is considered uncertain.

7.1.2 Cost-effectiveness

A systematic search of the literature found four economic evaluations of biologic DMARDs for patients with JIA. Two of the studies were presented as cost-utility-studies, one was a cost-effectiveness study and the other was cost-consequence study. The evaluations were published between 2002 and 2012 in the UK, the Netherlands, Russia and Canada. One of the studies was the assessment report which informed the previous NICE appraisal of etanercept (NICE TA35).¹²² The studies varied in design and structure, time horizons and the comparators included. There were limitations in the methodological quality in all the studies identified, and limited reporting of model parameters and assumptions.

A systematic search of the literature found two HRQoL studies in children and adolescents with JIA. One study assessed the effectiveness of a foot care programme in a RCT setting, while the other evaluated the quality of life in a cohort of patients from the Dutch ABC registry before and after treatment with etanercept.

Four drug companies submitted evidence to be considered as part of the NICE appraisal. Only one company (Roche) constructed a cost-utility analysis that included both costs and outcomes. Two companies (BMS and Pfizer) submitted cost analyses and assume that the biologic DMARDs were equivalent in effectiveness, whilst AbbVie did not submit an economic analysis due to suggested methodological limitations with any potential analysis. Roche submitted a Markov state-transition model with health states for uncontrolled / off treatment, on treatment and dead. The model compared treatment with adalimumab to tocilizumab. The base case results from the submission conclude that tocilizumab is of similar effectiveness and is less expensive than adalimumab.

We developed an independent cost-utility model comparing the biologic DMARDs to methotrexate only. From the model, the incremental cost-effectiveness versus methotrexate only for adalimumab, etanercept and tocilizumab is estimated at £38,127, £32,526 and £38,656 per QALY gained respectively. The incremental cost-effectiveness for abatacept as a second line biologic was £39,536 per QALY gained. The model results are most sensitive to changes to the HRQoL utility values.

The cost-effectiveness of biological DMARDs estimated in this report is associated with some uncertainty due to the limitations of the evidence base. For this reason, assumptions have had to be made to simplify the modelling. There was limited evidence on HRQoL, in particular with regard to disease progression. The HRQoL utility values were taken from a small Dutch registry study of patients receiving etanercept. The HRQoL values for patients treated with methotrexate were assumed to be constant over time. Patients with JIA who do not receive a biologic may experience disease progression and so their HRQoL may decline over time. In the model, we have assumed a constant

HRQoL utility value for patients in methotrexate only and so it may be that the biologic DMARDs would be more cost-effective than estimated by the economic model.

The model has not considered the underlying disease progression in terms of joint damage for patients with JIA. These patients may have sustained permanent damage to one or more joints affecting physical function and HRQoL into adulthood, potentially requiring joint surgery. The model has not considered the cost of this surgery and this assumption implicitly implies that biologic DMARDs have no impact on the long-term disease progression in terms of joint damage. It is unclear if biologic DMARDs reduce the long-term disease progression in JIA, however the AbbVie CS suggests that the reduction in orthopaedic surgery in JIA patients has been due to the increase in use of immunomodulatory agents among children in recent decades and so DMARDs and biological agents may have successfully prevented end-stage joint damage, based upon historical data that has shown a reduction. In the case where biologic DMARDs reduce long-term damage compared to treatment with methotrexate, the biologic DMARDs would be more cost-effective than estimated by the independent economic model.

The cost-effectiveness of biologic DMARD treatment of patients with JIA-associated uveitis has not been formally estimated in this economic evaluation, due to lack of suitable input parameter data. The current evidence base comprises mainly small retrospective observational studies, and suggests that adalimumab and infliximab are clinically effective in terms of improving intraocular inflammation and vision impairment.⁹⁶ A US cohort of children with JIA-associated uveitis and without JIA uveitis reported that vision-related HRQoL was worse in uveitis patients, but general HRQoL was similar to JIA patients without uveitis.¹⁶⁸ It can be assumed that biologic DMARD treatment in JIA-associated uveitis patients will result in bigger overall HRQoL improvement (including vision-related HRQoL) and therefore would be more cost-effective in this group than in JIA patients without uveitis.

It was also reported that significant predictors of uveitis were persistent oligoarthritis, and younger age at JIA diagnosis.¹⁶⁸ As discussed earlier in this report (section 1.1), persistent oligoarthritis accounts for up to 48% of JIA cases in the UK and is regarded as a milder form of JIA. In contrast, extended oligoarthritis accounts for between 6% and 17% of JIA cases in the UK and results in more severe symptoms and disease progression. Only extended oligoarthritis was explicitly included in the NICE scope for this appraisal, and therefore it can be considered that uveitis is less likely to affect the patient sub-types that are relevant to the appraisal.

The economic model does not include the wider societal costs associated with JIA and these are described in more detail in section 6. In the base case analysis we have not included caregiver benefits

associated with biologic DMARD treatment. A scenario analysis showed an improvement in cost-effectiveness for the biologic DMARDs when incorporating a utility disutility for patient caregivers.

The base case analysis includes only one line of biologic DMARD treatment, however in clinical practice some patients may switch to second or third line DMARDs. A scenario analysis was included that included a sequence of biologic treatments that most resembles current clinical practice. The cost-effectiveness of multiple lines of biologic therapy is similar to that seen in the base case analysis for one line of biologic therapy. There are many other possible treatment sequences but these have not been modelled as they were considered to be less likely to occur in clinical practice and the results for these sequences are similar to those presented. In clinical practice, infliximab is often used but this has not been included as a treatment in the economic model as it is licensed for this indication.

The cost-effectiveness results in this report are consistent with those from an earlier NICE technology appraisal for etanercept for patients with JIA (NICE TA35⁴³). The previous appraisal used a discount rate for 6% for costs and 1% for benefits.¹²² We ran the analysis for etanercept using these discount rates and the cost-effectiveness of etanercept improved to £21,718 per QALY. Using these discount rates etanercept would be cost-effective at a willingness-to-pay threshold of £30,000 per QALY.

7.2 Strengths and limitations of the assessment

The systematic reviews and economic evaluation in this report have been carried out independent of any competing interest, and the results are presented in a consistent and transparent manner.

The systematic reviews of clinical-effectiveness, cost-effectiveness and health-related quality of life have been undertaken following established methodology and principles for conducting a systematic review.⁵² The methods used were reported in a research protocol, which defined the decision problem in line with the NICE scope, and set out the inclusion and quality assessment criteria, data extraction process and the other methods to be employed during the evidence synthesis.

A multi-disciplinary advisory group has informed the review from its initiation. The research protocol was informed by comments received from the advisory group. The group also commented on a draft of the final report.

A *de novo* economic model has been developed following recognised guidelines. The model structure and data inputs are clearly presented in this report. The economic model is based upon data identified from systematic searches for clinical-effectiveness, cost-effectiveness and quality of life evidence, and other best available data.

This report is subject to certain limitations. The lack of head-to-head trials meant performing an indirect comparison of the biologic DMARDs, which is subject to a number of caveats due to heterogeneity between the trials (e.g. patient characteristics; treatment duration).

Limited HRQoL data were available for children with JIA, with none of the RCTs of biologic DMARDs reporting health utility data. The model results were based upon one Dutch registry study for patients treated with etanercept. It was necessary to make assumptions about the quality of life of patients treated with other biologic DMARDs. Due to the scarcity of the HRQoL data, it was not possible to link effectiveness data from the RCTs, in terms of ACR Pedi or CHAQ score, to a HRQoL utility measure. Furthermore, no HRQoL data were identified to inform the estimate of disutility of disease flare or the caregiver burden.

There were limited data available for the long-term discontinuation rates for patients for some of the biologic DMARDs and it was necessary to assume that the discontinuation rates for the biologic DMARDs were the same as each other.

The economic analysis has compared biologic DMARDs against methotrexate only, for patients with an insufficient response to previous methotrexate. The NICE scope also includes best supportive care (e.g. NSAIDs, corticosteroids) as a comparator in patients who cannot tolerate a DMARD (e.g. methotrexate) but this has not been included in the analysis due to lack of available data to make a comparison with best supportive care. Such patients would likely be offered a biologic DMARD rather than receiving best supportive care, therefore this comparison is not necessarily clinical relevant.

The model consists of a simple structure that does not incorporate the natural history of the disease in terms of long-term disease progression. JIA causes joint disease requiring joint operations and is associated with other co-morbidities. It is unclear from the current evidence how biologic DMARDs affect the natural history of the disease and the occurrence of these outcomes.

7.3 Uncertainties

The RCTs included in our systematic review of clinical-effectiveness did not report the impact of treatment on extra-articular manifestations. Uveitis is the most common of these manifestations and if not identified and adequately controlled can lead to permanent vision loss. Current guidance is to treat JIA patients with uveitis that has not responded to steroids or methotrexate with anti-TNF drugs (of

which etanercept and adalimumab are the two anti-TNF drugs within the scope of this assessment).²⁹ Expert clinical opinion suggests that etanercept would rarely be used to treat JIA-associated uveitis.

Furthermore, no HRQoL utility values for the impact of uveitis on the health-related quality of life in children with JIA were identified in our systematic review of quality of life. The paucity of good quality evidence for the effectiveness of biologic DMARDs means that the clinical and cost-effectiveness of treating JIA patients with uveitis is currently uncertain. However, it could be assumed that if biologic DMARD treatment of uveitis is effective in reducing sight impairment in addition to improving general JIA symptoms then the cost-effectiveness estimates generated in the independent economic evaluation in this assessment report would be improved. The SYCAMORE RCT of adalimumab in combination with methotrexate for JIA associated uveitis (funded by the NIHR HTA programme and Arthritis Research UK) has recently completed recruitment and will include an assessment of cost-effectiveness.⁸⁷

The lack of available suitable published cost-utility models, necessitated building a new model which aimed to resemble clinical practice, but also utilise the effectiveness data from the RCTs. The design of the RCTs does not necessarily represent clinical practice (e.g. there wouldn't be a lead-in phase with a biologic DMARD).

The model has not incorporated the impact of biologic DMARD treatment on disease progression, and assumes that the HRQoL of patients treated with methotrexate is constant over time. The results therefore may under-estimate the cost-effectiveness of treatment.

The model has assumed that treatment is equally effective for subsequent lines of biologic DMARD treatment as for the first line of treatment. If effectiveness is seen to be reduced in subsequent lines of therapy for particular switching regimens then cost-effectiveness may be reduced compared to the results presented in this report (relating to abatacept as 2nd line treatment, and the scenario analysis of three lines of treatment).

The model has been modelled with a 30 year time horizon in the base case analysis. There is a lack of long-term outcome data for JIA patients. In addition, there are often differences in the management of JIA patients as adults, than as children which may affect patient outcomes. However, there are little empirical data available on the management of adult patients with JIA.

8 CONCLUSIONS

8.1 Implications for service provision

Given that biologic DMARDs are currently used in the treatment of JIA any recommendation supporting their use is unlikely to have significant implications for service provision (e.g. in terms of changes to infrastructure, staff training). However, further recruitment and training of staff is required to address workforce capacity shortages in some specialist centres.

8.2 Suggested research priorities

Randomised head-to-head comparisons of biologic DMARDs are necessary to establish comparative effectiveness. Currently they are assumed to be equivalent based on indirect comparisons of a small number of trials with relatively small patient numbers. Trials should be sufficiently powered, with long-term follow-up of safety and efficacy, and should include an economic evaluation to assess cost-effectiveness. Treatment response should be assessed at a threshold that is considered clinically significant (e.g. ACR Pedi 70 or higher) and should also include measures disease inactivity. Additional instruments to the ACR Pedi criteria should be used, such as the JADAS instrument.³⁵⁻³⁷ Future trials of biologic DMARDs should stratify by disease phenotype to assess the differential effects of each treatment.

RCTs are also required for sub-types of JIA where evidence is currently lacking, including ERA, and PA. As mentioned, the SYCAMORE trial of adalimumab in patients with JIA-associated uveitis⁸⁷ has recently closed for recruitment early, following interim analysis showing that adalimumab is favourable in the treatment of JIA-associated uveitis.

Further research is needed to establish the HRQoL benefits associated with biological treatment in children with JIA and their caregivers.

9 REFERENCES

1. Prakken B, Albani S, Martini A. Juvenile idiopathic arthritis. *Lancet* 2011;**377**:2138-49.
2. Ravelli A, Martini A. Juvenile idiopathic arthritis. *Lancet* 2007;**369**:767-78.
3. Rigante D, Bosco A, Esposito S. The Etiology of Juvenile Idiopathic Arthritis. *Clin Rev. Allergy Immunol.* 2014.

4. Duffy CM, Colbert RA, Laxer RM, Schanberg LE, Bowyer SL. Nomenclature and classification in chronic childhood arthritis: time for a change? *Arthritis Rheum.* 2005;**52**:382-5.
5. Moorthy LN, Peterson MG, Hassett AL, Lehman TJ. Burden of childhood-onset arthritis. *Pediatr.Rheumatol.Online.J.* 2010;**8**:20-8.
6. Petty RE, Southwood TR, Manners P, Baum J, Glass DN, Goldenberg J *et al.* International League of Associations for Rheumatology classification of juvenile idiopathic arthritis: second revision, Edmonton, 2001. *J Rheumatol.* 2004;**31**:390-2.
7. Arthritis Research UK. What are the different types of JIA? <http://www.arthritisresearchuk.org/arthritis-information/conditions/juvenile-idiopathic-arthritis/different-types.aspx> . 2014. 15-10-2014.
8. Arthritis Care. Arthritis in children. <http://www.arthritiscare.org.uk/aboutarthritis/conditions/arthritisinchildren> . 7-10-2014. Arthritis Care. 15-10-2014.
9. Arthritis Care. *My child has oligoarticular JIA. A guide to the condition and its treatment.* London: Arthritis Care; 2013.
10. Arthritis Care. *My child has polyarticular JIA. A guide to the condition and its treatment.* London: Arthritis Care; 2013.
11. Arthritis Care. *My child has systemic JIA. A guide to the condition and its treatment.* London: Arthritis Care; 2013.
12. National Rheumatoid Arthritis Society. What is JIA? <http://www.nras.org.uk/information-on-jia-what-is-jia--81> . 2014. 15-10-2014.
13. McErlane, F. Validation of JADAS in all ILAR Subtypes of Juvenile Idiopathic Arthritis in the Clinical Setting (oral presentation). *Rheumatology* 2012. 2012. 1-5-2012.
14. Ruperto N, Giannini EH, Pistorio A, Brunner HI, Martini A, Lovell DJ. Is it time to move to active comparator trials in juvenile idiopathic arthritis?: a review of current study designs. *Arthritis and rheumatism* 2010;**62**:3131-9.
15. Minden K, Kiessling U, Listing J, Niewerth M, Doring E, Meincke J *et al.* Prognosis of patients with juvenile chronic arthritis and juvenile spondyloarthritis. *J Rheumatol.* 2000;**27**:2256-63.
16. Bertilsson L, Andersson-Gare B, Fasth A, Petersson IF, Forsblad-D'elia H. Disease course, outcome, and predictors of outcome in a population-based juvenile chronic arthritis cohort followed for 17 years. *J Rheumatol.* 2013;**40**:715-24.
17. Thierry S, Fautrel B, Lemelle I, Guillemin F. Prevalence and incidence of juvenile idiopathic arthritis: a systematic review. *Joint, bone, spine : revue du rhumatisme* 2014;**81**:112-7.
18. Parsons, S., Ingram, M., Clarke-Cornwell, S. M., and Symmons, D. P. M. *A Heavy Burden. The occurrence and impact of musculoskeletal conditions in the United Kingdom today.* The University of Manchester and Arthritis Research UK; 2011.

19. Martin, K. Juvenile Idiopathic Arthritis (JIA) explained. <http://www.ccaa.org.uk/index.php?id=3> 2014. 2014. Children's Chronic Arthritis Association. 15-10-2014.
20. NHS England Clinical Reference Group for Paediatric Medicine. *Interim Clinical Commissioning Policy Statement: Biologic Therapies for the treatment of Juvenile Idiopathic Arthritis (JIA)*. Redditch : NHS England; 2015. NHS England E03/PS/a. Available at: https://www.engage.england.nhs.uk/consultation/specialised-services-policies/user_uploads/biolgcs-juvenl-idiop-arthrs-pol.pdf. Accessed 22-4-2015
21. Office of National Statistics, Population Estimates Uni. *Population Estimates for UK, England and Wales, Scotland and Northern Ireland, Mid-2013*. Office of National Statistics; 2014. Available at: <http://www.ons.gov.uk/ons/publications/re-reference-tables.html?edition=tcm%3A77-322718>. Accessed 22-4-2015
22. Thomson W, Barrett JH, Donn R, Pepper L, Kennedy LJ, Ollier WE *et al*. Juvenile idiopathic arthritis classified by the ILAR criteria: HLA associations in UK patients. *Rheumatology.(Oxford)*. 2002;**41**:1183-9.
23. Kearsley-Fleet, L, Davies, R, Baildam, E, Beresford, M, Foster, H, Southwood, T. R, Thomson, W, and Hyrich, K. Factors associated with choice of biologic among children with juvenile idiopathic arthritis: results from 2 UK paediatric biologic registers. 2015.
24. Vidqvist KL, Malin M, Varjolahti-Lehtinen T, Korpela MM. Disease activity of idiopathic juvenile arthritis continues through adolescence despite the use of biologic therapies. *Rheumatology.(Oxford)*. 2013;**52**:1999-2003.
25. Bechtold S, Simon D. Growth abnormalities in children and adolescents with juvenile idiopathic arthritis. *Rheumatol.Int*. 2014;**34**:1483-8.
26. Stagi S, Masi L, Capannini S, Cimaz R, Tonini G, Matucci-Cerinic M *et al*. Cross-sectional and longitudinal evaluation of bone mass in children and young adults with juvenile idiopathic arthritis: the role of bone mass determinants in a large cohort of patients. *J Rheumatol*. 2010;**37**:1935-43.
27. Heiligenhaus A, Heinz C, Edelsten C, Kotaniemi K, Minden K. Review for disease of the year: epidemiology of juvenile idiopathic arthritis and its associated uveitis: the probable risk factors. *Ocul.Immunol.Inflamm*. 2013;**21**:180-91.
28. British Society for Paediatric and Adolescent Rheumatology (BSPAR). Guidelines for Screening for Uveitis in Juvenile Idiopathic Arthritis (JIA) Produced jointly by BSPAR and the RCPOphth 2006. <https://www.bspar.org.uk/DocStore/FileLibrary/PDFs/BSPAR%20Guidelines%20for%20Eye%20Screening%202006.pdf> . 2006. 22-4-0015.
29. NHS England Specialised Services Clinical Reference Group for Specialised Ophthalmology. *Clinical Commissioning Policy: Adalimumab (Humira) and Infliximab (Remicade) as Anti-TNF Alpha Treatment Options for Paediatric Patients with Severe Refractory Uveitis* . NHS England; 22-4-2015. NHS England D12/P/a
30. Tong A, Jones J, Craig JC, Singh-Grewal D. Children's experiences of living with juvenile idiopathic arthritis: a thematic synthesis of qualitative studies. *Arthritis Care Res (Hoboken.)*. 2012;**64**:1392-404.

31. Giannini EH, Ruperto N, Ravelli A, Lovell DJ, Felson DT, Martini A. Preliminary definition of improvement in juvenile arthritis. *Arthritis and rheumatism* 1997;**40**:1202-9.
32. Prince FHM, Otten MH, van Suijlekom-Smit LWA. Diagnosis and management of juvenile idiopathic arthritis. *British Medical Journal* 2010;**341**.
33. Consolaro A, Ruperto N, Bazso A, Pistorio A, Magni-Manzoni S, Filocamo G *et al*. Development and validation of a composite disease activity score for juvenile idiopathic arthritis. *Arthritis and rheumatism* 2009;**61**:658-66.
34. McErlane F, Beresford MW, Baildam EM, Chieng SE, Davidson JE, Foster HE *et al*. Validity of a three-variable Juvenile Arthritis Disease Activity Score in children with new-onset juvenile idiopathic arthritis. *Annals of the rheumatic diseases* 2013;**72**:1983-8.
35. Consolaro A, Negro G, Chiara Gallo M, Bracciolini G, Ferrari C, Schiappapietra B *et al*. Defining criteria for disease activity States in nonsystemic juvenile idiopathic arthritis based on a three-variable juvenile arthritis disease activity score. *Arthritis care & research* 2014;**66**:1703-9.
36. Consolaro A, Ruperto N, Bracciolini G, Frisina A, Gallo MC, Pistorio A *et al*. Defining criteria for high disease activity in juvenile idiopathic arthritis based on the juvenile arthritis disease activity score. *Annals of the rheumatic diseases* 2014;**73**:1380-3.
37. Horneff G, Becker I. Definition of improvement in juvenile idiopathic arthritis using the juvenile arthritis disease activity score. *Rheumatology (Oxford, England)* 2014;**53**:1229-34.
38. Consolaro A, Negro G, Lanni S, Solari N, Martini A, Ravelli A. Toward a treat-to-target approach in the management of juvenile idiopathic arthritis. *Clinical and experimental rheumatology* 2012;**30**:S157-S162.
39. Wallace CA, Ruperto N, Giannini E, Childhood Arthritis, Rheumatology Research Alliance, Pediatric Rheumatology International Trials Organization *et al*. Preliminary criteria for clinical remission for select categories of juvenile idiopathic arthritis. *The Journal of Rheumatology* 2004;**31**:2290-4.
40. Wallace CA, Giannini EH, Huang B, Itert L, Ruperto N. American College of Rheumatology provisional criteria for defining clinical inactive disease in select categories of juvenile idiopathic arthritis. *Arthritis care & research* 2011;**63**:929-36.
41. Brunner HI, Lovell DJ, Finck BK, Giannini EH. Preliminary definition of disease flare in juvenile rheumatoid arthritis. *J Rheumatol* 2002;**29**:1058-64.
42. Lovell DJ, Giannini EH, Reiff A, Cawkwell GD, Silverman ED, Nocton JJ *et al*. Etanercept in children with polyarticular juvenile rheumatoid arthritis. Pediatric Rheumatology Collaborative Study Group. *New England Journal of Medicine* 2000;**342**:763-9.
43. National Institute for Health and Care Excellence. Guidance on the use of etanercept for the treatment of juvenile idiopathic arthritis. NICE technology appraisal (TA35). <https://www.nice.org.uk/guidance/ta35> . 2002. 22-10-2014.
44. National Institute for Health and Care Excellence. Tocilizumab for the treatment of systemic juvenile idiopathic arthritis. NICE technology appraisals (TA238). <http://www.nice.org.uk/guidance/ta238> . 2011. 27-10-2014.

45. European Medicines Agency. Orencia: EPAR - Product Information. http://www.ema.europa.eu/ema/index.jsp?curl=pages/medicines/human/medicines/000701/human_med_000958.jsp&mid=WC0b01ac058001d124 . 17-6-2014. 15-10-2014.
46. European Medicines Agency. Humira: EPAR - Product Information. http://www.ema.europa.eu/ema/index.jsp?curl=pages/medicines/human/medicines/000481/human_med_000822.jsp&mid=WC0b01ac058001d124 . 17-6-2014. 15-10-2014.
47. European Medicines Agency. Etanercept: EPAR - Product Information. http://www.ema.europa.eu/ema/index.jsp?curl=pages/medicines/human/medicines/000262/human_med_000764.jsp&mid=WC0b01ac058001d124 . 9-3-2014. 22-10-2014.
48. European Medicines Agency. Tocilizumab: EPAR - Product Information. http://www.ema.europa.eu/ema/index.jsp?curl=pages/medicines/human/medicines/000955/human_med_001042.jsp&mid=WC0b01ac058001d124 . 29-9-2014. 22-10-2014.
49. Sonnenberg FA, Beck JR. Markov models in medical decision making: a practical guide. *Medical Decision Making* 1993;**13**:322-39.
50. Berard RA, Laxer RM. Etanercept (Enbrel) in the treatment of juvenile idiopathic arthritis. *Expert Opinion on Biological Therapy* 2013;**13**:1623-30.
51. Horizon Scanning Centre. Tocilizumab (RoActemra) for active polyarticular juvenile idiopathic arthritis - second line. <http://www.hsc.nihr.ac.uk/topics/tocilizumab-roactemra-for-active-polyarticular-juv/> . 2012. 22-10-2014.
52. Higgins, J. P. and Green, S. *Cochrane Handbook for Systematic Reviews of Interventions Version 5.1.0 [updated March 2011]*. The Cochrane Collaboration; 2011. Available at: www.cochrane-handbook.org.
53. Bucher HC, Guyatt GH, Griffith LE, Walter SD. The results of direct and indirect treatment comparisons in meta-analysis of randomized controlled trials. *J Clin.Epidemiol.* 1997;**50**:683-91.
54. Donegan S, Williamson P, Gamble C, Tudur-Smith C. Indirect comparisons: a review of reporting and methodological quality. *PLoS One* 2010;**5**:e11054.
55. Glenny AM, Altman DG, Song F, Sakarovitch C, Deeks JJ, D'Amico R *et al*. Indirect comparisons of competing interventions. *Health Technol.Assess.* 2005;**9**:1-iv.
56. Smith JA, Thompson DJ, Whitcup SM, Suhler E, Clarke G, Smith S *et al*. A randomized, placebo-controlled, double-masked clinical trial of etanercept for the treatment of uveitis associated with juvenile idiopathic arthritis. *Arthritis & Rheumatism* 2005;**53**:18-23.
57. Ruperto N, Lovell DJ, Quartier P, Paz E, Rubio-Perez N, Silva CA *et al*. Abatacept in children with juvenile idiopathic arthritis: a randomised, double-blind, placebo-controlled withdrawal trial. *Lancet* 2008;**372**:383-91.
58. Ruperto N, Lovell DJ, Li T, Sztajn bok F, Goldenstein-Schainberg C, Scheinberg M *et al*. Abatacept improves health-related quality of life, pain, sleep quality, and daily participation in subjects with juvenile idiopathic arthritis. *Arthritis care & research* 2010;**62**:1542-51.
59. Ruperto N, Lovell DJ, Quartier P, Paz E, Rubio-Perez N, Silva CA *et al*. Long-term safety and efficacy of abatacept in children with juvenile idiopathic arthritis. *Arthritis & Rheumatism* 2010;**62**:1792-802.

