

Cochrane Database of Systematic Reviews

Omega-3 fatty acid addition during pregnancy (Review)

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[Intervention Review]

Omega-3 fatty acid addition during pregnancy

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ABSTRACT

Background

Higher intakes of foods containing omega-3 long-chain polyunsaturated fatty acids (LCPUFA), such as fish, during pregnancy have been associated with longer gestations and improved perinatal outcomes. This is an update of a review that was first published in 2006.

Objectives

To assess the effects of omega-3 LCPUFA, as supplements or as dietary additions, during pregnancy on maternal, perinatal, and neonatal outcomes and longer-term outcomes for mother and child.

Search methods

For this update, we searched Cochrane Pregnancy and Childbirth's Trials Register, ClinicalTrials.gov, the WHO International Clinical Trials Registry Platform (ICTRP) (16 August 2018), and reference lists of retrieved studies.

Selection criteria

Randomised controlled trials (RCTs) comparing omega-3 fatty acids (as supplements or as foods, stand-alone interventions, or with a cointervention) during pregnancy with placebo or no omega-3, and studies or study arms directly comparing omega-3 LCPUFA doses or types. Trials published in abstract form were eligible for inclusion.

Data collection and analysis

Two review authors independently assessed study eligibility, extracted data, assessed risk of bias in trials and assessed quality of evidence for prespecified birth/infant, maternal, child/adult and health service outcomes using the GRADE approach.

Main results

In this update, we included 70 RCTs (involving 19,927 women at low, mixed or high risk of poor pregnancy outcomes) which compared omega-3 LCPUFA interventions (supplements and food) compared with placebo or no omega-3. Overall study-level risk of bias was mixed, with selection and performance bias mostly at low risk, but there was high risk of attrition bias in some trials. Most trials were conducted in upper-middle or high-income countries; and nearly half the trials included women at increased/high risk for factors which might increase the risk of adverse maternal and birth outcomes.



Preterm birth < 37 weeks (13.4% versus 11.9%; risk ratio (RR) 0.89, 95% confidence interval (CI) 0.81 to 0.97; 26 RCTs, 10,304 participants; high-quality evidence) and **early preterm birth < 34 weeks** (4.6% versus 2.7%; RR 0.58, 95% CI 0.44 to 0.77; 9 RCTs, 5204 participants; high-quality evidence) were both lower in women who received omega-3 LCPUFA compared with no omega-3. **Prolonged gestation > 42 weeks** was probably increased from 1.6% to 2.6% in women who received omega-3 LCPUFA compared with no omega-3 (RR 1.61 95% CI 1.11 to 2.33; 5141 participants; 6 RCTs; *moderate-quality evidence*).

For infants, there was a possibly reduced risk of **perinatal death** (RR 0.75, 95% CI 0.54 to 1.03; 10 RCTs, 7416 participants; moderate-quality evidence: 62/3715 versus 83/3701 infants) and possibly fewer **neonatal care admissions** (RR 0.92, 95% CI 0.83 to 1.03; 9 RCTs, 6920 participants; *moderate-quality evidence* - 483/3475 infants versus 519/3445 infants). There was a reduced risk of **low birthweight** (LBW) babies (15.6% versus 14%; RR 0.90, 95% CI 0.82 to 0.99; 15 trials, 8449 participants; high-quality evidence); but a possible small increase in **large-for-gestational age** (LGA) babies (RR 1.15, 95% CI 0.97 to 1.36; 6 RCTs, 3722 participants; moderate-quality evidence, for omega-3 LCPUFA compared with no omega-3. Little or no difference in **small-for-gestational age or intrauterine growth restriction** (RR 1.01, 95% CI 0.90 to 1.13; 8 RCTs, 6907 participants; moderate-quality evidence) was seen.

For the **maternal outcomes**, there is insufficient evidence to determine the effects of omega-3 on **induction post-term** (average RR 0.82, 95% CI 0.22 to 2.98; 3 trials, 2900 participants; low-quality evidence), **maternal serious adverse events** (RR 1.04, 95% CI 0.40 to 2.72; 2 trials, 2690 participants; low-quality evidence), **maternal admission to intensive care** (RR 0.56, 95% CI 0.12 to 2.63; 2 trials, 2458 participants; low-quality evidence), or **postnatal depression** (average RR 0.99, 95% CI 0.56 to 1.77; 2 trials, 2431 participants; low-quality evidence). Mean **gestational length** was greater in women who received omega-3 LCPUFA (mean difference (MD) 1.67 days, 95% CI 0.95 to 2.39; 41 trials, 12,517 participants; moderate-quality evidence), and **pre-eclampsia** may possibly be reduced with omega-3 LCPUFA (RR 0.84, 95% CI 0.69 to 1.01; 20 trials, 8306 participants; low-quality evidence).

For the **child/adult outcomes**, very few differences between antenatal omega-3 LCPUFA supplementation and no omega-3 were observed in **cognition**, **IQ**, **vision**, **other neurodevelopment and growth outcomes**, **language and behaviour** (mostly low-quality to very low-quality evidence). The effect of omega-3 LCPUFA on **body mass index at 19 years** (MD 0, 95% CI -0.83 to 0.83; 1 trial, 243 participants; very low-quality evidence) was uncertain. No data were reported for development of **diabetes** in the children of study participants.

Authors' conclusions

In the overall analysis, **preterm birth < 37 weeks** and **early preterm birth < 34 weeks** were reduced in women receiving omega-3 LCPUFA compared with no omega-3. There was a possibly reduced risk of **perinatal death** and of **neonatal care admission**, a reduced risk of **LBW** babies; and possibly a small increased risk of **LGA** babies with omega-3 LCPUFA.

For our GRADE quality assessments, we assessed most of the important perinatal outcomes as high-quality (e.g. preterm birth) or moderate-quality evidence (e.g. perinatal death). For the other outcome domains (maternal, child/adult and health service outcomes) GRADE ratings ranged from moderate to very low, with over half rated as low. Reasons for downgrading across the domain were mostly due to design limitations and imprecision.

