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GRADE EVIDENCE PROFILES FOR CLINICAL EVIDENCE

1.1 PSYCHOSOCIAL INTERVENTIONS

1.1.1 Behavioural therapies aimed at communication

1.1.1.1 Natural language teaching compared with analog language teaching for communication in adults with autism

			Quality assessn	nent					Summary	of Findings	
Participants (studies) Follow up	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study event r	ates (%) With Natural	Relative effect (95% CI)	Anticipated at	Produte effects
							language teaching	language teaching		Analog language teaching	Natural language teaching (95% CI)
Communicat	tion (meas	sured with: Langu	age acquition n	neasured by nu	umber of noun	s generalized; Be	tter indicated b	y lower values)	1	
24 (1 study) 3 months	serious ¹	no serious inconsistency	no serious indirectness	very serious ^{2,3}	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, imprecision	11.5	11.5	-		The mean communication in the intervention groups was 0.71 standard deviations lower (1.55 lower to 0.13 higher)

¹ Non-randomised and non-blind so high risk of bias

² Study was designed to compare two alternative treatments and not to determine overall treatment efficacy

³ Small sample size

1.1.1.2 Observational studies of functional communication skills training in adults with autism

		(Quality assessm	ient				S	ummary o	f Findings	
Participants (studies) Follow up	Risk of bias	Inconsistency	Indirectness	-	Publication bias	Overall quality of evidence	Study ev With Control	vent rates (%) With Functional communication skills training	Relative effect (95% CI)	Anticipated a Risk with Control	Risk difference with Functional communication skills training (95% CI)
Communica	tion (meas	sured with: Vinel	and Adaptive B	ehaviour Scale	e (VABS) subs	cale of commun	ication; Be	etter indicated by lower	values)		
18 (1 study) 18 months	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, imprecision	-	18	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ Observational study and cannot extract efficacy data ² Small sample size

1.1.2 Facilitated communication

1.1.2.1 Observational studies of facilitated communication in adults with autism

		Ç	Quality assessm	ent				Su	mmary of	Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev	vent rates (%)	Relative effect	Anticipated	absolute effects
Follow up	and social	interaction resp	onses (measure	ed with: Behav		evidence ations; Better in	With Control	With Observational studies of facilitated communication for adults with autism	(95% CI)	Risk with Control	Risk difference with Observational studies of facilitated communication for adults with autism (95% CI)
		_						· · ·			
12 (1 study) 17 weeks	very serious ^{1,2}	no serious inconsistency	no serious indirectness	very serious ^{3,4}	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3,4} due to risk of bias, imprecision	-	12	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ No control group ² Efficacy data could not be extracted

³ Small sample size

⁴ Behavioural observations were non-blind

1.1.3 Behavioural therapies aimed at behaviour management

1.1.3.1 Independence training versus no-treatment control group in adults with intellectual disability

			Quality asses	sment					Summary o	of Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study even	t rates (%)	Relative effect	Anticipated	absolute effects
Follow up							With No treatment	With Behavioural therapies	(95% CI)	Risk with No treatment	Risk difference with Behavioural therapies (95% CI)
Activities of	daily livi	ng (showering) (r	neasured with	: Task-specifio	c checklist for	showering; Better in	dicated by lo	ower values)	•	•	
72 (1 study) 7 months	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	36	36	-		The mean activities of daily living (showering) in the intervention groups was 8.40 higher (6.99 to 9.81 higher)

¹ No attention-placebo control group so participants did not receive same care apart from intervention, and non-blind so risk of performance and detection bias

² Extrapolating from adults with learning disabilities

³ The outcome measure was designed specifically for this study and lacks formal assessments of reliability and validity

1.1.3.2 Observational studies of adaptive skills training in adults with intellectual disability

			Quality asses	ssment				:	Summary o	f Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	ent rates (%)	Relative effect	Anticipated	l absolute effects
Follow up							With Control	With Behavioural therapies	(95% CI)	Risk with Control	Risk difference with Behavioural therapies (95% CI)
Activities of	daily livin	ng (measured with	: Behaviour M	laturity Checklis	t II-1978 toileti	ng subscale; Better	indicated	by lower values)		<u> </u>	-
51 (1 study) 10 years	very serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, indirectness	-	51	-	See comment	See comment

¹ Observational study with no control group and efficacy data cannot be extracted ² Extrapolating from adults with learning disabilities

1.1.3.3 Behavioural weight control versus no treatment control in adults with intellectual disability

			Quality asses	ssment				S	ummary of	Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	-	Publication bias	Overall quality of evidence	Study even	t rates (%)	Relative effect	Anticipated	absolute effects
Follow up							With No treatment	With Behavioural therapies	(95% CI)	Risk with No treatment	Risk difference with Behavioural therapies (95% CI)
Self care (me	easured wi	th: Weight loss; B	etter indicated	by lower valu	ues)				•		
21 (1 study) 26 weeks	very serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	11	10	-		The mean self care in the intervention groups was 0.44 standard deviations higher (0.43 lower to 1.30 higher)

¹ Control group consisted of drop-outs from the experimental group so there was high risk for selection bias. The study was also non-randomised and non-blind increasing the risk of performance and detection bias

² Extrapolating from adults with learning disabilities

³ Small sample size

1.1.3.4 Observational studies of self-instructional pictorial child care manuals in adults with intellectual disability

			Quality asses	ssment					Summary	of Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	-	Publication bias	Overall quality of evidence	Study ev	ent rates (%)	Relative effect	Anticipated ab	solute effects
Follow up							With Control	With Behavioural therapies	(95% CI)	Risk with Control	Risk difference with Behavioural therapies (95% CI)
Parenting sk	ill (measu	red with: Target c	hild-care beha	viour checklis	t; Better indica	ated by lower values)	I			ļ	
10 (1 study) 3 years	very serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	10	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ Observational study and efficacy data cannot be extracted

² Extrapolating from adults with learning disabilities

³ Small sample size

1.1.4 Cognitive behavioural therapies

1.1.4.1 Cognitive behavioural therapy versus treatment-as-usual for coexisting conditions in adults with autism

			Quality assessn	nent				S	ummary of	Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study event	rates (%)	Relative effect	Anticipated a	bsolute effects
Follow up						evidence	With Treatment as usual	With Cognitive behavioural therapies	(95% CI)	Risk with Treatment as usual	Risk difference with Cognitive behavioural therapies (95% CI)
Severity of c	oexisting	condition (OCD)) (measured wit	h: Yale-Brown	Obsessive Co	mpulsive Scale ((YBOCS) seve	rity scale; Better ind	licated by l	ower values)	
24 (1 study) 16 months	serious ¹	no serious inconsistency	no serious indirectness	serious ²	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, imprecision	12	12	-		The mean severity of coexisting condition (ocd) in the intervention groups was 2.42 higher (3.6 lower to 8.44 higher)

¹ No attention-placebo control group so participants did not receive same care apart from intervention, and non-randomised and non-blind so risk of selection, performance and detection bias

² Small sample size

1.1.4.2 Cognitive behavioural therapy versus treatment-as-usual for anti-victimization skills in adults with intellectual disability

			Quality as	sessment					Summary o	of Findings	
()	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study event ra	ites (%)	Relative effect	Anticipated ab	solute effects
Follow up							With Treatment as usual	With Cognitive behavioural therapies	(95% CI)	Risk with Treatment as usual	Risk difference with Cognitive behavioural therapies (95% CI)
Anti-victimi	zation skil	lls (measured with	n: Self Social Int	erpersonal Decis	sion Making Sci	ale & The Protective Be	haviour Skills Ev	aluation; Better indic	ated by lowe	r values)	- <u>-</u>
80 (3 studies ¹) 3-9 weeks	serious ²	no serious inconsistency	serious ³	serious ⁴	undetected	⊕⊖⊖⊖ VERY LOW ^{2,3,4} due to risk of bias, indirectness, imprecision	40	40	-		The mean anti-victimization skills in the intervention groups was 1.07 standard deviations higher (0.58 to 1.56 higher)
	1	l ls (assessed with:		, 			- /10	- /			
38 (1 study)	serious ²	no serious inconsistency	serious ³	no serious imprecision	undetected	$\oplus \ominus \ominus \ominus$ VERY LOW ^{2,3}	7/18 (38.9%)	5/20 (25%)	RR 0.64 (0.25 to	Study populati	on
3 months		licolocieley		mprecision		due to risk of bias, indirectness		(2000)	1.67)	389 per 1000	140 fewer per 1000 (from 292 fewer to 261 more)
										Moderate	
										389 per 1000	140 fewer per 1000 (from 292 fewer to 261 more)

¹ 2 RCTs (KHEMKA2000 & KHEMKA2005) and 1 QE (MAZZUCCHELLI2001) combined

² No attention-placebo control group so participants did not receive same care apart from intervention, and non-blind so risk of performance and detection bias

³ Extrapolating from adults with learning disabilities

⁴ The precision of the outcome measures for KHEMKA2000 and KHEMKA2005 is unclear

1.1.4.3 Cognitive behavioural therapy versus waitlist control or treatment-as-usual for anger management in adults with intellectual disability

			Quality assess	sment				Su	mmary of 1	Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	quality of	Study event rat	tes (%)	Relative effect	Anticipated abs	olute effects
Follow up						evidence	With Waiting list or treatment as usual control	With Cognitive behavioural therapies	(95% CI)	Risk with Waiting list or treatment as usual control	Risk difference with Cognitive behavioural therapies (95% CI)
Anger mana	gement (r	neasured with: D	undee Provoc	ation Inventory	r, Anger Inven	tory, & Provoca	tion Inventory; B	Better indicated by	lower valu	es)	
169 (3 studies) 4-9 months	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, indirectness	70	99	-		The mean anger management in the intervention groups was 0.59 standard deviations lower (0.9 to 0.27 lower)

¹ No attention-placebo control group so participants did not receive same care apart from intervention, and non-randomised and non-blind so risk of selection, performance and detection bias

² Extrapolating from adults with learning disabilities

1.1.4.4 Cognitive behavioural therapy for anger management in adults with intellectual disability

			Quality asses	ssment				:	Summary o	of Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev		Relative effect	Anticipated al	osolute effects
Follow up							With Control	With Cognitive behavioural therapies	(95% CI)	Risk with Control	Risk difference with Cognitive behavioural therapies (95% CI)
Anger mana	gement (n	। neasured with: Ag	ı ggressive gestu	res on the vic	leotaped rolep	ı ılay test & Anger Inv	entory for	Mentally Retarded	Adults; Bet	ter indicated by	v lower values)
65 (2 studies) 19-27 weeks	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	65	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ Observational studies and cannot extract efficacy data

² Extrapolating from adults with learning disabilities

³ The precision of the outcome measure in BENSON1996 is unclear

1.1.5 Leisure programmes

1.1.5.1 Leisure programmes versus waitlist control in adults with autism

			Quality asses	sment			Summary of Findings				
Participants (studies) Follow up	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence		rent rates (%)	Relative effect (95% CI)	Anticipa Risk	ted absolute effects Risk difference with
ľ			Quality of Life Questionnaire-Spanish version (QOL); Better indicated by lower values) With With Leisure With With Leisure Control program versus waiting list contrination in adults with autism spectrum conditions	program versus waiting list control in adults with autism spectrum		with Control	Leisure program versus waiting list control in adults with autism spectrum conditions (95% CI)				
Quality of li	fe (measu	ured with: Quality	y of Life Questio	onnaire-Spanis	h version (QO	L); Better indicate	d by lowe	r values)			
71 (1 study) 1 years	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	undetected	$\oplus \oplus \oplus \bigoplus$ MODERATE ¹ due to risk of bias	34	37	-		The mean quality of life in the intervention groups was 8.33 higher (5.21 to 11.45 higher)
Emotion reco	ognition ((measured with: ⁷	The Facial Disci	rimination Batte	ery (FDB)-Spa	nish version - recc	ognition of	emotion subscale; Bett	er indicated	d by lower	values)
40 (1 study) 1 years	serious ¹	no serious inconsistency	no serious indirectness	serious ²	undetected	$\begin{array}{c} \bigoplus \bigoplus \bigoplus \bigoplus \bigoplus \\ \textbf{LOW}^{1,2} \\ \text{due to risk of} \\ \text{bias,} \\ \text{imprecision} \end{array}$	20	20	-		The mean emotion recognition in the intervention groups was 12.77 higher (2.12 to 23.42 higher)