60. Lovell DJ, Ruperto N, Mouy R, Paz E, Rubio-Perez N, Silva CA *et al.* Cumulative long-term safety and efficacy of abatacept in children with juvenile idiopathic arthritis: Results up to 7 years of follow-up. *Arthritis and rheumatism* 2012;**64**:S717-S718.
61. Lovell DJ, Ruperto N, Goodman S, Reiff A, Jung L, Jarosova K *et al.* Adalimumab with or without methotrexate in juvenile rheumatoid arthritis. *New England Journal of Medicine* 2008;**359**:810-20.
62. Lovell DJ, Ruperto N, Jarosova K, Nemcova D, Vargova V, Michels H *et al.* The Impact of Adalimumab On Growth in Patients with Juvenile Idiopathic Arthritis. *Arthritis and rheumatism* 2012;**64**:S329-S330.
63. Ruperto N, Lovell DJ, Jarosova K, Nemcova D, Vargova V, Michels H *et al.* PREs-FINAL-2001: The impact of adalimumab on growth in patients with juvenile idiopathic arthritis. *Pediatric Rheumatology* 2013;**11**.
64. Ruperto N, Lovell D, Quartier P, Ravelli A, Karunaratne M, Kalabic J *et al.* Treating to target of minimal disease activity and normal function in polyarticular juvenile idiopathic arthritis with adalimumab: Analysis from a phase 3 clinical trial. *Annals of the rheumatic diseases* 2014;**73**:130-1.
65. Lovell DJ, Giannini EH, Reiff A, Jones OY, Schneider R, Olson JC *et al.* Long-term efficacy and safety of etanercept in children with polyarticular-course juvenile rheumatoid arthritis: interim results from an ongoing multicenter, open-label, extended-treatment trial. *Arthritis & Rheumatism* 2003;**48**:218-26.
66. Lovell DJ, Reiff A, Jones OY, Schneider R, Nocton J, Stein LD *et al.* Long-term safety and efficacy of etanercept in children with polyarticular-course juvenile rheumatoid arthritis. *Arthritis & Rheumatism* 2006;**54**:1987-94.
67. Lovell DJ, Reiff A, Ilowite NT, Wallace CA, Chon Y, Lin SL *et al.* Safety and efficacy of up to eight years of continuous etanercept therapy in patients with juvenile rheumatoid arthritis. *Arthritis & Rheumatism* 2008;**58**:1496-504.
68. Brunner HI, Ruperto N, Zuber Z, Keane C, Harari O, Kenwright A *et al.* Efficacy and safety of tocilizumab in patients with polyarticular-course juvenile idiopathic arthritis: results from a phase 3, randomised, double-blind withdrawal trial. *Ann.Rheum.Dis.* 2014.
69. Brunner H, Ruperto N, Zuber Z, Cuttica RnJ, Xavier R, Calvo I *et al.* A4: Efficacy and Safety of Tocilizumab in Patients With Polyarticular-Course Juvenile Idiopathic Arthritis: 2-Year Data From CHERISH. *Arthritis & Rheumatology* 2014;**66**:S5-S6.
70. Baildam E, Ramanan A, Woo P, Brunner H, Ruperto N, Zuber Z *et al.* Efficacy and safety of tocilizumab in polyarticular-course juvenile idiopathic arthritis: 2 year data from cherish. *Rheumatology (United Kingdom)* 2014;**i52-i53**.
71. Brunner HI, Ruperto N, Zuber Z, Cuttica RJ, Xavier R, Calvo I *et al.* Efficacy and Safety Of Tocilizumab In Patients With Polyarticular-Course Juvenile Idiopathic Arthritis: 2-Year Data From Cherish. *Arthritis & Rheumatism* 2013;**65**:S335.
72. De Benedetti F, Ruperto N, Zuber Z, Keane C, Harari O, Kenwright A *et al.* Efficacy and Safety of Tocilizumab in Patients with Polyarticular Juvenile Idiopathic Arthritis: Data from A Phase 3 Trial. *Annals of the rheumatic diseases* 2013;**72**:70.

73. Baildam E, Ruperto N, Brunner H, Zuber Z, Keane C, Harari O *et al.* Efficacy and Safety of Tocilizumab in Polyarticular Juvenile Idiopathic Arthritis: Cherish Results at Week 40. *Rheumatology* 2013;**52**:35.
74. De Benedetti F., Ruperto N, Zuber Z, Cuttica R, Xavier R, Calvo I *et al.* PRoS-FINAL-2180: Efficacy and safety of tocilizumab (TCZ) in patients with polyarticular-course juvenile idiopathic arthritis (pcJIA): 2-year data from CHERISH. *Pediatric Rheumatology* 2013;**11**:O15.
75. Brunner H, Ruperto N, Zuber Z, Keane C, Harari O, Kenwright A *et al.* Efficacy and Safety of Tocilizumab in Patients with Polyarticular Juvenile Idiopathic Arthritis: Data From a Phase 3 Trial. *Arthritis & Rheumatism* 2012;**64**:S682.
76. Bharucha, K. N., Brunner, H. I., Ruperto, N., Cabral, D. A., Gedalia, A., Gerloni, V., Jorgensen, C., Ramanan, A., Lovell, D., Martin, A., Frane, J., Wells, C., and De Benedetti Sr, F. Growth during tocilizumab therapy for polyarticular-course juvenile idiopathic arthritis: 2-year data from a phase 3 clinical trial. American College of Rheumatology Annual Scientific Meeting 2014 in Boston, USA; November 14-19, 2014 . 2014.
77. abbvie. *Adalimumab Multiple technology appraisal (MTA): Etanercept, abatacept, adalimumab and tocilizumab for treating juvenile idiopathic arthritis (including review of TA35)*. 2015.
78. Roche. *Tocilizumab (RoActemra®) for the treatment of juvenile idiopathic polyarthritis. MTA Submission*. 2015.
79. Otten MH, Anink J, Spronk S, Van Suijlekom-Smit LWA. Efficacy of biological agents in juvenile idiopathic arthritis: A systematic review using indirect comparisons. *Annals of the rheumatic diseases* 2013;**72**:1806-12.
80. Nikolakopoulou A, Chaimani A, Veroniki AA, Vasiliadis HS, Schmid CH, Salanti G. Characteristics of networks of interventions: a description of a database of 186 published networks. *PLoS One* 2014;**9**:e86754.
81. Mills EJ, Thorlund K, Ioannidis JP. Demystifying trial networks and network meta-analysis. *BMJ* 2013;**346**:f2914.
82. Kearsley-Fleet, L, Davies, R, Lunt, M, Southwood, T, and Hyrich, K. Factors Associated with Improvement in Disease Activity Following Initiation of Etanercept in Children and Young People with Juvenile Idiopathic Arthritis: Results from the British Society for Paediatric and Adolescent Rheumatology Etanercept Cohort Study (BSPAR-ETN). 2015.
83. Geikowski T, Becker I, Horneff G. Predictors of response to etanercept in polyarticular-course juvenile idiopathic arthritis. *Rheumatology.(Oxford)*. 2014;**53**:1245-9.
84. Albarouni M, Becker I, Horneff G. Predictors of response to methotrexate in juvenile idiopathic arthritis. *Pediatr.Rheumatol.Online.J.* 2014;**12**.
85. Bristol-Myers Squibb. *Bristol-Myers Squibb (BMS) abatacept submission for NICE Multiple Technology Appraisal (MTA) ID738: "Abatacept, adalimumab, etanercept and tocilizumab for treating juvenile idiopathic arthritis (including review of TA35)"*. 2015.
86. Lovell, D., Ruperto, N., Mouy, R., Paz, E., Rubio-Perez, N., Silva, C. A., Abud-Mendoza, C., Burgos-Vargas, R., Gerloni, V., Melo-Gomes, J. A., Saad-Magalhaes, C., Chavez, J., Huemer, C., Kivitz, A., Blanco, F. J., Foeldvari, I., Hofer, M., Huppertz, H., Job, Deslandre C., Minden,

- K., Flores, Nunez A., Block, A. J., Martini, A., and On behalf of the Pediatric Rheumatology Collaborative Study Group and Paediatric Rheumatology International Trials Organisation Investigators. Cumulative Long-Term Safety and Efficacy of Abatacept in Children with Juvenile Idiopathic Arthritis: Results up to 7 Years of Follow-up. American College of Rheumatology (ACR) Congress. *Arthritis & Rheumatism* 64, 1679. 2012. 9-11-2012.
87. Ramanan AV, Dick AD, Benton D, Compeyrot-Lacassagne S, Dawoud D, Hardwick B *et al.* A randomised controlled trial of the clinical effectiveness, safety and cost-effectiveness of adalimumab in combination with methotrexate for the treatment of juvenile idiopathic arthritis associated uveitis (SYCAMORE Trial). *Trials* 2014;**15**:14.
 88. Burgos-Vargas R, Tse SM, Horneff G, Pangan AL, Unnebrink K, Anderson JK. A3: Efficacy and Safety of Adalimumab in Pediatric Patients With Enthesitis Related Arthritis. *Arthritis & Rheumatology* 2014;**66**:S4.
 89. Burgos-Vargas R, Tse S, Horneff G, Pangan A, Unnebrink K, Anderson J. Efficacy and Safety of Adalimumab in Pediatric Patients with Enthesitis Related Arthritis. *Journal of Rheumatology* 2014;**41**:1532-3.
 90. Burgos-Vargas R, Tse SML, Horneff G, Pangan AL, Unnebrink K, Anderson JK. Efficacy and safety of adalimumab in paediatric patients with enthesitis related arthritis. *Rheumatology (United Kingdom)* 2014;**53**:i51-i52.
 91. Burgos-Vargas R, Tse SM, Horneff G, Pangan AL, Unnebrink K, Anderson JK. Efficacy and Safety of Adalimumab in Pediatric Patients With Enthesitis Related Arthritis. *Arthritis & Rheumatism* 2013;**65**:S336.
 92. Burgos-Vargas R, Tse S, Horneff G, Pangan AL, Unnebrink K, Anderson JK. PRoS-FINAL-2179: Efficacy and safety of adalimumab in pediatric patients with enthesitis related arthritis. *Pediatric Rheumatology* 2013;**11**:O14.
 93. Tse SM, Burgos-Vargas R, Horneff G, Pangan AL, Kalabic J, Unnebrink K *et al.* Discontinuation of Concomitant Medication for Enthesitis-Related Arthritis during 52 Weeks of Treatment with Adalimumab. *Arthritis & Rheumatology* 2014;**66**:S113.
 94. Kingsbury DJ, Bader-Meunier B, Patel G, Arora V, Kalabic J, Kupper H. Safety, effectiveness, and pharmacokinetics of adalimumab in children with polyarticular juvenile idiopathic arthritis aged 2 to 4 years. *Clinical Rheumatology* 2014;**33**:1433-41.
 95. Imagawa T, Takei S, Umebayashi H, Yamaguchi K, Itoh Y, Kawai T *et al.* Efficacy, pharmacokinetics, and safety of adalimumab in pediatric patients with juvenile idiopathic arthritis in Japan. *Clinical Rheumatology* 2012;**31**:1713-21.
 96. Simonini G, Druce K, Cimaz R, Macfarlane GJ, Jones GT. Current Evidence of Anti-Tumor Necrosis Factor alpha Treatment Efficacy in Childhood Chronic Uveitis: A Systematic Review and Meta-Analysis Approach of Individual Drugs. *Arthritis care & research* 2014;**66**:1073-84.
 97. Pfizer Ltd. *Multiple technology appraisal (MTA): Etanercept (Enbrel®) for treating juvenile idiopathic arthritis (including review of TA35)*. Company evidence submission. 2015.
 98. Zeng H, Zeng P, Xie Y, Tang Y, Li F. A randomized controlled clinical trial to evaluate the efficacy of recombinant human tumor necrosis factor- α receptor type II fusion protein antibody in juvenile idiopathic arthritis. *Allergy: European Journal of Allergy and Clinical Immunology* 2013;**68**:11.

99. Horneff G, Burgos-Vargas R, Constantin T, Foeldvari I, Vojinovic J, Chasnyk VG *et al.* Efficacy and safety of open-label etanercept on extended oligoarticular juvenile idiopathic arthritis, enthesitis-related arthritis and psoriatic arthritis: part 1 (week 12) of the CLIPPER study. *Ann.Rheum.Dis.* 2014;**73**:1114-22.
100. Ravelli, A., Ruperto, N., Pederzoli, S., Burgos-Vargas, R., Kobusinska, K., Schmeling, H., Sztajn bok, F., Weller-Heinemann, F., Zholobova, E., Zulian, F., Allen, R., Chaitow, J., Keane, C., Wells, C., Martini, A., Lovell, D. J., and De Benedetti, F. Assessment of Radiographic Progression in Patients With Polyarticular-Course Juvenile Idiopathic Arthritis Treated With Tocilizumab: 2-Year Data From CHERISH. *Arthritis & Rheumatology* 66, S17-S18. 2014.
101. Dias S, Sutton AJ, Ades AE, Welton NJ. Evidence synthesis for decision making 2: a generalized linear modeling framework for pairwise and network meta-analysis of randomized controlled trials. *Med.Decis.Making* 2013;**33**:607-17.
102. Foeldvari I, Horneff G, Minden K, Trauzeddel R, Kuemmerle-Deschner JB, Tenbrock K *et al.* Remission Induction By Etanercept in Enthesitis Related Arthritis JIA-Patients (juvenile undifferentiated Spondylarthropathy). *Arthritis & Rheumatology* 2014;**66**:3536.
103. Constantin, T., Foeldvari, I., Vojinovic, J., Horneff, G., Burgos-Vargas, R., Nikishina, I., Akikusa, J., Avcin, T., Chaitow, J., Koskova, E., Lauwerys, B., Bukowski, J., Zang, C., Wajdula, J., Woodworth, D., Vlahos, B., Martini, A., and Ruperto, N. Long-term safety and efficacy of etanercept in paediatric subjects with extended oligoarticular juvenile idiopathic arthritis, enthesitis-related arthritis, or psoriatic arthritis. *Arthritis and Rheumatism Conference: American College of Rheumatology/Association of Rheumatology Health Professionals Annual Scientific Meeting* . 2013.
104. Cordero-Coma M, Yilmaz T, Onal S. Systematic Review of Anti-Tumor Necrosis Factor-alpha Therapy for Treatment of Immune-mediated Uveitis. *Ocular Immunology and Inflammation* 2013;**21**:12-20.
105. Garcia-De-Vicuna C, Diaz-Llopis M, Salom D, Bou R, Diaz-Cascajosa J, Cordero-Coma M *et al.* Usefulness of adalimumab in the treatment of refractory uveitis associated with juvenile idiopathic arthritis. *Mediators.Inflamm.* 2013;**2013**:560632.
106. Diaz-Llopis M, Salom D, Garcia-De-Vicuna C, Cordero-Coma M, Ortega G, Ortego N *et al.* Treatment of refractory uveitis with adalimumab: a prospective multicenter study of 131 patients. *Ophthalmology* 2012;**119**:1575-81.
107. Magli A, Forte R, Navarro P, Russo G, Orlando F, Latanza L *et al.* Adalimumab for juvenile idiopathic arthritis-associated uveitis. *Graefes Arch.Clin.Exp.Ophthalmol.* 2013;**251**:1601-6.
108. Zannin ME, Birolo C, Gerloni VM, Misero cchi E, Pontikaki I, Paroli MP *et al.* Safety and efficacy of infliximab and adalimumab for refractory uveitis in juvenile idiopathic arthritis: 1-year followup data from the Italian Registry. *J Rheumatol.* 2013;**40**:74-9.
109. Simonini G, Taddio A, Cattalini M, Caputo R, de LC, Parentin F *et al.* Superior efficacy of Adalimumab in treating childhood refractory chronic uveitis when used as first biologic modifier drug: Adalimumab as starting anti-TNF-alpha therapy in childhood chronic uveitis. *Pediatr.Rheumatol.Online J* 2013;**11**:16.
110. Haapasaari JE, Kauppi M, Hakala MS, Kautiainen H. Economic evaluation of etanercept therapy in the treatment of re-refractory JIA. *Arthritis and rheumatism* 2002;**46**:S480.

111. Brodzsky, V., Pentek, M., Majer, I., Karpati, K., and Gulacsi, L. [*Etanercept in patients with juvenile idiopathic arthritis: systematic review and economic evaluation*]. 2006. Available at: http://hecon.uni-corvinus.hu/download/english/publ/hecon_research_20_eng.pdf. Accessed
112. Prince FHM, de Bekker-Grob EW, Twilt M, Van Rossum MAJ, Hoppenreijns EPAH, ten Cate R *et al.* An analysis of the costs and treatment success of etanercept in Juvenile Idiopathic Arthritis. *Clinical and experimental rheumatology* 2011;**29**:443.
113. Simpson K, Hubert MM, On PV, Cifaldi M, Shaw J. Long-Term Cost-Effectiveness of Adalimumab Therapy in Juvenile Idiopathic Arthritis: From a Canadian Perspective. *Journal of Rheumatology* 2012;**39**:1712.
114. Luca N, Burnett H, Ungar W, Beukelman T, Feldman BM, Schwartz G *et al.* Cost-Effectiveness Analysis of Early Biologic Treatment in Polyarticular Juvenile Idiopathic Arthritis. *Arthritis & Rheumatism* 2012;**64**:S501.
115. Luca N, Burnett H, Ungar W, Beukelman T, Feldman B, Schwartz G *et al.* Cost-effectiveness analysis of early biologic treatment in polyarticular juvenile idiopathic arthritis. *Journal of Rheumatology* 2013;Conference:6.
116. Chang S, Sawyer L, Dejonckheere F, van Suijlekom-Smit LW, Anink J, Diamantopoulos A. Tocilizumab in Polyarticular Juvenile Idiopathic Arthritis - A Cost-Utility Model for the United Kingdom. *Value in Health* 2013;**16**:A564.
117. All Wales Medicines Strategy Group. *Adalimumab (Humira®)*. 2013.
118. All Wales Medicines Strategy Group. *Etanercept (Enbrel®)*. 2013.
119. All Wales Medicines Strategy Group. *Abatacept (Orencia®)*. 2014.
120. All Wales Medicines Strategy Group. *Tocilizumab (RoActemra®)*. 2014.
121. Canadian Agency for Drugs and Technologies in Health. *Tocilizumab (Actemra - Hoffmann-La Roche Limited) new indication: polyarticular juvenile idiopathic arthritis*. 2014.
122. Cummins C, Connock M, Fry-Smith A, Burls A. A systematic review of effectiveness and economic evaluation of new drug treatments for juvenile idiopathic arthritis: etanercept. *Health Technology Assessment* 2002;**9**:1-43.
123. Prince FH, de Bekker-Grob EW, Twilt M, van Rossum MA, Hoppenreijns EP, ten Cate R *et al.* An analysis of the costs and treatment success of etanercept in juvenile idiopathic arthritis: results from the Dutch Arthritis and Biologicals in Children register. *Rheumatology (Oxford, England)* 2011;**50**:1131-6.
124. Simpson, K, Marlow, N., Shaw, J., and Rudakova, A. Pharmacoeconomic issues of adalimumab therapy in juvenile idiopathic arthritis. (*Pediatric pharmacology*) 9, 48-52. 2012. 2-2-2015.
125. Costa, V., Ungar, W. J., and Hancock-Howard, R. L. *The use of biologic response modifiers in polyarticular-course juvenile idiopathic arthritis*. 2010. Available at: http://www.sickkids.ca/pdfs/Research/TASK/biologics/33870-biologics-report_JIA.pdf. Accessed

126. Ungar WJ, Costa V, Hancock-Howard R, Feldman BM, Laxer RM. Cost-effectiveness of biologics in polyarticular-course juvenile idiopathic arthritis patients unresponsive to disease-modifying antirheumatic drugs. *Arthritis care & research* 2011;**63**:111-9.
127. Philips Z, Ginnelly L, Sculpher M, Claxton K, Golder S, Riemsma R *et al.* Review of guidelines for good practice in decision-analytic modelling in health technology assessment. *Health Technol Assess* 2004;**8**:1-158.
128. Drummond MF, O'Brien B, Stoddart GL, Torrance GW. Methods for the economic evaluation of health care programmes (3rd edition). Oxford: Oxford University Press, 2005.
129. National Institute for Health and Care Excellence (NICE). *Guide to the methods of technology appraisal*. London: NICE; 2013.
130. National Institute for Health and Care Excellence (NICE). Guidance on the use of etanercept for the treatment of juvenile idiopathic arthritis. <http://www.nice.org.uk/guidance/ta35> . 2002. 19-5-2015.
131. Simpson, K and Shaw, J. Mapping the Childhood Health Assessment Questionnaire to the Health Utilities Index Mark 2 and Mark 3 in children with juvenile idiopathic arthritis. 19th Annual Conference of the International Society for Quality of Life Research. Quality of life research 21, Abstract 103.2. 2012.
132. Yagudina, R., Kulikov, A., and Zinchuk, I. PMS25 Evaluation of direct costs for the treatment of active juvenile rheumatoid arthritis using biologics. ISPOR 14th annual European congress. Value in Health 14, A306. 2011.
133. Pennington B, Davis S. Mapping from the Health Assessment Questionnaire to the EQ-5D: The Impact of Different Algorithms on Cost-Effectiveness Results. *Value Health* 2014;**17**:762-71.
134. Ruperto N, Lovell DJ, Cuttica R, Wilkinson N, Woo P, Espada G *et al.* A randomized, placebo-controlled trial of infliximab plus methotrexate for the treatment of polyarticular-course juvenile rheumatoid arthritis. *Arthritis Rheum.* 2007;**56**:3096-106.
135. Hendry GJ, Watt GF, Brandon M, Friel L, Turner DE, Lorgelly PK *et al.* The effectiveness of a multidisciplinary foot care program for children and adolescents with juvenile idiopathic arthritis: an exploratory trial. *Journal of rehabilitation medicine* 2013;**45**:467-76.
136. Prince FH, Geerdink LM, Borsboom GJ, Twilt M, van Rossum MA, Hoppenreijns EP *et al.* Major improvements in health-related quality of life during the use of etanercept in patients with previously refractory juvenile idiopathic arthritis. *Annals of the rheumatic diseases* 2010;**69**:138-42.
137. MIMS. Prescription drug database and drug prescribing guide. <http://www.mims.co.uk/> . 2014. MIMS online.
138. Stevenson, M, Archer, R, Tosh, J., Simpson, E, Everson-Hock, E, Stevens, J, Wailoo, A, Hernandez, M, Paisley, S, Williams, K, Scott, D, and Young, A. Adalimumab, etanercept, infliximab, certolizumab pegol, golimumab, tocilizumab and abatacept for the treatment of rheumatoid arthritis not previously treated with disease-modifying anti-rheumatic drugs and after the failure of conventional disease-modifying anti-rheumatic drugs only: systematic review and economic evaluation. <http://www.nice.org.uk/guidance/gid-tag313/documents/rheumatoid-arthritis-adalimumab-etanercept-infliximab-certolizumab-pegol-golimumab-abatacept-and-tocilizumab-review-assessment-report2> . 2013.

139. Barton P, Jobanputra P, Wilson J, Bryan S, Burls A. The use of modelling to evaluate new drugs for patients with a chronic condition: the case of antibodies against tumour necrosis factor in rheumatoid arthritis. *Health Technol. Assess.* 2004;**8**:iii, 1-iii,91.
140. Khan KA, Petrou S, Rivero-Arias O, Walters SJ, Boyle SE. Mapping EQ-5D utility scores from the PedsQL generic core scales. *Pharmacoeconomics* 2014;**32**:693-706.
141. Briggs AH, O'Brien BJ. The death of cost-minimization analysis? *Health Econ.* 2001;**10**:179-84.
142. Dakin H, Wordsworth S. Cost-minimisation analysis versus cost-effectiveness analysis, revisited. *Health Econ.* 2013;**22**:22-34.
143. British Society for Paediatric and Adolescent Rheumatology. Submission to NICE - MTA- Abatacept, Adalimumab, Etanercept and Tocilizumab for the treatment of Juvenile Idiopathic Arthritis. 2015. 20-5-0015.
144. Briggs A, Sculpher M, Claxton K. Decision modelling for health economic evaluation. 2006.
145. Tynjala P, Vahasalo P, Honkanen V, Lahdenne P. Drug survival of the first and second course of anti-tumour necrosis factor agents in juvenile idiopathic arthritis. *Ann. Rheum. Dis.* 2009;**68**:552-7.
146. Klein A, Kaul I, Foeldvari I, Ganser G, Urban A, Horneff G. Efficacy and safety of oral and parenteral methotrexate therapy in children with juvenile idiopathic arthritis: an observational study with patients from the German Methotrexate Registry. *Arthritis Care Res. (Hoboken.)* 2012;**64**:1349-56.
147. Bruns A, Hilario MO, Jennings F, Silva CA, Natour J. Quality of life and impact of the disease on primary caregivers of juvenile idiopathic arthritis patients. *Joint, bone, spine : revue du rhumatisme* 2008;**75**:149-54.
148. Joint Formulary Committee. British National Formulary for Children. <http://www.medicinescomplete.com> . 2015. BMJ Group, Pharmaceutical Press and RCPCH Publications. 2015.
149. Joint Formulary Committee. British National Formulary. 2014. www.bnf.org, British Medical Association & Royal Pharmaceutical Society.
150. Department of Health. *National Schedule of Reference Costs 2013-14 for NHS trusts and NHS foundation trusts*. Department of Health: 2014.
151. Curtis, L. *Unit Costs of Health and Social Care 2013*. <http://www.pssru.ac.uk/project-pages/unit-costs/2012/>: 2014.
152. Thornton J, Lunt M, Ashcroft DM, Baildam E, Foster H, Davidson J *et al*. Costing juvenile idiopathic arthritis: examining patient-based costs during the first year after diagnosis. *Rheumatology. (Oxford)* 2008;**47**:985-90.
153. Cummins, C., Connock, M., Fry-Smith, A., and Burls, A. *A rapid review of new drug treatments for juvenile idiopathic arthritis: Etanercept [TA35 assessment report]*. National Institute for Health and Care Excellence (NICE); 8-1-2001. Available at: <http://www.nice.org.uk/guidance/ta35/documents/assessment-report-for-etanercept-for-juvenile-idiopathic-arthritis-2>. Accessed 5-6-2015

154. Riemsma, Rob, Al, Maiwenn J., Lhachimi, Stefan K., Armstrong, Nigel, Misso, Kate, Manning, Nathan, Lang, Shona, Severens, Johan L., and Kleijnen, Jos. *Tocilizumab for the treatment of systemic juvenile idiopathic arthritis [TA238 ERG report]*. National Institute for Health and Care Excellence (NICE); 16-6-2011. Available at: <http://www.nice.org.uk/guidance/ta238/documents/arthritis-juvenile-idiopathic-systemic-tocilizumab-appraisal-consultation-evidence-review-group-report2>. Accessed 5-6-2015
155. Chen, Yen Fu, Jobanputra, Paresh, Barton, Pelham, Jowett, Sue, Bryan, Stirling, Clark, Wendy, Fry-Smith, Anne, and Burls, Amanda. *A systematic review of the effectiveness of adalimumab, etanercept and infliximab for the treatment of rheumatoid arthritis in adults and an economic evaluation of their cost-effectiveness [TA130 assesment report]*. National Institute for Health and Clinical Excellence (NICE); 3-10-2005. Available at: <http://www.nice.org.uk/guidance/ta130/documents/rheumatoid-arthritis-adalimumab-etanercept-and-infliximab-assessment-report2>. Accessed 5-6-2015
156. Connock, Martin, Tubeuf, Sandy, Malotki, Kinga, Uthman, Abdulrahman, Round, Jeff, Bayliss, Sue, Meads, Catherine, and Moore, David. *Certolizumab pegol (CIMZIA®) for the treatment of Rheumatoid Arthritis [TA130 ERG report]*. National Institute for Health and Care Excellence (NICE); 1-8-2009. Available at: <http://www.nice.org.uk/guidance/ta186/documents/evidence-review-group-report2>. Accessed 5-6-2015
157. Malotki, Kinga, Barton, Pelham, Tsourapas, Angelos, Uthman, Abdulrahman, Liu, Zulian, Routh, Kristina, Connock, Martin, Jobanputra, Paresh, Moore, David, Fry-Smith, Anne, and Chen, Yen Fu. *Adalimumab, etanercept, infliximab, rituximab and abatacept for the treatment of rheumatoid arthritis after the failure of a TNF inhibitor [TA195 assessment report]*. National Institute for Health and Care Excellence (NICE); 24-11-2009. Available at: <http://www.nice.org.uk/guidance/ta195/documents/drugs-for-the-treatment-of-rheumatoid-arthritis-after-the-failure-of-tnf-inhibitor-assessment-report2>. Accessed 5-6-2015
158. Lloyd Jones, Myfanwy, Stevenson, Matt, Stevens, John, and Sutton, Anthea. *Abatacept for the treatment of rheumatoid arthritis after the failure of conventional disease-modifying anti-rheumatic drugs [TA247 rapid review of TA198, ERG report]*. National Institute for Health and Care Excellence (NICE); 2015. Available at: <http://www.nice.org.uk/guidance/ta247/documents/rheumatoid-arthritis-abatacept-2nd-line-evidence-review-group-report3>. Accessed 5-6-2015
159. Otten MH, Prince FH, Armbrust W, ten CR, Hoppenreijns EP, Twilt M *et al.* Factors associated with treatment response to etanercept in juvenile idiopathic arthritis. *JAMA* 2011;**306**:2340-7.
160. Baszis K, Garbutt J, Toib D, Mao J, King A, White A *et al.* Clinical outcomes after withdrawal of anti-tumor necrosis factor alpha therapy in patients with juvenile idiopathic arthritis: a twelve-year experience. *Arthritis Rheum.* 2011;**63**:3163-8.
161. Wallace CA, Huang B, Bandeira M, Ravelli A, Giannini EH. Patterns of clinical remission in select categories of juvenile idiopathic arthritis. *Arthritis Rheum.* 2005;**52**:3554-62.
162. Kuhlthau K, Kahn R, Hill KS, Gnanasekaran S, Ettner SL. The well-being of parental caregivers of children with activity limitations. *Matern.Child Health J.* 2010;**14**:155-63.
163. Gani R, Giovannoni G, Bates D, Kemball B, Hughes S, Kerrigan J. Cost-effectiveness analyses of natalizumab (Tysabri) compared with other disease-modifying therapies for people with highly active relapsing-remitting multiple sclerosis in the UK. *Pharmacoeconomics.* 2008;**26**:617-27.