Omega-3 LCPUFA supplementation during pregnancy is an effective strategy for reducing the incidence of preterm birth, although it probably increases the incidence of post-term pregnancies. More studies comparing omega-3 LCPUFA and placebo (to establish causality in relation to preterm birth) are not needed at this stage. A further 23 ongoing trials are still to report on over 5000 women, so no more RCTs are needed that compare omega-3 LCPUFA against placebo or no intervention. However, further follow-up of completed trials is needed to assess longer-term outcomes for mother and child, to improve understanding of metabolic, growth and neurodevelopment pathways in particular, and to establish if, and how, outcomes vary by different types of omega-3 LCPUFA, timing and doses; or by characteristics of women.

PLAIN LANGUAGE SUMMARY

Omega-3 fatty acid addition during pregnancy

What is the issue?

Do omega-3 long chain polyunsaturated fatty acids (LCPUFA) taken during pregnancy - either as supplements or as dietary additions in food (such as some types of fish) - improve health outcomes for babies and their mothers? This is an update of a Cochrane Review that was first published in 2006.

Why is this important?

Preterm birth (babies born before 37 weeks pregnancy (gestation)) is a leading cause of disability or death in the first five years of life. Fish and fish oil contain omega-3 LCPUFA (particularly docosahexaenoic acid (DHA) and eicosapentaenoic acid (EPA)) and have been associated with longer pregnancies. So it is suggested that additional omega-3 LCPUFAs in pregnancy may reduce the number of babies born preterm and may improve outcomes for children and mothers. However, many pregnant women do not eat fish very often. Encouraging pregnant women to eat fatty fish (which generally have low toxin levels) or to use omega-3 LCPUFA supplements may improve children's and women's health. This is an update of a Cochrane Review that was first published in 2006.



What evidence did we find?

We searched for evidence in August 2018 and found 70 randomised controlled trials (RCTs; this type of trial provides the most reliable results) (involving 19,927 women). Most trials evaluated a group of women who received omega-3 LCPUFA and compared them with a group of women who received something that looked like omega-3 LCPUFA but did not contain it (placebo) or received no omega-3. The trials were mostly undertaken in upper-middle or high-income countries. Some studies included women at increased risk of preterm birth. The quality of the evidence from the included studies ranged from high to very low; this affected the certainty of the findings for different outcomes.

We found the incidence of preterm birth (before 37 weeks) and very preterm birth (before 34 weeks) was lower in women who received omega-3 LCPUFA compared with no additional omega-3. There were also fewer babies with low birthweight. However, omega-3 LCPUFA probably increased the incidence of pregnancies continuing beyond 42 weeks, although there was no difference identified in induction of labour for post-term pregnancies. The risk of the baby dying or being very sick and going to neonatal intensive care may be lower with omega-3 LCPUFA compared with no omega-3. We did not see any differences between groups for serious adverse events for mothers or in postnatal depression. Very few differences between the omega-3 LCPUFA groups and no omega-3 groups were observed in child development and growth.

Eleven trials reported that they had received industry funding. When we omitted these trials from the main outcomes (such as preterm birth and very preterm birth) it made very little, or no difference, to the results.

What does this mean?

Increasing omega-3 LCPUFA intake during pregnancy, either through supplements or in foods, may reduce the incidence of preterm birth (before 37 weeks and before 34 weeks) and there may be less chance of having a baby with a low birthweight. Women who take omega-3 LCPUFA supplements during pregnancy may also be more likely to have longer pregnancies. More studies are underway and their results will be included in a further update of this review. Future studies could consider if and how outcomes may vary in different populations of women, and could test different ways of increasing omega-3 LCPUFA during pregnancy.

SUMMARY OF FINDINGS

Summary of findings for the main comparison. Birth/infant outcomes

Omega-3 LCPUFA compared with no omega-3 during pregnancy: birth/infant outcomes

Population: pregnant women and their babies

Settings: Angola (1 RCT), Australia (1 RCT), Belgium (1 RCT), Canada (1 RCT), Chile (1 RCT), Croatia (1 RCT), Chile (1 RCT), Denmark (3 RCTs), Egypt (1 RCT), Germany (2 RCTs), India (1 RCT), Iran (3 RCTs), Italy (1 RCT), Mexico (1 RCT), Netherlands (3 RCTs), Norway (1 RCT), Russia (1 RCT), Sweden (1 RCT), Turkey (1 RCT), UK (4 RCTs), USA (8 RCTs)

Intervention: omega 3 Comparison: no omega-3

Outcomes	(007007)		Relative effect - (95% CI)	No of participants (studies)	Quality of the evidence	Comments
	Assumed risk	Corresponding risk	(33 / 0 Ci)	(Studies)	(GRADE)	
	Risk with no omega-3	Risk with omega-3				
Preterm birth < 37 weeks	134/1000	119 per 1000	RR 0.89 (0.81 to 0.97)	10,304 (26 RCTs)	$\oplus \oplus \oplus \oplus$	
Weeks		(109 to 130)			HIGH ¹	
Early preterm birth < 34 weeks	46/1000	27 per 1000	RR 0.58 (0.44 to 0.77)	5204 (9 RCTs)	$\oplus \oplus \oplus \oplus$	
		(20 to 35)			HIGH ²	
Perinatal death	20/1000	15 per 1000	RR 0.75 (0.54 to 1.03)	7416 (10 RCTs)	⊕⊕⊕⊝	
		(11 to 21)			MODERATE ³	
SGA/IUGR	129/1000	130 per 1000	RR 1.01 (0.90 to 1.13)	6907 (8 RCTs)	⊕⊕⊕⊝	
		(116 to 146)			MODERATE ³	
LBW	156/1000	140	RR 0.90 (0.82 to 0.99)	8449 (15 RCTs)	$\Theta \Phi \Phi \Phi$	
		(128 to 154)			HIGH	
LGA	117/1000	134 per 1000	RR 1.15 (0.97 to 1.36)	3722 (6 RCTs)	⊕⊕⊕⊝	
		(113 to 159)			MODERATE ⁴	

Serious adverse RR 0.72 (0.53 to 0.99) 63/1000 45 per 1000 (37 to 62) 2690 (2 RCTs) events for neonate/ low:5 infant

*The basis for the assumed risk (e.g. the median control group risk across studies) is provided in footnotes. The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: confidence interval; LBW: low birth weight LGA: large-for-gestational age;RCT: randomised controlled trial; RR: risk ratio; SGA/IUGR: small-for-gestational age/intrauterine growth restriction

GRADE Working Group grades of evidence

High quality: further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: we are very uncertain about the estimate.