¹ No attention-placebo control group which increases the risk of performance bias

² Small sample size

1.1.6 Social learning interventions

1.1.6.1 Emotion recognition training versus treatment-as-usual in adults with autism

			Quality assessn	nent			Summary of Findings					
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study event r	rates (%)	Relative effect	Anticipated at	osolute effects	
Follow up						evidence	With Treatment as usual	With Emotion recognition training	(95% CI)	Risk with Treatment as usual	Risk difference with Emotion recognition training (95% CI)	
Emotion reco	ognition (measured with: T	he Cambridge N	lindreading (CAM) Face tas	k; Better indicate	d by lower val	ues)		1		
40 (1 study) 15 weeks	serious ¹	no serious inconsistency	no serious indirectness	serious ²	undetected	⊕⊕⊖⊖ LOW ^{1,2} due to risk of bias, imprecision	22	18	-		The mean emotion recognition in the intervention groups was 2.70 higher (2.27 lower to 7.67 higher)	

¹ No attention-placebo control group so participants did not receive same care apart from intervention, and non-blind so risk of performance and detection bias ² Small sample size

1.1.6.2 Observational studies of social skills group in adults with autism

			Quality assessn	nent			Summary of Findings				
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	ent rates (%)	Relative effect	Anticipated abs	solute effects
Follow up							With Control	With Social skills group	(95% CI)	Risk with Control	Risk difference with Social skills group (95% CI)
Social intera	iction (mea	sured with: Empa	thy quotient and	role play 'part	y' scenario; Be	tter indicated by lov	wer values)			
23 (2 studies) 8-52 weeks	serious ¹	no serious inconsistency	no serious indirectness	serious ²	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, imprecision	-	23	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ Observational study and cannot extrapolate efficacy data

² Small sample size

1.1.6.3 Social skills group versus waitlist control group in adolescents with autism

			Quality asse	essment		Summary of Findings					
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study even	nt rates (%)	Relative effect	Anticipated a	absolute effects
Follow up							With With Waitlist Social control skills group		(95% CI)	Risk with Waitlist control	Risk difference with Social skills group (95% CI)
Social intera	ction (mea	asured with: Test o	of Adolescent S	Social Skills Ki	nowledge; Bet	ter indicated by lower	values)				
33 (1 study) 24 weeks	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	16	17	-		The mean social interaction in the intervention groups was 6.30 higher (4.32 to 8.28 higher)

¹ No attention-placebo control group so participants did not receive same care apart from intervention, and non-blind so risk of performance and detection bias

² Extrapolating from adolescents with autism spectrum conditions

³ Sample size is small

1.1.6.4 Observational studies of social skills groups for adolescents with autism

			Quality as	sessment					Summar	y of Findings	
Participants (studies) Follow up	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev (%)	ent rates	Relative effect (95% CI)	Anticipated at	osolute effects
							With Control	With Social skills group	-	Risk with Control	Risk difference with Social skills group (95% CI)
Social intera	ction (mea	sured with: Blind-	expert video r	ating and soci	al responsiver	ness/social skills rating scal	es; Better	indicated by	lower value	s)	
49 (3 studies) 2.5-11 months	serious ¹	serious ²	serious ³	serious ⁴	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3,4} due to risk of bias, inconsistency, indirectness, imprecision	-	49	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted
Challenging	behaviou	r (measured with:	Aberrant Beha	viour Checkli	st Irritability s	ubscale; Better indicated by	y lower va	lues)			
30 (1 study) 12 weeks	serious ¹	no serious inconsistency	serious ³	serious ⁴	undetected	⊕⊖⊖⊖ VERY LOW ^{1,3,4} due to risk of bias, indirectness, imprecision	-	30	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ Observational studies and efficacy data cannot be extracted

² HERBRECHT2009 and WEBB2004 found no significant treatment effects, while TSE2007 found a significant treatment effect (effect size 0.39)

³ Extrapolating from adolescents with autism spectrum conditions ⁴ Sample size is small

1.1.6.5 Social skills group versus treatment-as-usual in adults with intellectual disability

			Quality asses	ssment			Summary of Findings					
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study event r	ates (%)	Relative effect	Anticipated ab	solute effects	
Follow up							With Treatment as usual	With Social skills group	(95% CI)	Risk with Treatment as usual	Risk difference with Social skills group (95% CI)	
Challenging	; behaviou	r (measured with	: Part 2 of the A	AAMD Adapt	ive Behavior S	Scale; Better indicated	by lower value	es)				
44 (1 study) 10 weeks	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	24	20	-		The mean challenging behaviour in the intervention groups was 2.03 lower (11.79 lower to 7.73 higher)	

¹ No attention-placebo control group so participants did not receive same care apart from intervention, and non-blind so risk of performance and detection bias

² Extrapolating from adults with learning disabilities

³ Sample size is small

1.1.7 Supported employment programmes

1.1.7.1 Supported employment versus sheltered workshop in adults with autism

			Quality assessr	nent					Summary	of Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study event	rates (%)	Relative effect	Anticipated a	bsolute effects
Follow up Autistic behavior							With Sheltered workshop	With Supported work	(95% CI)	Risk with Sheltered workshop	Risk difference with Supported work (95% CI)
Autistic beh	aviours (n	neasured with: Cl	hildhood Autism	Rating Scale	(CARS); Better	indicated by low	er values)			<u> </u>	
51 (1 study) 3 years	serious ¹	no serious inconsistency	no serious indirectness	serious ²	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, imprecision	26	25	-		The mean autistic behaviours in the intervention groups was 6.07 lower (10.09 to 2.05 lower)
Quality of li	fe (measu	red with: Quality	of Life Survey (QLS); Better in	dicated by low	ver values)	I				
51 (1 study) 3 years	serious ¹	no serious inconsistency	no serious indirectness	serious ²	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, imprecision	26	25	-		The mean quality of life in the intervention groups was 5.20 higher (2.69 to 7.71 higher)

¹ Group allocation not randomised

² Sample size figures varied throughout the paper with no explanation as to the changing values. The sample sizes used for analysis were selected from the demographic table but not clear that this assumption valid or correct

1.1.7.2 Supported employment versus waitlist control in adults with autism

			Quality assessn	nent			Summary of Findings					
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study event	rates (%)	Relative effect	Anticipated a	bsolute effects	
Follow up							With Waiting list control	With Supported work	(95% CI)	Risk with Waiting list control	Risk difference with Supported work (95% CI)	
Executive fu	nction (me	easured with: Stoc	kings of Cambrid	dge (SOC) Pla	nning task from	n CANTAB; Bette	er indicated by	v lower values)	<u> </u>		-	
44 (1 study) 30 months	serious ¹	no serious inconsistency	no serious indirectness	very serious ^{2,3}	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, imprecision	22	22	-		The mean executive function in the intervention groups was 2.75 lower (4.41 to 1.09 lower)	

¹ Group allocation not randomised

² Sample size not reported for each group. Analysis based on assumption of equal numbers in each group but may be invalid.

³ Sample size is small

1.1.7.3 Supported employment versus treatment-as-usual control in adults with autism

			Quality assess	ment					Summary o	f Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study eve	nt rates (%)	Relative effect	Anticipated a	absolute effects
Follow up						evidence	With With Control Support group work	Supported	(95% CI)	Risk with Control group	Risk difference with Supported work (95% CI)
	nts (assess	ed with: Number o	of participants in	work)	T	1	1			1	
50 (1 study)	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	undetected	$ \bigoplus \Theta \Theta \Theta $ VERY LOW ¹	5/20 (25%)	19/30 (63.3%)	RR 2.53 (1.13 to	Study popula	ation
2 years		inconsistency	nunceries5	Imprecision		due to risk of bias	(20,0)	(00.070)	5.67)	250 per 1000	382 more per 1000 (from 32 more to 1000 more)
										Moderate	1
									250 per 10	250 per 1000	382 more per 1000 (from 32 more to 1000 more)

¹ Group allocation not randomised

1.1.7.4 Observational studies of supported employment in adults with autism

			Quality assessment					Summary of Findings					
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev	ent rates (%)	Relative effect	Anticipated ab	solute effects		
Follow up						evidence	With Control	With Supported work	(95% CI)	Risk with Control	Risk difference with Supported work (95% CI)		
Job placmen	ts (measur	red with: Number	of participants in	work; Better ind	dicated by lowe	er values)				•			
89 (1 study) 1 years	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	undetected	$\oplus \bigcirc \bigcirc$ VERY LOW ¹ due to risk of bias	-	89	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted		

¹ No control group and efficacy data cannot be extracted

1.1.8 Support for families and carers

1.1.8.1 Coping skills training programme versus treatment as usual for mothers of adolescents with autism

		Ģ	Quality assessm	ent				S	ummary of	f Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study event r	ates (%)	Relative effect	Anticipated at	solute effects
Follow up						evidence	With Treatment as usual	With Coping skills training program	(95% CI)	Risk with Treatment as usual	Risk difference with Coping skills training program (95% CI)
Social suppo	ort (measure	d with: Coping St	rategy Indicator	; Better indica	ted by lower v	values)	<u> </u>		<u> </u>	Į	
20 (1 study) 4 weeks	very serious ^{1.2,3}	no serious inconsistency	no serious indirectness	serious ⁴	undetected	$\oplus \ominus \ominus \ominus$ VERY LOW ^{1,2,3,4} due to risk of bias, imprecision	10	10	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted
Hopelessnes	s (measured	l with: Beck Hope	less Scale; Better	r indicated by	lower values)	1	1		Į	1	<u> </u>
20 (1 study) 4 weeks	very serious ^{1,2,3}	no serious inconsistency	no serious indirectness	serious ⁴	undetected	$\oplus \ominus \ominus \ominus$ VERY LOW ^{1,2,3,4} due to risk of bias, imprecision	10	10	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ Group allocation not randomised

² Efficacy data cannot be extracted

³ Short duration of follow-up

⁴ Small sample size

1.1.8.2 Psychoeducational group permanency planning intervention versus treatment as usual for mothers of adults with intellectual disability

		(Quality assess	ment				Sun	nmary of F	indings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study even	t rates (%)	Relative effect	Anticipated	absolute effects
Follow up			evidence With With Treatment Psychoeducation as usual group permanency planning intervention	(95% CI)	Risk with Treatment as usual	Risk difference with Psychoeducation group permanency planning intervention (95% CI)					
Knowledge	and awar	eness about plar	nning (measur	ed with: Clus	ter based on s	Landardized and	original scale	es; Better indicated by lo	ower value	s)	
27 (1 study) 6 weeks	very serious ¹	no serious inconsistency	serious ²	very serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	14	13	-		The mean knowledge and awareness about planning in the intervention groups was 0.99 standard deviations lower (1.79 to 0.19 lower)
Competence	and conf	idence to plan (1	measured with	n: Cluster base	ed on standard	lized and origina	al scales; Bette	er indicated by lower va	lues)		
27 (1 study) 6 weeks	very serious ¹	no serious inconsistency	serious ²	very serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	14	13	-		The mean competence and confidence to plan in the intervention groups was 1.36 standard deviations lower (2.20 to 0.53 lower)