164. Reeves BC, Langham J, Walker J, Grieve R, Chakravarthy U, Tomlin K *et al.* Verteporfin Photodynamic Therapy Cohort Study: Report 2: Clinical Measures of Vision and Health-Related Quality of Life. *Ophthalmology* 2009;**116**:2463-70.
165. National Rheumatoid Arthritis Society (NRAS). A Focus on Juvenile Idiopathic Arthritis. www.nras.org.uk/publications/a-focus-on-juvenile-idiopathic-arthritis-report . 2014. 20-5-0015.
166. Ungar WJ, Costa V, Burnett HF, Feldman BM, Laxer RM. The use of biologic response modifiers in polyarticular-course juvenile idiopathic arthritis: A systematic review. *Seminars in Arthritis and Rheumatism* 2013;**42**:597-618.
167. Kemper A, Van Mater H, Coeytaux R, Williams J, Sanders G. Systematic review of disease-modifying antirheumatic drugs for juvenile idiopathic arthritis. *BMC Pediatrics* 2012;**12**.
168. Angeles-Han ST, McCracken C, Yeh S, Jenkins K, Stryker D, Rouster-Stevens K *et al.* Characteristics of a cohort of children with Juvenile Idiopathic Arthritis and JIA-associated Uveitis. *Pediatr.Rheumatol.Online J* 2015;**13**:19.
169. Higgins JPT, Altman DG, Gøtzsche PC, Jüni P, Moher D, Oxman AD *et al.* The Cochrane Collaboration's tool for assessing risk of bias in randomised trials. *British Medical Journal* 2011;**343**.

10 APPENDICES

Appendix 1 Search dates and example Medline search strategies for clinical-effectiveness, cost-effectiveness and HRQoL

Databases searched for the systematic reviews of clinical-effectiveness, cost-effectiveness and HRQoL are presented below. Clinical-effectiveness searches were updated 5th May 2015, cost-effectiveness and HRQoL searches 6th May 2015.

Database searched (host)	Clinical-effectiveness searches	Cost-effectiveness searches HRQoL searches
BIOSIS Previews (Web of Science)	Searched to 29/10/2014	1956 - 11/11/2014 1956 - 02/12/2014
Cochrane Central, Cochrane CDSR, Cochrane DARE, Cochrane HTA, and Cochrane Methods (Cochrane Library)	Searched to 04/11/2014	
Cochrane Central, Cochrane DARE, Cochrane Economic Evaluations, and Cochrane Methods (Cochrane Library)		HRQoL: searched to 09/12/2014
Centre for Reviews and Dissemination databases: DARE, HTA, and NHS EED (CRD)	Searched to 04/11/2014	All available years to 11/11/2014 All available years to 09/12/2014
Conference Proceedings Citation Index-Science (CPCI-S) (Web of Science)	1990 - 29/10/2014	1970 - 11/11/2014 1970 - 02/12/2014
DELPHIS		Costs: searched to 10/11//2014
EMBASE (Ovid)	All available years to 29/10/2014	Searched to 10/11//2014 1974 – 1/12/2014
MEDLINE(R) (Ovid)	Searched to 29/10/2014	1946 to October Week 5 2014 1946 to November Week 2 2014
MEDLINE(R) In-Process (MEIP) & Other Non-Indexed Citations (Ovid)	Searched to 29/10/2014	Searched to 10/11//2014 Searched to 25/11//2014
PSYCHINFO - Ebsco		HRQoL: 1954 – 09/12/2014
Science Citation Index Expanded (SCI-EXPANDED) (Web of Science)	1970 - 29/10/2014	1970 - 11/11/2014 1970 - 02/12/2014
Zetoc (Mimas)	Searched to 04/11/2014	

Searched for ongoing trials (all searched on 13/05/2015)

National Institute for Health Research Clinical Research Network (NIHR CRN Portfolio, formally UKCRN website)

Clinical trials.gov

Medline search strategies for clinical-effectiveness, cost-effectiveness and HRQoL are shown here.

These were adapted for other databases and are available on request.

Clinical-effectiveness Medline search strategy

- 1 Arthritis, Juvenile/
- 2 JIA.tw.
- 3 exp Arthritis/
- 4 (arthriti* or oligoarthriti* or polyarthriti* or polyartacula*).tw.
- 5 Rheumatoid Factor/
- 6 "rheumatoid factor".tw.
- 7 or/3-6
- 8 (juvenile* or child* or teen* or adolescen* or youth* or "young person" or "young people" or pediatric* or paediatric*).tw.
- 9 exp Child/ or Adolescent/
- 10 7 and (8 or 9)
- 11 1 or 2 or 10
- 12 (etanercept or enbrel).mp.
- 13 (abatacept or orenca).mp.
- 14 (adalimumab or humira).mp.
- 15 (tocilizumab or toclizumab or RoActemra).mp.
- 16 or/12-15
- 17 11 and 16
- 18 limit 17 to English language
- 19 limit 18 to humans
- 20 (letter or editorial or comment).pt.
- 21 19 not 20

Cost-effectives Medline search strategy

- 1 Arthritis, Juvenile/
- 2 JIA.tw.
- 3 exp Arthritis/
- 4 (arthriti* or oligoarthriti* or polyarthriti* or polyartacula*).tw.
- 5 Rheumatoid Factor/
- 6 "rheumatoid factor".tw.
- 7 or/3-6
- 8 (juvenile* or child* or teen* or adolescen* or youth* or "young person" or "young people" or pediatric* or paediatric*).tw.
- 9 exp Child/ or Adolescent/
- 10 7 and (8 or 9)
- 11 1 or 2 or 10
- 12 (etanercept or enbrel).mp.
- 13 (abatacept or orenca).mp.
- 14 (adalimumab or humira).mp.
- 15 (tocilizumab or toclizumab or RoActemra).mp.
- 16 or/12-15
- 17 11 and 16
- 18 limit 17 to English language
- 19 limit 18 to humans
- 20 (letter or editorial or comment).pt.

- 21 19 not 20
- 22 exp economics/
- 23 exp economics hospital/
- 24 exp economics pharmaceutical/
- 25 exp economics nursing/
- 26 exp economics medical/
- 27 exp "Costs and Cost Analysis"/
- 28 Cost Benefit Analysis/
- 29 exp models economic/
- 30 exp fees/ and charges/
- 31 exp budgets/
- 32 (economic* or cost or costs or costly or costing or price or prices or pricing or pharmaco-economic*).tw.
- 33 (value adj1 money).tw.
- 34 budget\$.tw.
- 35 or/22-34
- 36 ((energy or oxygen) adj cost).tw.
- 37 (metabolic adj cost).tw.
- 38 ((energy or oxygen) adj expenditure).tw.
- 39 or/36-38
- 40 35 not 39
- 41 (letter or editorial or comment or historical article).pt.
- 42 40 not 41
- 43 21 and 42

HRQoL Medline search strategy

- 1 Arthritis, Juvenile/
- 2 JIA.tw.
- 3 exp Arthritis/
- 4 (arthriti* or oligoarthriti* or polyarthriti* or polyarticular*).tw.
- 5 Rheumatoid Factor/
- 6 "rheumatoid factor".tw.
- 7 or/3-6
- 8 (juvenile* or child* or teen* or adolescent* or youth* or "young person" or "young people" or pediatric* or paediatric*).tw.
- 9 exp Child/ or Adolescent/
- 10 7 and (8 or 9)
- 11 1 or 2 or 10
- 12 CHAQ.tw.
- 13 childhood health assessment questionnaire.tw.
- 14 child health questionnaire.tw.
- 15 CHQ.tw.
- 16 CHU 9D.tw.
- 17 PedsQL.tw.
- 18 "Paediatric Quality of Life Inventory".tw.
- 19 "Pediatric Quality of Life Inventory".tw.
- 20 "juvenile arthritis disease activity score".tw.
- 21 JADAS*.tw.
- 22 value of life/
- 23 quality adjusted life year/
- 24 quality adjusted life.ti,ab.
- 25 (qaly* or qald* or qale* or qtime*).tw.
- 26 disability adjusted life.ti,ab.
- 27 daly*.ti,ab.
- 28 health status indicators/

29 eq 5d 3l.tw.
 30 (euroqol or euro qol or eq5d or eq 5d).tw.
 31 (hql or hqol or "h qol" or hrqol or "hr qol").tw.
 32 (hye or hyes).tw.
 33 health* year* equivalen*.ti,ab.
 34 health utilit*.ab.
 35 (hui or hui1 or hui2 or hui3).ti,ab.
 36 disutil*.ti,ab.
 37 rosser.ti,ab.
 38 "quality of well being".tw.
 39 "quality of wellbeing".tw.
 40 qwb.tw.
 41 "willingness to pay".tw.
 42 "standard gamble*".tw.
 43 "time trade off".tw.
 44 "time tradeoff".tw.
 45 tto.tw.
 46 (index adj2 "well being").mp.
 47 (quality adj2 "well being").mp.
 48 (health adj3 utilit*).mp.
 49 ((multiattribute* or "multi attribute*") adj3 ("health ind*" or theor* or "health state*" or
 utilit* or analys*)).mp.
 50 "quality adjusted life year*".mp.
 51 (15D or "15 dimension*").mp.
 52 (12D or "12 dimension*").mp.
 53 "rating scale*".mp.
 54 "linear scal*".mp.
 55 "linear analog".mp.
 56 "visual analog*".mp.
 57 (categor* adj2 scal*).mp.
 58 or/12-57
 59 11 and 58
 60 (comment or editorial or letter).pt.
 61 59 not 60
 62 limit 61 to English language

Appendix 2 Screening Phase 1 – Titles and abstracts for systematic review of clinical-effectiveness

Language	
Non- English language	Exclude
Intervention	
- Abatacept (Orencia) (with or without methotrexate) - Adalimumab (Humira) (with or without methotrexate) - Etanercept (Enbrel) - Tocilizumab (RoActemra) (with or without methotrexate)	Can be either with or without methotrexate (will check usage is as per licenced indication at full paper screen)
Participants	
Juvenile idiopathic arthritis - extended oligoarthritis - poly arthritis (onset or course) - enthesitis-related - psoriatic - undifferentiated	For mixed populations e.g. including systemic or oligo-arthritis, include only if the proportion of the unwanted type(s) is <33% (i.e. 2/3 rd of the population should meet the inclusion criteria) Exclude systemic arthritis (unless NO active systemic symptoms in the previous 6 months); exclude persistent oligoarthritis
Comparators	
- DMARDs e.g. methotrexate, azathioprine, cyclosporin, penicillamine, sulphasalazine and gold preparations. - Best supportive care if DMARDs not tolerated (e.g. NSAIDs, corticosteroids) - Interventions compared with each other	
Outcomes	
One or more of: • Disease activity • Disease flares • Physical function • Joint damage • Pain • Corticosteroid reducing regimens • Extra-articular manifestations (e.g. uveitis) • Body weight and height • Mortality • Adverse effects of treatment • Health-related quality of life	Don't exclude at TIAB screening stage on outcome. Get full paper to check.
Design	
RCT	If NO but data may not be available from RCTs e.g. long-term adverse events, height and growth)
Systematic review	If YES (or possibly Yes) & can't exclude on P, I or C, RETRIEVE for full paper screen & possible ref list check if meets criteria
Abstracts/conference presentations	
Published 2011 or earlier	Exclude
Published 2012 or later: are sufficient details presented to allow appraisal of methodology and assessment of results?	If can't definitely exclude on P,I, C or D RETRIEVE (for full text screen & possible tie up with full papers or ongoing studies)

Appendix 3 Screening Phase 2 – Full papers for systematic review of clinical-effectiveness

Study name or Number:

Design																				
RCT			Yes ↓ Next Q			Unclear ↓ Next Q			No ↓ Exclude											
Abstracts/conference presentations																				
Published 2012 or later			Yes ↓ Next Q			Unclear ↓ Next Q			No ↓ Exclude											
Intervention												Comments								
<ul style="list-style-type: none"> ▪ abatacept (Orencia) ▪ adalimumab (Humira) ▪ etanercept (Enbrel) ▪ tocilizumab (RoActemra) 			Yes (Y) ↓ To drug specific section (row below)			Unclear (U) ↓ Skip to Comparator section (Comp)			No (N) ↓ Exclude (Ex)			etanercept should be monotherapy (not with methotrexate)								
Abatacept (Orencia)			Adalimumab (Humira)			Etanercept (Enbrel)						Tocilizumab (RoActemra)								
JIA sub-type polyarthritis (poly) with insufficient response to other DMARDs including at least 1 TNF inhibitor ^a			JIA sub-type poly with inadequate response to 1 or more DMARDs ^a			JIA sub-type enthesitis-related arthritis (ERA) with inadequate response/intolerance to conventional treatment ^a			JIA sub-type poly (including RF +ve or -ve and extended oligo-arthritis) with inadequate response/intolerance to methotrexate ^a			JIA sub-type psoriatic arthritis (PA) with inadequate response/intolerance to methotrexate ^a			JIA sub-type enthesitis-related arthritis (ERA) with inadequate response/intolerance to conventional treatment ^a			JIA sub-type poly (including RF +ve or -ve and extended oligo-arthritis) in patients not responding to other NSAIDs or corticosteroids ^a		
Y ↓ Next Q	U ↓ Next Q	N ↓ Ex	Y ↓ Next Q	U ↗ Next Q	N ↗ Next Q	Yes ↓ Next Q	U ↓ Age Qs	N ↓ Ex	Y ↓ Next Q	U ↗ Next Q	N ↗ Next Q	Y ↓ Next Q	U ↗ Next Q	N ↗ Next Q	Y ↓ Next Q	U ↓ Age Qs	N ↓ Ex	Y ↓ Next Q	U ↓ Next Q	N ↓ Ex
Participant age 6- years			Participant age 2- years			Participant age 6- years -			Participant age 2- years			Participant age 12- years			Participant age 2- years					
Y ↓ Comp	U ↓ Comp	N ↓ Ex	Y ↓ Comp	U ↓ Comp	N ↓ Ex	Y ↓ Comp	U ↓ Comp	N ↓ Ex	Y ↓ Comp	U ↓ Comp	N ↓ Ex	Y ↓ Comp	U ↓ Comp	N ↓ Ex	Y ↓ Comp	U ↓ Comp	N ↓ Ex	Y ↓ Comp	U ↓ Comp	N ↓ Ex
Comments:			Comments:			Comments:			Comments:			Comments:			Comments:					
Comparators (Comp)												Comments								
<ul style="list-style-type: none"> ▪ A DMARD (e.g. methotrexate, azathioprine, cyclosporin, penicillamine, sulphasalazine and gold preparations) ▪ Best supportive care if DMARDs not tolerated (e.g. NSAIDs, corticosteroids) ▪ Interventions compared with each 			Yes ↓ Next Q			Unclear ↓ Next Q			No ↓ Exclude			Note what the comparator is								

other				
Outcomes				
Any one or more from the list below: Disease activity; Disease flares; Physical function; Joint damage; Pain Corticosteroid reducing regimens; Extra-articular manifestations (e.g. uveitis); Body weight and height; Mortality; Adverse effects of treatment; HR-QoL.	Yes ↓ Next Q	Unclear ↓ Next Q	No ↓ Exclude	
Abstracts/conference presentations				
Published with sufficient detail to allow appraisal of methodology and assessment of results	Yes ↓ Make final decision	Unclear ↓ Make final decision	No ↓ Exclude	
Final Decision	INCLUDE	UNCLEAR (Discuss)	EXCLUDE	

Appendix 4 Table of excluded and unclear studies from systematic review of clinical-effectiveness

Excluded study	Primary reason for exclusion (comments)
Amarilyo G, Tarp S, Foeldvari I, Cohen N, Pope TD, Woo JMP <i>et al.</i> Efficacy and safety of biologic agents in patients with poly-articular juvenile idiopathic arthritis: Network meta-analysis of randomized controlled withdrawal trials. <i>Arthritis and Rheumatism</i> 2013;65(S10):S922-S923.	Design (NMA)
Anink J, Otten MH, Spronk S, van Suijlekom-Smit LW. Efficacy of Biologic Agents in Juvenile Idiopathic Arthritis: A Systematic Review Using Indirect Comparisons. <i>Arthritis & Rheumatism</i> 2012;64(S10):S490.	Design (SR and indirect comparison)
Canadian Agency for Drugs and Technologies in Health. Common drug review: clinical review report for tocilizumab (Actemra, intravenous) for the treatment of signs and symptoms of active polyarticular juvenile idiopathic arthritis. 2014. Available at: http://www.cadth.ca/media/cdr/clinical/SR0343_Actemra%20pJIA_CL_Report_e.pdf .	Design (SR)
Cummins C, Connock M, Fry-Smith A, Burls A. A systematic review of effectiveness and economic evaluation of new drug treatments for juvenile idiopathic arthritis: etanercept. <i>Health Technology Assessment</i> 2002;6(17):1-43.	Design (SR and economic evaluation)
Decelle K, Horton ER. Tocilizumab for the treatment of juvenile idiopathic arthritis. [Review]. <i>Annals of Pharmacotherapy</i> 2012;46(6):822-9.	Design (SR)
Foster CS, Tufail F, Waheed NK, Chu D, Miserocchi E, Baltatzis S <i>et al.</i> Efficacy of etanercept in preventing relapse of uveitis controlled by methotrexate. <i>Arch.Ophthalmol.</i> 2003;121(4):437-40.	Population (adults)
Gartlehner G, Hansen RA, Jonas BL, Thieda P, Lohr KN. Biologics for the treatment of juvenile idiopathic arthritis: A systematic review and critical analysis of the evidence. <i>Clinical Rheumatology</i> 2008;27(1):67-76.	Design (SR)
Kemper AR, Van Mater HA, Coeytaux RR, Williams JW, Jr., Sanders GD. Systematic review of disease-modifying antirheumatic drugs for juvenile idiopathic arthritis. [Review]. <i>BMC Pediatrics</i> 2012;1229.	Design (SR)
Kingsbury D, Quartier P, Arora V, Kalabic J, Kupper H, Mozaffarian N. Safety and Effectiveness of Adalimumab in Children with Polyarticular Juvenile Idiopathic Arthritis Aged 2 to < 4 Years Or >= 4 Years Weighing < 15 Kg. <i>Annals of the Rheumatic Diseases</i> 2013;72(S3):A729.	Design
Kingsbury D, Quartier P, Arora V, Kalabic J, Kupper H, Mozaffarian N. PRoS-FINAL-2161: Safety and effectiveness of adalimumab in children with polyarticular juvenile idiopathic arthritis aged 2 to <4 years or >=4 years weighing <15 kg. <i>Pediatric Rheumatology</i> 2013;11(S2):P173.	Design

Kingsbury DJ, Quartier P, Arora V, Kalabic J, Kupper H, Mozaffarian N. Safety and Effectiveness Of Adalimumab In Children With Polyarticular Juvenile Idiopathic Arthritis Aged 2 To < 4 Years Or >= 4 Years Weighing < 15 Kg. <i>Arthritis and Rheumatism</i> 2013;65:S117.	Design
Maneiro JR, Salgado E, Gomez-Reino JJ. Immunogenicity of monoclonal antibodies against tumor necrosis factor used in chronic immune-mediated Inflammatory conditions: systematic review and meta-analysis. [Review]. <i>JAMA Internal Medicine</i> 2013;173(15):1416-28.	Design (SR and MA)
Martini A. Etanercept improves active polyarticular juvenile rheumatoid arthritis. <i>Clinical & Experimental Rheumatology</i> 2001;19(2):122-4.	Design (commentary)
Mease P, Genovese MC, Gladstein G, Kivitz AJ, Ritchlin C, Tak PP <i>et al.</i> Abatacept in the treatment of patients with psoriatic arthritis: results of a six-month, multicenter, randomized, double-blind, placebo-controlled, phase II trial. <i>Arthritis & Rheumatism</i> 2011;63(4):939-48.	Population (adults)
Mori M, Takei S, Imagawa T, Imanaka H, Nerome Y, Kurosawa R <i>et al.</i> Etanercept in the treatment of disease-modifying anti-rheumatic drug (DMARD)-refractory polyarticular course juvenile idiopathic arthritis: experience from Japanese clinical trials. <i>Modern Rheumatology</i> 2011;21(6):572-8.	No comparator
Mori M, Takei S, Imagawa T, Imanaka H, Nerome Y, Higuchi R <i>et al.</i> Safety and efficacy of long-term etanercept in the treatment of methotrexate-refractory polyarticular-course juvenile idiopathic arthritis in Japan. <i>Modern Rheumatology</i> 2012;22(5):720-6.	Design (open-label part)
Otten MH, Anink J, Spronk S, van Suijlekom-Smit LWA. Efficacy of biological agents in juvenile idiopathic arthritis: A systematic review using indirect comparisons. <i>Annals of the Rheumatic Diseases</i> 2013;72(11):1806-12.	Design (review)
Pato E, Munoz-Fernandez S, Francisco F, Abad MA, Maese J, Ortiz A <i>et al.</i> Systematic Review on the Effectiveness of Immunosuppressants and Biological Therapies in the Treatment of Autoimmune Posterior Uveitis. <i>Seminars in Arthritis and Rheumatism</i> 2011;40(4):314-23.	Design (review)
Sawyer L, Diamantopoulos A, Brunner HI, Benedetti F, Ruperto N, Dejonckheere F <i>et al.</i> PRoS-FINAL-2070: Efficacy of biologic treatments in juvenile idiopathic arthritis with a polyarticular course: An indirect comparison. <i>Pediatric Rheumatology</i> . 2013;11(S2):P82.	Design (indirect comparison)
Sawyer L, Diamantopoulos A, Brunner H, De Benedetti F, Ruperto N, Dejonckheere F <i>et al.</i> Efficacy of Biologic Treatments in Juvenile Idiopathic Arthritis with A Polyarticular Course: An Indirect Comparison. <i>Annals of the Rheumatic Diseases</i> 2013;72(S3):740-1.	Design (indirect comparison)
Sawyer L, Diamantopoulos A, Brunner HI, De BF, Ruperto N, Dejonckheere F <i>et al.</i> Efficacy of biologic treatments in juvenile idiopathic arthritis with a polyarticular course: An indirect comparison. <i>Arthritis and Rheumatism</i> 2013;65(S10):S119.	Design (indirect comparison)
Simonini G, Druce K, Cimaz R, Macfarlane GJ, Jones GT. Current Evidence of Anti-Tumor Necrosis Factor alpha Treatment Efficacy in Childhood Chronic Uveitis: A Systematic Review and Meta-Analysis Approach of Individual Drugs. <i>Arthritis care & research</i>	Design (review)

2014;66(7):1073-84.	
Simonini G, Katie D, Cimaz R, Macfarlane GJ, Jones GT. Does switching anti-TNFalpha biologic agents represent an effective option in childhood chronic uveitis: The evidence from a systematic review and meta-analysis approach. <i>Seminars in Arthritis and Rheumatism</i> 2014;44(1):39-46.	Design (review)
Ungar WJ, Costa V, Burnett HF, Feldman BM, Laxer RM. The use of biologic response modifiers in polyarticular-course juvenile idiopathic arthritis: A systematic review. <i>Seminars in Arthritis and Rheumatism</i> 2013;42(6):597-618.	Design (review)
Wallace CA, Giannini EH, Spalding SJ, Hashkes PJ, O'Neil KM, Zeff AS <i>et al.</i> The effects of early aggressive therapy in JIA: Results of the TREAT study. <i>Pediatric Rheumatology</i> 2012;10(S1):32.	Abstract (methods)
Wallace CA, Giannini EH, Spalding SJ, Hashkes PJ, O'Neil KM, Zeff AS <i>et al.</i> Trial of early aggressive therapy in polyarticular juvenile idiopathic arthritis. <i>Arthritis & Rheumatism</i> 2012;64(6):2012-21.	Unclear population
Wallace CA, Giannini EH, Spalding SJ, Hashkes PJ, O'Neil KM, Zeff AS <i>et al.</i> Predictors and sustainability of clinical inactive disease in polyarticular juvenile idiopathic arthritis given aggressive therapy very early in the disease course. <i>Arthritis and Rheumatism</i> 2013;65(10):S334-S335.	Abstract (methods)
Wallace CA, Giannini EH, Spalding SJ, Hashkes PJ, O'Neil KM, Zeff AS <i>et al.</i> Clinically inactive disease in a cohort of children with new-onset polyarticular juvenile idiopathic arthritis treated with early aggressive therapy: time to achievement, total duration, and predictors. <i>Journal of Rheumatology</i> 2014;41(6):1163-70.	Unclear population
Wallace CA, Bonsack J, Spalding SJ, Brunner H, O'Neil KM, Milojevic D <i>et al.</i> Results Of a 24 Month Extension Study In PatientsWhoParticipated In The Trial Of Early Aggressive Therapy In Polyarticular Juvenile Idiopathic Arthritis. <i>Arthritis & Rheumatism</i> 2013;65 (S10, Sp. Iss. SI):S116.	Abstract (methods)

MA, meta-analysis. NMA, network metal-analysis. SR, Systematic review.

Unclear studies

Smith JA, Thompson DJ, Whitcup SM, Suhler E, Clarke G, Smith S *et al.* A randomized, placebo-controlled, double-masked clinical trial of etanercept for the treatment of uveitis associated with juvenile idiopathic arthritis. *Arthritis & Rheumatism* 2005;53(1):18-23.