- 1 Design limitations: larger studies of high quality, but some smaller studies with unclear risk of selective reporting and some smaller studies with unclear or high attrition bias at the time of birth (not downgraded for study limitations)
- ² Design limitations: larger studies of higher quality, but several studies with unclear or high attrition bias at the time of birth, or baseline imbalances (not downgraded for study limitations)
- ³ Imprecision (-1): downgraded one level due to crossing line of no effect and/or wide confidence intervals
- ⁴ Imprecision (-1): downgraded one level due to wide confidence intervals
- ⁵ Design limitations (-2): downgraded two levels; one study with unclear allocation concealment and attrition bias; specific adverse events not detailed in this study

Summary of findings 2. Maternal outcomes

Omega-3 LCPUFA compared with no omega-3 during pregnancy: maternal outcomes

Population: pregnant women

Settings: Angola (1 RCT), Australia (2 RCTs), Belgium (1 RCT), Brazil (1 RCT), Chile (1 RCT), Croatia (1 RCT), Denmark (3 RCTs), Egypt (1 RCT), Germany (3 RCTs), Hungary (1 RCT), Iran (5 RCTs), India (1 RCT), Italy (2 RCTs), Mexico (1 RCT), Netherlands (4 RCTs), Norway (2 RCTs), Russia (1 RCT), Scotland (2 RCTs), Spain (4 RCTs) Sweden (2 RCTs), Turkey (1 RCT), UK (3 RCTs) USA (12 RCTs), Venezuela (1 RCT)

Intervention: omega-3 Comparison: no omega-3

	Outcomes	Illustrative comparative risks* (95% CI)	Relative effect (95% CI)	No of partici- pants	Quality of the evidence	Comments
		Assumed risk Corresponding risk	(55 /0 C1)	(studies)	(GRADE)	
	Risk with no Risk with omega-3 omega-3					

Prolonged gestation > 42	16/1000	26/1000	RR 1.61 (1.11 to 2.33)	5141 (6)	000 0	
weeks	,	(18 to 37)	, ,	,	MODERATE ⁶	
Induction post-term	83/1000	68/1000	Average RR 0.82 (0.22 to 2.98)	2900 (3)	⊕⊕⊝⊝	
		(18 to 247)			LOW ⁷	
Pre-eclampsia	53/1000	44/1000	RR 0.84 (0.69 to 1.01)	8306 (20)	⊕⊕⊝⊝	Defined as hy-
		(37 to 53)			LOW ⁷	pertension with proteinuria
Gestational length		al age in the intervention group	Average MD 1.67 days (0.95 to	12,517 (41)	⊕⊕⊕⊝	
	was 1.67 days greate greater)	r (0.95 greater to 2.39 days	2.39)		MODERATE ⁸	
Maternal serious ad-	6/1000	6/1000	RR 1.04 (0.40 to 2.72)	2690 (2)	⊕⊕⊝⊝	
verse events		(2 to 16)			LOW ⁹	
Maternal admission to	1/1000	1/1000	RR 0.56 (0.12 to 2.63)	2458 (2)	⊕⊕⊝⊝	
intensive care		(0 to 3)			LOW ⁹	
Postnatal depression	112/1000	100	Average RR 0.99 (0.56 to 1.77)	2431 (2)	⊕⊕⊝⊝	
		(80 to 125)			LOW ¹⁰	

^{*}The basis for the assumed risk (e.g. the median control group risk across studies) is provided in footnotes. The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: confidence interval; RR: risk ratio

GRADE Working Group grades of evidence

High quality: further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: we are very uncertain about the estimate.

⁶ Design limitations (-1): downgraded one level due to some studies with attrition bias and some selective reporting bias; and some imprecision (not downgraded)

⁷ Design limitations (-1): downgraded one level for combined study limitations (mostly attrition bias and selective reporting bias); Imprecision (-1): downgraded one level due to confidence intervals including line of no effect

⁸ Design limitations (-1): downgraded one level for study limitations (mainly attrition bias): heterogeneity I² = 54%, but not downgraded due to use of a random-effects model

⁹ Imprecision (-2): downgraded two levels for wide confidence intervals and only 2 studies

10 Design limitations (-1): downgraded one level for study limitations (unclear randomisation in 1 study); downgraded one level for imprecision (wide confidence intervals; 2 studies only)

Summary of findings 3. Child/adult outcomes

Omega-3 LCPUFA compared with no omega-3 during pregnancy: child/adult outcomes

Population: children of women randomised to omega-3 or no omega-3 during pregnancy

Settings: Australia (2 RCTs), Bangladesh (1 RCT), Canada (1 RCT), Denmark (1 RCT), Hungary (1 RCT), Germany (1 RCT), Spain (2 RCTs), Mexico (1 RCT), Netherlands (1 RCT)