Appraisals	of the plan	nning process (n	neasured with	: Cluster base	d on standard	ized and original	scales; Bett	er indicated by lov	wer values)	
27 (1 study) 6 weeks	very serious ¹	no serious inconsistency	serious ²	very serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	14	13	-	The mean appraisals of the planning process in the intervention groups was 0.61 standard deviations lower (1.39 lower to 0.1 higher)
Intermedia	te plannin	g behaviours (m	neasured with	Cluster based	l on standardi	zed and original	scales; Bette	er indicated by low	ver values)	
27 (1 study) 6 weeks	very serious ¹	no serious inconsistency	serious ²	very serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	14	13	-	The mean intermediate planning behaviours in the intervention groups was 0.49 standard deviations lower (1.25 lower to 0.28 higher)
Residentia	l and legal	planning (meas	ured with: Clu	ıster based on	standardized	and original scal	es; Better ir	dicated by lower	values)	
27 (1 study) 6 weeks	very serious ¹	no serious inconsistency	serious ²	very serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	14	13	-	The mean residential and legal planning in the intervention groups was 1.02 standard deviations lower (1.82 to 0.21 lower)

¹ Non-blind allocation, administration and assessment. Randomisation methods are unclear. It is not clear that the control group received the same care apart from the intervention. There was also a relatively short duration of follow-up and concerns regarding the reliability and validity of outcome measures

² Extrapolating from adults with intellectual disability

³ Small sample size and group N not clear (assumed N=13 in experimental and N=14 in control but not clear that this assumption is correct

1.2 BIOMEDICAL INTERVENTIONS

1.2.1 Antipsychotics: grade profiles

1.2.1.1 Risperidone versus placebo for behaviour management in adults with autism

			Quality assessme	ent					Sun	nmary of Fin	dings
(studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study eve	ent rates (%)	Relative effect	Anticipated	absolute effects
Follow up							With Placebo	With Risperidone	(95% CI)	Risk with Placebo	Risk difference with Risperidone (95% Cl)
Challenging	behaviour (m	easured with: Aber	rant Behaviour Ch	ecklist and SII	3-Q (Aggressio	n); Better indicated by	y lower val	ues)			
66 (2 studies) 12-22 weeks	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	undetected	⊕⊕⊕⊖ MODERATE ¹ due to imprecision	33	33	-		The mean challenging behaviour in the intervention groups was 0.79 standard deviations lower (1.29 to 0.28 lower)
Autistic beha	aviours (meas	ured with: Ritvo-Fr	eeman Real-life Ra	ting Scale; Bet	ter indicated by	v lower values)			•		
31 (1 study) 12 weeks	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	undetected	⊕⊕⊕⊖ MODERATE ¹ due to imprecision	16	15	-		The mean autistic behaviours in the intervention groups was 0.72 standard deviations lower (1.45 lower to 0.01 higher)
Core ASC sy	mptom (repet	itive behaviour) (m	neasured with: Yal	e-Brown Obse	ssive Compuls	on Scale; Better indic	ated by low	ver values)	I	<u>.</u>	
31 (1 study) 12 weeks	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	undetected	⊕⊕⊕⊖ MODERATE ¹ due to imprecision	16	15	-		The mean core asc symptom (repetitive behaviour) in the intervention groups was 0.94 standard deviations lower (1.68 to 0.19 lower)

Symptom sev	verity/improvo	ement (measured v	with: Clinical Glob	al Impression	(CGI) scale; Bet	ter indicated by lowe	er values)			
			no serious indirectness	serious ¹		⊕⊕⊕⊖ MODERATE ¹ due to imprecision	16	15	-	The mean symptom severity/improvement in the intervention groups was 1.40 standard deviations lower (2.18 to 0.61 lower)

¹ Sample size is small

1.2.1.2 Risperidone versus placebo for behaviour management in adults with intellectual disability

			Quality asse	essment					Summ	ary of Findi	ings
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	rent rates (%)	Relative effect	Anticipate	d absolute effects
Follow up							With Placebo	With Risperidone	(95% CI)	Risk with Placebo	Risk difference with Risperidone (95% CI)
Challenging	behaviou	ı ır (measured with	n: Aberrant Be	haviour Checkl	ist score (chall	enging behaviour); I	Better indi	cated by lower	values)	I	<u>,</u>
58 (1 study) 26 weeks	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊕⊖⊖ LOW ^{1,2} due to risk of bias, indirectness	29	29	-		The mean challenging behaviour in the intervention groups was 4.77 lower (18.38 lower to 8.84 higher)
Aggression (measured	with: Modified (Overt Aggress	ion Scale (MOA	S); Better indi	cated by lower value	es)				
58 (1 study) 26 weeks	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊕⊖⊖ LOW ^{1,2} due to risk of bias, indirectness	29	29	-		The mean aggression in the intervention groups was 0.58 higher (4.90 lower to 6.06 higher)

Symptom s	everity/im	provement (mea	sured with: Cl	inical Global In	pressions (CC	GI) Scale; Better indic	ated by lo	ower values)			
132 (2 studies) 4-26 weeks	serious ¹	serious ³	very serious ^{2,4}	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3,4} due to risk of bias, inconsistency, indirectness	66	66	-		The mean symptom severity/improvement in the intervention groups was 0.30 standard deviations lower (0.64 lower to 0.04 higher)
Quality of	l ife (measu	red with: Qualit	y of life question	onnaire; Better i	ndicated by lo	wer values)	Į		_	1	
58 (1 study) 26 weeks	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊕⊖⊖ LOW ^{1,2} due to risk of bias, indirectness	29	29	-		The mean quality of life in the intervention groups was 2.88 higher (2.56 lower to 8.32 higher)
Challengin	g behaviot	ır (narrative rep	orting) (measu	red with: Aberr	rant Behaviour	Checklist total score	e; Better i	ndicated by lov	ver values)	1	1
38 (1 study) 8 weeks	serious⁵	no serious inconsistency	serious ²	serious ⁶	undetected	⊕⊖⊖⊖ VERY LOW ^{2,5,6} due to risk of bias, indirectness, imprecision	19	19	-	See comment	See comment
Symptom s	everity/im	provement (narr	ative reporting	g) (measured w	ith: Clinical G	lobal Impressions (C	GI) scale;	Better indicate	ed by lower	values)	
	serious ⁵	no serious	serious ²	serious ⁶	undetected	$\oplus \ominus \ominus \ominus$	19	19	-	See	See comment

¹ Data is skewed in TYRER2008

² Extrapolating from a learning disabilities population

³ GAGIANO2005 found significant differences whereas TYRER2008 did not

⁴ Participants in GAGIANO2005 had co-existing conditions including conduct disorder, disruptive behaviour disorder, intermittent explosive disorder, oppositional defiant disorder, and antisocial personality disorder

⁵ The data reported does not allow for a calculation of effect size

⁶ Small sample size

1.2.1.3 Open-label risperidone for behaviour management in adults with intellectual disability

			Quality ass	sessment					Summary	y of Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study eve	ent rates (%)	Relative effect	Anticipated a	bsolute effects
Follow up							With Control	With Open-label risperidone	(95% CI)	Risk with Control	Risk difference with Open- label risperidone (95% CI)
Challenging	behaviour	(narrative reportin	g) (measured w	ith: Aberrant I	Behaviour Chee	L cklist (ABC); Better indicated	by lower v	values)			
24 (1 study) 76.4 days	very serious ¹	no serious inconsistency	very serious ^{2,3}	serious ⁴	undetected	$\oplus \Theta \Theta \Theta$ VERY LOW ^{1,2,3,4} due to risk of bias, indirectness, imprecision	-	24	-	See comment	See comment
Symptom sev	verity/outc	ome (narrative repo	orting) (measure	ed with: Clinic	al Global Impr	essions (CGI) scale; Better in	dicated by	lower values)	1	1	Į
24 (1 study) 76.4 days	very serious ¹	no serious inconsistency	very serious ^{2,3}	serious ⁴	undetected	$\bigcirc \bigcirc \bigcirc \bigcirc$ VERY LOW ^{1,2,3,4} due to risk of bias, indirectness, imprecision	-	24	-	See comment	See comment
Quality of lif	fe (measure	ed with: Composite	Autonomic Syn	nptom Scale (C	COMPASS) mo	dified version; Better indicate	ed by lower	r values)	1	1	<u> </u>
24 (1 study) 76.4 days	very serious ¹	no serious inconsistency	very serious ^{2,3}	serious ⁴	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3,4} due to risk of bias, indirectness, imprecision	-	24	-	See comment	See comment

¹ Observational study with open-label treatment and data extracted did not allow for calculation of effect sizes

² Extrapolating from adults with learning disabilities

³ Learning disabilities population also have co-existing psychiatric conditions including epilepsy and organic behaviour disorder

⁴ Small sample size

1.2.1.4 Haloperidol versus placebo for behaviour management in adults with autism

	_		Quality asse	ssment					Summary	of Findings	;
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	ent rates (%)	Relative effect	Anticipate	d absolute effects
Follow up							With Placebo	With Haloperidol	(95% CI)	Risk with Placebo	Risk difference with Haloperidol (95% CI)
Autistic beh	aviours (n	l neasured with: Chi	ldhood Autisr	n Rating Scale	: ; Better indica	ted by lower values)	1			<u> </u>	
33 (1 study) 21 weeks	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	$\oplus \ominus \ominus \ominus$ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	16	17	-		The mean autistic behaviours in the intervention groups was 2.70 lower (7.19 lower to 1.79 higher)
Side effects	(measured	with: Dosage Trea	atment Emerge	ent Symptom	Scale; Better ir	l ndicated by lower value	es)		<u> </u>	ļ	1
33 (1 study) 21 weeks	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	16	17	-		The mean side effects in the intervention groups was 1.50 higher (0.28 lower to 3.28 higher)

¹ High risk of attrition bias due to higher dropout as a consequence of side effects in the haloperidol group

² Sample is of adolescents with autism

³ Sample size is small

1.2.1.5 Haloperidol ve	ersus placebo for beha	viour management in adu	lts with intellectual disability
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			Quality asso	essment					S	ummary of F	indings
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study eve	ent rates (%)	Relative effect	Anticipated	l absolute effects
Follow up							With Placebo	With Haloperidol	(95% CI)	Risk with Placebo	Risk difference with Haloperidol (95% CI)
Challenging	behaviour	(measured with: A	Aberrant Behav	iour Checklist (A	BC); Better indic	cated by lower values	3)				
57 (1 study) 26 weeks	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊕⊖⊖ LOW ^{1,2} due to risk of bias, indirectness	29	28	-		The mean challenging behaviour in the intervention groups was 4.30 lower (19.30 lower to 10.70 higher)
Aggression ((measured	with: Modified Ov	ert Aggression	Scale; Better indic	ated by lower v	values)	<u> </u>		1	1	
57 (1 study) 26 weeks	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	$\oplus \oplus \ominus \ominus$ LOW ^{1,2} due to risk of bias, indirectness	29	28	-		The mean aggression in the intervention groups was 4.12 lower (8.53 lower to 0.29 higher)
Symptom se	verity/imp	rovement (measur	ed with: Clinica	l Global Impress	ions Scale (CGI)	- Improvement; Bett	er indicated	d by lower value	s)	1	1
57 (1 study) 26 weeks	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊕⊖⊖ LOW ^{1,2} due to risk of bias, indirectness	29	28	-		The mean symptom severity/improvement in the intervention groups was 0.88 lower (1.57 to 0.19 lower)
Quality of lit	fe (measure	ed with: Quality of	life questionna	ire; Better indicat	ed by lower val	ues)	ļ		1	<u> </u>	1
57 (1 study) 26 weeks	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊕⊖⊖ LOW ^{1,2} due to risk of bias, indirectness	29	28	-		The mean quality of life in the intervention groups was 1.87 lower (7.38 lower to 3.64 higher)