Appendix 5 Clinical-effectiveness data extraction tables

Data extraction - Abatacept

Reference and design	Intervention and Comparator	Participants	Outcome measures
<p><i>Study identifier:</i> Ruperto 2008⁵⁷, Ruperto 2010⁵⁸, Ruperto 2010⁵⁹, Lovell 2012⁶⁰</p> <p><i>Study acronym:</i> AWAKEN (Abatacept Withdrawal study to Assess efficacy and safety in Key Endpoints)</p> <p><i>Study design:</i> withdrawal RCT (4 months open-label lead-in phase, 6 months double- blind randomised phase, open-label extension phase)</p> <p><i>Country or countries:</i> Europe (none appear from UK), Latin America, USA.</p> <p><i>Number of centres:</i> 45</p> <p><i>Recruitment dates:</i> February 2004 - June 2006 (date of last treatment, recruitment likely finished before then)</p> <p><i>Funding:</i> Bristol-Myers Squibb</p>	<p><i>Intervention:</i> 4 month open-label lead- in phase (days 1–113) abatacept (ABA) (10 mg/kg according to weight max dose 1000 mg) on days 1, 15, 29, 57, and 85.</p> <p><i>Double-blind phase:</i> Abatacept (ABA) given at doses of 10 mg/kg at randomisation and at about 28-day intervals thereafter for 6 months (days 114–283), or until a flare of arthritis</p> <p><i>Comparator:</i> Matching placebo</p> <p><i>Other interventions used:</i> All disease-modifying antirheumatic drugs except MTX (stable dose) withdrawn and prohibited during the trial (wash-out period of at least 4 weeks for any DMARD other than MTX, before the first dose of study medication). Oral corticosteroids were stabilised 4 weeks before enrolment. Non-steroidal anti- inflammatory drugs or analgesics permitted for pain control. Folic acid or folic acid permitted. 140/190 (74%) received MTX concomitantly.</p>	<p><i>(Open-label (OL) lead-in phase, number enrolled:</i> <i>n =190. Those achieving ACR Pedi 30 response randomised in double blind phase).</i> (NB. Limited data extracted for this phase)</p> <p><i>Double-blind withdrawal phase</i> <i>Number of randomised participants:</i> ABA: n= 60 Placebo: n= 62</p> <p>Open-label extension (OLE) study (up to day 1681 (year 5.5) efficacy, and up to 7 years safety)^{59;60}</p> <p>Non-responders to ABA during OL phase: n=36 ABA treated patients in double blind phase: n=58 Placebo treated patients in double blind phase: n=59 Total in OLE: n=153</p> <p><i>Inclusion criteria:</i> Juvenile idiopathic arthritis (extended oligoarticular, polyarticular positive or negative for rheumatoid factor, or systemic without systemic manifestations); Aged 6–17 years; At least 5 active joints (those with swelling or, in the absence of swelling, limited range of motion, accompanied by either pain or tenderness) and active disease (at least 2 active joints and 2 joints with a limited range of motion); Inadequate response to, or intolerance to, at least one DMARD</p>	<p><i>Primary outcome(s):</i> Time to disease flare</p> <p><i>Secondary outcomes:</i> proportion of patients at the end of six months double-blind phase who had disease flare; changes from baseline in each of the 6 ACR core variables; pain; assessment of safety and tolerability; HRQoL. (sleep and missed school days reported but not extracted here)</p> <p><i>Method of assessing outcomes:</i> Disease flare defined as worsening of 30% or more in at least 3 of the 6 ACR core-response variables for JIA, and at least 30% improvement in no more than 1 variable during the double-blind period. If a global assessment by either physician or parent was used, flare was defined as a worsening of 20 mm or more on the 100 mm visual analogue scale. If the number of active joints or joints with limited range of motion was used for assessment, it was defined as worsening in 2 or more joints.</p> <p>Improvement defined as an improvement of 30% or more in at least 3 of 6 ACR core-response variables and at least 30% worsening in not more than 1 variable. Improvements were also defined by 50%, 70%, and 90% improvements in the ACR paediatric criteria.</p> <p>Child Health Assessment Questionnaire (CHAQ) used to assess physical, emotional, and social aspects of HRQoL. Higher scores indicate better HRQoL, 0- 100 scale. Childhood Health Assessment Questionnaire (CHAQ) disability index is scored 0 to 3, with a higher score indicating greater disability. CHQ used to assess pain on 100mm visual analogue scale. Higher score indicates more severe pain.</p> <p><i>Length of follow-up:</i></p>

		including biological agents (e.g. etanercept, infliximab, & adalimumab). (Previous anti-TNF therapy reported in 57/190 patients during open-label lead in) <i>Exclusion criteria:</i> Active uveitis, major concurrent medical conditions, or were pregnant or lactating.	End of double-blind period (day 169), plus assessments made for OLE at ≥21 months (day 589) (efficacy and safety), and at day 1681 (study year 5.5 - efficacy and safety, and study year 7 (safety)). It is presumed that these time points are in relation to the start of the OL lead-in.	
Baseline characteristics (double-blind period)		ABA (n=60)	Placebo (n=62)	Comments
Age mean years (SD)		12.6 (3)	12.0 (3)	
Sex female, n (%)		43 (72)	45 (73)	
Ethnic origin, n (%)				
White		46 (77)	49 (79)	
Black		5 (8)	4 (7)	
Other		9 (15)	9 (15)	
Type of JIA				
Persistent oligoarthritis		0	2 (3)	
Extended oligoarthritis		9 (15)	7 (11)	
Polyarthritis (RF +ve)		14 (23)	12 (19)	
Polyarthritis (RF -ve)		26 (43)	28 (45)	
Systemic		11 (18)	12 (19)	
Rheumatoid factor +ve, n (%)		19 (32)	12 (19)	
Rheumatoid factor -ve, n (%)		41 (68)	50 (81)	
Duration of JIA mean years (SD)		3.8 (3.7)	3.9 (3.5)	
Previous anti-TNF therapy discontinued, n (%)		8 (13)	13 (21)	
Lack of efficacy		7 (12)	11 (18)	
For financial reasons		1 (2)	2 (3)	
Results (for double-blind period ⁵⁷ , unless otherwise stated)				
Primary Outcome		ABA (n=60)	Placebo (n=62)	p-value
Time to flare (median, months)		Not reached	6	0.0002
Comments: Kaplan-Meier survival curves are presented, but the survival probabilities can only be read off from the curves and have not been extracted here. Inter-quartile range could not be calculated for the placebo group as there were too few events.				
Secondary Outcomes		ABA (n=60)	Placebo (n=62)	p-value
Disease activity, n (%) ¹				
ACR Pedi 30		49 (82)	43 (69)	0.1712
ACR Pedi 50		46 (77)	32 (52)	0.0071
ACR Pedi 70		32 (53)	19 (31)	0.0185
ACR Pedi 90		24 (40)	10 (16)	0.0062
Inactive disease ²		18 (30)	7 (11)	0.0195
Disease flares, n (%)		12 (20)	33 (53)	0.0003
Disease flares, hazard ratio		0.31 (95% CI 0.16 to 0.59)		NR
Core-response variables, mean (SD)				
Physician's global assessment (VAS: 100mm)		14.7 (18.9)	23.2 (21.8)	0.0004
Parent's global assessment (VAS: 100mm)		17.9 (22.2)	23.9 (21.6)	0.6992
Physical function (CHAQ disability index: 0-3)		0.8 (0.9)	0.8 (0.7)	0.0388
No. of active joints		4.4 (7.0)	6.0 (5.8)	0.0245
No. of joints with limited range of motion		8.8 (12.8)	8.6 (12.0)	0.0128
ERS (mm per hour)		25.1 (26.4)	30.7 (30.1)	0.9562
CRT (mg/L)		0.16 (0.25)	0.29 (0.54)	0.0255
Pain (mean parent global assessment of pain, CHAQ 100mmVAS)		15 ³	21 ³	0.105
Corticosteroid reducing regimens		NR		

Extra-articular manifestations	NR		
Body weight and height	NR		
Mortality	NR		
Health-related quality of life ⁵⁸³			
CHQ Physical summary score	43.6	41 ⁴	p=0.666
CHQ Psychosocial summary score	51.7	47 ⁴	p=0.056
Adverse Events (for double-blind period, unless otherwise stated)(number & % of patients experiencing event)			
Total serious adverse events (SAEs), n (%)	0	2 (3)	0.50
Total serious adverse events, OLE ⁵ , n (%)	23 (15)		
Total adverse events, ⁶ n (%)	37 (62)	34 (55)	0.47
Infections and infestations, n (%)	27 (45%)	27 (44%)	1.00
Gastrointestinal disorders, n (%)	10 (17%)	9 (15%)	0.81
General disorders & administration site conditions, n (%)	4 (7%)	9 (15%)	0.24
Nervous system disorders, n (%)	3 (5%)	2 (3%)	0.68
Respiratory, thoracic & mediastinal disorders, n (%)	6 (10%)	3 (5%)	0.32
<p>Comments:</p> <p>P-values for the core-response variables were based on the difference in the adjusted mean % change from day 113 to day 282.</p> <p>¹ NB. After 6 months of double-blind treatment, or at the time of flare for patients who did not complete this period. In addition to the ACR Pedi overall response, data for the respective 6 ACR Pedi core response variables are reported at the start and end of the double blind period. Only the number of active joints, number of joints with limited range of motion and the CHAQ disability index (physical function) are data extracted here.</p> <p>² Defined as no joints with active arthritis, a physician's assessment of 10 or less on a 100 mm visual analogue scale, and a normal erythrocyte sedimentation rate.</p> <p>³ Read off from graph by reviewer. Number of patients in the trial arms not clear. Abatacept-treated patients (n=52) had improved scores for 14 of the 15 subscales and placebo-treated patients (n=34) for 6 of the 15 subscales (p >0.05 for abatacept versus placebo for all subscales; details not data extracted).</p> <p>⁴ Abatacept treated patients (n=52) had improved scores for 14 of the 15 subscales of the CHQ from start to end of double blind period (P >0.05 for abatacept versus placebo for all subscales). Placebo treated patients (n=34) had improved scores for 6 of the 15 subscales.</p> <p>⁵ SAEs during the OLE (by day 589) occurred in 23/153 patients including an arthritis flare (n =6), arthralgia (n =2), foot deformity (n =2), pyrexia (n =2), and vomiting (n=2). At 7 year follow up 30/153 (19.6%) patients had SAEs. Most were unrelated and were primarily musculoskeletal or infectious events. The incidence rate (per 100 patient-years) of SAEs in the OLE (5.6/100 patient years) did not increase versus the 6 month double blind rate (6.8/100 patient years).</p> <p>⁶ Adverse events that occurred in at least 5% of patients in the open-label and double-blind phases.</p>			
Results OLE			
Original group sizes in double-blind phase	ABA (n=60)	Placebo (n=62)	
ACR Pedi Outcomes OLE at day 589⁵⁹	Received ABA in double blind phase (n= 51)	Received Placebo in double blind phase (n= 47)	
ACR Pedi 30	46/51 (90%)	41/47 (87%)	
ACR Pedi 50	45/51 (88%)	39/47 (83%)	
ACR Pedi 70	38/51 (75%)	35/47 (75%)	
ACR Pedi 90	29/51 (57%)	19/47 (40%)	
ACR Pedi 100	20/51 (39%)	9/47 (19%)	
Inactive disease ¹	22/51 (43%)	11/47 (23%)	
<p>Comments: Patients treated with abatacept during the double-blind phase had in total (lead-in, double-blind & OLE phases) received continuous abatacept therapy for a minimum of 31 months (those recruited to the study earliest had been treated longer, the maximum was 52 months at the time of database lock) whereas those who received placebo during the double blind phase usually received abatacept for a shorter period (length not stated). An analysis according to prior exposure to biologic agents, ACR-Pedi data for those in the OLE who had not taken part in the double-blind phase and information on anti-abatacept and anti-CTLA-4 antibody production is presented but has not been extracted.</p> <p>¹ Inactive disease was defined as having no joint with active disease, a physician's global assessment of disease severity score <10 mm, and an ESR ≤ 20 mm/hour.</p>			
Methodological comments			
<ul style="list-style-type: none"> <i>Allocation to treatment groups:</i> Patients randomly assigned (1:1) to receive either abatacept or placebo. The 			

<p>sequential number for each patient was allocated according to a computer-generated randomisation schedule.</p> <ul style="list-style-type: none"> • <i>Blinding</i>: The main phase of the trial was described as double-blind. Responder and flare status were determined by independent blinded evaluators at the coordinating centres. • <i>Comparability of treatment groups</i>: Appear similar on most variables, though placebo group had greater proportion of rheumatoid factor negative patients than ABA group (81% versus 68%). • <i>Method of data analysis</i>: Kaplan–Meier survival curves used to estimate the distribution of time to disease flare for each group in the 6-month double-blind phase. Log-rank test used to compare the time to disease flare between groups. A Cox proportional-hazards model, with treatment as the only covariate, was used to compare the hazard ratio and 95% CIs for flare of arthritis between the two groups. Missing values in the double-blind phase imputed with the last-observation carried forward method in the analysis of the individual components of the 6 ACR paediatric response variables, the ACR responses, and inactive disease status. HRQoL analysis (CHQ) based on available data at each time point. • <i>Sample size/power calculation</i>: estimated to need 200 patients into the open-label phase to have a sufficient sample size to compare the time to flare over 6 months between the abatacept and placebo groups (with two-sided log-rank tests at 5% significance). Assuming that 64% of patients would respond to treatment (based on experience with rheumatoid arthritis in adults), a sample size of 128 patients would yield 95% power to detect a difference of 35%, assuming a flare rate of 65% in placebo controls and a dropout rate of 10% for the double-blind phase. (The actual flare rate for placebo was 53% and the drop-out rate was 34%, with a difference of 33% between abatacept and placebo in percentage of patients experiencing a flare). • <i>Attrition/drop-out</i>: 42 (34%) patients discontinued during the double-blind period (31 (50%) in the placebo group, and 11 (18%) in the abatacept group); all but one (abatacept-treated patient) did so because the treatment was not effective. Eight patients (2 ABA, 6 placebo) did not receive treatment according to protocol during the double-blind phase but were included in end-point analysis.
<p>General comments</p> <ul style="list-style-type: none"> • <i>Generalisability</i>: Results applicable to patients aged 6 to 17 years with JIA (extended oligoarthritis, polyarthritis, or systemic without systemic manifestations) with an inadequate response to, or intolerance to, at least one DMARD (including biological agents), receiving background MTX. • <i>Outcome measures</i>: Appear appropriate. • <i>Inter-centre variability</i>: Not reported, but to minimise variability in joint assessments each centre had at least 2 certified joint assessors who underwent specific and standardised joint assessment training. • <i>Conflict of interests</i>: The first two authors have received funding for research activity from a variety of pharmaceutical companies including Bristol-Myers Squibb, though have not received funding from companies as personal contribution for assistance during the trial. Three other authors are employees of Bristol-Myers Squibb.

NR = not reported; VAS = visual analogue scale

Quality criteria (Cochrane Collaboration Risk of Bias tool) RCTs¹⁶⁹

Criteria	Judgement ¹	Support for judgement
Random sequence generation (selection bias)	Yes	Computer generated randomisation sequence
Allocation concealment (selection bias)	Yes	Centres were informed of the random allocation of patients by an interactive voice-randomisation system run by the central drug management group
Blinding of participants and personnel (performance bias)	Yes	Double blind phase of the study (after open-label lead in)
Blinding of outcome assessment (detection bias)	Yes	Responder and flare status determined by independent blinded evaluators at the coordinating centres
Incomplete outcome data addressed (attrition bias)		Larger proportion of patients dropped out of the placebo group in the double blind phase than the ABA group (50% versus 18%). Main reason for drop-out was lack of efficacy.
ACR responses, inactive disease status	Yes	Missing values imputed with LOCF method
HRQoL ⁵⁸	No	Analyses based on available data (observed analysis)
Selective reporting (reporting bias)	Yes	All outcomes reported on.
Other sources of bias	Unclear	Inter-centre variability not discussed

¹Yes (low risk of bias); No (high risk of bias); Unclear (uncertain risk of bias).

Data extraction – Adalimumab

Reference and design	Intervention and Comparator	Participants	Outcome measures
<p><i>Study identifier:</i> Lovell 2008⁶¹, Lovell 2012⁶², Ruperto 2013⁶³ & 2014⁶⁴</p> <p><i>Study acronym:</i> None</p> <p><i>Study design:</i> medication- withdrawal RCT (16 weeks randomised open- label, 32 weeks double-blind randomised withdrawal phase, open-label extension phase)</p> <p><i>Country or countries:</i> not specially stated but appear to be Belgium, Czech Republic, France, Germany, Italy, Spain, Slovak Republic and USA.</p> <p><i>Number of centres:</i> 31 (not specified by country)</p> <p><i>Recruitment dates:</i> 19/9/2002- 13/1/2005</p> <p><i>Funding:</i> research grant from Abbott Laboratories</p>	<p>(16 weeks open-label phase: 24 mg of adalimumab (ADA) per square metre (max. of 40 mg) subcutaneously every other week)</p> <p>Licence indication: polyarticular onset JIA – presumed to be the same as polyarticular-course JIA</p> <p><u>Double-blind phase:</u> ADA: as per open- label phase for 32 weeks</p> <p><i>Comparator:</i> placebo</p> <p><i>Other interventions used:</i> MTX: ≥10 mg per square metre per week for 3-month prior screening, same dosage during open- label lead-in and double-blind phase.</p> <p>No MTX: have never received MTX or had discontinued it ≥2 weeks prior to study drug.</p> <p>Stable dosages of NSAIDs and low-dose corticosteroids (≤0.2 mg of prednisone or prednisone equivalent per kilogram of body weight per day to a max. of 10 mg per day) were permitted. Pain medications were allowed except for the 12 hours preceding assessment of the joints.</p>	<p>(1st randomisation, open- label lead-in phase, number randomised: n =171 MTX: n=85 No-MTX: n=86) (NB. Limited data extracted for this phase)</p> <p>2nd randomisation, double-blind withdrawal phase: No. randomised: n=133 MTX/placebo: n=37 MTX/ADA: n=38 No data for ADA (n=30) and Placebo (n=28) group not receiving MTX extracted.</p> <p>Loss to follow-up: n=4 (5.3%) ADA: n=3 (7.9%) Placebo: n=1 (2.7%)</p> <p><i>Inclusion criteria:</i></p> <ul style="list-style-type: none"> • Age 4 - 17 years • Polyarticular-course JIA (with any type of onset) with active disease (≥ 5 swollen joints and ≥ 3 joints with limitation of motion) and without adequate response to NSAIDs • Either no previous treatment with MTX or previous treatment with MTX and AEs or an inadequate response. <p><u>Had to have an ACR Pedi 30 response at week 16 to enter double-blind phase.</u></p> <p><i>Exclusion criteria:</i></p> <ul style="list-style-type: none"> • Clinically significant deviations in clinically hematologic, hepatic, or renal indicators • Ongoing infection or recent major infection requiring hospitalisation or 	<p><i>Primary outcome(s):</i> % of pts not receiving MTX with disease flares (week 16-48)</p> <p><i>Secondary outcomes:</i> adverse events (AEs)</p> <p><i>Method of assessing outcomes:</i> every 12 weeks.</p> <p>Disease flare (ACR Pedi responses): worsening of ≥30% in ≥3 of the 6 core criteria for JIA and an improvement of ≥30% in ≤1 of the criteria. If the no. of joints with active arthritis was used as a criterion of flare, an increase in the no. of active joints to ≥2 was required if there were no initial active joints or only 1 active joint – same approach if the no. of joints with loss of motion was used as a criterion of flare. If either of the global assessments was used as a criterion of flare, any increase of >30% in the VAS of 0 to 100 was sufficient and no minimum clinically important increase was required (e.g. an increase from 2 - 4 would qualify for use of that criterion in the determination of flare).</p> <p>ACR Pedi criteria: physician’s and patient’s/ parent’s global assessment of overall well-being (both measured with the use of a 100-mm VAS: 0 = no disease activity or “very well” for overall well-being, 100 = most disease activity or “very poor” for overall well-being) the no. of joints with active arthritis (defined as joints with swelling not caused by deformity or joints, in the absence of swelling, with limitation of passive motion accompanied by pain tenderness, or both), the no. of joints with limitation of passive motion, physical function measured by the Disability Index of the Childhood Health Assessment Questionnaire (CHAQ-DI), and a laboratory assessment of inflammation (C-reactive protein concentrations). ACR Pedi 50, 70, 90, and 100 levels of response were evaluated, defined as improvements of 50% or more, 70% or more, 90% or more, and 100%, respectively, in ≥3 of the 6 core criteria for JIA, with worsening of 30% or more in only 1 criterion.</p>

		intravenous antibiotics <ul style="list-style-type: none"> • Recent live or attenuated vaccines • Previously treated with other biologic agents at any time or recent treatment with intravenous immune globulin, cytotoxic agents, investigational agents, DMARDs other than MTX, or corticosteroids administered by the intra-articular, intramuscular or intravenous route. 	Safety: physical examinations, laboratory results, vital signs and AEs. Post-hoc analysis: Clinical outcomes: 27-joint Juvenile Arthritis Disease Activity Score (JADAS27) based on C-creative protein; Functional outcome: CHAQ-DI. Minimal disease activity (MDA) defined as JADAS27 <3.8 and normal function defined as CHAQ-DI <0.5. Higher scores indicate higher disease activity. <i>Length of follow-up:</i> 70 days after last dose for AEs for all patients who discontinued study medication. Those enrolled in the double-blind phase were eligible to receive open-label treatment with ADA in an extension phase of the study (duration not specified).
--	--	--	--

Baseline characteristics	ADA (n=38)	Placebo (n=37)	Comments
Age, mean years (SD)	11.7 (3.3)	10.8 (3.4)	
Age group, n (%)			
4-8 years	6 (16)	12 (32)	
9-12 years	17 (45)	10 (27)	
13-17 years	15 (40)	15 (41)	
Sex – Female, n (%)	30 (79)	30 (81)	
Race, n (%)			Determined by the patient or parent
White	36 (95)	36 (97)	
Black	0	0	
Other	2 (5)	1 (3)	
Body weight, mean kg (SD)	42.1 (17.9)	44.3 (18.9)	
Type of JIA			Reported as poly-articular-course JIA
RF -ve, n/N (%)	27/37 (73)	30/36 (83)	
Duration of JIA, mean years (SD)	4.3 (4.1)	4.0 (3.5)	
Previous medication use, n (%)			
Methotrexate (MTX)	38 (100)	37 (100)	
Other DMARDs	1 (3)	7 (19)	
Methylprednisolone	2 (5)	2 (5)	
Results (double-blind phase, weeks 16-48)			
Primary Outcome	ADA (n=38)	Placebo (n=37)	p-value
Disease flares, n/N (%)	14/ 38 (37)	24/37 (65)	p=0.02
Comments:			
Secondary Outcomes			
Disease Activity			

ACR Pedi response ¹ week 48 (%)					
30	63	38		p=0.03	
50	63	38		p=0.03	
70	63	27		p=0.002	
90	42	27		p=0.17	
Physical function	NR	NR			
Joint damage	NR	NR			
Pain	NR	NR			
Corticosteroid reducing regimens	NR	NR			
Extra-articular manifestations (such as uveitis)	NR	NR			
Body weight and height	NR	NR			
Mortality	See comments AEs				
Health-related quality of life	NR	NR			
Comments: NR, not reported.					
¹ A patient who had a flare according to the protocol definition was classified as having no response (ACR Pedi <30) from that point forward, regardless of the patient's ACR Pedi response at that time.					
Adverse Events, no. of events (no. of events per patient-year)	ADA (+MTX) (n=37; 15 Patient-yrs)	Placebo (+MTX) (n=38; 18.3 Patient-yrs)	p-value		
Any AE	155 (10.3)	234 (12.8)			
Most frequently reported AEs					
Related to injection-site reaction	57 (3.8)	73 (4.0)			
Contusion	7 (0.5)	12 (0.7)			
Nasopharyngitis	6 (0.4)	5 (0.3)			
Upper respiratory tract infection	5 (0.3)	6 (0.3)			
Viral infection	3 (0.2)	7 (0.4)			
Vomiting	2 (0.1)	4 (0.2)			
Excoriation	1 (0.1)	10 (0.6)			
Serious AEs, possibly related to study drug ¹	1 (0.1) - Gastroduodenitis	0			
AEs leading to the discontinuation of the drug	0	0			
Comments: no occurrence of deaths, opportunistic infections, malignant conditions, demyelinating diseases or lupus-like reactions. ¹ Serious adverse events were death or any event that was life-threatening; required hospitalisation or prolongation of existing hospitalisation; resulted in persistent or significant disability, congenital anomaly, or spontaneous or elective abortion; or required medical or surgical intervention to prevent another serious outcome.					
(27/171 (16%) had at least 1 positive test for anti-ADA antibody during the open-label and double-blind phases (MTX: 5/85 (6%), No MTX: 22/86 (26%), but development of anti-ADA antibody did not lead to a greater rate of discontinuation of the study drug, nor did it increase the incidence of serious AEs)					
First 104 weeks of open-label extension phase⁶¹					
ACR Pedi at 104 weeks of OLE, %	OLE phase (n=128)¹				
30	89% ²				
50	86% ²				
70	77% ²				
90	59% ²				
100	40%				
Comments:					
¹ Only 71/128 (55%) of this group received methotrexate during the open-label and double-blind phases of the study and meet the licenced indication for adalimumab (ie. the 128 includes participants not receiving methotrexate in the two study arms that do not meet the licenced indication).					
² Data extracted from figure using Engauge software. Data available for earlier time points in OLE (weeks 8, 16, 24, 56, 104) but not data extracted. For missing values the last observation was carried forward.					
Post-hoc analysis - OLE,⁶⁴ n(%)	ADA		Placebo		p-value²
	week 48	week 88	week 48	week 88	
Minimal disease activity ³	19 (76)	26 (83.9)	15 (62.5)	14.0 (50.0)	
Minimal disease activity ³ with normal	17 (68.0)	24 (77.4)	15 (62.5)	14 (50.0)	

function ⁴					
<p>Comments: ² statistical comparison between ADA with or without MTX vs Placebo with or without MTX only. Post-hoc analysis on growth in patients with JIA^{62,63} reported per MTX or non -MTX group only (not data extracted). ³ JADAS27 <3.8. ⁴ CHAQ-DI <0.5.</p> <p>The OLE⁶¹ was ongoing at the time the key effectiveness paper was published and the time period for which events were reported is not clear.⁶¹ Serious AEs considered possibly related to study drug occurred in 7 patients during the OLE (a table in the published paper⁶¹ suggests none were receiving methotrexate, in which case they were not receiving adalimumab treatment according to the licenced indication). Three patients discontinued treatment due to adverse events during the OLE.</p>					
<p>Methodological comments</p> <ul style="list-style-type: none"> • <i>Allocation to treatment groups</i>: randomisation at a 1:1 ratio within patients' previous respective strata (stratified according to MTX use), no further details reported. • <i>Blinding</i>: double-blind (investigators, study coordinators, assessors, patients and parents were unaware of the treatment assignment during the double-blind phase of the study). • <i>Comparability of treatment groups</i>: states no significant differences in baseline characteristics between the Placebo and the ADA group within either stratum (MTX or No MTX) - no statistical comparison between the ADA-MTX and Placebo-MTX group reported. There were some baseline differences between the later treatment groups: the ADA group had a higher percentage of children in the 9-12 year age group compared to placebo; conversely mean body weight was slightly lower. Mean negative for rheumatoid factor was 10% lower in this group compared to placebo, duration of JIA was slightly lower and previous medication use for other DMARDs was 16% lower compared to the placebo group. • <i>Method of data analysis</i>: Efficacy analyses ITT, however this was defined as all patients who received at ≥1 dose of the study drug during the phase of the study for which the analysis was being conducted. For the primary efficacy end point and for all secondary analyses of disease flare, missing values were treated as disease flares. For secondary analyses of ACR Pedi 30, 50, 70, and 90 responses during the open-label lead-in and double-blind phases, missing values were imputed as nonresponses. In addition, patients in whom a flare occurred according to the protocol definition during the double-blind phase were classified as having no response (ACR Pedi <30) at week 48, regardless of their actual ACR Pedi responses. • <i>Sample size/power calculation</i>: assumption of a 70% response rate to ADA reported, requiring 42 patients in the open-label lead-in phase to yield the 29 patients needed for each group in the double-blind phase. This estimate was based on a 40% difference in the rate of flare between the Placebo and the ADA groups and provided a power of 80% at an alpha level of 0.05. However, states that study was not statistically powered to detect differences between patients receiving and those not receiving MTX. • <i>Attrition/drop-out</i>: Double-blind: n=4 (5.3%); ADA, n=3 (withdrew for other reasons, no further details), Placebo, n=1 (withdrew consent). [Loss to follow-up, total: n= 43 (25%); Open-label, all: n=38 (22%); Double-blind all n=5 (3.8%)] 					
<p>General comments</p> <ul style="list-style-type: none"> • <i>Generalisability</i>: limited to polyarticular-course JIA patients aged 4 to 17 years, who have previously received 16 wks ADA treatment and treated with MTX, and had an ACR Pedi 30 response at wk 16. • <i>Outcome measures</i>: appear to be appropriate. • <i>Inter-centre variability</i>: Not discussed. • <i>Conflict of interests</i>: Authors received various financial support and/or unrestricted continued medical education grants from various pharmaceutical companies including the drug manufacturer. States 'no other potential conflict of interests relevant to this article was reported'. Individuals at JK Associates and Abbott Laboratories provided editorial support, while individuals at Abbott Laboratories helped with data management and statistical analysis. 					

Quality criteria (Cochrane Collaboration Risk of Bias tool) RCTs¹⁶⁹

Criteria	Judgement ¹	Support for judgement
Random sequence generation (selection bias)	Unclear	No details reported
Allocation concealment (selection bias)	Unclear	No details reported
Blinding of participants and personnel (performance bias)	Yes	Double-blind, details reported
Blinding of outcome assessment (detection bias)	Yes	Double-blind, details reported
Incomplete outcome data addressed (attrition bias)	Yes	Details reported
Selective reporting (reporting bias)	Yes	All outcomes stated were reported
Other sources of bias	Unclear	Inter-centre variability not discussed

¹ Yes (low risk of bias); No (high risk of bias); Unclear (uncertain risk of bias).