Intervention: omega-3

Comparison: no omega-3

Outcomes	Illustrative comparative risks* (9	Relative effect (95% CI)	No of partici- pants (studies)	Quality of the evidence	Comments	
	Assumed risk Corresponding risk			(3370 CI)	(GRADE)	
	Risk with no omega-3	Risk with omega-3				
Cognition:		ns in the intervention group was 0.37	MD -0.37 (-1.49 to 0.76)	1154 (4)	⊕⊕⊝⊝	
BSID II score at < 24 months	points lower in the intervention group (1.47 lower to 0.76 higher)		10 0.76)		LOW ¹¹	
Cognition:	The mean BSID III score at 24 months in the intervention group was 0.04		MD 0.04 (-1.59 to 1.68)	809 (2)	⊕⊕⊙⊙	
BSID III score at < 24 months	points nigher (1.59 lower to 1.68 ni	ints higher (1.59 lower to 1.68 higher)			LOW ¹²	
IQ: WASI at 7 years	, , ,		MD 1.00 (-0.79 to 2.79)	543 (1)	⊕⊕⊝⊝	
	mean in the control group (0.79 po	mean in the control group (0.79 points lower to 2.79 higher)			LOW ¹²	
IQ: WISC-IV at 12	The WISC-IV at 12 years in the intervention group was identical to in the control group (5.16 points lower to 7.16 higher)		MD 1.00 (-5.16	50 (1)	⊕⊝⊝⊝	
years			to 7.16)		VERY LOW ¹³	
Behaviour: BSID	-	our score in the intervention group at	MD -1.20 (-3.12	809 (2)	⊕⊕⊝⊝	At 12 months
III adaptive behav- iour score at 12-18 months	12-18 months was 1.20 points lower (3.12 lower to 0.72 higher)		to 0.72)		LOW ¹⁴	(one study), 18 months (one study)

Behaviour: SDQ To- tal Difficulties at 7 years	The mean SDQ total difficulties score at 7 years in the intervention group was 1.08 higher (0.18 higher to 1.98 higher)	MD 1.08 (0.18 to 1.98)	543 (1)	⊕⊕⊝⊝ LOW ¹²
BMI at 19 years	The mean BMI at 19 years in the intervention group was identical to that in the control group (0.83 lower to 0.83 higher)	MD 0 (-0.83 to 0.83)	243 (1)	⊕⊝⊝⊝ VERY LOW ¹⁵
Diabetes	Not reported			

^{*}The basis for the assumed risk (e.g. the median control group risk across studies) is provided in footnotes. The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

BMI: body mass index; BSID: Bayley Scales of Infant Development; CI: confidence interval; IQ: Intelligence Quotient; MD: mean difference; SDQ: Strengths and Difficulties Ouestionnaire; WASI: Weschler Abbreviated Scale of Intelligence; WISC: Weschler Intelligence Scale for Children

GRADE Working Group grades of evidence

High quality: further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: we are very uncertain about the estimate.

- 11 Design limitations (-1): downgraded one level due to unclear randomisation in 3 studies (that contributed 40% to meta-analysis) and some studies at high risk of attrition bias; Imprecision (-1): downgraded one level for wide confidence intervals including line of no effect
- 12 Imprecision (-2): downgraded one level for confidence intervals including line of no effect; and one level for small number of studies/single study
- 13 Design limitations (-1): downgraded one level for unclear selection bias (not clear if random sequence generated), possible attrition and/or reporting bias: Imprecision (-2): downgraded two levels for wide confidence intervals including line of no effect and 1 study with small number of participants
- 14 Design limitations (-1): downgraded one level for unclear randomisation (possible lack of allocation concealment), possible attrition and/or selective bias in 1 of the trials (contributing 15% to analysis); Imprecision (-1): downgraded one level for confidence intervals including line of no effect and few studies

Design limitations (-1): downgraded one level for unclear sequence generation and unclear blinding: Imprecision (-2): downgraded two levels for confidence intervals including line of no effect and 1 study with small number of participants

Summary of findings 4. Health service outcomes

Omega-3 compared with no omega-3 during pregnancy: health services outcomes

Population: pregnant women and their infants

Settings: Australia (1 RCT), Belgium (1 RCT), Denmark (2 RCTs), Egypt (1), Iran (2 RCTs), Italy (1 RCT), Netherlands (1 RCT), Norway (1 RCT), Russia (1 RCT), Scotland (1 RCT), UK (1 RCT), USA (5 RCTs)

Intervention: omega-3

Comparison: no omega-3

Outcomes	Illustrative comparative risks* (95% CI)		Relative effect - (95% CI)	No of partici- pants	Quality of the evidence	Comments
	Assumed risk	Corresponding risk	- (33% CI)	(studies)	(GRADE)	
	no omega-3	omega-3				
Maternal hospital ad-	273/1000	251/1000	RR 0.92 (0.81 to	2876 (5)	⊕⊕⊝⊝	
mission (antenatal)		(221 to 284)	1.04)		LOW ¹⁶	
Infant admission to	151/1000	139/1000	RR 0.92 (0.83 to	6920 (9)	⊕⊕⊕⊝	
neonatal care		(125 to 156)	1.03)		MODERATE ¹⁷	
Maternal length of hos-		the intervention group was 0.18	MD 0.18 (-0.20 to	2290 (2)	⊕⊕⊝⊝	
pital stay (days)	days greater (0.20 less to 0.	.57 days greater)	0.57)		LOW ⁸	
Infant length of hospital		the intervention group was 0.11	MD 0.11 (-1.40 to	2041 (1)	⊕⊕⊝⊝	
stay (days)	days greater (1.40 less to 1.	.oz uays greater)	1.62)		LOW ⁸	
Costs	Not reported					

^{*}The basis for the assumed risk (e.g. the median control group risk across studies) is provided in footnotes. The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: confidence interval; MD: mean difference; RR: risk ratio

GRADE Working Group grades of evidence

High quality: further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: we are very uncertain about the estimate.

¹⁶ Design limitations (-1): downgraded one level due to some studies with possible risk of attrition bias; Imprecision (-1): downgraded one level for confidence intervals including line of no effect

¹⁷ Imprecision (-1): downgraded one level for confidence intervals including line of no effect

¹⁸ Imprecision (-2): downgraded one level for confidence intervals including line of no effect and once for small number of studies



BACKGROUND

Description of the condition

Complications of pregnancy such as preterm birth, fetal growth restriction, postnatal depression and pre-eclampsia are relatively common and are associated with poorer outcomes for both the mother and child.

Of these, preterm birth has the highest burden of mortality and morbidity. Worldwide, approximately 15 million infants are born preterm (<37 weeks completed gestation) every year (World Health Organization 2017). National rates range from 5% to 18%, and are rising in most countries (World Health Organization 2017). Preterm birth is the leading cause of death in newborns, accounting for more than 85% of all perinatal complications and death (Thornton 2008). Preterm birth is also the leading cause of deaths in children under five years of age, with 1 million of the 5.9 million child deaths each year due to preterm birth complications (Liu 2016).