Appendix 19

¹ Data is skewed in TYRER2008

² Extrapolating from adults with learning disabilities

1.2.1.6 Zuclopenthixol versus placebo for behaviour management in adults with intellectual disability

			Quality assess	ment					Summar	y of Findir	ngs
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	ent rates (%)	Relative effect	Anticipa	ted absolute effects
Follow up							With Placebo	With Zuclopenthixol	(95% CI)	Risk with Placebo	Risk difference with Zuclopenthixol (95% CI)
Challenging	; behaviour	(aggression)									
39 (1 study)	no serious	no serious inconsistency	serious ¹	serious ²	undetected		1/20 (5%)	7/19 (36.8%)	RR 7.37 (1.2 to	Study po	pulation
18 weeks	risk of bias	licononocity				due to indirectness, imprecision		(2010 /2)	16.85)	50 per 1000	319 more per 1000 (from 10 more to 793 more)
						Imprecision				Moderate	2
										50 per 1000	319 more per 1000 (from 10 more to 793 more)
Challenging	; behaviour	(irritability) cha	nge from base	line (measured	with: Nurse's	Observation Scal	e for In-pa	atient Evaluation (1	NOISE-30);	Better ind	icated by lower values)
85 (1 study) 12 weeks	serious ³	no serious inconsistency	very serious ^{1,4}	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,3,4} due to risk of bias, indirectness	40	45	-		The mean challenging behaviour (irritability) change from baseline in the intervention groups was 2.20 lower (3.86 to 0.54 lower)

43 (1 study)	serious ³	no serious inconsistency	very serious ^{1,4}	serious ²	undetecte d	⊕⊖⊖⊖ VERY LOW ^{1,2,3,4}	1/19 5/24 (5.3 (20.8		R 3.96 .51 to	Study p	opulation
18 weeks						due to risk of bias, indirectness,	%)	, .	.47)	53 per 1000	156 more per 1000 (from 26 fewer to 656 more)
						imprecision				Modera	te
										50 per 1000	148 more per 1000 (from 25 fewer to 624 more)
Symptom	severity/imp	provement (change	e from baselin	e) (measured w	ith: Clinical (Global Impression	(CGI) scale;	; Better indicated by	v lower va	lues)	
85	serious ³	no serious inconsistency	very serious ^{1,4}	no serious imprecision	undetecte d		40 45	-			The mean symptom severity/improvement (change from baseline) in th

¹ Extrapolating from a learning disabilities population
 ² Sample size is small
 ³ Higher attrition rate in the placebo group
 ⁴ Study is very old

1.2.1.7 Prothipendyl versus placebo for behaviour management in adults with intellectual disability

			Quality asse	essment					Summary o	f Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	ent rates (%)	Relative effect	Anticipate	d absolute effects
Follow up							With Placebo	With Prothipendyl	(95% CI)	Risk with Placebo	Risk difference with Prothipendyl (95% CI)
Symptom sev	verity/imp	provement (assesse	ed with: Clinica	al observation	rating scale)	L					
39 (1 study)	serious ¹	no serious inconsistency	very serious ^{2,3}	serious ⁴	undetected	$ \bigoplus \ominus \ominus \ominus \\ \mathbf{VERY} \ \mathbf{LOW}^{1,2,3,4} $	9/19 (47.4%)	16/20 (80%)	RR 1.69 (1.04 to	Study pop	ulation
16 weeks						due to risk of bias, indirectness, imprecision	()		1.99)	474 per 1000	327 more per 1000 (from 19 more to 469 more)
										Moderate	<u> </u>
										50 per 1000	35 more per 1000 (from 2 more to 49 more)

¹ Pre-trial differences between experimental and control groups in IQ

² Extrapolating from adults with learning disabilities

³ Study is very old

⁴ Sample size is small

1.2.1.8 Pipamperone versus placebo for behaviour management in adults with intellectual disability

Quality assessment							Summary of Findings				
Participants (studies) Follow up	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study event rates (%)		Relative effect	Anticipated absolute effects	
							With Placebo	With Pipamperone	(95% CI)	Risk with Placebo	Risk difference with Pipamperone (95% CI)
Challenging behaviour (narrative reporting) (measured with: Experiment-specific behaviour checklist; Better indicated by lower values)											
20 (1 study) 4 months	very serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	10	10	-	See comment	See comment

¹ Data reported did not allow for calculation of effect size

² Extrapolating from a learning disabilities population

³ Small sample size

	_		Quality assess	sment					Summary	of Findings	
studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study event ra	ates (%)	Relative effect	Anticipated ab	solute effects
Follow up							With Haloperidol	With Cis(z)- clopenthixol	(95% CI)	Risk with Haloperidol	Risk difference with Cis(z)-clopenthixol (95% CI)
Symptom sev	verity/improv	ement (assessed w	ith: Clinical Glo	bal Impression (CGI) scale)						
98 1 study)	no serious risk of bias	no serious inconsistency	very serious ^{1,2}	no serious imprecision	undetected	$\oplus \oplus \ominus \ominus$ LOW ^{1,2}	7/49 (14.3%)	24/49 (49%)	RR 3.43 (1.86 to	Study populati	on
2 weeks				1		due to indirectness			5.02)	143 per 1000	347 more per 1000 (from 123 more to 574 more)
										Moderate	
					143 per 1000	347 more per 1000 (from 123 more to 575 more)					
Side effects (assessed with	: Clinical Global In	npression (CGI)	scale)							
98 1 study)	no serious risk of bias	no serious inconsistency	very serious ^{1,2}	no serious imprecision	undetected	$\oplus \oplus \ominus \ominus$ LOW ^{1,2}	39/49 (79.6%)	33/49 (67.3%)	RR 0.85 (0.57 to	Study populati	on
2 weeks	TISK OF DIAS	intonsistenty	Scrieus			due to indirectness	(12.0.10)	(07.070)	1.05)	796 per 1000	119 fewer per 1000 (from 342 fewer to 40 more)
										Moderate	
									796 per 1000	119 fewer per 1000 (from 342 fewer to 40 more)	

1.2.1.9 Cis(z)-clopenthixol versus haloperidol for behaviour management in adults with intellectual disability

¹ Extrapolating from a learning disabilities population

² Study is very old

1.2.1.10 Open-label olanzapine for behaviour management in adults with intellectual disability

			Quality asse	ssment			Summary of Findings					
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	ent rates (%)	Relative effect	Anticipated	absolute effects	
Follow up		Control label		With Open- label olanzapine	(95% CI)	Risk with Control	Risk difference with Open-label olanzapine (95% CI)					
Challenging	; behaviou	r (narrative report	i ng) (measured	d with: Aberra	ant Behaviour	Checklist (ABC); Better	r indicated	by lower values)		I		
16 (1 study) 8 weeks	very serious ¹	no serious inconsistency	very serious ^{2,3}	serious ⁴	undetected	$\oplus \ominus \ominus \ominus$ VERY LOW ^{1,2,3,4} due to risk of bias, indirectness, imprecision	-	16	-	See comment	See comment	
Symptom se	everity/out	come (narrative re	porting) (meas	sured with: Cl	inical Global I	mpressions (CGI) scale	; Better in	dicated by lower v	values)	1	<u></u>	
16 (2 studies) 8-11 weeks	very serious ¹	no serious inconsistency	very serious ^{2,3}	serious ⁴	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3,4} due to risk of bias, indirectness, imprecision	-	16	-	See comment	See comment	

¹ Observational studies with open-label treatment and data extracted did not allow for calculation of effect sizes

² Extrapolating from adults with learning disabilities

³ Learning disabilities population also have co-existing psychiatric conditions including disruptive behaviour disorder, attention-deficit/hyperactivity disorder, oppositional defiant disorder , stereotypic movement disorder, conduct disorder, impulse control disorder, epilepsy, and organic behaviour disorder ⁴ Small sample size

1.2.2 Anticonvulsants

1.2.2.1 Valproate versus placebo for behaviour management in children with autism

			Quality asses	sment					Sun	ımary of F	indings
Participants (studies) Follow up	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev (%)	rent rates	Relative effect (95% CI)	Anticipat	ted absolute effects
							With Placebo	With Valproate		Risk with Placebo	Risk difference with Valproate (95% CI)
Challenging	; behaviou	r (irritability) (me	easured with:	ABC Irritabilit	y and CGI-Irr	itability; Better indic	ated by lo	wer values)		<u> </u>	
57 (2 studies) 8-12 weeks	no serious risk of bias	serious ¹	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to inconsistency, indirectness, imprecision	25	32	-		The mean challenging behaviour (irritability) in the intervention groups was 0.05 standard deviations lower (0.58 lower to 0.48 higher)
Challenging	; behaviou	r (irritability) (as	sessed with: C	GI-Irritability))	I	I		I	I	
27 (1 study)	no serious	no serious inconsistency	serious ²	serious ³	undetected	$\oplus \oplus \ominus \ominus$ LOW ^{2,3}	1/11 (9.1%)	10/16 (62.5%)	RR 6.87 (1.59 to	Study po	pulation
12 weeks	risk of bias					due to indirectness, imprecision			10.36)	91 per 1000	534 more per 1000 (from 54 more to 851 more)
						Imprecision				Moderate	2
										91 per 1000	534 more per 1000 (from 54 more to 852 more)

(1 Study)	risk of	1				due to		. /	1.26)	786 per	149 more per 1000
30 (1 study)	no serious	no serious inconsistency	serious	serious ³	undetected	⊕⊕⊝⊝ LOW ³	11/14 (78.6%)	15/16 (93.8%)	RR 1.19 (0.73 to	Study po	-
Side effect	s (assessed	with: Checklist de	erived from F	'hysicians' De	sk Reference)	I					
8 weeks	risk of bias					due to indirectness, imprecision					improvement) in the intervention groups was 0.37 lower (0.97 lower to 0.23 higher)
30 (1 study)	no serious	no serious inconsistency	serious ²	serious ³	undetected	$ \bigoplus \bigoplus \ominus \ominus \\ LOW^{2,3} $	14	16	-		The mean symptom severity/improvement (cgi-
Symptom	severity/im	provement (CGI-	Improvemer	t) (measured	with: Clinical G	lobal Impressions	- Improvem	ent scale; B	etter indicate	ed by lowe	r values)
8 weeks	risk of bias	inconsistency				due to indirectness, imprecision					(aggression) in the intervention groups was 0.14 higher (2.93 lower to 3.21 higher)
(1 study)	no serious	no serious inconsistency	serious ²	serious ³	undetected	$\oplus \oplus \ominus \ominus$ LOW ^{2,3}	14	16	-		The mean challenging behaviou (aggression) in the intervention

¹ HELLINGS2005 found a negative response and HOLANDER2010 found a positive response for valproate on ABC irritability scores

² Extrapolation from children with ASC ³ Small sample sizes

1.2.2.2 Lamotrigine versus placebo for behaviour management in children with autism

			Quality asse	essment					Summary o	of Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	ent rates (%)	Relative effect	Anticipated	absolute effects
Follow up							With Placebo	With Lamotrigine	(95% CI)	Risk with Placebo	Risk difference with Lamotrigine (95% CI)
Autistic beh	aviours (n	arrative reporting)	(measured wi	ith: Childhood	l Autism Ratin	g Scale; Better indicated	l by lower	values)			
28 (1 study) 18 weeks	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	14	14	-	See comment	See comment
Challenging	behaviou	r (narrative reporti	i ng) (measured	d with: Aberra	ant Behaviour (l Checklist - Irritability; B	Better indica	ated by lower va	lues)		1
28 (1 study) 18 weeks	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	14	14	-	See comment	See comment

¹ Efficacy data could not be extracted

² Extrapolating from children with ASC

1.2.2.3 Open-label topiramate for behaviour management in children with autism

			Quality asse	ssment			Summary of Findings					
Participants (studies)	Risk of bias	Inconsistency	Indirectness	-	Publication bias	Overall quality of evidence	Study event rates (%)		Relative effect	t i		
Follow up							With Control	With Open- label topiramate	(95% CI)	Risk with Control	Risk difference with Open-label topiramate (95% CI)	
Challenging	; behaviou	r (narrative report	ing) (measure	ı d with: Conne	ers Parent Scale	e - Conduct subscale; B	etter indic	ated by lower valu	ıes)	1		
15 (1 study ¹) 25 weeks	very serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	15	-	See comment	See comment	