Data extraction – Etanercept

Reference and design	Intervention & Comparator	Participants	Outcome measures
<p><i>Study identifier:</i> Lovell 2000,⁴² Lovell 2003,⁶⁵ Lovell 2006⁶⁶ and Lovell 2008⁶⁷</p> <p><i>Study acronym:</i> none</p> <p><i>Study design:</i> medication-withdrawal RCT (3 months open-label lead-in phase, 4 months double-blind randomised withdrawal phase, open-label extension phase)</p> <p><i>Country or countries:</i> Canada and USA</p> <p><i>Number of centres:</i> not specifically stated, but appears to be 9⁶⁵</p> <p><i>Recruitment dates:</i> not reported</p> <p><i>Funding:</i> Immunex Corporation¹⁶⁹</p>	<p><i>Intervention:</i> 0.4 mg of etanercept per kilogram subcutaneously twice weekly until disease flare occurred or four months elapsed</p> <p><i>(Includes 33% of patients with systemic onset JIA)</i></p> <p><i>Comparator:</i> Placebo</p> <p><i>Other interventions used:</i> Stable doses of NSAIDs, low doses of corticosteroids (≤ 0.2 mg of prednisone per kilogram per day, with a max. 10 mg per day) or both. Pain medications were allowed except during the 12 hrs before a joint assessment. MTX discontinued 14 days and other DMARDs 28 days before receipt of etanercept.</p> <p>Open-label extension (OLE): 0.4mg/kg etanercept twice weekly (max. dose 25 mg per injection) or 0.8 mg/kg once weekly (max. dose of 50 mg/week) subcutaneously.</p>	<p><i>(3 months open-label (OL) lead-in phase: n=69)</i> (NB. Limited data extracted for this phase)</p> <p><i>Double-blind withdrawal phase</i> <i>No. of randomised participants:</i> n = 51 Etanercept: n=25 Placebo: n=26</p> <p>Open-label extension (OLE): n=58</p> <p>Loss to follow-up: [Part 1: n= 5 (7%)] Part 2-RCT: Etanercept: n=6 (24%); Placebo: n=19 (73%) Part 3: OLE: n=38 (66%)</p> <p><i>Inclusion criteria:</i> children aged 4 - 17 years with JIA, with active disease (≥ 5 swollen joints and ≥ 3 joints with limitation of motion and pain, tenderness, or both) despite treatment with NSAIDs and with MTX at doses of at least 10 mg per square metre of body-surface area per week.</p> <p><i>Exclusion criteria:</i> No intra-articular and soft-tissue corticosteroid injections for 1 month before the trial. Patients with major concurrent medical conditions; Pregnant and lactating patients.</p>	<p><i>Primary outcome(s):</i> number of patients with disease flare</p> <p><i>Secondary outcomes:</i> not specifically stated</p> <p><i>Method of assessing outcomes for Part 2:</i> Physical examinations, measures of disease activity and lab. tests (hematologic analysis, serum chemical analysis, and urinalysis) on day 1 (before etanercept or placebo) and day 15 and at the end of each month. Final safety assessments 30 days after discontinuation of study drug for withdrawals or at next scheduled visit if withdrawal due to disease flare. Serum at the end of 7 months for testing for autoantibodies (antinuclear antibodies, antibodies to double-stranded DNA, IgG and IgM anticardiolipin antibodies, and antibodies to extractable nuclear antigens), and on day 1 before the administration of the study drug and at the end of months 7 for testing for antibodies to etanercept.</p> <p>Physician's global assessment of disease severity 0-10 (best-worst); Patient's or parent's global assess of overall well-being - 0-10 (best-worst); Childhood Health Assessment Questionnaire (CHAQ) – 0-3 (best-worst); Erythrocyte sedimentation rate (ESR) - normal ranges 1 to 30 mm per hour for females and 1 to 13 mm per hour for males; Articular severity score - 0 (best) to 962 (worst); Pain – VAS 0 cm (best) to 10 cm (worst); C-reactive protein - normal range is 0 to 0.79 mg per decilitre. Other: 73 joints were evaluated for the total active-joint count; 71 for limitation of motion with pain, tenderness, or both; 66 for swollen joints; and 71 for limitation of motion.</p> <p>Definition of disease flare: Change in the core set of response variables from the beginning of the double-blind study - worsening of $\geq 30\%$ in 3/6 response variables and a minimum of 2 active joints. Could also have improvement of $\geq 30\%$ in no more than 1/6 six response variables. Global assessments, if used to define flare, had to change by at least 2 units on a scale from 0 to 10.</p> <p>Definition of improvement of disease response was based on changes from baseline values at enrolment, whereas flare was measured from beginning of the double-blind study. For example, 28 active joints at baseline, but only 2 active joints at the time of randomisation - a change to 3 active joints would be considered a flare (at least 30 percent worse than the</p>

			condition at the time of randomisation) but would also still be considered improvement (at least 30 percent improved from base line). <i>Length of follow-up:</i> 4 months (double-blind only); OLE: 8 years ⁶⁷
--	--	--	---

Baseline characteristics: double-blind study	Etanercept (n=25)	Placebo (n=26)	Comments
Mean age, year	8.9	12.2	
Age group, n (%)			p<0.02
4-8 years	13 (52)	5 (19)	
9-12 years	5 (20)	4 (15)	
13-17 years	7 (28)	17 (65)	
Sex, n (%)			
Female	19 (76)	15 (58)	
Male	6 (24)	11 (42)	
Ethnicity, n (%)			p<0.02
White	14 (56)	23 (88)	
Black	3 (12)	1 (4)	
Hispanic	6 (24)	2 (8)	
Other	2 (8)	0	
Type of JIA, n (%)			
Pauciarticular	2 (8)	1 (4)	
Polyarticular	14 (56)	17 (65)	
Systemic	9 (36)	8 (31)	
RF +ve, n (%)	4 (16)	8 (31)	
Mean duration of JIA, year	5.3	6.4	
Previous medication, n (%)			
MTX	25 (100)	26 (100)	
DMARDs at washout, n (%)			
Methotrexate	16 (64)	19 (73)	
Hydroxychloroquine	2 (8)	7 (27)	
Concomitant therapy at washout, n (%)			
Corticosteroids	6 (24)	13 (50)	
NSAIDs	25 (100)	24 (92)	
Mean dose of corticosteroids — mg/day	6.5	5.5	

Results: double-blind study

Primary Outcome	Etanercept (n=25)	Placebo (n=26)	p-value
Disease flare, n (%)	7 (28)	21 (81)	p=0.003 ¹
Corticosteroid use at baseline ²			
Yes	3/6 (50)	12/13 (92)	p=0.05
No	4/19 (21)	9/13 (69)	
Time to flare, median days	>116	28	p<0.001

Comments: ¹ p<0.001 after adjustment for baseline characteristics in logistic regression model.

² Authors state that with the exception of corticosteroid use at base line (p=0.05), none of the baseline characteristics were significant predictors of flare rates (p>0.15).

Because 13/25 patients were still receiving etanercept at the end of the study (day 116) without disease flare, the median time to flare was greater than 116 days.

Secondary Outcomes – 7 months, median	Etanercept (n=25)	Placebo (n=26)	p-value
30% improvement, n (%)	20 (80)	9 (35)	p<0.01
50% improvement, n (%)	18 (72)	6 (23)	
70% improvement, n (%)	11 (44)	5 (19)	
JIA core set criteria, median			
Total number of active joints (out of 73 joints)	7.0	13.0	
No. of joints with limitation of motion and with pain, tenderness, or both (out of 71 joints)	1.0	4.5	
Physician's global assessment of disease severity	2	5	
Patient/parent's global assess. of overall well-being	3	5	

Score on CHAQ (disability domain)	0.8	1.2	
ESR	18	30	
Articular severity score, median	38	66	
Duration of stiffness (min), median	5	38	
Pain (on a visual-analogue scale), median	1.5	3.5	
C-reactive protein, median	0.4	3.0	
No. of swollen joints, median	4.0	11.0	
No. of joints with limitation of motion, median	9	22	
Corticosteroid reducing regimens	NR	NR	
Extra-articular manifestations (such as uveitis)	NR	NR	
Body weight and height	NR	NR	
Mortality	See AEs		
Comments: NR, not reported. Authors state that in the double-blind study as compared with the end of the open-label study, a significant proportion of patients who received placebo had shifts from normal levels of C-reactive protein and erythrocyte sedimentation rates to above-normal values ($p \leq 0.003$ for each variable). Last observation carried forward approach for missing data and visits and for early termination (LOCF).			
Adverse Events	Etanercept (n=25)	Placebo (n=26)	p-value
Death, n	0	0	
Urticaria, n	1 ¹	0	
Hospitalisation for serious AEs, n			
Depression and personality disorder	1	0	
Gastroenteritis-flu syndrome	1	0	
Injection-site reactions, n	1	1	
Tested positive for non-neutralising antibody to etanercept, n	2	N/A	
Comments: ¹ after 1 st dose of etanercept (responded to oral antihistamines). Other AEs were reported to be of mild-to-moderate intensity, with no significant difference in the frequency of AEs between the treatment groups. There were no laboratory abnormalities requiring urgent treatment in the etanercept group. No patient had persistent elevations in autoantibodies or had signs or symptoms of another autoimmune disease.			
Baseline characteristics: OPE, 8 year follow-up⁶⁷	OLE (n=58)	8th year of OLE (n=26)	
Age, mean years (SD)	10.4 (3.8)	10.8 (3.9)	
JRA onset type, n (%)			
Pauciarticular	5 (9)	2 (8)	
Polyarticular	34 (58)	19 (73)	
Systemic	19 (33)	5 (19)	
Duration of JRA, mean years (SD)	5.9 (3.2)	6.4 (3.4)	
RF+ve, n (%)	13 (23) (n=56)	6 (24) (n=25)	
Concomitant therapy at enrolment, n (%)			
NSAIDs	56 (97)	25 (96)	
Corticosteroids	22 (38)	8 (31)	
Corticosteroid dosage, mg/day mean (SD)	5.7 (3.2)	4.1 (2.3)	
Comments: After 1 year of the OLE, the dosages and the use of other medications for JRA (including corticosteroids, intra-articular injections of steroids and NSAIDs) could be adjusted or added at the discretion of the treating physician, without restriction. MTX could be added to the regimen (dosage limited to 10–20 mg/m ² /week)			
Results: OLE year 8, mean (SEM)²		Completed year 8 (n=16)	
Total no. of joints with active arthritis (n=11)		2.2 (0.9)	
Total no. of joints with LOM and tenderness and/or pain on motion (n=11)		0	
Total no. of joints with LOM (n=11)		11.8 (4.4)	
Physician's global assessment		1.6 (0.3)	
Patient's/parent's global assessment		2.0 (0.6)	
Pain score		1.8 (0.5)	
CHAQ score (n=11)		0.6 (0.2)	
C-reactive protein ³		1.1 (0.5)	
Comments: ² 74 joints were assessed for tenderness and/or pain on motion, 71 for limitation of motion (LOM), and 66 for swelling. ³ New high-sensitivity method of analysing C-reactive protein levels for year 8 (old method: normal range 0–0.79 mg/dl; new method: normal range 0–0.287 mg/dl).			
ACR Pedi response , 8 years (LOCF), % n/N			

ACR Pedi 30	83 (40/48)			
ACR Pedi 50	77 (36/47)			
ACR Pedi 70	61 (28/46)			
ACR Pedi 90	41 (19/46)			
ACR Pedi 100	18 (8/45)			
Year of etanercept treatment from RCT (excluding gaps between RCT and OLE)	SAE⁴		MII⁵	
	No. of events	No. of events/patient year	No. of events	No. of events/patient year
1 (n=69; 57 patient-years of drug exposure)	5	0.09	2	0.04
9 (n=14; 4 patient-years of drug exposure)	0	0	0	0
Total for all years (n=69; 318 patient-years of drug exposure)	39	0.12	9	0.03
<p>Comments: SAEs defined as events that were fatal or life-threatening, required hospitalisation or prolonged an existing hospitalisation, resulted in a persistent or significant disability or incapacity, or resulted in a congenital anomaly or birth defect.</p> <p>⁴ SAEs occurring during the study or within 30 days of the last dose of etanercept.</p> <p>⁵ Defined as medically important infections resulting in the need for intravenous antibiotic therapy or hospitalisation.</p> <p>Only 1 MII reported by patients since report at 4 years (pyelonephritis). The most common new SAEs reported beyond 4 years of drug exposure were a flare or worsening of disease [6/9 SAEs (67%)].</p>				
Methodological comments				
<ul style="list-style-type: none"> • <i>Allocation to treatment groups:</i> A blocked randomisation scheme with stratification according to study centre and number of active joints (≤ 2 vs. > 2) at the end of month 3 (in the open-label study) was used to assign patients to their treatment group. • <i>Blinding:</i> Double-blind (no further details). • <i>Comparability of treatment groups:</i> Authors state that the groups were well balanced in the double-blind study, except for age group and race ($p < 0.02$) and corticosteroid use at baseline ($p = 0.05$) and that the unequal randomisation did not affect the study results. The etanercept group had a significantly higher number of younger patients compared to the placebo group (4-8yr old: E 52% vs 19% placebo) and a greater ethnic mix (white: E 56% vs 88% placebo), while the placebo group had a significantly larger use of corticosteroid use at washout (placebo 50% vs 24% E). • <i>Method of data analysis:</i> Statistical methods employed were reported. All tests were two-sided, with a significance level of 0.05. Patients who withdrew early without disease flare were counted in the analysis with those who continued to have a response - a LOCF approach was used for missing data and visits and for early termination. To evaluate any bias introduced by the withdrawal assumption in the primary analysis, an analysis of time to flare (by the log-rank test) was undertaken in which data on patients who withdrew without flare were censored at the time of withdrawal. The effect of baseline characteristics on flare rates was assessed by main-effects logistic regression. <p>OLE: Data from patients who reached the age of 18 and discontinued the study and who therefore no longer had valid childhood efficacy measures were not included in efficacy analysis (summary of the last visit using the LOCF method). Adult-specific measures of disease for patients ≥ 18 years of age were not included in analyses (n=5 each at years 7 and 8).</p> <ul style="list-style-type: none"> • <i>Sample size/power calculation:</i> none reported. • <i>Attrition/drop-out:</i> Part 1 OL: 64/69 (93%) urticaria with the 1st dose of etanercept n=1; refusal of treatment n=2; lack of response n=2. Part 2 - RCT: Etanercept 6/25 (24%): disease flare n=6; Placebo: 19/26 (73%): parental refusal to allow continuation n=1, disease flare n=18. Part 3 OLE: 38/58 (66%): lack of efficacy n=7 (12%); AEs n=4 (7%); physician decision n=5 (9%); protocol issue n=3 (5%); lost to follow-up n=3 (5%); patient/guardian refusal n=5 (9%); other n= 8 (14%). 36% patients (n=21) discontinued during the first 4 years. 				
General comments				
<ul style="list-style-type: none"> • <i>Generalisability:</i> limited to pauciarticular, polyarticular and systemic onset JIA patients aged 4 to 17 years, who did not tolerate or had an inadequate response to MTX and had received 3 months of etanercept treatment. • <i>Outcome measures:</i> appear to be appropriate. • <i>Inter-centre variability:</i> not discussed. • <i>Conflict of interests:</i> 2 authors had served as ad hoc consultants to Immunex. 				

Quality criteria (Cochrane Collaboration Risk of Bias tool) RCTs¹⁶⁹

Criteria	Judgement¹	Support for judgement
Random sequence generation (selection bias)	Unclear	Blocked randomisation scheme with stratification, no details about how randomisation was performed.
Allocation concealment (selection bias)	Unclear	No details reported
Blinding of participants and personnel (performance bias)	Yes	Double-blind phase of study (after open-label lead in)
Blinding of outcome assessment (detection bias)	Yes	Paper states that study site-staff who were not involved in patient assessments constituted the contents of the vials (etanercept or placebo).
Incomplete outcome data addressed (attrition bias)	Yes	Details reported, but drop-outs are nearly three times higher in the placebo group. Incomplete data appears to have been address with the LOCF method (used for missing data/visits and early terminations).
Selective reporting (reporting bias)	Yes	All outcomes reported on
Other sources of bias	Unclear	Inter-centre variability not discussed

¹ Yes (low risk of bias); No (high risk of bias); Unclear (uncertain risk of bias).

Data extraction – Tocilizumab

Reference and design	Intervention and Comparator	Participants	Outcome measures
<p><i>Study identifier:</i> Brunner 2014,⁶⁸ Brunner 2014,⁶⁹ Baildam 2014,⁷⁰ Baildam 2013,⁷³ DeBenedetti 2013,⁷² DeBenedetti 2013,⁷⁴ Brunner 2013,⁷¹ and Brunner 2012⁷⁵</p> <p><i>Study acronym:</i> CHERISH</p> <p><i>Study design:</i> medication-withdrawal RCT (16-week randomised open-label, 24-week double-blind randomised withdrawal phase, open-label extension phase)</p> <p><i>Country or countries:</i> Australia, Canada, Europe, Latin America, Russia and USA</p> <p><i>Number of centres:</i> 58</p> <p><i>Recruitment dates:</i> 14/10/2009 – 31/1/2011</p> <p><i>Funding:</i> funding for manuscript preparation by H Hoffmann-La Roche Ltd</p>	<p><i>Intervention:</i> intravenous tocilizumab (TCZ) at 8 mg/kg (8 mg/kg for <30 kg group) or 10 mg/kg (10 mg/kg for <30 kg group) every 4 weeks (based on pharmacokinetic modelling and simulation, doses of 10 mg/kg for patients weighing <30 kg achieved TCZ exposure comparable to that of 8 mg/kg for patients weighing ≥30 kg).</p> <p><i>Comparator:</i> Placebo</p> <p><i>Other interventions used:</i> Stable doses of NSAIDs and low-dose glucocorticoids (≤ 0.2 mg/kg/day prednisone; daily max. 10 mg) and MTX (10–20 mg/m² body surface area/ week).</p>	<p><i>(Randomised, open-label (OL) lead-in phase: TCZ every 4 weeks until week 16, n=188 - patients randomised to <30kg TCZ (n=69) or ≥30kg TCZ (n=119) (NB. Limited data extracted for this phase)</i></p> <p><i>Double-blind withdrawal phase</i> <i>No. of randomised participants:</i> n =166 <i>TCZ:</i> n=82: T 10mg/kg <30kg BW: n= 16; T 8mg/kg <30 kg BW: n=11; T 8 mg/kg ≥30kg BW: n=55 <i>Placebo:</i> n= 84: P 10mg/kg <30kg BW: n=15; P 8mg/kg <30kg BW: n=13; P 8 mg/kg ≥30kg BW: n=56</p> <p><i>Loss to follow-up:</i> OL: n=22/188 (11.7%) RCT: TCZ n=3/82 (3.7%), Placebo 3/84 (3.6%). OL-Extension (OLE): n=5/160 (3.1%)</p> <p><i>Inclusion criteria:</i></p> <ul style="list-style-type: none"> • 2-17 years old; • Diagnoses of RF+ve or RF-ve polyarticular-course JIA or extended oligoarticular JIA; • Disease duration ≥6months; • Inadequate responses to or intolerant of MTX; • ≥5 more active joints, with LOM present in ≥3 of the active joints; • No MTX ≥4 weeks before and including baseline visit or had been taking MTX ≥12 weeks immediately before and including baseline visit and on stable dose of 10-20 mg/m² for ≥8 weeks before and including baseline visit together with either folic acid or folinic acid; • No oral glucocorticoids at baseline visit or had been taking oral glucocorticoids at a stable dose for ≥4 weeks before and including baseline visit (n≤10 mg/day or 0.2mg/kg/day); • No NSAIDs at baseline or more than 1 type of NSAID at a stable dose (less than or equal 	<p><i>Primary outcome(s):</i> Proportion of patients in whom a JIA-flare occurred during part 2 (up to and including wk40) compared with wk16</p> <p><i>Secondary outcomes wk40:</i> JIA-ACR 30/50/70/90 responses, change from baseline in JIA-core response variables (CRVs) and clinically inactive disease.</p> <p><i>Method of assessing outcomes:</i> 4 weekly assessments.</p> <p>JIA-ACR30 response: defined as ≥30% improvement of ≥3/6 JIA core response variables (JIA-CRVs) without >30% worsening in ≥1 of the remaining JIA-CRVs compared with baseline. (Part 1. Patients who had at least 1 JIA-ACR30 response entered part 2).</p> <p>Active joints: defined as the presence of swollen joints (or, in the absence of swelling, joints with limitation of movement (LOM) plus pain on motion and/or tenderness with palpation).</p> <p>Clinically inactive disease was defined as PGA, indicating no disease activity plus absence of all the following: joints with active arthritis, uveitis and ESR greater than 20 mm/h.</p> <p>Serious infections were defined in accordance with the definition of serious adverse events (SAEs) in the International Conference on Harmonisation guidelines (reference provided).</p> <p>Physician global assessment (PGA) of disease activity: VAS 0–100 (0=inactive disease); Assessment of patient overall well-being: VAS 0–100 (0=very poor); Physical function measured by</p>

		<p>to the recommended daily dose) ≥ 2 weeks before and including the baseline visit;</p> <ul style="list-style-type: none"> Never been treated with biologics or had been previously treated with biologics and discontinued them for at least the following periods: anakinra: 1 week; etanercept: 2 weeks; rilonacept: 5 weeks; infliximab or adalimumab: 8 weeks; abatacept: 12 weeks; canakinumab: 20 weeks, before and including the baseline visit. <p><i>Exclusion criteria:</i> None in addition to the above.</p>	<p>the Childhood Health Assessment Questionnaire-Disability Index (CHAQ-DI): 0–3 (0=no disability).</p> <p>Patients continued RCT until week 40 unless JIA-flare ($>30\%$ worsening in 3/6 JIA-CRVs without $> 30\%$ improvement in ≥ 1 of the remaining JIA-CRV) compared with week 16. They then entered the OL-extension study.</p> <p><i>Length of F-up:</i> OL 16 weeks; RCT 24 weeks; total 40 weeks. OLE 64 weeks (total 104 weeks).</p>
--	--	---	---

Baseline characteristics ¹	TCZ 8mg/kg <30kg (n=34)	TCZ 10mg/kg <30kg (n=35)	TCZ 10mg/kg ≥ 30 kg (n=119)
Age, years	7.6 (2.71)	6.9 (3.02)	13.1 (2.78)
Sex, females n (%)	24 (71)	30 (86)	90 (76)
Ethnicity	Not reported		
Type of JIA	Not reported		
Weight (kg)	22.4 (5.3)	20.7 (5.7)	50.0 (12.6)
RF +ve, n (%)	2 (6)	4 (11)	48 (40)
Duration of JIA, years	3.5 (2.57)	3.4 (2.39)	4.7 (4.16)
Previous medication, n (%)			
DMARD	26 (76)	21 (60)	87 (73)
Biological agent ²	6 (18)	8 (23)	47 (39)
Joints with active arthritis, n	21.2 (13.6)	23.9 (18.3)	18.9 (13.0)
Joints with LOM, n	17.3 (13.3)	23.1 (19.2)	16.0 (12.7)
Assessment of patient overall well-being, VAS	59.1 (26.2)	51.5 (26.9)	51.6 (24.1)
PGA of JIA activity, VAS	64.7 (18.5)	64.7 (20.5)	59.4 (21.3)
CRP (mg/L) (standard ref. range 0–10 mg/L)	26.6 (33.6)	21.8 (32.3)	22.8 (38.8)
CHAQ-DI score	1.8 (0.68)	1.7 (0.71)	1.2 (0.69)
ESR (mm/h) (standard ref. range 0–18 mm/h)	36.6 (23.0)	35.1 (24.1)	34.2 (26.7)
Concurrent MTX use, n (%)	30 (88)	29 (83)	89 (75)
Dose (mg/m ² /week)	13.8 (2.9)	16.5 (11.1)	11.6 (2.7)
Concurrent glucocorticoid use, n (%) ³	18 (53)	15 (43)	54 (45)
Dose (mg/kg/day) ³	0.15 (0.038)	0.15 (0.033)	0.12 (0.052)

Comments: ¹ all patients randomised in part 1. 15/188 (7.9%) did not achieve JIA-ACR30 response and were not randomised in part 2. ² 9% of patients previously received ≥ 3 biological agents. TNF inhibitors: n=56, anakinra n=5, abatacept n=5, canakinumab n=1. ³ measured in prednisone equivalents.
NB – baseline characteristic not given the TCZ vs placebo groups in Part 2.

Results, week 40 double-blind study ^{68;73;75}			
Primary Outcome	TCZ ⁴ (n=82)	Placebo (n=81)	Difference ⁴ TCZ vs Placebo (95% CI); p value
Proportion with JIA-ACR30 flare (compared with week 16), n (%)	21 (25.6%)	39 (48.1)	-0.21 (-0.35, 0.08); 0.0024
Secondary Outcomes	TCZ ⁴ (n=82)	Placebo (n=81)	Difference ⁴ TCZ vs Placebo (95% CI); p value
Proportion of patients with JIA-ACR30 improvement relative to baseline, n (%)	61 (74.4)	44 (54.3)	0.09 (0.05, 0.33); 0.0084
Proportion of patients with JIA-ACR50 improvement relative to baseline, n (%)	60 (73.2)	42 (51.9)	0.20 (0.06, 0.34); 0.0050
Proportion of patients with JIA-ACR70 improvement	53 (64.6)	34 (42.0)	0.22 (0.07, 0.37); 0.0032

relative to baseline, n (%)			
Change from baseline in number of active joints, adjusted mean	-14.3	-11.4	-2.9 (-5.7, -0.1); 0.0435
Change from baseline in PGA (VAS), adjusted mean	-45.2	-35.2	-9.9 (-16.5, -3.4); 0.0031
Change from baseline in the pain (VAS), adjusted mean	-32.4	-22.3	-10.2 (-17.6, -2.7); 0.0076
Change from baseline in number of joints with LOM, adjusted mean	-9.5	-7.7	-1.8 (-4.1, 0.5) ; 0.1229
Change from baseline in ESR (mm/h), adjusted mean	-26.3	-12.0	-14.3 (-19.6, -9.0) ⁵
CHAQ-disability score	-0.8	-0.6	-0.2 (-0.4, 0.0) ⁵
Proportion with JIA-ACR90 improvement, n (%)	37 (45.1)	19 (23.5)	0.21 (0.07, 0.35) ⁵
Proportion with inactive disease, n (%)	30 (36.6)	14 (17.3)	0.18 (0.05, 0.32) ⁵
Corticosteroid reducing regimens	Not reported		
Extra-articular manifestations (such as uveitis)	Not reported		
Body weight and height	Not reported		
Mortality	0	0	
Change from baseline in patient global assessment of well-being adjusted mean	-32.1	-24.7	-7.4 (-14.8, 0.0) ⁵

Comments: ⁴ Adjusted for baseline stratification factors (background use of MTX and oral glucocorticoids). ⁵ p values were not provided because they fell below a non-significant parameter in the hierarchical chain to address multiplicity.

Time to JIA-ACR30 flare reported in a Kaplan Meier curve, but not presented here.

Proportion of patients in the ITT population with JIA-ACR70 and JIA-ACR90 response at wk40 by background methotrexate, glucocorticoid and previous biological agent use at baseline⁶

Concomitant therapies and previous exposure to biological agent, n/N (% N)	Response level	TCZ (n=82)		Placebo (n=81)	
		Yes	No	Yes	No
Background MTX	JIA-ACR70	45/67 (67.2)	8/15 (53.3)	30/64 (46.9)	4/17 (23.5)
	JIA-ACR90	32/67 (47.8)	5/15 (33.3)	18/64 (28.1)	1/17 (5.9)
Background glucocorticoid	JIA-ACR70	23/33 (69.7)	30/49 (61.2)	4/38 (36.8)	20/43 (46.5)
	JIA-ACR90	16/33 (48.5)	21/49 (42.9)	5/38 (13.2)	14/43 (32.6)
Previous biological agent	JIA-ACR70	13/27 (48.1)	40/55 (72.7)	2/23 (8.7)	32/58 (55.2)
	JIA-ACR90	5/27 (18.5)	32/55 (58.2)	2/23 (8.7)	17/58 (29.3)

Comments: ⁶ Patients who withdrew or escaped to O-L TCZ or for whom the end point could not be determined were classified as non-responders.