Advances in perinatal and neonatal care mean more preterm babies are surviving, but many of these infants go on to suffer the short-and long-term consequences of being born before their organs are mature (Saigal 2008). Infants born before 34 weeks often require intensive care and are at increased risk of respiratory distress syndrome, intraventricular haemorrhage, necrotising enterocolitis, blindness and cerebral palsy (Saigal 2008). In early childhood, developmental difficulties may emerge, with later societal and economic impacts due to low educational achievement, high unemployment, and deficits in social and emotional well-being (Westrupp 2014).

For mothers, postnatal depression is the most prevalent mood disorder associated with childbirth; symptoms include mood disturbances, sleep disturbances (not related to the infant), appetite disturbances or weight loss, and suicidal ideation. Systematic reviews report that nearly 20% of women experience depression within 12 weeks of giving birth (Gaynes 2005), with symptoms persisting beyond the first year in 8% of affected women (Dennis 2012). Postnatal depression impairs maternal social and psychological functioning with possible subsequent adverse effects on child development outcomes (Conroy 2012; Zhu 2014).

Fetal growth restriction is associated with stillbirth, neonatal death and perinatal morbidity and an increased risk of adverse health outcomes into adulthood (Stillbirth CRE 2018). Pre-eclampsia, characterised by high blood pressure and protein in the urine, can affect the kidneys, liver and blood-clotting systems and have serious life-threatening complications for the mother, such as eclampsia and can also result in preterm birth and fetal growth restriction (Mol 2016).

Description of the intervention

Maternal diet, including type and quantity of fat consumed, can have profound effects on pregnancy outcomes (Nordgren 2017). Omega-3 long chain polyunsaturated fatty acid status (LCPUFA) in pregnancy was first linked to longer gestation, higher birthweight and less preterm birth by researchers observing longer pregnancies among Faroe Islanders (who consume a diet high in fish) than the Danish population (Olsen 1985; Olsen 1986; Olsen 1991).

A prospective observational study in 8729 Danish women showed that reporting low consumption of fish in pregnancy was a strong risk factor for preterm and early preterm birth (Olsen 2002; Olsen 2006) particularly if low intake occurred during a prolonged period of pregnancy (Olsen 2006). A study pooling results from 19 European birth cohorts with over 150,000 mother-child pairs has subsequently shown an association between consumption of fish more than once a week by the mother and lower risk of preterm birth (Leventakou 2014), while a later study from Norway of over 67,000 women has also shown an association between increased fish consumption (particularly lean fish) and a lower prevalence of preterm birth (Brantsaeter 2017). Brantsaeter 2017 also examined the effect of omega-3 LCPUFA in the form of supplements, and found an association with reduced early, but not later, preterm birth. Observational studies have also shown links between fish consumption in pregnancy and child neurodevelopment (Hibbeln 2007).

In this review we have taken a comprehensive approach and specified any form or dose of omega-3 fatty acid as eligible, whether as fish or algal oil supplements, as food, or advice to consume particular foods rich in omega-3 LCPUFA (such as fish). We have also specified any type of omega-3 fatty acid (e.g. docosahexaenoic acid (DHA); eicosapentaenoic acid (EPA)); and any combination of omega-3 LCPUFAs as eligible. We have also included the omega-3 PUFA alpha-linolenic acid for completeness, although it is not a LCPUFA.

How the intervention might work

Consumption of omega-3 fatty acids during pregnancy and lactation, particularly those forms derived from fish or marine sources, are thought to influence a wide range of maternal, fetal, neonatal, and later outcomes. These include child growth and development outcomes (Borge 2017; Jensen 2006), preventing childhood allergies (see separate Cochrane Review - Gunaratne 2015), prevention of pre-eclampsia, decreasing maternal depression and anxiety (Golding 2009; Vaz Jdos 2013), and increasing gestational length (as discussed above).

When consumed in the diet, the essential fatty acid alpha-linolenic acid (ALA; 18:3 omega-3) can be converted to biologically active derivatives including eicosapentaenoic acid (EPA; 20:5 omega-3), docosapentaenoic acid (DPA; 22:5 omega-3) and docosahexaenoic acid (DHA; 22:6 omega-3). These fatty acids are precursors to a range of compounds that are known to minimise and help resolve inflammatory responses and oxidative stress (Leghi 2016). Pregnancy outcomes with an inflammatory component, such as preterm birth, are thought to be reduced by increasing omega-3 LCPUFA concentrations through including fish in the maternal diet or taking fish oil supplements. Maintaining a balance between the metabolites of omega-3 LCPUFA and the often pro-inflammatory omega-6 arachidonic acid is important in maintaining normal gestation length and is a critical element in cervical ripening and the initiation of labour (Zhou 2017). Adequate DHA, in particular, is thought to be crucial in fetal and early-life brain development (Shulkin 2018).

Fish and seafood are the richest dietary sources of DHA (Greenberg 2008). However, fish consumption is low in many countries, and women of childbearing age may be reluctant to increase their fish intake due to perceptions that mercury and other pollutants in fish may affect their unborn child (Oken 2018). For example only 10%



of women of childbearing age in Australia meet the recommended docosahexaenoic acid (DHA) intake (Koletzko 2007; Meyer 2016), which includes fish as well as fish oil supplementation. Many pregnant women are likely to have low concentrations of omega-3 LCPUFA and may benefit from increasing DHA in their diet, either from food sources or as supplements.

Why it is important to do this review

Over the last 40 years, a slew of observational studies, randomised trials and reviews addressing omega-3 fatty acids and pregnancy (e.g. Newberry 2016), involving hundreds of thousands of women, have been published. However many of these studies and reviews have concentrated on a particular focus such as allergy or child development, and reported only a selection of outcomes. Some outcomes such as preterm birth have not always been reported, despite the growing realisation that omega-3 LCPUFA supplementation may have a role in preventing it. Furthermore, studies and reviews on omega-3 LCPUFAs in pregnancy have differed in their findings and conclusions (e.g. Saccone 2016), sometimes due to selective reporting and other methodological issues.