¹ Obervational case series and efficacy data could not be extracted

² Extrapolating from children with ASC ³ Small sample size

1.2.3 Drugs affecting cognition

1.2.3.1 Donepezil hydrochloride versus placebo for behaviour management in children with autism

		Ç	Juality assessi	nent			Summary of Findings					
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	ent rates (%)	Relative effect	Anticipate	d absolute effects	
Follow up							With Placebo	With Donepezil hydrochloride	(95% CI)	Risk with Placebo	Risk difference with Donepezil hydrochloride (95% CI)	
Autistic beh	aviours (me	asured with: Mod	ified parent-co	ompleted Chil	dhood Autism	Rating Scale (CAI	RS); Better	indicated by lower v	values)	1		
34 (1 study) 6 weeks	no serious risk of bias	no serious inconsistency	serious ¹	serious ²	undetected	⊕⊕⊖⊖ LOW ^{1,2} due to indirectness, imprecision	17	17	-		The mean autistic behaviours in the intervention groups was 0.40 higher (4.88 lower to 5.68 higher)	

¹ Extrapolating from children with autism spectrum conditions

1.2.3.2 Amantadine hydrochloride versus placebo for behaviour management in children with autism

l		Ç	Quality assessi	nent			Summary of Findings				
Participants (studies)	Risk of bias	Inconsistency	Indirectness	-	Publication bias	Overall quality of evidence	Study ev	ent rates (%)	Relative effect	Anticipate	d absolute effects
Follow up							With Placebo	With Amantadine hydrochloride	(95% CI)	Risk with Placebo	Risk difference with Amantadine hydrochloride (95% CI)
	[(irritability) (asse	ssed with: Abe		our Checklist (. undetected	ABC) parent-comp ⊕⊕⊝⊝	7/19	9/19	RR 1.29	Study pop	ulation
	risk of	inconsistency	serious	serious-	undelected	LOW ^{1,2}	(36.8%)	(47.4%)	(0.60 to	Study pop	
5 weeks	bias					due to indirectness, imprecision			2.74)	368 per 1000	107 more per 1000 (from 147 fewer to 641 more)
										Moderate	1
										368 per 1000	107 more per 1000 (from 147 fewer to 640 more)

¹ Extrapolating from children with autism spectrum conditions

1.2.3.3 Open-label memantine for behaviour management in children with autism

			Qualit	y assessment					Summar	y of Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study eve	ent rates (%)	Relative effect	Anticipated a	bsolute effects
Follow up						With With Control Memanti		With Memantine	(95% CI)	Risk with Control	Risk difference with Memantine (95% CI)
ASC core syn	ntpoms (co	mmunication) (mea	sured with: Cli	inical Global II	mpression Impr	 ovement Scale (CGI-Language); Bett	er indicated	d by lower values	5)		
	very serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	151	-	See comment	See comment
Challenging	behaviour	(measured with: CG	I-Improvemen	t Behaviour Sc	cale and Abbera	 nt Behaviour Checklist (ABC) Irritab	oility subsca	ale; Better indicat	ed by lower	values)	
	very serious ¹	no serious inconsistency	serious ²	serious ³	undetected	$\bigoplus \ominus \ominus \ominus$ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	165	-	See comment	See comment
Symptom sev	verity/impr	ovement (measured	with: Clinical	Global Impres	sions - Severity	scale (CGI-S); Better indicated by lov	wer values)	1			
	very serious ¹	serious ⁴	serious ²	serious ⁵	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,4,5} due to risk of bias, inconsistency, indirectness, imprecision	-	32	-	See comment	See comment

¹ No control group and efficacy data cannot be extracted

² Extrapolating from children with autism spectrum conditions

³ Clinical Global Impressions (CGI) scale usually used to rate symptom severity/improvement and not clear it is a precise enough scale to evaluate and differentiate language and behaviour scores as used in this study

⁴ ERICKSON2007 reports large treatment effect and OWLEY2006 reports non-significant treatment effect

1.2.3.4 Open-label galantamine for behaviour management in children with autism

			Quality ass	sessment					Summary	of Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study eve	ent rates (%)	Relative effect	Anticipated	absolute effects
Follow up							With Control	With Galantamine	(95% CI)	Risk with Control	Risk difference with Galantamine (95% CI)
Challenging	behaviour	(measured with: A	l. berrant Behavi	l iour Checklist	(ABC) Irritabil	l ity subscale; Better indicate	d by lower	r values)			
13 (1 study) 12 weeks	very serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	13	-	See comment	See comment
Autistic Beh	aviours (m	easured with: Child	lren's Psychiati	ric Rating Scal	e Autism Facto	r; Better indicated by lowe	r values)	•		<u> </u>	<u> </u>
13 (1 study) 12 weeks	very serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	13	-	See comment	See comment
Symptom se	verity/imp	rovement (measure	ed with: Clinica	l Global Impre	essions - Severi	ity scale (CGI-S); Better ind	icated by l	ower values)	1		
13 (1 study) 12 weeks	very serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	13	-	See comment	See comment

¹ No control group and efficacy data could not be extracted

² Extrapolating from children with autism spectrum conditions

1.2.4 Adrenocorticotrophic hormones

1.2.4.1 Adrenocorticotrophic hormone (ORG 2766) versus placebo for behaviour management in children with autism

			Quality asses	sment				Su	mmary of F	indings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	vent rates (%)	Relative effect	Anticipa	ted absolute effects
Follow up							With Placebo	With Adrenocorticotrophic hormone (ORG 2766)	(95% CI)	Risk with Placebo	Risk difference with Adrenocorticotrophic hormone (ORG 2766) (95% CI)
Challenging	behaviou	ır (social withdr	awal) (assessed	d with: Aberra	nt Behaviour	Checklist)	1		1	<u> </u>	
47 (1 study)	serious ¹	serious ²	serious ³	serious ⁴	undetected		4/18 (22.2%)	10/29 (34.5%)	RR 1.55 (0.57 to	Study po	opulation
6 weeks						due to risk of bias, inconsistency, indirectness,		`	4.22)	222 per 1000	122 more per 1000 (from 96 fewer to 716 more)
						imprecision				Moderat	e
										222 per 1000	122 more per 1000 (from 95 fewer to 715 more)
Challenging	behaviou	ır (social isolatio	on) (measured	with: General	Assessment F	arents Scale; Bette	r indicated	l by lower values)		<u> </u>	
20 (1 study) 36 weeks	no serious risk of bias	serious ²	serious ³	serious ⁴	undetected	⊕⊖⊖⊖ VERY LOW ^{2,3,4} due to inconsistency, indirectness, imprecision	10	10	-		The mean challenging behaviour (social isolation) in the intervention groups was 0.92479 standard deviations lower (1.82 to 0.02 lower)

Symptom se	everity/imj	provement (mea	sured with: Cl	inical Global I	mpression Sca	ale; Better indicated	l by low	er values)		
69 (2 studies) 6-36 weeks	serious ¹	no serious inconsistency	serious ³	no serious imprecision	undetected	⊕⊕⊖⊖ LOW ^{1,3} due to risk of bias, indirectness	29	40	-	The mean symptom severity/improvement in the intervention groups was 0.97 standard deviations lower (1.48 to 0.45 lower)

¹ Randomisation methods were unclear in BUITELAAR1996 (authors state 'randomised in principle' and there was a trend for group differences in age and CARS score at baseline ² BUITELAAR1992 found statistically significant treatment effects for challenging behaviour as measured by social isolation on the GAP, whereas BUITELAAR1996 found no

significant differences for social withdrawal as measured by ABC

³ Extrapolating from children with autism spectrum conditions

1.2.5 Secretin

1.2.5.1 Secretin versus placebo for autistic behaviours in children with autism

			Quality	assessment					:	Summary of I	indings
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study eve	nt rates (%)	Relative effect	Anticipated	absolute effects
Follow up							With Placebo	With Secretin	(95% CI)	Risk with Placebo	Risk difference with Secretin (95% CI)
Core ASC sy	mptom of	communication (1	neasured with:	Communication	and Symbolic B	ehaviour Scale and Presc	hool Language	Scale-3; Bett	er indicated	l by lower val	ues)
157 (2 studies) 3-8 weeks	serious ¹	serious ²	serious ³	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, inconsistency, indirectness	79	78	-		The mean core asc symptom of communication in the intervention groups was 0.29 standard deviations lower (0.77 lower to 0.2 higher)
Autistic beha	aviours (m	easured with: Chi	ldhood Autism	Rating Scale & R	eal Life Ritvo Be	ehaviour Scale; Better indi	icated by lowe	r values)	[<u> </u>
86 (2 studies) 3-8 weeks	serious ¹	no serious inconsistency	serious ³	no serious imprecision	undetected	$\oplus \oplus \ominus \ominus$ LOW ^{1,3} due to risk of bias, indirectness	43	43	-		The mean autistic behaviours in the intervention groups was 0.24 standard deviations lower (0.67 lower to 0.18 higher)
Challenging	behaviou	r (measured with:	Parent-complete	ed Global Behavi	our Rating Scale	es; Better indicated by low	ver values)		<u> </u>		1
62 (1 study) 8 weeks	serious ¹	no serious inconsistency	serious ³	no serious imprecision	undetected	$\oplus \oplus \bigcirc \bigcirc$ LOW ^{1,3} due to risk of bias, indirectness	31	31	-		The mean challenging behaviour in the intervention groups was 0.13678 standard deviations lower (0.64 lower to 0.36 higher)

¹ For LEVY2003 there was a significant difference between the groups in baseline CARS total score

² The studies found modest but non-significant effect sizes in different directions

³ Extrapolating from children with autism spectrum conditions

1.2.6 Melatonin

1.2.6.1 Open-label melatonin for insomnia in children with autism

			Quality asse	ssment		Summary of Findings					
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	ent rates (%)	Relative effect	Anticipated abs	olute effects
Follow up							With Control	With Melatonin	(95% CI)	Risk with Control	Risk difference with Melatonin (95% CI)
Sleep patter	ns (measure	ed with: Actigraph;	Better indicat	ed by lower v	alues)					L	
15 (1 study) 5 weeks	very serious ^{1,2}	no serious inconsistency	serious ³	serious ²	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	15	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ Open-label study with no control group and efficacy data cannot be extracted

² Small sample size

³ Extrapolating from children with autism spectrum conditions

1.2.7 Stimulants

1.2.7.1 Methylphenidate versus placebo for coexisting hyperactivity in children with autism

			Quality asses	sment					Summary	of Finding	s
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	ent rates (%)	Relative effect	Anticipat	ed absolute effects
Follow up							With Placebo	With Methylphenidate	(95% CI)	Risk with Placebo	Risk difference with Methylphenidate (95% CI)
Hyperactivit	y (measur	ed with: Aberran	l t Behaviour Cl	l necklist Hypera	ctivity subscal	le (parent-report);	Better indi	icated by lower value	es)		
62 (1 study) 5 weeks	no serious risk of bias	no serious inconsistency	serious ¹	no serious imprecision	undetected	⊕⊕⊕⊖ MODERATE ¹ due to indirectness	30	32	-		The mean hyperactivity in the intervention groups was 8.80 lower (13.72 to 3.88 lower)
Social intera	ction (init	iating joint atten	tion) (measure	ed with: Joint A	ttention Meas	ure from the EScs (Early Soci	al Communication S	Scales) (JAN	/IES); Bette	r indicated by lower values)
34 (1 study) 5 weeks	no serious risk of bias	no serious inconsistency	serious ¹	serious ²	undetected	⊕⊕⊖⊖ LOW ^{1,2} due to indirectness, imprecision	17	17	-		The mean social interaction (initiating joint attention) in the intervention groups was 6.50 higher (2.85 lower to 15.85 higher)
Repetitive b	ehaviour (measured with: (L Children's Yale	e-Brown Obsess	ive Compulsiv	ve Scales-PDD (CY	BOCS-PD	D); Better indicated	by lower v	alues)	
63 (1 study) 5 weeks	no serious risk of bias	no serious inconsistency	serious ¹	no serious imprecision	undetected	⊕⊕⊕⊖ MODERATE ¹ due to indirectness	31	32	-		The mean repetitive behaviour in the intervention groups was 0.92 lower (2.82 lower to 0.98 higher)