Authors report an ad hoc analysis of patients who received TCZ continuously in parts 1 and 2 (not data extracted).

AEs and SAEs			
SAEs and AEs occurring $\geq 5\%$ of patients, n (%)	TCZ ⁷ (n=82)	Placebo ⁷ (n=81)	p value
Duration in study (years)	32.33	27.41	
Patients with ≥ 1 AE	58 (70.7)	60 (74.1)	
Total no. of AEs ⁸	147	141	
Rate of AEs per 100 patient-years	454.7	514.4	
Most frequent AEs			
Nasopharyngitis	14 (17.1)	9 (11.1)	
Headache	3 (3.7)	0	
Upper respiratory infection	4 (4.9)	2 (2.5)	
Cough	2 (2.4)	1 (1.2)	
Pharyngitis	3 (3.7)	3 (3.7)	
Nausea	2 (2.4)	2 (2.5)	
Diarrhoea	2 (2.4)	3 (3.7)	
Rhinitis	2 (2.4)	1 (1.2)	
Vomiting	3 (3.7)	1 (1.2)	
Abdominal pain	2 (2.4)	2 (2.5)	
Oropharyngeal pain	1 (1.2)	5 (6.2)	
Rash	4 (4.9)	1 (1.2)	
SAEs			
Patients with ≥ 1 SAE	3 (3.7)	3 (3.7)	

Rate of SAEs per 100 patient-years	9.3	10.9		
Patients with ≥ 1 infectious SAE	1 (1.2)	0		
Rates of infectious SAEs per 100 patient-years	3.1	0		
Pneumonia	1 (1.2)	0		
Upper limb fracture	1 (1.2)	0		
Uveitis	0	1 (1.2)		
Psychosomatic disease	1 (1.2)	0		
Enterocolitis	0	1 (1.2)		
Complicated migraine	0	1 (1.2)		
AEs leading to study drug discontinuation				
Increased blood bilirubin level ⁹	1 (1.2)			
Gastroenteritis		1 (1.2) ¹⁰		
<p>Comments: ⁷ AE data on open-label TCZ escape therapy were excluded. ⁸ Multiple occurrences of the same AE in one individual were counted. ⁹ Highest total bilirubin reading, 50μmol/L (normal range, 3–24μmol/L); 2 consecutive readings >51μmol/L mandated withdrawal per protocol. The event resolved without sequelae. ¹⁰ Occurred 46 days after the last of five doses of placebo.</p> <p>Exposure to TCZ varied for individual patients, depending on the period from the first dose of TCZ to the date of data cut or withdrawal (max. exposure 1.8 years). The safety population consisted of all patients who received ≥ 1 dose of study medication. Safety data included full exposure data for each patient.</p>				
Results for OLE, 104 weeks				
Efficacy endpoints and percentage change from baseline in JIA ACR components¹¹ (continuous TCZ, n=82)^{69;71;74}	Baseline	Week 40	Week 104 (n=160)	Change from baseline to week 104, %
JIA ACR70 responders, ¹² n (%)	-	65 (79.3)	71 (86.6)	-
JIA ACR90 responders, ¹² n (%)	-	41 (50.0)	58 (70.7)	-
Active joints (0–71), mean (SD)	19.7 (14.0)	4.7 (9.1)	3.3 (9.1)	-87.7 (27.1)
Joints with LOM (0–67), mean (SD)	16.5 (13.8)	5.6 (10.1)	3.6 (7.3)	-81.3 (31.7)
Patient global ¹³ (VAS 0–100 mm), mean (SD)	45.5 (23.1)	12.2 (19.0)	9.1 (18.4)	-75.4 (43.8)
PGA (VAS 0–100 mm), mean (SD)	57.8 (20.3)	8.8 (10.9)	5.0 (10.5)	-89.7 (23.7)
CHAQ-DI (0–3), mean (SD)	1.2 (0.7)	0.4 (0.5)	0.2 (0.4)	-76.7 (34.7)
ESR (mm/h), mean (SD)	31.7 (22.9)	5.4 (6.3)	5.1 (5.6)	-76.2 (27.3)
Inactive disease, ¹⁴ n (%)	-	33 (40.2)	52 (63.4)	-
Remission, ¹⁵ n (%)	-	5 (6.1)	31 (37.8)	-
Minimal disease activity (JADAS-71 <3.8), n (%)	0 (0)	49 (59.8)	60 (73.2)	-
Inactive disease (JADAS-71 <1), n (%)	0 (0)	24 (29.3)	48 (58.5)	-
<p>Comments: ¹¹ Patients who withdrew because of non-safety reasons are non-responders. Patients who withdrew because of safety are included using last observation carried forward. ¹² Two abstracts^{69;74} contain a table with a footnote to indicate patients who withdrew were excluded, however in the third abstract⁷¹ the table footnote states that patients who withdrew due to non-safety reasons are non-responders whereas patients who withdrew due to safety are included using LOCF. ¹³ Parent-rated. ¹⁴ No active joints, no active uveitis, ESR <20 mm/h and physician global assessment VAS ≤ 10. ¹⁵ Met criteria for inactive disease at each visit for 6 preceding months.</p>				
AEs and SAEs^{69-71;74}	(Safety population=188 with 307 patient-years)			
AEs, rates/100 patient-years	406.5			
SAEs, rates/100 patient-years	11.1			
Most common AE - infections	151.4			
Infections - SAE	5.2			
ALT elevations ≥ 3 x upper limit of normal, %	6.4			
AST elevations ≥ 3 x upper limit of normal, %	2.7			
Grade 3 lowest neutrophil count, %	5.9			
Grade 2/3/4 thrombocytopenia, %	1.6			
Low-density lipoprotein cholesterol ≥ 110 mg/dL, %	16.2			
Comments: ALT – alanine aminotransferase; AST – aspartate aminotransferase.				
Methodological comments				
<ul style="list-style-type: none"> <i>Allocation to treatment groups:</i> Randomly assigned 1:1, stratified by background use of MTX and oral glucocorticoid use. O-L (Part 1): patients weighing <30kg were randomly assigned 1:1 to receive intravenous TCZ 8 mg/kg or 10mg/kg, while patients weighing ≥ 30kg received 8 mg/kg. Double-blind RCT (Part 2): each of the 3 previous groups was randomised to the received either the existing dose of TCZ or placebo, equating to 6 groups in total. No further details about randomisation procedure were reported. 				

- *Blinding*: Double-blind, no further details of procedure reported. States that JIA-ACR response rates and clinically inactive disease status were performed in real time by independent masked evaluators at the coordinating centres of PRINTO and PRCSSG, according to validated criteria.
- *Comparability of treatment groups*: no baseline characteristics for Part 2 (RCT) reported. States that disease characteristics at baseline for the O-L (Part 1) were generally similar across the 3 groups with the exception of body weight based dosing regime, but no details are reported for the 6 groups in the RCT Part 2.
- *Method of data analysis*: ITT, however 3/166 patients from the placebo group were excluded as they discontinued without receiving the study drug therefore modified ITT. To control for the type 1 error rate, secondary endpoints were tested in a hierarchical fixed-sequence approach provided the primary endpoint was found to be statistically significant. The robustness of the results of the statistical procedure used for the primary endpoint analysis was assessed by logistic regression analysis of the proportion of patients with JIA-flare in the ITT during Part 2, showed a statistically significant treatment difference in favour of TCZ and was consistent with the primary analysis.
Primary endpoint analysis was conducted with the Cochran–Mantel–Haenszel test (also used for secondary endpoints), adjusted for stratification factors; patients who withdrew or for whom the endpoint could not be determined were considered to have experienced JIA-flare. Patients who escaped or withdrew or for whom the end point could not be determined were considered non-responders. Continuous variables were evaluated using analysis of variance, adjusted for baseline differences between groups and stratification variables. Ad hoc analysis was conducted in patients continuously treated with TCZ up to week 40, including those who escaped from blinded to O-L TCZ, using an ITT approach.
- *Sample size/power calculation*: Sample size estimation was reported. States that recruitment was planned to ensure that a sufficient number of patients were available for randomisation in Part 2, needing 60 patients in each group to achieve 80% power to detect a significant difference in assumed JIA-flare rates (35% TCZ, 65% placebo) between groups using a 2-sided significance test with $\alpha=0.05$. For the results, the 3 TCZ groups were combined and so were the 3 placebo groups, giving each combined group sufficient power.
- *Attrition/drop-out*: Part 1 lead in – states 10.6% discontinued (n=20), but flow chart shows n=22 (11.7%): lack of JIA-ACR30 response n=15, withdrew consent, n=3, AEs n=3, failure to return n=1.
Part 2 RCT: TCZ 3/82 (3.7% (10mg/kg <30kg BW group: n=1; 8mg/kg \geq 30kg BW: n=2) – AEs n=1, insufficient therapeutic response n=1, withdrew consent n=1. Placebo n=3/84 (3.6%) – AEs n=2, insufficient therapeutic response n=1.
OL-Extension: 5/160 (3.1%) – reasons not reported.⁶⁹

General comments

- *Generalisability*: To patients aged 2 to 17 years, with diagnoses of rheumatoid factor-positive or rheumatoid factor-negative pcJIA or extended oligoarticular JIA, with a minimum disease duration of at least 6 months and had inadequate responses to or were intolerant of MTX, and experienced at least one (JIA-ACR30) response to TCZ.
- *Outcome measures*: appear to be appropriate
- *Inter-centre variability*: not discussed.
- *Conflict of interests*: various authors received funding/support from a variety of pharmaceutical companies.

Quality criteria (Cochrane Collaboration Risk of Bias tool) RCTs¹⁶⁹

Criteria	Judgement ¹	Support for judgement
Random sequence generation (selection bias)	Unclear	Insufficient information.
Allocation concealment (selection bias)	Unclear	Insufficient information.
Blinding of participants and personnel (performance bias)	Yes	Double-blind.
Blinding of outcome assessment (detection bias)	Yes	JIA-ACR response rates and clinically inactive disease status were performed in real time by independent masked evaluators at 2 coordinating centres according to validated criteria.
Incomplete outcome data addressed (attrition bias)	Yes	Details reported and similar between groups.
Selective reporting (reporting bias)	Yes	All outcomes reported on.
Other sources of bias	Unclear	Inter-centre variability not discussed

¹ Yes (low risk of bias); No (high risk of bias); Unclear (uncertain risk of bias).

Appendix 6 Table of excluded studies for systematic review of cost-effectiveness

Excluded study	Primary reason for exclusion
Haapasaari JE, Kauppi M, Hakala MS, Kautiainen H. Economic evaluation of etanercept therapy in the treatment of re-fractory JIA. <i>Arthritis and Rheumatism</i> 2002; 46(9):S480.	Abstract
Brodzsky V, Pentek M, Majer I, Karpati K, Gulacsi L. [Etanercept in patients with juvenile idiopathic arthritis: systematic review and economic evaluation]. 2006.	Abstract
Prince FHM, de Bekker-Grob EW, Twilt M, Van Rossum MAJ, Hoppenreijs EPAH, ten Cate R et al. An analysis of the costs and treatment success of etanercept in Juvenile Idiopathic Arthritis. <i>Clinical and experimental rheumatology</i> 2011; 29(2):443.	Abstract
Simpson K, Hubert MM, On PV, Cifaldi M, Shaw J. Long-Term Cost-Effectiveness of Adalimumab Therapy in Juvenile Idiopathic Arthritis: From a Canadian Perspective. <i>Journal of Rheumatology</i> 2012; 39(8):1712.	Abstract
Luca N, Burnett H, Ungar W, Beukelman T, Feldman BM, Schwartz G et al. Cost-Effectiveness Analysis of Early Biologic Treatment in Polyarticular Juvenile Idiopathic Arthritis. <i>Arthritis & Rheumatism</i> 2012; 64(10, Suppl. S):S501.	Abstract
Luca N, Burnett H, Ungar W, Beukelman T, Feldman B, Schwartz G et al. Cost-effectiveness analysis of early biologic treatment in polyarticular juvenile idiopathic arthritis. <i>Journal of Rheumatology</i> 2013; Conference(var.pagings):6.	Abstract
Chang S, Sawyer L, Dejonckheere F, van Suijlekom-Smit LW, Anink J, Diamantopoulos A. Tocilizumab in Polyarticular Juvenile Idiopathic Arthritis - A Cost-Utility Model for the United Kingdom. <i>Value in Health</i> 2013; 16(7):A564.	Abstract
All Wales Medicines Strategy Group (. Adalimumab (Humira®). 2013.	Not economic evaluation
All Wales Medicines Strategy Group (. Etanercept (Enbrel®). 2013.	Not economic evaluation
All Wales Medicines Strategy Group (. Abatacept (Orencia®). 2014.	Not economic evaluation
All Wales Medicines Strategy Group (. Tocilizumab (RoActemra®). 2014.	Not economic evaluation
CADTH. Tocilizumab (Actemra - Hoffmann-La Roche Limited) new indication: polyarticular juvenile idiopathic arthritis. 2014.	Not economic evaluation

Appendix 7 Table of excluded studies for systematic review of health-related quality of life

Identified studies from titles/abstracts & full papers	Reason for exclusion
Anink J, Prince FHM, Dijkstra M, Otten H, Twilt M, Ten CR, et al. Long term functional outcome and quality of life of patients with refractory juvenile idiopathic arthritis treated with etanercept: Results of the Dutch arthritis and biologicals in children register. <i>Pediatr Rheumatol. Paediatric Rheumatology</i> , 2014; Conference proceedings.	Abstract
Duarte-Salazar C, Guzman-Vazquez S, Soto-Molina H, Chaidez-Rosales P, Ilizaliturri-Sanchez V, Nieves-Silva J, et al. Disability impact on quality of life in Mexican adults with juvenile idiopathic arthritis and juvenile ankylosing spondylitis. <i>Clin Exp Rheumatol</i> . 2007	No utilities reported
Hendry GJ, Gardner-Medwin J, Turner DE, Woodburn J, Lorgelly PK. Self- vs proxy-reported health-related quality of life of patients with juvenile idiopathic arthritis: Implications for a cost-utility analysis of multidisciplinary foot care. <i>Rheumatology Aust</i> ; 2011;Conference Available from: http://rheumatology.oxfordjournals.org/content/50/suppl_1/i2.full.pdf+html	Abstract
Hendry GJ, Gardner-Medwin J, Steultjens MPM, Woodburn J, Sturrock RD, Turner DE. Frequent discordance between clinical and musculoskeletal ultrasound examinations of foot disease in juvenile idiopathic arthritis. <i>Arthritis Care Res.</i> ; 2012;64(3):441–7.	No utilities reported
Janse AJ, Uiterwaal CS, Gemke RJ, Kimpen JL, Sinnema G. A difference in perception of quality of life in chronically ill children was found between parents and pediatricians. <i>J Clin Epidemiol</i> . 2005 May;58	Irelevant population
Janse AJ, Sinnema G, Uiterwaal CS, Kimpen JL, Gemke RJ. Quality of life in chronic illness: perceptions of parents and paediatricians. <i>Arch Dis Child</i> . 2005 May;90(England PT - Journal Article LG - English DC - 20050426):486–91.	Irelevant population
Janse A, Sinnema G, Uiterwaal C, Kimpen J, Gemke R. Quality of life in chronic illness: children, parents and paediatricians have different, but stable perceptions. <i>Acta Paediatr [Internet]</i> . 2008;97(8):1118–24.	Irelevant population
Angeles-Han ST, Griffin KW, Lehman TJA, Rutledge JR, Lyman S, Nguyen JT, et al. The importance of visual function in the quality of life of children with uveitis. <i>J AAPOS</i> ; 2010;14(2):163–8. 91-85	Irelevant population
Céspedes-Cruz A, Gutierrez-Suarez R, Pistorio A, Ravelli A, Loy A, Murray KJ, et al. Methotrexate improves the health-related quality of life of children with juvenile idiopathic arthritis. <i>Ann Rheum Dis</i> . 2008 Mar;67:309–14.	No utilities reported
Feinstein AB, Forman EM, Masuda A, Cohen LL, Herbert JD, Moorthy LN, et al. Pain intensity, psychological inflexibility, and acceptance of pain as predictors of functioning in adolescents with juvenile idiopathic arthritis: a preliminary investigation. <i>J Clin Psychol Med Settings</i> 2011 18,3,:291–8.	No utilities reported
Maetzel, A., Strand, V., Tugwell, P., Wells, G., & Bombardier, C. 2002. Economic comparison of leflunomide and methotrexate in patients with rheumatoid arthritis: an evaluation based on a 1-year randomised controlled trial. <i>Pharmacoeconomics</i> , 20, (1) 61-70	Irelevant population
Matza, L.S., Boye, K.S., Feeny, D.H., Johnston, J.A., Bowman, L., & Jordan, J.B. 2014. Impact of caregiver and parenting status on time trade-off and standard gamble utility scores for health state descriptions. <i>Health & Quality of Life Outcomes</i> , 12, 48	Irelevant population
McTaggart-Cowan, H.M., Brazier, J.E., & Tsuchiya, A. 2010. Clustering Rasch results: A novel method for developing rheumatoid arthritis states for use in valuation studies. <i>Value in Health</i> , 13, (6) 787-795	Irelevant population
Medrare, L., Ngeuleu, A., Rkain, M., Bouaddi, I., Znat, F., El, K.S., Lakhdar, T., Benslama, I., Rkain, H., Allali, F., Khattab, M., El, K.M., & Hajjaj-Hassouni, N. 2014. Is there any relationship between the children health assessment questionnaire (CHAQ) and the european quality of life (EUROQOL) in children suffering from chronic haemophilic arthropathy? <i>Annals of the Rheumatic Diseases</i> , Conference,	Irelevant population

Mo, F., Choi, B.C., Li, F.C., & Merrick, J. 2004. Using Health Utility Index (HUI) for measuring the impact on health-related quality of Life (HRQL) among individuals with chronic diseases. <i>TheScientificWorldJournal</i> , 4, 746-757	Irelevant population
Nordvag, B.-Y., Bernklev, T., Slevolden, E., Myhr, K.-M., & Stensland, E. 2012. Norwegian quality registry for biological drugs: The NOKBIL project. <i>Scandinavian Journal of Rheumatology</i> , Conference, (var.pagings) 50	Abstract
Osnes-Ringen, H., Kvien, T.K., Henriksen, J.E., Mowinckel, P., & Dagfinrud, H. 2009. Orthopaedic surgery in 255 patients with inflammatory arthropathies: Longitudinal effects on pain, physical function and health-related quality of life. <i>Annals of the Rheumatic Diseases</i> , 68, (10) 1596-1601	Irelevant population
Osnes-Ringen, H., Kvien, T.K., Henriksen, J.E., & Dagfinrud, H. 2010. Patients with inflammatory arthropathies undergo feet surgery later in the disease course than hand surgery. <i>Clinical and Experimental Rheumatology</i> , 28, (5) 702-707	Irelevant population
Osnes-Ringen, H., Kvamme, M.K., Kristiansen, I.S., Thingstad, M., Henriksen, J.E., Kvien, T.K., & Dagfinrud, H. 2011. Cost-effectiveness analyses of elective orthopaedic surgical procedures in patients with inflammatory arthropathies. <i>Scandinavian Journal of Rheumatology</i> , 40, (2) 108-115	Irelevant population
Shelepina, T.A., Stepanenko, N.Y., & Fedorov, E.S. 2011. Comparative characteristic of quality of life with patients suffering from jUVENILE idiopathic arthritis (JIA), attending school and taught at home. <i>Pediatric Rheumatology</i> , Conference, (var.pagings)	Abstract
Simpson, K., Hubert, M.M., On, P.V., Cifaldi, M., & Shaw, J. 2012. Long-term cost-effectiveness of adalimumab therapy in jUVENILE idiopathic arthritis: From a canadian perspective. <i>Journal of Rheumatology</i> ,	Abstract
Solari, N., Viola, S., Pistorio, A., Magni-Manzoni, S., Vitale, R., Ruperto, N., Ullmann, N., Filocamo, G., Martini, A., & Ravelli, A. 2008. Assessing current outcomes of juvenile idiopathic arthritis: a cross-sectional study in a tertiary center sample. <i>Arthritis & Rheumatism</i> , 59, (11) 1571-1579	No utilities reported
Sparsa, L., Job, D.C., Quartier, P., Kahan, A., & Wipff, J. 2013. Quality of life of juvenile idiopathic arthritis cohort at adulthood in a transition program. <i>Annals of the Rheumatic Disease</i> , , (var.pagings)	Abstract
Wade AG. Crawford GM. Pumford N. Koscielny V. Maycock S. McConnachie A. Baseline characteristics and patient reported outcome data of patients prescribed etanercept: web-based and telephone evaluation.	No utilities reported
Wang, H.-M., Beyer, M., Gensichen, J., & Gerlach, F.M. 2008. Health-related quality of life among general practice patients with differing chronic diseases in Germany: Cross sectional survey. <i>BMC Public Health</i> , 8 , 2008. Article Number,	Irelevant population

Appendix 8 Cost-effectiveness studies – data extraction forms

1	Study	Cummins, 2002 ¹²²															
2	Research question	To provide background info and systematic review of JIA, including economic evidence of etanercept compared to other treatment options															
3	Country/setting	Not stated															
4	Funding source	Not stated															
5	Analysis type	Cost-utility analysis															
6	Study type	Industry submission CU model using results from one JIA trial (Lovell et al., 2000). The model assumes response related to health assessment (HAQ) and mortality.															
7	Perspective	Health care system															
8	Time horizon	Model cycle length: 3m, 6m, and 1y, then yearly intervals over the life-course.															
9	Model assumptions	It is an adaptation of a rheumatoid arthritis model for adults using strong and questionable assumptions, related to health assessment, utility, mortality and costs.															
10	Discounting (rate)	Yes, costs 6% per annum and benefits 1%															
11	Costing year, currency	2001, £															
12	Population	Etanercept in children with polyarticular juvenile rheumatoid arthritis. Definition of condition: JIA, heterogeneous group of painful conditions involving persistent swelling of the joints with variable presentation and course.															
13	Intervention(s), comparator(s)	Etanercept vs placebo (placebo effect assumed to last 3m)															
14	Intervention effect	Effect size measured in terms of CHAQ and mortality. Cost offset per HAQ point £860 38% increase in mortality per point change in HAQ Relative risk of mortality in JIA was 2.98 Placebo and etanercept HAQ progression: responders 0-4 years 0, responders >4 years 0.034, non-responders 0.0669 Annual withdrawal from responders to non-responders: placebo 50%, etanercept 13%.															
15	Health state utilities	EQ-5D adults															
16	Intervention cost	Adult cost															
17	Indirect costs	n/a															
18	Results	<table border="1"> <thead> <tr> <th>Discounted/ undiscounted</th> <th>Intervention</th> <th>Comparator</th> <th>Incremental</th> <th>ICER</th> </tr> </thead> <tbody> <tr> <td>Costs</td> <td>£40,624</td> <td>£12,602</td> <td>£28,022</td> <td></td> </tr> <tr> <td>QALY</td> <td>15.0</td> <td>13.3</td> <td>1.7</td> <td>£16,082</td> </tr> </tbody> </table>	Discounted/ undiscounted	Intervention	Comparator	Incremental	ICER	Costs	£40,624	£12,602	£28,022		QALY	15.0	13.3	1.7	£16,082
Discounted/ undiscounted	Intervention	Comparator	Incremental	ICER													
Costs	£40,624	£12,602	£28,022														
QALY	15.0	13.3	1.7	£16,082													
19	Sensitivity analysis	Sensitivity analysis varied from £3,900 to £34,000 (SF-36 regression used)															
20	Author's conclusions	Insufficient data to construct a model for JIA, and little is known about HRQoL in JIA. The ICER should be viewed with caution.															
21	Reviewer's comments	Limited relevance in cost/utilities; for adult RA cannot be assumed to be the same for JIA in children.															

Critical appraisal checklist for economic evaluations (based on Drummond et al)

Item	Y/N/?
1. Is the decision problem (including interventions compared and patient group) relevant to the UK?	Y
2. Is the setting comparable to the UK?	Y
3. Is the analytical and modelling methodology appropriate?	N
4. Are all the relevant costs and consequences for each alternative identified?	N
5. Are the data inputs for the model described and justified?	Y
6. Are health outcomes measured in QALYs?	Y

7. Is the time horizon considered appropriate?	Y
8. Are costs and outcomes discounted?	Y
9. Is an incremental analysis performed?	Y
10. Is uncertainty assessed?	Y
<i>Y – yes, N – no, ? – nuclear</i>	
Comments	
Out of date, not children specific, relying on very strong assumptions due to lack of evidence. Informative in general terms but not relevant.	

1	Study	Prince, 2011 ¹²³
2	Research question	To analyse and report the costs and effects of etanercept therapy in patients with JIA
3	Country/setting	Netherlands/ national Arthritis and Biologicals in Children register
4	Funding source	The Dutch Board of Health Insurances and Wyeth International
5	Analysis type	Cost consequence analysis
6	Study type	Trial-based: Prospective etanercept effectiveness and safety add-on study with JIA patients in 7 of 9 Dutch paediatric rheumatology centres.
7	Perspective	Health care system
8	Time horizon	27 months
9	Model assumptions	N/A
10	Discounting (rate)	No
11	Costing year, currency	2008, € (euro)
12	Population	Dutch JIA patients younger than 18 years are eligible for treatment with etanercept if the disease has a polyarticular course and the response to the maximum (tolerated) dose of methotrexate (MTX) is not sufficient. Onset subtype JIA (n=49): <ul style="list-style-type: none"> - Systemic (22%) - Polyarticular RF+ (8%) - Polyarticular RF- (37%) - Oligoarticular extended (22%) - Entesitis-related arthritis (4%) - Juvenile arthritis psoriatica (6%) Concomitant drug use at start of etanercept: <ul style="list-style-type: none"> - NSAID (92%) - Glucocorticoids systemic (47%) - MTX (80%) - Other DMARD (10%)
13	Intervention(s), comparator(s)	Intervention: Etanercept (add-on to conventional treatment) Comparator: conventional treatment with synthetic disease-modifying anti-rheumatoid drugs (DMARD), mostly MTX, if required accompanied by anti-inflammatories or systemic glucocorticoids.
14	Intervention effect	Effect size measured in the study in terms of change in disease activity response variables of the JIA core set and HUI3
15	Health state utilities	HUI3 – preference-based HRQoL measure completed by the parents of study participants (8 domains in 15-item parent questionnaire). Valuation using value scores obtained by Fenny et al (2002) from the Canadian general population.
16	Intervention cost	Unit costs for medication were retrieved from the Pharmacotherapeutic Compass provided by the Dutch Board of Health Insurances, and treatment costs were calculated with the exact dose of medication and administration period as reported in the patients' files. Etanercept unit cost ~ €10,478/year

17	Indirect costs	N/A		
18	Results:			
	Undiscounted	Intervention	Comparator	Incremental
	Costs (€, 27 months)	28,075	8,370	19,705
	Utility	0.78 (27 months)	0.53 (0 months)	0.25
19	Sensitivity analysis: N/A			
20	Author's conclusions	"Although etanercept is expensive, the major utility gain justifies the costs."		
21	Reviewer's comments	Sound trial-based evaluation of costs and consequences (including disease activity improvement and utility) associated to adding etanercept to conventional care. Full incremental cost-effectiveness/utility analysis not performed and there is no indication of the variation from the mean estimates reported nor assessment of uncertainty.		