Therefore, a comprehensive systematic review of omega-3 fatty acids in pregnancy that covers all relevant maternal, perinatal and child outcomes (except allergy which is covered in Gunaratne 2015), all forms of omega-3 fatty acids, and comparisons of doses, timing and types of omega-3 fatty acids is required.

OBJECTIVES

To assess the effects of omega-3 LCPUFA, as supplements or as dietary additions, during pregnancy on maternal, perinatal, and neonatal outcomes and longer-term outcomes for mother and child.

METHODS

Criteria for considering studies for this review

Types of studies

Randomised controlled trials (RCTs), including quasi-randomised trials, and trials published in abstract form were eligible for inclusion.

We intended to include RCTs that use a cluster-randomised design but identified none for inclusion in this update. Cross-over trials are not eligible for inclusion in this review.

Types of participants

Pregnant women, regardless of their risk for pre-eclampsia, preterm birth or intrauterine growth restriction (IUGR).

Types of interventions

- Omega-3 fatty acids (usually fish or algal oils) compared with placebo or no omega-3 fatty acids
- Trials that assessed omega-3 fatty acid co-interventions (e.g. omega-3 with another agent)
- Studies or study arms that compared omega-3 doses or types of omega-3 (e.g. DHA versus EPA) directly

Types of outcome measures

Primary outcomes

- Preterm birth < 37 weeks
- Early preterm birth < 34 weeks
- Prolonged gestation (> 42 weeks)

Secondary outcomes

For the woman

- Hypertension
- · Pre-eclampsia
- Eclampsia
- Admission to hospital (antenatal or postnatal)
- Caesarean section
- · Caesarean section (post-term)
- Induction (post-term)
- · Haemorrhage; blood loss
- Serious morbidity/mortality
- Length of gestation
- · Adverse effects
- Gestational diabetes
- Depression
- Anxiety
- · Stress (scale or response to challenge)
- Gestational weight gain
- Miscarriage

For babies

- Stillbirths
- Neonatal deaths
- Perinatal deaths
- Birthweight
- Birth length
- Head circumference
- Low birthweight (< 2.5 kg)
- Small-for-gestational age (SGA) (< 10th percentile)/IUGR
- Large-for-gestational age
- Intraventricular haemorrhage (and grade)
- Respiratory distress syndrome
- Necrotising enterocolitis
- Jaundice requiring phototherapy
- Sepsis
- · Retinopathy of prematurity
- Neonatal convulsion
- Admission to a neonatal intensive care unit

Longer term infant/child follow-up

- Physical growth
- Mental and emotional health
- Behaviour
- Neurological/neurosensory and developmental outcomes (including cognitive domains: attention, executive function, language, memory, visuospatial and motor development)



• Neurological disorders (e.g. cerebral palsy)

For health service resources

- Admission and length of stay in hospital and intensive care facilities
- · Use of community health services

Search methods for identification of studies

The following methods section of this review is based on a standard template used by Cochrane Pregnancy and Childbirth.

Electronic searches

For this update, we searched Cochrane Pregnancy and Childbirth's Trials Register by contacting their Information Specialist (16 August 2018)

The Register is a database containing over 25,000 reports of controlled trials in the field of pregnancy and childbirth. It represents over 30 years of searching. For full current search methods used to populate Pregnancy and Childbirth's Trials Register including the detailed search strategies for CENTRAL, MEDLINE, Embase and CINAHL; the list of handsearched journals and conference proceedings, and the list of journals reviewed via the current awareness service, please follow this link

Briefly, Cochrane Pregnancy and Childbirth's Trials Register is maintained by their Information Specialist and contains trials identified from:

- monthly searches of the Cochrane Central Register of Controlled Trials (CENTRAL);
- 2. weekly searches of MEDLINE (Ovid);
- 3. weekly searches of Embase (Ovid);
- 4. monthly searches of CINAHL (EBSCO);
- handsearches of 30 journals and the proceedings of major conferences;
- 6. weekly current awareness alerts for a further 44 journals plus monthly BioMed Central email alerts.

Search results are screened by two people and the full text of all relevant trial reports identified through the searching activities described above is reviewed. Based on the intervention described, each trial report is assigned a number that corresponds to a specific Pregnancy and Childbirth review topic (or topics), and is then added to the Register. The Information Specialist searches the Register for each review using this topic number rather than keywords. This results in a more specific search set that has been fully accounted for in the relevant review sections (Included studies; Excluded studies; Studies awaiting classification; Ongoing studies).

In addition, we searched ClinicalTrials.gov and the World Health Organization (WHO) International Clinical Trials Registry Platform (ICTRP) for unpublished, planned and ongoing trial reports (29 August 2017) using the search terms given in Appendix 1.

Searching other resources

We searched the reference lists of retrieved studies.

We did not apply any language or date restrictions.

Data collection and analysis

Selection of studies

Two review authors independently assessed all the potential studies we identified as a result of the search strategy for inclusion. We resolved any disagreement through discussion or, if required, we consulted a third review author.

Data extraction and management

We designed a form to extract data. For eligible trials, two review authors extracted the data using the agreed form. We resolved discrepancies through discussion or, if required, we consulted a third review author. We entered data into Review Manager 5 software (Review Manager 2014), and checked for accuracy.

When information regarding any of the above was unclear, we attempted to contact authors of the original reports to request further details.

Assessment of risk of bias in included studies

Two review authors independently assessed risk of bias for each study using the criteria outlined in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). We resolved any disagreement by discussion or by involving a third assessor.

(1) Random sequence generation (checking for possible selection bias)

For each included study we described the method used to generate the allocation sequence in sufficient detail to allow an assessment of whether it should produce comparable groups.

We assessed the method as:

- low risk of bias (any truly random process, e.g. random number table; computer random number generator);
- high risk of bias (any non-random process, e.g. odd or even date of birth; hospital or clinic record number);
- unclear risk of bias.

(2) Allocation concealment (checking for possible selection bias)

For each included study we described the method used to conceal allocation to interventions prior to assignment and assessed whether intervention allocation could have been foreseen in advance of, or during recruitment, or changed after assignment.