 1 Extrapolating from children with autism spectrum conditions 2 Small sample size

1.2.8 Antidepressants

1.2.8.1 Clomipramine versus placebo for autistic behaviours in adolescents with autism

			Quality assess	sment				Sur	nmary of F	indings	
-	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	vent rates (%)	Relative effect	Anticipa	ted absolute effects
Follow up							With Control	With Clomipramine versus placebo for behaviour management in adults with autism	(95% CI)	Risk with Control	Risk difference with Clomipramine versus placebo for behaviour management in adults with autism (95% CI)
Autistic beh	aviours (1	measured with: C	Childhood Aut	ism Rating Sca	ale (CARS); Be	etter indicated by	lower val	ues)		•	
32 (1 study) 21 weeks	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	16	16	-		The mean autistic behaviours in the intervention groups was 1.60 lower (7.07 lower to 3.87 higher)
Side effects	(global) (i	measured with: I	Dosage Treatm	ent Emergent	Symptom Sca	le (DOTES); Bette	r indicate	d by lower values)		<u> </u>	
32 (1 study) 21 weeks	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	16	16	-		The mean side effects (global) in the intervention groups was 1.20 higher (0.45 lower to 2.85 higher)

¹ Risk of attrition bias due to high drop out in the clomipramine group

² Sample includes children and adolescents with autism and mean age is 16 years

Appendix 19

			Quality assess	ment					Summary	of Findin	gs
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev	vent rates (%)	Relative effect	Anticipa	ted absolute effects
Follow up						evidence	With Control	With Fluvoxamine versus placebo for behaviour management in adults with autism	(95% CI)	Risk with Control	Risk difference with Fluvoxamine versus placebo for behaviour management in adults with autism (95% CI)
Core autistic	c sympton	n (repetitive beh	aviour) (measu	red with: Yale	e-Brown Obse	ssive Compulsiv	e Scale (Y-	BOCS); Better indicate	ed by lowe	r values)	
30 (1 study) 12 weeks	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ^{1,2}	undetected	⊕⊕⊖⊖ LOW ^{1,2} due to imprecision	15	15	-		The mean core autistic symptom (repetitive behaviour) in the intervention groups was 8.20 lower (13.92 to 2.48 lower)
Autistic beh	aviours (r	neasured with: F	Ritvo-Freeman I	Real-Life Ratir	ng Scale; Bette	r indicated by lov	ver values	3)			
30 (1 study) 21 weeks	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	undetected	⊕⊕⊕⊖ MODERATE ¹ due to imprecision	15	15	-		The mean autistic behaviours in the intervention groups was 0.82 standard deviations lower (1.56 to 0.07 lower)
Challenging	g behaviou	ur (aggression) c	hange-from-ba	seline (measu	red with: Bro	wn Aggression S	cale; Bette	r indicated by lower v	values)		1
30 (1 study) 21 weeks	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	undetected	⊕⊕⊕⊖ MODERATE ¹ due to imprecision	15	15	-		The mean challenging behaviour (aggression) change- from-baseline in the intervention groups was 0.92 standard deviations lower

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]	(1.68 to 0.17 lower)
Maladaptiv	ve behavio	ur (change from	baseline) (mea	sured with: V	ineland Adap	tive Behaviour Sc	ale; Bett	er indicated by lower v	values)		
30 (1 study) 21 weeks	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	undetected	⊕⊕⊕⊖ MODERATE ¹ due to imprecision	15	15	-		The mean maladaptive behaviour (change from baseline) in the intervention groups was 1.61 standard deviations lower (2.43 to 0.79 lower)
Symptom s	everity/im	provement (dich	notomous) (asse	essed with: Cl	inical Global I	mpressions (CGI) scale)				
30 (1 study)	no serious	no serious inconsistency	no serious indirectness	serious ¹	undetected	$\oplus \oplus \oplus \ominus$ MODERATE ¹	0/15 (0%)	8/15 (53.3%)	RR 17 (1.07 to	Study po	opulation
21 weeks	risk of bias					due to imprecision	()	· · ·	270.41)	0 per 1000	-
										Moderat	e
										0 per 1000	-
Symptom s	everity/im	provement (cont	t inuous) (meas	ured with: Cli	nical Global Ir	mpressions (CGI)	scale; B	etter indicated by lowe	r values)		1
30 (1 study) 21 weeks	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	undetected	$\oplus \oplus \oplus \bigcirc$ MODERATE ¹ due to imprecision	15	15	-		The mean symptom severity/improvement (continuous) in the intervention groups was 1.94 standard deviations lower (2.8 to 1.07 lower)

¹ Small sample size ² Y-BOCS scale valid and reliable for assessing severity of obsessive-compulsive symptoms in individuals with OCD but reliability and validity for assessing repetitive thoughts in autism is unknown

1.2.8.3 Open-label fluoxetine for behaviour management in adolescents with autism

			Quality assess	sment				S	Summary o	f Findings	
-	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	vent rates (%)	Relative effect	Anticipated	absolute effects
Follow up							With Control	With Open-label fluoxetine for behaviour management in adults with autism	(95% CI)	Risk with Control	Risk difference with Open-label fluoxetine for behaviour management in adults with autism (95% CI)
Symptom se	verity/im	provement (mea	sured with: Cl	inical Global I	mpressions (C	CGI) scale; Better i	ndicated l	by lower values)	I		
23 (1 study) 189 days	very serious ¹	no serious inconsistency	very serious ^{2,3}	serious ⁴	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3,4} due to risk of bias, indirectness, imprecision	-	23	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted
Compulsive	behaviou	ur (measured wit	h: Clinical Glo	bal Impression	ns (CGI) scale,	; Better indicated l	by lower v	values)			
23 (1 study) 189 days	very serious ¹	no serious inconsistency	very serious ^{2,3}	serious ⁴	undetected	$\bigcirc \bigcirc \bigcirc$ VERY LOW ^{1,2,3,4} due to risk of	-	23	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ No control group and efficacy data cannot be extracted

² The mean age is above 15 years but this is predominantly a child and adolescent sample

³ Participants also had coexisting psychiatric disorders

			Quality assessn	nent				Su	immary of	Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev	rent rates (%)	Relative effect	Anticipate	d absolute effects
Follow up						evidence	With Control	With Open-label sertraline for behaviour management in adults with autsim	(95% CI)	Risk with Control	Risk difference with Open-label sertraline for behaviour management in adults with autsim (95% CI)
Core autistic	sympton	n (repetitive beha	aviour) (measu	red with: Yale	Brown Obses	sive Compulsiv	e Scale (Y	-BOCS); Better indicated	by lower v	alues)	
37 (1 study) 12 weeks	very serious ¹	no serious inconsistency	no serious indirectness	very serious ²	undetected	$\oplus \ominus \ominus \ominus$ VERY LOW ^{1,2} due to risk of bias, imprecision	-	37	-	See comment	See comment
Autistic beh	aviours (r	neasured with: R	itvo-Freeman R	eal-Life Rating	g Scale; Better	indicated by lo	wer value	s)	1	1	
37 (1 study) 12 weeks	very serious ¹	no serious inconsistency	no serious indirectness	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,3} due to risk of bias, imprecision	-	37	-	See comment	See comment
Maladaptive	e behavior	ur (measured wit	h: Vineland Ad	aptive Behavi	our Scale; Bet	ter indicated by	lower valı	ues)	•	•	
37 (1 study) 12 weeks	very serious ¹	no serious inconsistency	no serious indirectness	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,3} due to risk of bias, imprecision	-	37	-	See comment	See comment

1.2.8.4 Open-label sertraline for autistic behaviours in adults with autism

Appendix 19

Symptom se	wmptom severity/improvement (measured with: Clinical Global Impressions global improvement item; Better indicated by lower values)														
37 (1 study) 12 weeks	2		no serious indirectness	serious ³		⊕⊖⊖⊖ VERY LOW ^{1,3} due to risk of bias, imprecision	-	37		See comment	See comment				

¹ No control group and efficacy data cannot be extracted ² Y-BOCS scale valid and reliable for assessing severity of obsessive-compulsive symptoms in individuals with OCD but reliability and validity for assessing repetitive thoughts in autism is unknown

1.2.9 Restrictive diets, vitamins, minerals and supplements

1.2.9.1 Gluten-and-casein-free diet versus treatment as usual for autistic behaviours in children with autism

			Quality asses	sment					Summar	Summary of Findings					
(studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study event	rates (%)	Relative effect	Anticipated a	bsolute effects				
Follow up							With Treatment as usual	With Gluten- and-casein- free diet	(95% CI)	Risk with Treatment as usual	Risk difference with Gluten-and-casein-free diet (95% CI)				
Autistic beha	aviours (s	ocial isolation an	d bizarre beh	aviours) (mea	sured with: D	iagnosis of Psychot	ic Behaviour	in Children; Be	tter indicat	ed by lower va	lues)				
20 (1 study) 1 years	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	10	10	-		The mean autistic behaviours (social isolation and bizarre behaviours) in the intervention groups was 5.60 lower (9.04 to 2.16 lower)				

¹ Risk of performance bias as unclear if intervention groups received same care apart from treatment, and participants receiving care and individuals administering care were not blind to group allocation

² Extrapolating from children with autism spectrum conditions

1.2.9.2 Open-label ketogenic diet for autistic behaviours in children with autism

			Quality asse	essment		Summary of Findings					
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	ent rates (%)	Relative effect	Anticipated ab	solute effects
Follow up							With Control	With Ketogenic diet	(95% CI)	Risk with Control	Risk difference with Ketogenic diet (95% CI)
Autistic beh	aviours (m	neasured with: Chi	ldhood Autisn	n Rating Scale	; Better indicat	ed by lower values)				1	
30 (1 study) 6 months	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	30	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ Observational study with no control group so high potential for bias and not possible to extract efficacy data

² Extrapolating from children with autism spectrum conditions

1.2.9.3 L-carnosine versus placebo for autistic behaviours in children with autism

			Quality asse	ssment					Summar	y of Findin	gs
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study eve	ent rates (%)	Relative effect	Anticipate	d absolute effects
Follow up							With Placebo	With L- Carnosine	(95% CI)	Risk with Placebo	Risk difference with L- Carnosine (95% CI)
Autistic beh	aviours (n	neasured with: Chi	l Idhood Autisr	n Rating Scale	; Better indica	ted by lower values)	Į	·		I	
31 (1 study) 8 weeks	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	17	14	-		The mean autistic behaviours in the intervention groups was 4.01 lower (9.03 lower to 1.01 higher)
Symptom in	nproveme	nt (measured with	: Clinical Glob	al Impressions	s improvemen	t scale; Better indicated	l by higher	values)	1		1
31 (1 study) 8 weeks	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	17	14	-		The mean symptom improvement in the intervention groups was 2.14 higher (0.99 lower to 5.27 higher)

¹ Baseline group differences in autistic behaviours as measured by the Gilliam Autism Rating Scale (GARS)