Critical appraisal checklist for economic evaluations (based on Drummond et al)

Item	Y/N/?
1. Is the decision problem (including interventions compared and patient group) relevant to the UK?	Y
2. Is the setting comparable to the UK?	?
3. Is the analytical and modelling methodology appropriate?	Y ^a
4. Are all the relevant costs and consequences for each alternative identified?	Y
5. Are the data inputs for the model described and justified?	N/A
6. Are health outcomes measured in QALYs?	N
7. Is the time horizon considered appropriate?	N
8. Are costs and outcomes discounted?	N
9. Is an incremental analysis performed?	? ^a
10. Is uncertainty assessed?	N
<i>Y – yes, N – no, ? – unclear</i>	
Comments	
^a The methodology is appropriate for a cohort-based evaluation; however, a full incremental cost utility analysis has not been performed	

1	Study	Simpson, 2012 ¹²⁴
2	Research question	To evaluate the cost-effectiveness of adalimumab <i>versus</i> non-biologic therapy for the treatment of JIA in Russian children and adolescents.
3	Country/setting	Russia / health care system
4	Funding source	Not stated
5	Analysis type	Cost-utility analysis
6	Study type	Markov model; Mutually exclusive health states: <ul style="list-style-type: none"> - Base model (children under 18 years): (1) mild disease activity, (2) moderate disease activity, (3) severe disease activity, (4) remission without movement limitations, (5) remission with movement limitations - Second part of the model (adults from 18 years to death): (6) remission, (7) active mild disability, (8) active moderate disability, and (9) active severe disability - (10) Death (not clear if children mortality is included)
7	Perspective	Health care system and Society
8	Time horizon	Lifetime. Model cycle length – 4 months
9	Model assumptions	<ul style="list-style-type: none"> - Adult patients with moderate to severe disability are assumed to have hip and knee prosthetic surgery at the frequency observed in patients (Packham and Hall 2002 Rheumatology); - Patients who do not achieve remission after 1 year of treatment had a median time on treatment of 3 years (as observed in DE038); - Mean age of 11 years at start of therapy

Comments

Limited assessment of uncertainty

1	Study	Ungar, 2011 ¹²⁶																				
2	Research question	To determine the incremental costs of biologics per additional responder compared to conventional treatment (methotrexate)																				
3	Country/setting	Canada, secondary care																				
4	Funding source	Ontario Ministry of Health and Long-term Care Drug Innovation Fund																				
5	Analysis type	CEA																				
6	Study type	Decision analysis model																				
7	Perspective	Societal																				
8	Time horizon	1 year time horizon with 2 consecutive 6 month cycles																				
9	Model assumptions	Model incorporated probabilities that patients would, based on their response at 6 months, either continue with the same treatment or switch.																				
10	Discounting (rate)	Not included																				
11	Costing year, currency	2008 Canadian dollars																				
12	Population	In the base case, a 40 kg patient was assumed, similar to the mean weight in 2 paediatric RCTs. Patients had JIA with a prior inadequate response or intolerance to disease modifying antirheumatic drugs (DMARDs).																				
13	Intervention(s), comparator(s)	Etanercept, adalimumab, abatacept and infliximab vs. MTX																				
14	Intervention effect	<p>The effectiveness measure was the proportion of patients who had a reduction in symptoms at 1 year according to the ACR Pedi 30 criteria. Effect size were taken from RCTs: etanercept (Lovell et al, 2000), adalimumab (Lovell et al, 2008), infliximab (Ruperto et al 2007) and abatacept (Ruperto et al 2008).</p> <p>For the base case, patients achieving ACR Pedi 30, %</p> <table border="1"> <thead> <tr> <th>Time</th> <th>Etanercept</th> <th>Adalimumab</th> <th>Abatacept</th> <th>Infliximab</th> <th>DTX</th> </tr> </thead> <tbody> <tr> <td>6 months</td> <td>79</td> <td>80</td> <td>82</td> <td>80</td> <td>30</td> </tr> <tr> <td>12 months</td> <td>79</td> <td>63</td> <td>82</td> <td>79</td> <td>30</td> </tr> </tbody> </table>	Time	Etanercept	Adalimumab	Abatacept	Infliximab	DTX	6 months	79	80	82	80	30	12 months	79	63	82	79	30		
Time	Etanercept	Adalimumab	Abatacept	Infliximab	DTX																	
6 months	79	80	82	80	30																	
12 months	79	63	82	79	30																	
15	Health state utilities	No utility values included.																				
16	Intervention cost	Total annual costs for treatment were: abatacept (\$14,733), infliximab (\$17,259), etanercept (\$18,966), adalimumab (\$18,654), methotrexate (\$952). Treatment costs included medication costs, preparation and administration costs and concomitant medications.																				
17	Indirect costs	The costs for abatacept and infliximab included parental time losses of \$1875 and \$1071. There were no indirect costs for the other treatments.																				
18	Results Intervention vs MTX	<table border="1"> <thead> <tr> <th>Undiscounted</th> <th>Etanercept</th> <th>Adalimumab</th> <th>Abatacept</th> <th>Infliximab</th> </tr> </thead> <tbody> <tr> <td>Incremental costs, \$</td> <td>11,090</td> <td>13,107</td> <td>7,873</td> <td>12,167</td> </tr> <tr> <td>Incremental effectiveness, %</td> <td>47.6</td> <td>29.4</td> <td>49.4</td> <td>43.2</td> </tr> <tr> <td>ICER</td> <td>26,061</td> <td>46,711</td> <td>16,204</td> <td>31,209</td> </tr> </tbody> </table>	Undiscounted	Etanercept	Adalimumab	Abatacept	Infliximab	Incremental costs, \$	11,090	13,107	7,873	12,167	Incremental effectiveness, %	47.6	29.4	49.4	43.2	ICER	26,061	46,711	16,204	31,209
Undiscounted	Etanercept	Adalimumab	Abatacept	Infliximab																		
Incremental costs, \$	11,090	13,107	7,873	12,167																		
Incremental effectiveness, %	47.6	29.4	49.4	43.2																		
ICER	26,061	46,711	16,204	31,209																		
19	Sensitivity analysis	<p>Deterministic analysis was performed for extreme efficacy with biologic high efficacy and MTX low efficacy and vice versa.</p> <p>Probabilistic sensitivity analyses were conducted for each treatment vs MTX and cost-effectiveness acceptability curves were calculated. If a decision maker was willing to pay no more than \$30,000 to gain a responder, then the probability that etanercept would demonstrate a net economic benefit would be 95%. The willingness to pay points at which the biologic had a 50% probability of cost-effectiveness were \$45,000,</p>																				

	\$17,000 and \$27,500 for adalimumab, abatacept and infliximab respectively.	
20	Author's conclusions	JIA patients with a prior suboptimal response or intolerance to MTX may benefit from treatment with biologic for at least 1 year.
21	Reviewer's comments	Results not present in QALYs, which makes results difficult to interpret. Short time horizon used (1 year). Unclear how ICERs are calculated.

Critical appraisal checklist for economic evaluations (based on Drummond et al)

Item	Y/N/?
1. Is the decision problem (including interventions compared and patient group) relevant to the UK?	Y
2. Is the setting comparable to the UK?	N
3. Is the analytical and modelling methodology appropriate?	Y
4. Are all the relevant costs and consequences for each alternative identified?	Y
5. Are the data inputs for the model described and justified?	Y
6. Are health outcomes measured in QALYs?	N
7. Is the time horizon considered appropriate?	N
8. Are costs and outcomes discounted?	N
9. Is an incremental analysis performed?	Y
10. Is uncertainty assessed?	Y
<i>Y – yes, N – no, ? – unclear</i>	

Appendix 9 Health related quality of life systematic review– data extraction forms

Reference

Hendry, 2013, ¹³⁵

Study Characteristics

Research question

What are the stated objectives of the study?

To evaluate the effectiveness of multidisciplinary foot-care, and to evaluate the methodological considerations of a trial of multidisciplinary care in juvenile idiopathic arthritis.

Describe the type of study and study design.

Exploratory randomised controlled trial.

Was the sample from i) the general population, ii) patients with the disease of interest, iii) individuals with knowledge of the disease, iv) other?

Are inclusion/exclusion criteria clearly described? Do these exclude any individuals that may be relevant (eg >80 years)?

The sample was drawn from patients with the disease of interest, i.e. children and adolescents with a definitive diagnosis of JIA and inflammatory joint disease affecting the foot/ankle.

The inclusion/exclusion criteria were clearly stated; however, might exclude a proportion of individuals with the disease of interest but whose disease has not affected the foot/ankle.

Patients were included if they satisfied at least one of the following: (i) previously documented arthritis in the foot including small joints derived from medical case notes, (ii) previously documented foot arthritis in one or more large joints derived from medical case notes, or (iii) current widespread polyarthritis involving large and small foot joints derived from clinical examination by a consultant paediatric rheumatologist. Patients with an unconfirmed diagnosis of JIA, and/or only upper limb, jaw, or neck involvement were excluded.

What are the characteristics of the baseline cohort for the evaluation?

Age, years, mean(SD), intervention arm; control arm	10.1(4.22); 10.0(3.39)
Male/Female, n, intervention arm; control arm	7/14; 6/17
Race (if appropriate)	nr
Disease subtypes, n (%), intervention arm; control arm:	
- Persistent oligoarthritis	7 (33); 4 (17)
- Extended oligoarthritis	4 (19); 5 (22)
- Polyarthritis rheumatoid factor negative	6 (29); 10 (43)
- Polyarthritis rheumatoid factor positive	0 (0); 2 (9)
- Psoriatic arthritis	2 (10); 1 (4)
- Enthesitis related arthritis	2 (10); 0 (0)
- Undifferentiated	0 (0); 1 (4)
Sample size, n, intervention arm; control arm:	21; 23
Pharmacological management, n (%), intervention arm; control arm:	
- Analgesics	2 (9); 3 (13)
- NSAIDs	2 (9); 3 (13)

<ul style="list-style-type: none"> - Methotrexate - Etanercept - Sulphasalazine - Rituximab - Combination methotrexate & etanercept 	<p>18 (86); 16 (70) 7 (33); 5 (22) 1 (5); 0 (0) 0 (0); 1 (4) 5 (24); 5 (22)</p>
QoL instrument	EQ-5D-Y (patients) and EQ-5D-3L (parents/guardians) questionnaires
Utility values, (Y/N)	Y
Treatment effect, if reported	Both the treatment groups appeared to improve by one point on the JAFI impairment scale between baseline and 12 months follow up, however, the differences between groups for change scores did not reach statistical significance.

Country/ setting

What is the country and setting for the evaluation?

Royal Hospital for Sick Children. Glasgow, UK

Data Sources

Effectiveness

Were the QoL data derived from: a single (observational) study, a review / synthesis or combination of previous studies, expert opinion?

This single exploratory RCT

Results

Summarise the results

There were no significant differences between treatment groups for secondary outcomes at final follow up.

	Intervention arm	Control arm
Baseline		
Self EQ-5D utility index, mean (SD)	0.57 (0.31)	0.58 (0.35)
Self EQ-5D utility index, median (IQR)	0.62 (0.52 to 0.76)	0.66 (0.52 to 0.75)
Proxy EQ-5D utility index, mean (SD)	0.69 (0.29)	0.60 (0.33)
Proxy EQ-5D utility index, median (IQR)	0.69 (0.58 to 1)	0.62 (0.55 to 0.82)
Change at 12 months		
Self EQ-5D utility index, median (IQR)	0 (-0.1 to 0.01)	0 (-0.04 to 0.04)
Proxy EQ-5D utility index, median (IQR)	0 (0 to 0.11)	0 (0 to 0.1)

Were the methods for deriving these data adequately described (give sources if using data from other published studies)? (Was a valid preference based instrument used to describe health states, such as EQ-5D? Was the valuation of health states from the UK general population?)

A valid preference-based instrument was used – EQ-5D-Y and EQ-5D-3L

Are the levels of missing data reported? How are they dealt with?

For missing data identified at the end of the study, a sensitivity analysis was performed in order to identify the most appropriate method to address this problem (LOCF, mean value imputation, maximum value imputation, minimum value imputation, and random value imputation). LOCF was found to be the most conservative method while being less labour intensive, thus it was subsequently used to impute all missing data at final follow-up.

Mapping

If a model was used, describe the type of model (eg. regression) or other conversion algorithm

Not applicable

Conclusions/ Implications

Give a brief summary of the author's conclusions from their analysis

Integrated multidisciplinary foot care did not result in a significant reduction in disease-related foot impairments and disability.

What are the implications of the study for the model

In both arms, a proportion of participants received etanercept, so the utility values reported cannot be used in the model for baseline HRQoL with standard of care.

Reference

Prince, 2010^{123,136}

Study Characteristics

Research question

What are the stated objectives of the study?

To evaluate changes in health related quality of life in patients with refractory juvenile idiopathic arthritis who are being treated with etanercept

Describe the type of study and study design.

Prospective study

Was the sample from i) the general population, ii) patients with the disease of interest, iii) individuals with knowledge of the disease, iv) other?

Are inclusion/exclusion criteria clearly described? Do these exclude any individuals that may be relevant (eg >80 years)?

JIA patients younger than 18 years old treated with etanercept

What are the characteristics of the baseline cohort for the evaluation?

Study	Prince 2010	Prince 2011
Age	11.9 years (IQR 8.1 – 14.9)	11.6 years (IQR 7.9 – 14.9)
Sex, n (%)	Male 20 (38), female 33 (62)	Male 20 (41), female 33 (59)
Race (if appropriate)		
Indication / disease, n (%)	Systemic 14 (26) Polyarticular rheumatoid factor positive 5 (9) Polyarticular rheumatoid factor negative 18 (34) Oligoarticular extended 11 (21) Enthesitis-related arthritis 2 (4) Juvenile psoriatic arthritis 3 (6)	Systemic 11 (22) Polyarticular rheumatoid factor positive 4 (8) Polyarticular rheumatoid factor negative 18 (37) Oligoarticular extended 11 (22) Enthesitis-related arthritis 2 (4) Juvenile psoriatic arthritis 3 (6)
Other characteristics (sample size)	Sample size 53 Median disease duration JIA (years) at start of etanercept 3.0	Sample size 49 Median disease duration JIA (years) at start of etanercept 3.6
QoL instrument	HUI3	HUI3
Utility values, (Y/N)	Yes	Yes

Treatment effect, if reported	Significant improvements were shown after 3 months and these continued at least up to 27 months	Significant improvements were shown after 3 months and these continued at least up to 27 months
-------------------------------	---	---

Country/ setting

What is the country and setting for the evaluation?

The Netherlands

Data Sources

Effectiveness

Were the QoL data derived from: a single (observational) study, a review / synthesis or combination of previous studies, expert opinion?

Single prospective study

Results

Summarise the results

Prince (2010)	Baseline	3 months	15 months	27 months
HUI3 mean (SE)	0.53 (0.04)	0.69 (0.05)	0.74 (0.06)	0.78 (0.07)

Were the methods for deriving these data adequately described (give sources if using data from other published studies)? (Was a valid preference based instrument used to describe health states, such as EQ-5D? Was the valuation of health states from the UK general population?)

Yes

Are the levels of missing data reported? How are they dealt with?

Not reported

Mapping

If a model was used, describe the type of model (eg. regression) or other conversion algorithm

Mapping was not used

Conclusions/ Implications

Give a brief summary of the author's conclusions from their analysis

This study shows that the HRQoL of patients with refractory JIA can be substantially improved by the use of etanercept

What are the implications of the study for the model

This is a potential source of HRQoL for the SHTAC economic model

Appendix 10 Cost-effectiveness data extraction forms for the company submissions

1 Reference

AbbVie, 2015,⁷⁷

1.1 Health technology

Adalimumab

1.2 Interventions and comparators

What interventions/ strategies were included?

No economic evaluation was conducted; however, reasons for not conducting an economic evaluation were discussed and included the following interventions: adalimumab, etanercept, abatacept, tocilizumab, and methotrexate.

Was a no treatment/ supportive care strategy included?

No

Describe interventions/ strategies

p.13 CS: “The aim of drug therapy in JIA patients is to induce and maintain remission of symptoms, and thus allow a child to achieve normal growth, development, and allow full participation in school, career, sport and all other aspects of normal life.^{Error! Bookmark not defined.} The initial aim is induction of complete disease remission using corticosteroids – either intravenously (IV) or intra-articular. Oral corticosteroids are avoided where possible to avoid side effects (can affect growth or increase risk of osteoporosis) but may be needed for short time periods.”

p. 14 EMA licence: “Adalimumab in combination with methotrexate is indicated for the treatment of active polyarticular juvenile idiopathic arthritis, in patients from the age of 2 years who have had an inadequate response to one or more disease-modifying anti-rheumatic drugs (DMARDs). Adalimumab can be given as monotherapy in case of intolerance to methotrexate or when continued treatment with methotrexate is inappropriate. Adalimumab has not been studied in patients aged less than 2 years.”

Adalimumab is delivered as 24mg/m² body surface area (with varying maximum doses dependent on weight and study protocol) subcutaneous injection with concomitant methotrexate for 24 weeks or more. Patients were allowed to take NSAIDs, and prednisone or equivalents to prednisone.

1.3 Research question

What are the stated objectives of the evaluation?

No economic evaluation was conducted.

1.4 Study type Cost-effectiveness/ cost-utility/ cost-benefit analysis?

No economic evaluation was conducted.

1.5 Study population

What definition was used for [condition]? What are the characteristics of the baseline cohort for the evaluation?

From pp 9-10 of the CS: “Juvenile Idiopathic Arthritis (JIA) is the most common rheumatic disease of childhood and describes a group of conditions that involve joint inflammation which lasts for more than 6 weeks in people under 16 years of age. ... JIA is an ‘umbrella’ term which covers a number of different sub-types listed below that were proposed by the International League of Associations for Rheumatology (ILAR) in 1995 for the classification of JIA:

- Oligoarticular JIA - Oligoarthritis is the most common type of JIA, accounting for up to 50% of new diagnoses in Europe each year. It is diagnosed when four or fewer joints are affected in the

first 6 months of disease.

- Extended Oligoarticular JIA - If oligoarthritis progresses and affects more than four joints during the first 6 months, it is called extended oligoarthritis.
 - Poly-Articular JIA (RF –ve or RF +ve) - Polyarticular JIA is diagnosed when five or more joints are affected at presentation, and can be further divided into rheumatoid factor positive arthritis and rheumatoid factor negative disease.
 - Systemic-Onset JIA - Systemic JIA accounts for 5-10% of new diagnoses and is diagnosed when arthritis is part of a general illness involving features such as fever, lymphadenopathy, hepatosplenomegaly and serositis. This patient group was not included in the NICE Scope for this project.
 - Psoriatic JIA - Psoriatic arthritis accounts for 2-15% of new diagnoses and is diagnosed when there is joint swelling associated with psoriasis, or a family history of psoriasis.
 - Entesitis-Related Arthritis (ERA) - ERA accounts for 2-10% of new diagnoses and is diagnosed in the presence of arthritis or inflammation of tendon attachments to the bones (entheses), in association with two or more other features of spondyloarthritis.”
- No economic evaluation was conducted, so there is no relevant base-line cohort.

1.6 Institutional setting Where is/are the intervention(s) being evaluated usually provided?

Paediatric secondary care

1.7 Country/ currency

Has a country setting been provided for the evaluation? What currency are costs expressed in and does the publication give the base year to which those costs relate?

No economic evaluation was conducted

1.8 Funding source

AbbVie

1.9 Analytical perspective

What is the perspective adopted for the evaluation (health service, health and personal social services, third party payer, societal (i.e. including costs borne by individuals and lost productivity)?

No economic evaluation was conducted

2 Effectiveness

Were the effectiveness data derived from: a single study, a review/ synthesis of previous studies or expert opinion? Give the definition of treatment effect used in the evaluation. Give the size of the treatment effect used in the evaluation

No economic evaluation was conducted

3 Intervention Costs

Were the cost data derived from: a single (observational) study, a review/ synthesis of previous studies expert opinion? Were the methods for deriving these data adequately described (give sources if using data from other published studies)? List the direct intervention costs and other direct costs used in the evaluation – include resource estimates (and sources for these estimates, if appropriate) as well as sources for unit costs used.

No economic evaluation was conducted

3.1 Indirect Costs (costs due to lost productivity, unpaid inputs to patient care)

Were indirect costs included:

No economic evaluation was conducted

4 Health state valuations/ utilities (if study uses quality of life adjustments to outcomes)

Were the utility data derived from: a single (observational) study, a review/ synthesis of previous studies expert opinion. Were the methods for deriving these data adequately described (give sources if using data from other published studies)?

No economic evaluation was conducted

4.1 List the utility values used in the evaluation

No economic evaluation was conducted

5 Modelling

If a model was used, describe the type of model used (e.g. Markov state transition model, discrete event simulation). Was this a newly developed model or was it adapted from a previously reported model? If an adaptation, give the source of the original. What was the purpose of the model (i.e. why was a model required in this evaluation)? What are the main components of the model (e.g. health states within a Markov model)? Are sources for assumptions over model structure (e.g. allowable transitions) reported – list them if reported.

No modelling was undertaken.

5.1 Extract transition probabilities for [natural history/disease progression] model and show sources (or refer to table in text).

No modelling was undertaken.

5.2 What is the model time horizon?

No modelling was undertaken.

5.3 What, if any, discount rates have been applied in the model? Same rate for costs and outcomes?

No modelling was undertaken.

5.4 If no economic evaluation was conducted, state the manufacturer's reasons for this.

No economic evaluation was conducted due to heterogeneity in study methods and populations between the interventions that complicated indirect comparisons, and a lack of appropriate utility data for HRQoL and lack of long-term data.

6 Results/ Analysis

What measure(s) of benefit were reported in the evaluation?

No economic evaluation was undertaken.

6.1 Provide a summary of the clinical outcome/ benefits estimated for each intervention/ strategy assessed in the evaluation

No economic evaluation was undertaken.

6.2 Provide a summary of the costs estimated for each intervention/ strategy assessed in the evaluation

No economic evaluation was undertaken.

6.3 Synthesis of costs and benefits – are the costs and outcomes reported together (e.g. as cost-effectiveness ratios)? If so, provide a summary of the results.

No economic evaluation was undertaken.

6.4 Give results of any statistical analysis of the results of the evaluation.

No economic evaluation was undertaken.

6.5 Was any sensitivity analysis performed – if yes, what type(s) (i.e. deterministic (one-way, two-way etc) or probabilistic).

No economic evaluation was undertaken.

6.6 What scenarios were tested in the sensitivity analysis? How do these relate to structural uncertainty (testing assumptions over model structure such as relationships between health states), methodological uncertainty (such as choices of discount rate or inclusion of indirect costs) or parameter uncertainty (assumptions over values of parameters in the model, such as costs, quality of life or disease progression rates)?

No economic evaluation was undertaken.

6.7 Give a summary of the results of the sensitivity analysis – did they differ substantially from the base case analysis. If so, what were the suggested causes?

No economic evaluation was undertaken.

7 Conclusions/ Implications

Give a brief summary of the author’s conclusions from their analysis

No economic evaluation was undertaken.

7.1 What are the implications of the evaluation for practice?

No economic evaluation was undertaken.

8 SHTAC Commentary

Selection of comparators:

While no economic evaluation was conducted, the comparators listed by the manufacturer were appropriate and in accord with the NICE Scope.

Validity of estimate of measure of benefit:

No economic evaluation was undertaken.

Validity of estimate of costs:

No economic evaluation was undertaken.

1 Reference

Bristol Myers Squibb (BMS), 2015, ⁸⁵

1.1 Health technology

Abatacept

1.2 Interventions and comparators

What interventions/ strategies were included?

Abatacept, adalimumab, etanercept and tocilizumab

Was a no treatment/ supportive care strategy included?

No.

Describe interventions/ strategies

Abatacept is a biological DMARD that prevents T-cell activation, thus down-regulating the immune response of inflammatory disease. (p.9). Abatacept is administered intravenously.

p. 9 CS: “The recommended dose of abatacept for polyarticular JIA patients aged 6 to 17 years (who

weigh less than 75kg) is 10mg/kg, calculated based on the patient's body weight at each administration. Paediatric patients weighing 75kg or more should follow the abatacept adult dosing regimen and should not exceed a maximum dose of 1,000mg. Abatacept should be administered as a 30-minute intravenous infusion. Following the initial administration, abatacept should be given at 2 and 4 weeks after the first infusion, and every 4 weeks thereafter.”

Etanercept is a biological DMARD that inhibits TNF activation. It is administered subcutaneously. For patients aged 2-18 400µg/kg (max 25mg twice a week) or 800µg/kg, (max 50mg once a week) was administered. For patients above aged 18 or above the dose was 50mg.

Adalimumab is a biological DMARD that inhibits TNF activation. It is administered subcutaneously. For patients aged 4-13 24mg/m² (max 40mg) was administered every other week. For patients aged 13 and above the dose was 40mg.

Tocilizumab is a biological DMARD. It is a humanised monoclonal antibody that inhibits the cytokine interleukin-6. It is administered by intravenous infusion. Tocilizumab dose was based on weight. For patients weighing under 30kg the dose was 10mg/kg. For patients 30kg and above the dose was 8mg/kg (max 800mg). Doses were administered every four weeks. All tocilizumab patients were greater than two years of age.

All drugs were administered with subcutaneous methotrexate. BMS indicated that methotrexate was given every four weeks at a dosage of 13.5mg/m².

1.3 Research question

What are the stated objectives of the evaluation?

From CS model: “The model evaluates the cost-effectiveness of abatacept against other biological disease modifying anti rheumatic drugs (bDMARDs) in moderate-to-severe active polyarthritis in paediatric patients from the age of 6 years and who have shown insufficient response to other DMARDs, including at least one anti-TNF.”

BMS reports that this is not in perfect agreement with the NICE Scope, but is in accord with the drug license. The NICE Scope is broader with no specifications for patient age, or insufficient response on other DMARDs (including failure of at least one TNF inhibitor). The NICE Scope also includes enthesitis-related arthritis.

1.4 Study type Cost-effectiveness/ cost-utility/ cost-benefit analysis?

BMS has conducted a cost-minimisation analysis (CMA) with an assumption that there is no difference between the bDMARDs in effectiveness. BMS indicated that effectiveness evidence was not considered because it “would lead to uncertainty within the model.”

1.5 Study population

What definition was used for [condition]? What are the characteristics of the baseline cohort for the evaluation?

p.8 CS: “JIA encompasses all forms of arthritis of unknown aetiology that persist for at least 6 weeks and begin in patients younger than 16 years.1 JIA comprises several heterogeneous subtypes (oligoarthritis, polyarthritis, systemic, psoriatic, enthesitis-related and undifferentiated), all presenting with different clinical signs and symptoms.1,3,14 Overall, JIA is characterised by persistent joint swelling, pain and limitation of movement and has an estimated incidence in the UK of 1 per 10,000 children and a prevalence in the order of 1 per 1,000 children.2 Polyarticular JIA (classifiable as polyarthritis [rheumatoid factor-positive or -negative]) is characterised by arthritis affecting five or more joints during the first 6 months of the disease1,3,14,15, and it affects 13%–37% of patients with JIA.3

JIA causes functional impairment due to joint and back pain, heel pain, swelling of joints and morning

stiffness, contractures, pain and anterior uveitis leading to blindness.¹⁶ This leads to suboptimal health-related quality of life (HRQL) in patients and parents or carers alike.^{17,18} Moreover, as JIA patients reach adulthood, they face possible continuing disease activity, medication-associated morbidity, life-long disability and the risk of emotional and social dysfunction.¹⁶

The baseline cohort population was defined as 12 year old “moderate-to-severe active polyarticular JIA [patients] who have had an insufficient response to other DMARD, including at least one TNF inhibitor” in the decision problem stated by BMS.

1.6 Institutional setting Where is/are the intervention(s) being evaluated usually provided?

The institutional setting appears to be paediatric secondary care, but this isn’t entirely clear. The delivery environment is not specifically referenced.

1.7 Country/ currency

Has a country setting been provided for the evaluation? What currency are costs expressed in and does the publication give the base year to which those costs relate?

The country setting given is UK. Costs are expressed in pounds sterling (£). Costs were derived from the MIMS database (accessed Nov. 2014). The price year was not explicitly stated.