We assessed the methods as:

- low risk of bias (e.g. telephone or central randomisation; consecutively numbered sealed opaque envelopes);
- high risk of bias (open random allocation; unsealed or nonopaque envelopes, alternation; date of birth);
- unclear risk of bias.

(3.1) Blinding of participants and personnel (checking for possible performance bias)

For each included study we described the methods used, if any, to blind study participants and personnel from knowledge of which intervention a participant received. We considered that studies were at low risk of bias if they were blinded, or if we judged that



the lack of blinding unlikely to affect results. We assessed blinding separately for different outcomes or classes of outcomes.

We assessed the methods as:

- · low, high or unclear risk of bias for participants;
- low, high or unclear risk of bias for personnel.

(3.2) Blinding of outcome assessment (checking for possible detection bias)

For each included study we described the methods used, if any, to blind outcome assessors from knowledge of which intervention a participant received. We assessed blinding separately for different outcomes or classes of outcomes.

We assessed methods used to blind outcome assessment as:

• low, high or unclear risk of bias.

(4) Incomplete outcome data (checking for possible attrition bias due to the amount, nature and handling of incomplete outcome data)

For each included study we described, and for each outcome or class of outcomes, the completeness of data including attrition and exclusions from the analysis. We stated whether attrition and exclusions were reported and the numbers included in the analysis at each stage (compared with the total randomised participants), reasons for attrition or exclusion where reported, and whether missing data were balanced across groups or were related to outcomes. Where sufficient information was reported, or could be supplied by the trial authors, we planned to reinclude missing data in the analyses which we undertook.

We assessed methods as:

- low risk of bias (e.g. no missing outcome data; missing outcome data balanced across groups);
- high risk of bias (e.g. numbers or reasons for missing data imbalanced across groups; 'as treated' analysis done with substantial departure of intervention received from that assigned at randomisation);
- · unclear risk of bias.

(5) Selective reporting (checking for reporting bias)

For each included study we described how we investigated the possibility of selective outcome reporting bias and what we found.

We assessed the methods as:

- low risk of bias (where it is clear that all of the study's prespecified outcomes and all expected outcomes of interest to the review were reported);
- high risk of bias (where not all the study's prespecified outcomes were reported; one or more reported primary outcomes was not prespecified; outcomes of interest are reported incompletely and so cannot be used; study failed to include results of a key outcome that would have been expected to have been reported);
- unclear risk of bias.

(6) Other bias (checking for bias due to problems not covered by (1) to (5) above)

For each included study we described any important concerns we had about other possible sources of bias.

(7) Overall risk of bias

We made explicit judgements about whether studies were at high risk of bias, according to the criteria given in the *Handbook* (Higgins 2011).

Assessment of the quality of the evidence using the GRADE approach

For this update, we evaluated the quality of the evidence for the outcomes below using the GRADE approach as outlined in the GRADE handbook. The GRADE approach uses five considerations (study limitations, consistency of effect, imprecision, indirectness and publication bias) to assess the quality of the body of evidence for each outcome. In randomised controlled trials, the evidence can be downgraded from 'high quality' by one level for serious (or by two levels for very serious) limitations, depending on assessments for risk of bias, indirectness of evidence, serious inconsistency, imprecision of effect estimates or potential publication bias.

Baby/infant

- Preterm birth < 37 weeks
- Preterm birth < 34 weeks
- Perinatal death
- SGA/IUGR
- Low birthweight
- Large-for-gestational age

Mother

- Prolonged gestation (> 42 weeks)
- Induction post-term
- Pre-eclampsia
- Length of gestation
- Maternal adverse events
- Maternal morbidity composite (serious morbidity)
- Depression and/or anxiety (postnatal)

Child/adult

- Cognition
- Vision (neurosensory outcome)
- Neurodevelopment
- Behaviour
- BMI (long-term growth outcome)
- Diabetes (long-term development outcome)

Health services

- Maternal hospital admission (antenatal; postnatal)
- NICU admission
- Maternal length of hospital stay
- · Infant length of hospital stay
- Resource use



'Summary of findings' table

We used the GRADEpro Guideline Development Tool to import data from Review Manager 5 in order to create 'Summary of findings' tables for maternal, baby/infant, child and health service outcomes (Review Manager 2014). We created 'Summary of findings' tables for the main comparison: omega-3 LCPUFA versus no omega-3 (e.g. placebo or no supplement). We have presented summaries of the intervention effect and measures of quality according to the GRADE approach in the 'Summary of findings' tables.

Measures of treatment effect

Dichotomous data

For dichotomous data, we have presented results as summary risk ratios with 95% confidence intervals.

Continuous data

For continuous data, we have used the mean differences if outcomes were measured in the same way between trials. In future updates, we plan to use the standardised mean difference to combine trials that measure the same outcome, but use different methods.

Unit of analysis issues

Cluster-randomised trials

We did not identify any cluster-randomised trials.

In future updates of this review, if cluster-randomised trials are included, we will adjust their sample sizes and event rates using the methods described in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011), using an estimate of the intra-cluster correlation co-efficient (ICC) derived from the trial (if possible), from a similar trial or from a study of a similar population. If we use ICCs from other sources, we will report this and conduct sensitivity analyses to investigate the effect of variation in the ICC. If we identify both cluster-randomised trials and individually-randomised trials, we plan to synthesise the relevant information. We will consider it reasonable to combine the results from both if there is little heterogeneity between the study designs and the interaction between the effect of intervention and the choice of randomisation unit is considered to be unlikely.

We will also acknowledge heterogeneity in the randomisation unit and we will perform a subgroup analysis to investigate the effects of the randomisation unit.

Cross-over trials

We considered cross-over designs to be an inappropriate design for this research question.

Multi-arm trials

For included multi-arm trials, we used methods described in the *Cochrane Handbook for Systematic Reviews of Interventions* to overcome possible unit-of analysis errors (Higgins 2011), by combining groups to make a single pair-wise comparison (where appropriate), or by splitting the 'shared' group into two (or more) groups with smaller sample sizes, and including the two (or more) comparisons (see Included studies text for details of how this was done for each of the 10 multi-arm trials we included).