² Extrapolating from children with autism spectrum conditions

1.2.9.4 Micronutrient versus standard medication for autistic behaviours in children with autism

			Quality assess	sment					Summa	ary of Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study event r	ates (%)	Relative effect	Anticipated at	psolute effects
Follow up						evidence	With Standard medication	With Micronutrient	(95% CI)	Risk with Standard medication	Risk difference with Micronutrient (95% CI)
Autistic beha	aviours (m	neasured with: Cl	nildhood Autis	m Rating Scale;	; Better indicat	ed by lower valu	es)			<u> </u>	
88 (1 study ¹) 3-98 months	serious ²	no serious inconsistency	serious ³	no serious imprecision	undetected	$\bigcirc \bigcirc \bigcirc$ VERY LOW ^{2,3} due to risk of bias, indirectness	44		-		The mean autistic behaviours in the intervention groups was 0.50 higher (5.62 lower to 6.62 higher)
Challenging	behaviou	r (irritability) (m	easured with:	Aberrant Behav	viour Checklist	; Better indicated	by lower value	es)	Į	Į	
88 (1 study ¹) 3-98 months	serious ²	no serious inconsistency	serious ³	no serious imprecision	undetected	$\textcircled{\basis} \bigcirc \bigcirc \bigcirc$ VERY LOW ^{2,3} due to risk of bias, indirectness	44		-		The mean challenging behaviour (irritability) in the intervention groups was 7.40 lower (9.91 to 4.89 lower)
Symptom se	verity (me	asured with: Clir	nical Global Im	pressions sever	ity scale; Bette	r indicated by low	wer values)		ł	1	
88 (1 study ¹) 3-98 months	serious ²	no serious inconsistency	serious ³	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{2,3} due to risk of bias, indirectness	44		-		The mean symptom severity/improvement in the intervention groups was 1.38 lower (2.04 to 0.72 lower)

¹ case-control

² This is a non-randomized and non-blinded study so there is a high risk of bias ³ Extrapolating from children with autism spectrum conditions

1.2.9.5 Open-label iron supplementation for coexisting sleep problems in children with autism

			Quality asse	ssment					Summary	of Findings	
Participants (studies)	udies) bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	ent rates (%)	Relative effect	Anticipated ab	solute effects
Follow up							With Control	With Iron supplement	(95% CI)	Risk with Control	Risk difference with Iron supplement (95% CI)
Sleep patter	ns (measu:	red with: Restless	Sleep score; Be	tter indicated	by lower valu	ues)	I			1	
33 (1 study) 8 weeks	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	33	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted
Challenging	; behaviou	r (measured with:	Clinical Globa	ll Impressions	s - Irritability; I	Better indicated by low	er values)			I	
33 (1 study) 8 weeks	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	33	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ Observational study with no control group, no blinding, and a high attrition rate so there is potential for bias. It is also not possible to extract efficacy data ² Extrapolating from children with autism spectrum conditions

1.2.9.6 Open-label magnesium-vitamin B6 supplementation for core autistic symptoms in children with autism

			Quality asses	sment					Summary	of Findings	
Participants (studies)	Risk of bias	bias evidence	Study ev	ent rates (%)	Relative effect	Anticipated absolute effects					
Follow up							With Control	With Magnesium- vitamin B6	(95% CI)	Risk with Control	Risk difference with Magnesium-vitamin B6 (95% CI)
Core ASC sy	mptoms (s	ocial interaction,	communicatio	on, stereotype	d behaviour)	(measured with: DSN	I-IV clinic	al evaluation; Bette	er indicated	by lower value	s)
33 (1 study) 24 months	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	33	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted
Symptom se	everity/imp	rovement (measu:	red with: Beha	viour Summa	rized Evaluati	on; Better indicated b	y lower v	alues)	Į	<u> </u>	
11 (1 study) 14 weeks	very serious ^{1,4}	no serious inconsistency	very serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3,4} due to risk of bias, indirectness, imprecision	-	11	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ No control group results in high risk of bias and efficacy data cannot be extracted

² Extrapolating from children with autism spectum conditions

³ Small sample size

⁴ Sample selected for their previous sensitivity to the treatment

		Ç	Quality assessi	ment					Summary of	f Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev	rent rates (%)	Relative effect	Anticipat	ted absolute effects
Follow up						evidence	With Placebo	With Digestive enzyme supplementation	(95% CI)	Risk with Placebo	Risk difference with Digestive enzyme supplementation (95% CI)
Autsitic core	symptom	(communication) (measured w	rith: Language	e Developmen	t Survey (LDS) \	/ocabulary	score; Better indicate	d by lower v	alues)	
43 (1 study) 6 months	no serious risk of bias	no serious inconsistency	serious ¹	serious ²	undetected	⊕⊕⊖⊖ LOW ^{1,2} due to indirectness, imprecision	22	21	-		The mean autsitic core symptom (communication) in the intervention groups was 1.36 higher (15.74 lower to 18.46 higher)
Gastrointest	inal sympt	toms (measured v	with: Parent-ra	ted Additiona	l Rating Scale	e (ARS) gastrointe	estinal sym		er indicated l	oy lower va	alues)
43 (1 study) 6 months	no serious risk of bias	no serious inconsistency	serious ¹	serious ²	undetected	$\oplus \oplus \ominus \ominus$ LOW ^{1,2} due to indirectness, imprecision	22	21	-		The mean gastrointestinal symptoms in the intervention groups was 0.18 higher (0.27 lower to 0.63 higher)
Challenging	behaviou	r (measured with	: Parent-rated	Global Behavi	our Rating Sc	ale (GBRS); Bette	er indicated	l by higher values)		1	
43 (1 study) 6 months	no serious risk of bias	no serious inconsistency	serious ¹	serious ²	undetected	$\oplus \oplus \ominus \ominus$ LOW ^{1,2} due to indirectness, imprecision	22	21	-		The mean challenging behaviour in the intervention groups was 0.14 higher (0.19 lower to 0.47 higher)

1.2.9.7 Digestive enzyme supplementation versus placebo for behaviour management in children with autism

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 $^1\,\rm Extrapolating$ from children with autism spectrum conditions $^2\,\rm Small$ sample size

1.3 SETTINGS FOR CARE

1.3.1 Community based teams

1.3.1.1 Current living compared to developmental group home training environment for adults with intellectual disability

			Quality assess	sment				S	bummary o	of Findings	
-	Risk of bias	Inconsistency	Indirectness	-	Publication bias	Overall quality of evidence	Study event rates (%))	Relative effect	Anticipated absolute	effects
Follow up							With Developmental centre group home training environment	With Current living	(95% CI)	Risk with Developmental centre group home training environment	Risk difference with Current living (95% CI)
Community	living sk	ills (measured w	ith: Average n	umber of skil	ls gained acro	ss community liv	ing skills behavioural	domains; B	etter indica	ated by lower values)	
20 (1 study) 1 years	very serious ¹	no serious inconsistency	serious ²	very serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	10	10	-		The mean community living skills in the intervention groups was 8.90 higher (8.06 to 9.74 higher)

¹ Non-randomised allocation and non-blind assessment of outcome increasing the risk of selection and detction bias

² Extrapolating from adults with learning disabilities

³ The precision and reliability and validity of the outcome measure is unclear as under-specified and the sample size is small

1.3.1.2 Specialist behaviour therapy team compared with treatment as usual for adults with intellectual disability

			Quality asses	sment				St	ummary of	Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study event r	rates (%)	Relative effect	Anticipated al	bsolute effects
Follow up							With Treatment as usual	With Specialist behaviour therapy team	(95% CI)	Risk with Treatment as usual	Risk difference with Specialist behaviour therapy team (95% CI)
Challenging	behaviou	ır (lethargy/hype	ractivity) (mea	sured with: Ab	errant Behavic	ur Checklist (AB	C); Better indic	ated by lower value	ues)		
63 (1 study) 6 months	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	$\oplus \oplus \bigcirc \bigcirc$ LOW ^{1,2} due to risk of bias, indirectness	31	32	-	See comment	See comment
Challenging	behaviou	ur (irritability) (m	neasured with:	Aberrant Behav	viour Checklist	: (ABC); Better ind	licated by lowe	er values)	<u> </u>	1	
63 (1 study) 6 months	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	$\oplus \oplus \bigcirc \bigcirc$ LOW ^{1,2} due to risk of bias, indirectness	31	32	-	See comment	See comment

¹ Cannot extract data for efficacy as median values and interquartile ranges were reported. This may also imply that the data was skewed. We are thus restricted to analysing the results from this study via narrative review

² Extrapolating from adults with learning disabilities

1.3.1.3 Observational studies of specialist assessment and treatment units for adults with intellectual disability

			Quality asses	sment			Summary of Findings				
Participants (studies) Follow up	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev With Control	ent rates (%) With Specialist assessment and treatment unit	Relative effect (95% CI)	Anticipated a Risk with Control	Risk difference with Specialist assessment and treatment unit (95% CI)
Challenging	; behaviou	r (measured with	: Adaptive Beł	naviour Scale	Part II violent	behaviour domain; l	Better indi	icated by lower value	es)		
16 (1 study) 6 months	very serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	16	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ Small sample size and ABS data only available for half of the participants. There was also no control group and efficacy data could not be extracted

² Extrapolating from adults with learning disabilities

1.3.1.4 Liaison worker compared with treatment as usual for adults with intellectual disability

			Quality asse	ssment					Summary o	of Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study event ra	ntes (%)	Relative effect	Anticipated ab	solute effects
Follow up							With Treatment as usual	With Liason worker	(95% CI)	Risk with Treatment as usual	Risk difference with Liason worker (95% CI)
Access to ser	rvices (me	asured with: Num	ber of contacts	with services	; Better indicat	ted by lower values)	I		1	1	_
26 (1 study) 9 months	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	14	12	-	See comment	See comment

¹ Erfficacy data could not be extracted

² Extrapolating from adults with learning disabilities

1.3.2 Residential accommodation and related services

1.3.2.1 Residential institution compared with community housing for adults with intellectual disability

			Quality asses	ssment				S	Summary o	f Findings	
Participants (studies)	Risk of bias	bias of evidence With With		ates (%)	Relative effect	Anticipated at	osolute effects				
Follow up							With Community housing	With Residential institution	(95% CI)	Risk with Community housing	Risk difference with Residential institution (95% CI)
Residential	satisfactio	n - social life (m	easured with:	Satisfaction Qu	stionnaire of S	l Geltzer and Seltzer's	s (1978) Comm	unity Adjustmen	it Scale; Bet	ter indicated by	lower values)
29 (1 study) 0.1-8 years	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	15	14	-		The mean residential satisfaction - social life in the intervention groups was 5.80 higher (3.14 to 8.46 higher)
Residential	satisfactio	n - autonomy (m	leasured with:	Satisfaction Qu	stionnaire of s	Seltzer and Seltzer'	s (1978) Comm	unity Adjustmer	nt Scale; Be	tter indicated by	v lower values)
29 (1 study) 0.1-8 years	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	$\oplus \ominus \ominus \ominus$ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	15	14	-		The mean residential satisfaction - autonomy in the intervention groups was 1.20 lower

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										(2.28 to 0.12 lower)
Residential	satisfactio	n - total (measur	ed with: Satis	faction Qustion	naire of Seltze	r and Seltzer's (1978	8) Commun	ity Adjustment	Scale; Better ind	icated by lower values)
29 (1 study) 0.1-8 years	serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	15	14	-	The mean residential satisfaction - total in the intervention groups was 5.60 higher (1.1 to 10.1 higher)
Adaptive be indicated by	,		Adaptive Beha	viour Scale (AB	S), Vineland A	Adaptive Behaviour	Scales or a	modified version	on of the Behavio	our Development Survey; Better
224 (3 studies) 12-48 months	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, indirectness	103	121	-	The mean adaptive behaviour in the intervention groups was 0.48 standard deviations lower (0.75 to 0.2 lower)
Social skills	(measure	d with: Staff-rate	d social skills;	Better indicated	l by lower val	ues)			I	
100 (1 study) 30 months	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, indirectness	50	50	-	The mean social skills in the intervention groups was 5.10 lower (14.31 lower to 4.11 higher)
Quality of li	fe (measu	red with: Behavi	oural observat	ions of quality	of life ; Better i	indicated by lower	values)			
100 (1 study) 30 months	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, indirectness	50	50	-	The mean quality of life in the intervention groups was