1.8 Funding source

BMS

1.9 Analytical perspective

What is the perspective adopted for the evaluation (health service, health and personal social services, third party payer, societal (i.e. including costs borne by individuals and lost productivity)?

The model reports that it adopts an NHS and PSS perspective; however, it appears that only drug and administration costs have been included in the model. The NHS and PSS perspective generally includes costs associated with the disease. This would routinely include hospitalisation costs, costs for physician visits and nurse time, as well as costs for managing adverse events. This could also include reductions in costs due to temporary dose reductions and interruptions. The NHS and PSS perspective presented by BMS is much more limited than what is commonly presented in NHS economic evaluations.

2 Effectiveness

Were the effectiveness data derived from: a single study, a review/ synthesis of previous studies or expert opinion? Give the definition of treatment effect used in the evaluation. Give the size of the treatment effect used in the evaluation

No effectiveness data were used.

3 Intervention Costs

Were the cost data derived from: a single (observational) study, a review/ synthesis of previous studies expert opinion? Were the methods for deriving these data adequately described (give sources if using data from other published studies)? List the direct intervention costs and other direct costs used in the evaluation – include resource estimates (and sources for these estimates, if appropriate) as well as sources for unit costs used.

Intervention costs were derived from MIMS. A patient access scheme (PAS) was incorporated for abatacept ([REDACTED]). Sensitivity analyses were run for the price of tocilizumab using various percentage price discounts as a CIC PAS has been agreed for tocilizumab.

Drug	Cost	Dose	PAS Discount	PAS Cost
Abatacept	302.40	250mg	[REDACTED]	[REDACTED]

Etanercept	35.75	10mg		
	89.38	25mg		
	178.75	50mg		
Adalimumab	352.14	40mg		
Tocilizumab	102.40	80mg		
	256.00	200mg		
	512.00	400mg		
Methotrexate	14.85	7.5mg		
	15.29	10mg		
	16.50	12.5mg		
	16.57	15mg		
	17.50	17.5mg		
	17.84	20mg		
Administration Method		Costs		
Infusion		154.00		
Subcutaneous Injection		3.05		

3.1 Indirect Costs (costs due to lost productivity, unpaid inputs to patient care)

Were indirect costs included:

No.

4 Health state valuations/ utilities (if study uses quality of life adjustments to outcomes)

Were the utility data derived from: a single (observational) study, a review/ synthesis of previous studies expert opinion. Were the methods for deriving these data adequately described (give sources if using data from other published studies)?

No health valuations were undertaken.

4.1 List the utility values used in the evaluation

No health valuations were undertaken.

5 Modelling

If a model was used, describe the type of model used (e.g. Markov state transition model, discrete event simulation). Was this a newly developed model or was it adapted from a previously reported model? If an adaptation, give the source of the original. What was the purpose of the model (i.e. why was a model required in this evaluation)? What are the main components of the model (e.g. health states within a Markov model)? Are sources for assumptions over model structure (e.g. allowable transitions) reported – list them if reported.

The model is essentially a one state model where child height and weight change as they age which only affects the cost of drug doses. There are no health states.

5.1 Extract transition probabilities for [natural history/disease progression] model and show sources (or refer to table in text).

There was no natural history modelling.

5.2 What is the model time horizon?

The base-case model has a six year time-horizon. The time-horizon is user adjustable within the model by setting different exit ages.

5.3 What, if any, discount rates have been applied in the model? Same rate for costs and outcomes?

A discount rate of 3.5% annually has been applied to costs, in accordance with the NICE Reference Case.

5.4 If no economic evaluation was conducted, state the manufacturer's reasons for this.

A Cost minimisation analysis was conducted.

6 Results/ Analysis

What measure(s) of benefit were reported in the evaluation?

No benefit measure was evaluated.

6.1 Provide a summary of the clinical outcome/ benefits estimated for each intervention/ strategy assessed in the evaluation

No clinical outcomes nor benefit measures were evaluated.

6.2 Provide a summary of the costs estimated for each intervention/ strategy assessed in the evaluation

Discounted results for the base case (Table 13 in company submission)

Results for the base-case model (12 year olds, 6 year time horizon, from Excel model)				
	Abatacept	Adalimumab	Etanercept	Tocilizumab
Drug costs				
Administration costs	£11,797	£871	£871	£11,646
Total costs				
Cost savings with abatacept				

Undiscounted results for the base case (Table 12 in company submission)

Results for the base-case model (12 year olds, 6 year time horizon, from Excel model)				
	Abatacept	Adalimumab	Etanercept	Tocilizumab
Drug costs				
Administration costs	£13,040	£964	£964	£12,889
Total costs				
Cost savings with abatacept				

6.3 Synthesis of costs and benefits – are the costs and outcomes reported together (e.g. as cost-effectiveness ratios)? If so, provide a summary of the results.

A cost minimisation analysis was undertaken, so there was no synthesis of costs and benefits.

6.4 Give results of any statistical analysis of the results of the evaluation.

There were no statistical analyses of the results of the evaluation.

6.5 Was any sensitivity analysis performed – if yes, what type(s) (i.e. deterministic (one-way, two-way etc) or probabilistic).

A probabilistic sensitivity analysis was undertaken and scenario analyses were undertaken.

6.6 What scenarios were tested in the sensitivity analysis? How do these relate to structural uncertainty (testing assumptions over model structure such as relationships between health states), methodological uncertainty (such as choices of discount rate or inclusion of indirect costs) or parameter uncertainty (assumptions over values of parameters in the model, such as costs, quality of life or disease progression rates)?

- A sensitivity analysis was undertaken wherein the infusion costs for tocilizumab were increased due to the longer infusion time. This evaluates parameter uncertainty.
- The starting age of patients in the model was varied between 6 and 16 years. This related to the structural assumption of starting age. In the biological DMARD trials the mean age was 11 years at baseline, but the drug licenses were for much younger ages.
- The time horizon of the model was varied between 6 months and 20 years. Longer time horizons

were meant to represent that a third of children with JIA will have it continue into adulthood. It was unclear why shorter time horizons were tested.

d) PAS discount for tocilizumab. This represents parameter uncertainty in the cost of tocilizumab.

e) Exclude methotrexate from the etanercept arm. This scenario dabbles across all the types of uncertainty. NICE specifies that etanercept probably benefits from methotrexate being given concurrently, but the license is not for etanercept plus methotrexate, so there is some uncertainty in the appropriateness of methodology recommended by NICE. Changing a comparator is a structural modification and requires new parameter estimates.

6.7 Give a summary of the results of the sensitivity analysis – did they differ substantially from the base case analysis. If so, what were the suggested causes?

Adjusting the starting age of patients downward favoured etanercept with a starting age of 10 or under resulting in etanercept being cost saving compared to abatacept. Applying a PAS discount of █████ to tocilizumab makes the costs for the two drugs identical. Excluding methotrexate costs from the etanercept arm made etanercept cost-saving compared to abatacept. There were no suggested causes for any of the analyses, only a statement of the analysis results.

7 Conclusions/ Implications

Give a brief summary of the author's conclusions from their analysis

In the base case, abatacept is the least costly bDMARD and has similar efficacy and safety to other bDMARDs. These results remain stable for a wide range of scenarios.

7.1 What are the implications of the evaluation for practice?

There are no implications, because the economic evaluation did not evaluate practice, it only evaluated drug pricing. The model assumes that the drugs will have identical discontinuation rates, adverse events, and onset of effectiveness and duration of effectiveness. These assumptions are unlikely to be true. Even in an analysis that assumes there is no difference in effectiveness, differences in how the drugs behave in practice should be reflected in the costs.

8 SHTAC Commentary

Selection of comparators:

The comparators were consistent with the NICE Scope.

Validity of estimate of measure of benefit:

The assumption of equivalent efficacy was of unclear validity. While BMS provided justification for assuming equivalence, the nature of the data available may not justify this approach. The trials were small, but generally share many characteristics, and indirect comparisons were referenced by BMS and conducted by BMS. A full evaluation of adverse events using data beyond the clinical trials was not undertaken, and no comparisons of discontinuation rates were undertaken. The data from the trials was of insufficient quantity to make equivalency assumptions on event rates over time.

Given that there is a large amount of uncertainty in the effectiveness data, it may have been more appropriate to conduct a full economic evaluation with that uncertainty incorporated.

Validity of estimate of costs:

The costs were derived from appropriate sources, but BMS has assumed that the drugs will have identical adverse event costs and discontinuation rates (and identical everything else), both of which would lead to costs that have not been captured here. The strong assumption that there are no differences in the behaviour of the drugs in spite of different licenses and mechanisms of action lacks face validity.

Table: Critical appraisal checklist of economic evaluation (Questions in this checklist based on Philips et al and Drummond et al.)

	Item	MS 1
1	Is there a clear statement of the decision problem?	Yes
2	Is the comparator routinely used in UK NHS?	Yes
3	Is the patient group in the study similar to those of interest in UK NHS?	Yes
4	Is the health care system comparable to UK?	Yes
5	Is the setting comparable to the UK?	Yes
6	Is the perspective of the model clearly stated?	Yes
7	Is the study type appropriate?	No
8	Is the modelling methodology appropriate?	No
9	Is the model structure described and does it reflect the disease process?	? ^a
10	Are assumptions about model structure listed and justified?	Yes
11	Are the data inputs for the model described and justified?	Yes
12	Is the effectiveness of the intervention established based on a systematic review?	N/A
13	Are health benefits measured in QALYs?	N/A
14	Are health benefits measured using a standardised and validated generic instrument?	N/A
15	Are the resource costs described and justified?	Yes
16	Have the costs and outcomes been discounted?	No. ^b
17	Has uncertainty been assessed?	Yes
18	Has the model been validated?	Yes

1 Reference

Pfizer, 2015⁹⁷

1.1 Health technology

Etanercept

1.2 Interventions and comparators

What interventions/ strategies were included?

No economic evaluations were conducted, however a cost analysis was conducted comparing etanercept, adalimumab and tocilizumab

Was a no treatment/ supportive care strategy included?

No

Describe interventions/ strategies

No economic evaluation was conducted. The cost analysis included etanercept, adalimumab and tocilizumab. Etanercept is administered by subcutaneous injection at a recommended dose of 0.4 mg/kg (up to a maximum of 25 mg per dose), given twice weekly as a subcutaneous injection with an interval of 3-4 days between doses or 0.8 mg/kg (up to a maximum of 50 mg per dose) given once weekly.

1.3 Research question

What are the stated objectives of the evaluation?

No economic evaluation was conducted

1.4 Study type Cost-effectiveness/ cost-utility/ cost-benefit analysis?

No economic evaluation was conducted, however a cost analysis was conducted

1.5 Study population

What definition was used for [condition]? What are the characteristics of the baseline cohort for the evaluation?

The cost analysis was undertaken for a cohort with polyarticular JIA

1.6 Institutional setting Where is/are the intervention(s) being evaluated usually provided?

NHS outpatient setting

1.7 Country/ currency

Has a country setting been provided for the evaluation? What currency are costs expressed in and does the publication give the base year to which those costs relate?

No economic evaluation. The cost analysis was conducted in pounds sterling (£) but does not state the price year.

1.8 Funding source

Pfizer

1.9 Analytical perspective

What is the perspective adopted for the evaluation (health service, health and personal social services, third party payer, societal (i.e. including costs borne by individuals and lost productivity)?

No economic evaluation was conducted

2 Effectiveness

Were the effectiveness data derived from: a single study, a review/ synthesis of previous studies or expert opinion? Give the definition of treatment effect used in the evaluation. Give the size of the treatment effect used in the evaluation

No economic evaluation was conducted

3 Intervention Costs

Were the cost data derived from: a single (observational) study, a review/ synthesis of previous studies expert opinion? Were the methods for deriving these data adequately described (give sources if using data from other published studies)? List the direct intervention costs and other direct costs used in the evaluation – include resource estimates (and sources for these estimates, if appropriate) as well as sources for unit costs used.

No economic evaluation was conducted. The cost analysis used drug costs and administration costs for 1st year of treatment for different patient ages and weights. Cost sources are not given.

3.1 Indirect Costs (costs due to lost productivity, unpaid inputs to patient care)

Were indirect costs included:

No economic evaluation was conducted

4 Health state valuations/ utilities (if study uses quality of life adjustments to outcomes)

Were the utility data derived from: a single (observational) study, a review/ synthesis of previous studies expert opinion. Were the methods for deriving these data adequately described (give sources if using data from other published studies)?

No economic evaluation was conducted

4.1 List the utility values used in the evaluation

No economic evaluation was conducted

5 Modelling

If a model was used, describe the type of model used (e.g. Markov state transition model, discrete event simulation). Was this a newly developed model or was it adapted from a previously reported model? If an adaptation, give the source of the original. What was the purpose of the model (i.e. why was a model required in this evaluation)? What are the main components of the model (e.g. health states within a Markov model)? Are sources for assumptions over model structure (e.g. allowable transitions) reported – list them if reported.

No economic evaluation was conducted

5.1 Extract transition probabilities for [natural history/disease progression] model and show sources (or refer to table in text).

No economic evaluation was conducted

5.2 What is the model time horizon?

No economic evaluation was conducted

5.3 What, if any, discount rates have been applied in the model? Same rate for costs and outcomes?

No economic evaluation was conducted

5.4 If no economic evaluation was conducted, state the manufacturer's reasons for this.

The company submission notes the limitation raised in previous NICE submission TA35 and TA238. These relate to the limitations in the HRQoL data and the limited evidence on the long term outcomes and the effectiveness of the treatments. The company states that any cost-effectiveness evidence would be associated with considerable and unresolvable uncertainty and have therefore not submitted a cost-effectiveness model for this appraisal.

6 Results/ Analysis

What measure(s) of benefit were reported in the evaluation?

No economic evaluation was conducted

6.1 Provide a summary of the clinical outcome/ benefits estimated for each intervention/ strategy assessed in the evaluation

No economic evaluation was conducted

6.2 Provide a summary of the costs estimated for each intervention/ strategy assessed in the evaluation

No economic evaluation was conducted; The cost analysis shows the costs were similar between etanercept, adalimumab and tocilizumab.

	Etanercept†	Adalimumab	Tocilizumab‡
2 years	£1,859.00	£9,155.64	£5,000.19
3 years	£3,718.00	£9,155.64	£5,000.19
4 years	£3,718.00	£9,155.64	£5,665.79
5 years	£3,718.00	£9,155.64	£5,665.79
6 years	£3,718.00	£9,155.64	£6,331.39
7 years	£3,718.00	£9,155.64	£6,331.39
8 years	£3,718.00	£9,155.64	£6,996.99
9 years	£4,647.76	£9,155.64	£6,996.99

10 years	£4,647.76	£9,155.64	£6,996.99
11 years	£9,295.00	£9,155.64	£6,996.99
12 years	£9,295.00	£9,155.64	£8,328.19
13 years	£9,295.00	£9,155.64	£8,993.79
14 years	£9,295.00	£9,155.64	£8,993.79
15 years	£9,295.00	£9,155.64	£8,993.79
16 years	£9,295.00	£9,155.64	£10,324.99
17 years	£9,295.00	£9,155.64	£10,324.99

To reflect clinical practice and avoidance of drug wastage, doses were rounded down to the nearest available combination of vial strengths to a maximum of 10% variation from estimated dose.

†Where relevant the cheapest dosage regimen was assumed to be used in selecting between once weekly and twice weekly options. ‡Includes cost of administration in hospitals.

6.3 Synthesis of costs and benefits – are the costs and outcomes reported together (e.g. as cost-effectiveness ratios)? If so, provide a summary of the results.

No economic evaluation was conducted

6.4 Give results of any statistical analysis of the results of the evaluation.

No economic evaluation was conducted

6.5 Was any sensitivity analysis performed – if yes, what type(s) (i.e. deterministic (one-way, two-way etc) or probabilistic).

No economic evaluation was conducted

6.6 What scenarios were tested in the sensitivity analysis? How do these relate to structural uncertainty (testing assumptions over model structure such as relationships between health states), methodological uncertainty (such as choices of discount rate or inclusion of indirect costs) or parameter uncertainty (assumptions over values of parameters in the model, such as costs, quality of life or disease progression rates)?

No economic evaluation was conducted

6.7 Give a summary of the results of the sensitivity analysis – did they differ substantially from the base case analysis. If so, what were the suggested causes?

No economic evaluation was conducted

7 Conclusions/ Implications

Give a brief summary of the author's conclusions from their analysis

No economic evaluation was conducted

7.1 What are the implications of the evaluation for practice?

No economic evaluation was conducted

8 SHTAC Commentary

Selection of comparators:

No economic evaluation was conducted, the cost analysis did not include abatacept.

Validity of estimate of measure of benefit:

No economic evaluation was conducted

Validity of estimate of costs:

No economic evaluation was conducted. Costs used in the cost analysis appear reasonable.

1 Reference

Roche 2015⁷⁸

1.1 Health technology

Tocilizumab

1.2 Interventions and comparators

What interventions/ strategies were included?

Tocilizumab vs adalimumab

Was a no treatment/ supportive care strategy included?

No

Describe interventions/ strategies

Tocilizumab + MTX vs Adalimumab + MTX;

Tocilizumab only vs adalimumab only

1.3 Research question

What are the stated objectives of the evaluation?

To demonstrate the cost-effectiveness of tocilizumab when used in patients with pJIA who had an inadequate response to DMARDs

1.4 Study type Cost-effectiveness/ cost-utility/ cost-benefit analysis?

Cost utility

1.5 Study population

What definition was used for [condition]? What are the characteristics of the baseline cohort for the evaluation?

Patients entering the model have active JIA and have previously experienced an inadequate response to, or were intolerant of methotrexate (MTX). The modelled population is in line with the CHERISH trial population.

1.6 Institutional setting Where is/are the intervention(s) being evaluated usually provided?

NHS outpatient care

1.7 Country/ currency

Has a country setting been provided for the evaluation? What currency are costs expressed in and does the publication give the base year to which those costs relate?

UK. Costs have been taken from sources from year 2011-2015 with some costs taken from the Netherlands. Costs have not been inflated to a common base year.

1.8 Funding source

Roche

1.9 Analytical perspective

What is the perspective adopted for the evaluation (health service, health and personal social services, third party payer, societal (i.e. including costs borne by individuals and lost productivity)?

UK NHS and PSS

2 Effectiveness

Were the effectiveness data derived from: a single study, a review/ synthesis of previous studies or expert opinion? Give the definition of treatment effect used in the evaluation. Give the size of the treatment effect used in the evaluation

The company completed a systematic review of biologics in the treatment of JIA. The effectiveness data were derived from a WinBUGS indirect comparison with an order probit model. The results were in terms of level of ACR response.

		JIA ACR30	JIA ACR50	JIA ACR70	JIA ACR90
Without MTX	Placebo	31%	28%	25%	12%
	Tocilizumab	62%	59%	54%	35%
	Adalimumab	52%	49%	44%	26%
With MTX	MTX	52%	51%	41%	25%
	Tocilizumab	72%	70%	61%	44%
	Adalimumab	76%	75%	66%	49%

3 Intervention Costs

Were the cost data derived from: a single (observational) study, a review/ synthesis of previous studies expert opinion? Were the methods for deriving these data adequately described (give sources if using data from other published studies)? List the direct intervention costs and other direct costs used in the evaluation – include resource estimates (and sources for these estimates, if appropriate) as well as sources for unit costs used.

The costs associated with each health state was obtained from Prince et al (2011), who report costs data from the Dutch ABC register for the year before and after starting etanercept. The total 6 month health state cost for patients on treatment is £912.33 and off treatment is £1591.43.

The source of the treatment acquisition costs was not stated (assumed to be BNF).

Treatment	Dose1	Frequency	Unit cost (list price)
Adalimumab (40mg)	40 mg (assume wastage and all children receive 40mg vial)	Every 2 weeks	£352.14
Etanercept (10mg)	0.4mg/kg (max 25mg)	Twice a week	£35.75
Etanercept (25mg)			£89.38
Methotrexate (10mg, oral)	10mg	Every week	£0.56
Tocilizumab (80mg)	10 mg/kg for patients < 30 kg;	Every 4 weeks	£102.40
Tocilizumab (200mg)	8 mg/kg for patients ≥ 30 kg)		£256.00
Tocilizumab (400mg)			£512.00

3.1 Indirect Costs (costs due to lost productivity, unpaid inputs to patient care)

Were indirect costs included:

Indirect costs are not included

4 Health state valuations/ utilities (if study uses quality of life adjustments to outcomes)

Were the utility data derived from: a single (observational) study, a review/ synthesis of previous studies expert opinion. Were the methods for deriving these data adequately described (give sources if using data from other published studies)?

The company conducted a literature review that identified one study that reported utility values suitable for use in the model (Prince et al 2011).¹²³ This study reported utility scores obtained using the HUI3 questionnaire to JIA patients starting treatment with etanercept in the Dutch ABC register.

Based on these data, the company used values at time 0 for the patients who are off treatment, and used values at time 1 year for patients on treatment.

4.1 List the utility values used in the evaluation

On treatment: 0.7275

Off treatment: 0.53

Dead: 0

5 Modelling

If a model was used, describe the type of model used (e.g. Markov state transition model, discrete event simulation). Was this a newly developed model or was it adapted from a previously reported model? If an adaptation, give the source of the original. What was the purpose of the model (i.e. why was a model required in this evaluation)? What are the main components of the model (e.g. health states within a Markov model)? Are sources for assumptions over model structure (e.g. allowable transitions) reported – list them if reported.

A de novo Markov state transition model with 3 health states (uncontrolled disease / off treatment, on treatment and dead) was developed. The model has 6 month cycles. Patients start with uncontrolled disease at cycle 0 then move to first line treatment. Patients discontinue from treatment at a rate proportional to their response. Once all lines of treatment are exhausted, patients move into uncontrolled disease health state.

The model uses a 1% 6-month mortality rate.

5.1 Extract transition probabilities for [natural history/disease progression] model and show sources (or refer to table in text).

The 6-month discontinuation rate is 0.126 for no response, 0.09 for moderate response, and 0.042 for good response.

5.2 What is the model time horizon?

25 year time frame. The company states that this reflects the chronic nature of the disease and allows for all relevant costs and benefits to be included in the analysis.

5.3 What, if any, discount rates have been applied in the model? Same rate for costs and outcomes?

3.5% for costs and benefits

5.4 If no economic evaluation was conducted, state the manufacturer's reasons for this

Not applicable

6 Results/ Analysis

What measure(s) of benefit were reported in the evaluation?

Cost per QALY gained

6.1 Provide a summary of the clinical outcome/ benefits estimated for each intervention/ strategy assessed in the evaluation

Combination therapy			
	Adalimumab + MTX	Tocilizumab + MTX	
Total QALYs	18.76	18.72	
Monotherapy			
	Adalimumab	Tocilizumab	
Total QALYs	18.65	18.7	

6.2 Provide a summary of the costs estimated for each intervention/ strategy assessed in the evaluation

Combination therapy			
	Adalimumab + MTX	Tocilizumab + MTX	
Total Cost	£81,827	£70,707	
Monotherapy			
	Adalimumab	Tocilizumab	
Total Cost	£74,576	£68,560	

6.3 Synthesis of costs and benefits – are the costs and outcomes reported together (e.g. as cost-effectiveness ratios)? If so, provide a summary of the results.

	Combination therapy	Monotherapy
Incremental QALYs	(0.03)	0.0455
Incremental cost	(£11,120)	(6,015)
Incremental ICER	£280,370	Tocilizumab dominant

6.4 Give results of any statistical analysis of the results of the evaluation.

None reported

6.5 Was any sensitivity analysis performed – if yes, what type(s) (i.e. deterministic (one-way, two-way etc) or probabilistic).

None reported

6.6 What scenarios were tested in the sensitivity analysis? How do these relate to structural uncertainty (testing assumptions over model structure such as relationships between health states), methodological uncertainty (such as choices of discount rate or inclusion of indirect costs) or parameter uncertainty (assumptions over values of parameters in the model, such as costs, quality of life or disease progression rates)?

An exploratory analysis has been performed for tocilizumab versus etanercept. The analysis assumes a class effect across TNFs in pJIA. The analysis found that tocilizumab was a cost-effective alternative to etanercept.

6.7 Give a summary of the results of the sensitivity analysis – did they differ substantially from the base case analysis. If so, what were the suggested causes?

None reported

7 Conclusions/ Implications

Give a brief summary of the author's conclusions from their analysis

Adalimumab and tocilizumab have similar outcomes for patients with JIA, however tocilizumab is a less expensive alternative to adalimumab.

7.1 What are the implications of the evaluation for practice?

None

8 SHTAC Commentary

Selection of comparators:

Results not presented for tocilizumab compared to methotrexate only

Validity of estimate of measure of benefit:

Based on only utility estimates available for this population.

Validity of estimate of costs:

Based on relevant dataset of costs for patients on etanercept in Holland. May be differences in costs between countries.

Table: Critical appraisal checklist of economic evaluation (Questions in this checklist based on Philips et al and Drummond et al.)

	Item	Roche
1	Is there a clear statement of the decision problem?	Yes
2	Is the comparator routinely used in UK NHS?	Yes
3	Is the patient group in the study similar to those of interest in UK NHS?	Yes
4	Is the health care system comparable to UK?	? ^a
5	Is the setting comparable to the UK?	Yes
6	Is the perspective of the model clearly stated?	Yes
7	Is the study type appropriate?	Yes
8	Is the modelling methodology appropriate?	Yes
9	Is the model structure described and does it reflect the disease process?	Yes
10	Are assumptions about model structure listed and justified?	Yes
11	Are the data inputs for the model described and justified?	Yes
12	Is the effectiveness of the intervention established based on a systematic review?	Yes
13	Are health benefits measured in QALYs?	Yes
14	Are health benefits measured using a standardised and validated generic instrument?	Yes
15	Are the resource costs described and justified?	Yes
16	Have the costs and outcomes been discounted?	Yes
17	Has uncertainty been assessed?	No ^b
18	Has the model been validated?	No
^a Costs and utilities have been taken from a Dutch registry study ^b An exploratory analysis was conducted against etanercept		

Appendix 11 Parameters used in the independent model PSA

Parameter	Mean	Higher CI	Lower CI	standard error	distribution
Utility values					
No treatment	0.13	0.15	0.11	0.010	beta
Treatment 3 months	0.17	0.20	0.15	0.013	beta
Treatment 15 months phase	0.19	0.21	0.16	0.015	beta
Tx long term 27+ months	0.20	0.23	0.16	0.018	beta
Disease flare disutility	0.03	0.04	0.02	0.006	beta
Disease Flare					
Placebo	0.25	0.34	0.16	0.046	beta
Abatacept	0.09	0.16	0.05	0.021	beta
Adalimumab	0.14	0.23	0.09	0.028	beta
Etanercept	0.09	0.17	0.04	0.021	beta
Tocilizumab	0.14	0.20	0.09	0.025	beta
Adverse events 1st cycle					
Abatacept	0.53%	1.51%	0.00%	0.005	beta
Adalimumab	1.75%	3.71%	0.00%	0.010	beta
Etanercept	1.45%	4.19%	0.00%	0.014	beta
tocilizumab	1.60%	3.36%	0.00%	0.009	beta
Loss of efficacy					
Abatacept	9.47%	13.59%	5.36%	0.021	beta
Adalimumab	3.51%	6.25%	0.76%	0.014	beta
Etanercept	2.90%	6.82%	0.00%	0.020	beta
tocilizumab	7.98%	11.90%	4.06%	0.020	beta
Further line treatment					
Adverse events biological DMARD	0.43%	0.82%	0.04%	0.002	beta
Loss of efficacy biological DMARD	2.00%	2.59%	1.41%	0.003	beta
Adverse events MTX	0.58%	0.82%	0.34%	0.001	beta
Loss of efficacy MTX	0.42%	0.79%	0.05%	0.002	beta
Costs					
On bDMARD cost	£724	£724	£941	£507	gamma
Off bDMARD cost	£724	£724	£941	£507	gamma
Serious Adverse event cost	£1,533	£1,533	£1,993	£1,073	gamma
Disease flare cost	£430	£430	£559	£301	gamma