											12.90 lower (16.05 to 9.75 lower)
Activity out	side the h	ome (measured v	with: Diary self	f-report on the 1	number of trip	s outside the home	e; Better ind	icated by lower va	lues)		•
	1	1					1			1	
36	serious1	no serious	serious ²	serious ³	undetected	$\oplus \Theta \Theta \Theta$	18	18	-		The mean activity
(1 study)		inconsistency				VERY LOW ^{1,2,3}					outside the home in
18 months						due to risk of					the intervention
						bias,					groups was
						indirectness,					3.00 lower
						imprecision					(6.99 lower to 0.99
						-					higher)

¹ Non-randomised allocation and non-blind assessment of outcome increases the risk of selection and detection bias

² Extrapolating from adults with learning disabilities ³ Small sample size

1.3.2.2 Small residential homes compared with institutions for adults with intellectual disability

			Quality asses	sment					Summary o	of Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study even	t rates (%)	Relative effect	Anticipated	absolute effects
Follow up							With Institution	With Small residential homes	(95% CI)	Risk with Institution	Risk difference with Small residential homes (95% CI)
Quality of li	fe (measu	red with: Quality	of Life Questio	onnaire (QOLQ)	; Better indicat	ted by lower value	es)			<u> </u>	
179 (1 study)	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, indirectness	76	103	-		The mean quality of life in the intervention groups was 11.40 higher (8.79 to 14.01 higher)
Choice maki	i ng (measi	ured with: Reside	nce Choice Ass	sessment Scale (RCAS); Better	indicated by lowe	r values)	•	_	<u> </u>	
179 (1 study)	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, indirectness	76	103	-		The mean choice making in the intervention groups was 36.60 higher (30.89 to 42.31 higher)
Community	inclusion	(measured with:	Use of Comm	unity Facilities S	Gcale (UCFS); B	l Setter indicated by	lower values	5)	_	ļ	
179 (1 study)	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, indirectness	76	103	-		The mean community inclusion in the intervention groups was 7.40 higher (4.86 to 9.94 higher)

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Contact with	1 family (r	neasured with: Fr	requency of fac	e-to-face visits;	Better indicate	d by lower values)			
179 (1 study)	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, indirectness	76	103	-	The mean contact with family in the intervention groups was 0.60 higher (0.36 to 0.84 higher)

¹ Non-randomised allocation of participants and significant group differences in adaptive/maladaptive behaviour

² Extrapolating from adults with learning disabilities

1.3.2.3 Dispersed supported living compared with residential homes for adults with intellectual disability

			Quality asses	sment				S	ummary of	Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study event 1	rates (%)	Relative effect	Anticipated ab	solute effects
Follow up						evidence	With Residential homes	With Dispersed supported living	(95% CI)	Risk with Residential homes	Risk difference with Dispersed supported living (95% CI)
Social inclus	sion (meas	sured with: Numb	per of commun	ity amenities us	sed in past mo	nths; Better indic	ated by lower	values)		I	
241 (1 study)	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, indirectness	138	103	-		The mean social inclusion in the intervention groups was 0.90 higher (0.43 to 1.37 higher)

¹ Limited data could be extracted from the study as a measure of variation (SD) was only reported for one scale item. Non-randomised allocation and non-blind assessment of outcome also increases the risk of selection and detection bias

² Extrapolating from adults with learning disabilities

1.3.2.4 Group homes compared with semi-independent apartments for adults with intellectual disability

			Quality asses	sment					Summary	v of Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study event rate	es (%)	Relative effect	Anticipated absol	ute effects
Follow up						evidence	With Semi- independent apartments	With Group home	(95% CI)	Risk with Semi- independent apartments	Risk difference with Group home (95% CI)
Resident sat	isfaction	measured with: I	Lifestyle satisfa	action scale (LSS	5); Better indic	ated by lower val	ues)		1		
204 (1 study) 1 years	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, indirectness	147	57	-		The mean resident satisfaction in the intervention groups was 8.72 lower (12.61 to 4.83 lower)

¹ Differences in sample sizes across groups, and significant differences in demographic factors found between groups, e.g. group home residents oldest, and participants in independent apartments had the highest mean score for adaptive behaviour and the lowest mean score for challenging behaviour which were not controlled for in statistical analysis. Non-randomisation and non-blind assessment of outcome also increases the risk of selection and detection bias

² Extrapolating from adults with learning disabilities

1.3.2.5 Intermediate care placement between institution and community compared with direct community placement for adults with intellectual disability

		Ģ	Quality assess	ment				Sun	nmary of F	indings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study event r	ates (%)	Relative effect	Anticipated at	osolute effects
Follow up	haviour (measured with: 4	AMD Adapti	ve Behaviour 9	Scala: Better in	evidence	With Direct community placement	With Intermediate care placement between institution and community	(95% CI)	Risk with Direct community placement	Risk difference with Intermediate care placement between institution and community (95% CI)
Adaptive be		measured with. <i>I</i>	Алири Ацари	ve benaviour a	Scale, Detter II	luicated by low	er values)				
57 (1 study) 1 years	serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, indirectness	39	18	-		The mean adaptive behaviour in the intervention groups was 5.89 higher (12.24 lower to 24.02 higher)

¹ Discrepancy in sample size between groups. Also non-randomised allocation and non-blind assessment of outcomes increases the risk of selection and detection bias ² Extrapolating from adults with learning disabilities

1.3.2.6 Person-centred compared with system-centred planning for adults with intellectual disability

			Quality asses	ssment				S	ummary of	Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study event ra	ates (%)	Relative effect	Anticipated a	bsolute effects
Follow up							With System- centred planning	With Person- centred planning	(95% CI)	Risk with System- centred planning	Risk difference with Person- centred planning (95% CI)
Movement in 37 (1 study)	nto comm serious ¹	unity (assessed w no serious inconsistency	rith: Number o serious ²	f participants serious ³	moving into c undetected	<pre>ommunity) ⊕⊖⊖⊖ VERY LOW^{1,2,3}</pre>	5/18 (27.8%)	18/19 (94.7%)	RR 3.41 (1.61 to	Study popula	tion
3 years		y				due to risk of bias, indirectness, imprecision		(*****)	7.24)	278 per 1000	669 more per 1000 (from 169 more to 1000 more)
										Moderate	-

¹ Allocation was not randomised increasing the risk of selection bias

² Extrapolating from adults with learning disabilities

³ Small sample size

1.3.2.7 Observational studies of the TEACCH approach in a residential setting for adults with autism

			Quality assess	ment					Summary o	of Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of	Study ev	ent rates (%)	Relative effect	Anticipated a	bsolute effects
Follow up						evidence	With Control	With TEACCH approach in residential setting	(95% CI)	Risk with Control	Risk difference with TEACCH approach in residential setting (95% CI)
Social abiliti	ies (measu	ured with: staff-re	port questionnai	re (based on Vi	neland Adapti	ve Behaviour Sca	les) and o	bservation checklist	; Better indi	icated by lower	values)
12 (1 study) 6 months	very serious ¹	no serious inconsistency	no serious indirectness	serious ²	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, imprecision	-	12	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted
Functional c	ommunic	ation (measured v	with: staff-report	questionnaire	(based on Vine	eland Adaptive B	ehaviour S	Scales) and observat	ion checklis	st; Better indica	ted by lower values)
12 (1 study) 6 months	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	undetected	$\bigoplus \bigcirc \bigcirc \bigcirc$ VERY LOW ¹ due to risk of bias	-	12	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ No control group and efficacy data could not be extracted. This study also used a small sample size ² Small sample size

1.3.2.8 Observational studies of the move from institutional to community settings for adults with intellectual disability

			Quality asse	ssment				5	Summary o	f Findings	
(studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	vent rates (%)	Relative effect	Anticipated a	bsolute effects
Follow up							With Control	With Move from institutional to community settings	(95% CI)	Risk with Control	Risk difference with Move from institutional to community settings (95% CI)
Challenging	behaviou	I Ir (measured with	n: Modified Ov	ert Aggression	Scale (MOAS)) and Problems Que	estionnair	e (PQ); Better indicate	ed by lower	values)	
329 (3 studies) 12-24 months	very serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2} due to risk of bias, indirectness	-	329	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted
Quality of Li	ife (measu	ured with: The Qu	lestionnaire or	Quality of Life	e; Better indica	ted by lower value	s)		<u> </u>	Į	
29 (1 study) 53 months	very serious ¹	no serious inconsistency	serious ²	serious ³	undetected	\bigcirc \bigcirc \bigcirc VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	29	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted
Family conta	i ct (measu	red with: Develo	pmental Disab	ilities Quality A	Assurance Que	estionnaire (DDQA)	Q); Better	indicated by lower v	alues)	ļ	
177 (1 study) 5 years	very serious ¹	no serious inconsistency	serious ²	no serious imprecision	undetected	$\begin{array}{c} {}{}{}{}{}{}{$	-	177	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

						bias, indirectness			
Adaptive B	ehaviour (1	neasured with: A	AMD's Adapt	ive Behaviour S	Scale (ABS) Pa	rt I total score; Bette	er indicated by lower values)	ł	L
32 (1 study) 5.5 years	very serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	- 32	5	Efficacy data cannot be extracted

¹ No control group and efficacy data could not be extracted ² Extrapolating from adults with learning disabilities ³ Small sample size

1.3.2.9 Observational studies of the move from more restrictive to less restrictive work or living environments for adults with intellectual disability

			Quality assess	sment				Si	ummary of	Findings	
Participants (studies)	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall quality of evidence	Study ev	rent rates (%)	Relative effect	Anticipated	absolute effects
Follow up							With Control	With Move from more restrictive to less restrictive work or living environments	(95% CI)	Risk with Control	Risk difference with Move from more restrictive to less restrictive work or living environments (95% CI)
Self-determi	nation (m	l neasured with: A	rcs's Self-Dete	rmination Sca	le: Adult Vers	ion; Better indicat	ed by low	er values)		1	
31 (1 study) 1 years	very serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	31	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted
Autonmous	functioni	ng (measured wi	ith: Autonomo	us Functionin	g Checklist (A	AFC); Better indica	ited by lov	ver values)		1	
31 (1 study) 1 years	very serious ¹	no serious inconsistency	serious ²	serious ³	undetected	⊕⊖⊖⊖ VERY LOW ^{1,2,3} due to risk of bias, indirectness, imprecision	-	31	-	Efficacy data cannot be extracted	Efficacy data cannot be extracted

¹ No control group and efficacy data could not be extracted

² Extrapolating from adults with learning disabilities

³ Sample size is small

GRADE EVIDENCE PROFILES FOR ECONOMIC EVIDENCE

Evide	ence prof	ile - econo	mic evidence				
Study & countr y	Limitatio ns	Applicabilit y	Other comments	Incrementa l cost (£)	Increment al effect	ICER (£/week employed)	Uncertainty
Guide line econo mic analys is	Potentiall y serious limitation s ¹	Directly applicable	Cost effectiveness analysis of employment support programme.Public sector perspective.	£6,921	44 weeks in employme nt	£158 per additional week of employm ent in the supported group	One-way sensitivity analysis to all parameters: Range of ICER: £80 to £338 per week in employment

1.3.2.10 Clinical / economic question: Employment support scheme versus standard care

¹ The data on rates of different types of accommodation of employed and unemployed is assumed. The model is extended to third year under certain assumptions. The standard service received by the control group not reported in MAWHOOD1999, it is assumed same as day service and adult education.

Evidence profile - economic evidence							
Study & countr y	Limitatio ns	Applicabilit y	Other comments	Incrementa l cost (£)	Incrementa l effect (QALYs)	ICER (£/QALY)	Uncertainty
Mawh ood and Howli n, 1999, UK	Potentially serious limitations 2	Directly applicable	RCT. NHS Perspective. Intervention costs of employment support only included – intervention costs of control group not estimated	Cost of intervention of additional person in the programme £25,451	38% NA	Cost of interventi on per additional adult employed: £33,474N A	Not reported

1.3.2.11 Clinical / economic question: Employment support scheme versus standard care

² Short time horizon, simple cost analysis, resource use or cost of standard service received by the control group not reported.