Dealing with missing data

For included trials, we noted levels of attrition.

For all outcomes, we carried out analyses, as far as possible, on an intention-to-treat basis, that is, we have attempted to include all participants randomised to each group in the analyses, and all participants were analysed in the group to which they were allocated, regardless of whether or not they received the allocated intervention. The denominator for each outcome in each trial was the number randomised minus any participants whose outcomes were known to be missing.

Assessment of heterogeneity

We assessed statistical heterogeneity in each meta-analysis using the Tau², I² and Chi² statistics. We regarded heterogeneity as substantial if I² was greater than 30% and either Tau² was greater than zero, or there was a low P value (less than 0.10) in the Chi² test for heterogeneity.

Assessment of reporting biases

Where there were 10 or more trials in a meta-analysis we investigated reporting biases (such as publication bias) using funnel plots. We assessed funnel plot asymmetry visually.

Data synthesis

We carried out statistical analysis using Review Manager 5 software (Review Manager 2014). We used fixed-effect meta-analysis for combining data where it was reasonable to assume that trials were estimating the same underlying treatment effect, that is, where trials were examining the same intervention, and the trials' populations and methods were judged to be sufficiently similar. Where there was clinical heterogeneity sufficient to expect that the underlying treatment effects differed between trials, or where substantial statistical heterogeneity was detected ($I^2 > 30\%$), we used random-effects meta-analysis to produce an overall summary, if an average treatment effect across trials was considered clinically meaningful. The random-effects summary was treated as the average of the range of possible treatment effects and we have discussed the implications of treatment effects differing between trials. If the average treatment effect was not clinically meaningful, we would not have combined trials.

Where we have used random-effects analyses, the results have been presented as the average treatment effect with 95% confidence intervals, and the estimates of T^2 and I^2 .

Subgroup analysis and investigation of heterogeneity

We investigated substantial heterogeneity using subgroup analyses and sensitivity analyses.

We carried out the following subgroup analyses.

1. Type of intervention

All the following interventions compared with each other:

- omega-3 LCPUFA supplements only;
- omega-3 supplements plus omega enriched food or dietary advice;
- omega enriched food only;
- omega-3 LCPUFA supplements plus advice and/or other agents.



2. Dose of omega-3 LCPUFA

The following doses compared to each other:

- low (< 500 mg/day);
- mid (500 mg to 1 g/day);
- high (> 1 g/day).

3. Timing

Comparison of the following gestational ages when omega-3 LCPUFA supplements commenced:

- ≤ 20 weeks' gestation;
- > 20 weeks' gestation.

4. Type of omega-3

Comparison of the following types of omega-3:

- DHA/largely DHA;
- mixed EPA/DHA;
- mixed DHA/EPA/other

5. Risk of poorer maternal/perinatal outcomes

Comparison of the following risk levels with each other:

- · increased or high risk
- · low risk
- any or mixed risk

For subgroup 1 type of intervention (Analysis 2) we did not restrict this analysis to the selected group of outcomes used in the other subgroup analyses. This was done to help readers to see results across all outcomes by type of omega-3 intervention (except for longer term outcomes or other outcomes reporting multiple time points (analyses 1.63 to 1.92) which were sparsely reported).

The following outcomes were used in the other four subgroup analyses (analyses 2-5):

- preterm birth < 37 weeks;
- early preterm birth < 34 weeks;
- prolonged gestation (> 42 weeks);
- · pre-eclampsia;
- · caesarean section;
- length of gestation;
- perinatal death;
- stillbirth;
- · neonatal death;
- low birthweight;
- SGA/IUGR;
- birthweight.

We assessed subgroup differences by interaction tests available within Review Manager 5 (Review Manager 2014). We reported the results of subgroup analyses quoting the Chi² statistic and P value, and the interaction test I² value.

Sensitivity analysis

We carried out sensitivity analyses (Analysis 9) to explore the effects of trial quality assessed by sequence generation and concealment of allocation, and inadequate blinding, by omitting trials rated as 'high risk of bias' or 'unclear risk of bias' for any one or more of these sources of bias, to assess whether this made any difference to the overall result. We restricted this analysis to 12 outcomes:

- preterm birth < 37 weeks;
- early preterm birth < 34 weeks;
- prolonged gestation > 42 weeks;
- pre-eclampsia;
- · caesarean section;
- birthweight;
- perinatal death;
- stillbirth;
- neonatal death;
- · gestational age;
- low birthweight;
- SGA/IUGR

These outcomes are this review's three primary outcomes, plus nine secondary outcomes that were selected for use in subgroup analyses 3, 4 and 5).

RESULTS

Description of studies

Results of the search

For this update, we assessed 447 trial reports in total. This included 406 new reports, plus we reassessed the six included studies (17 reports), 15 excluded studies (20 reports), three ongoing studies and one awaiting further classification in the previous version of the review (Makrides 2006Makrides 2006).

Where required, we reclassified some of the studies/records which were listed as excluded, ongoing or awaiting classification in the previous version of this review (Makrides 2006).

Overall, we have included 70 trials (374 reports). The six trials originally included are still included. The three trials originally listed as ongoing have reported results and are now included. Eight trials that were previously excluded are now included (either due to the enlarged scope of the review or changes in review methodology (e.g. fulfilling inclusion criteria, even if the trial does not report any of the review's prespecified outcomes)).

As of August 2018, we have:

- 15 excluded studies (25 reports) (Escobar 2008; Fievet 1985; Gholami 2017; Herrera 1993; Herrera 1998; Herrera 2004; Lauritzen 2004; Marangell 2004; Morrison 1984; Morrison 1986; Nishi 2016; Starling 1990; Valentine 2013; Velzing-Aarts 2001; Yelland 2016).
- 23 ongoing studies (28 reports) (Albert 2017; Carlson 2017 ADORE; Carvajal 2014; de Carvalho 2017; Dos Santos 2018; Dragan 2013; FOPCHIN; Garg 2017; Garmendia 2015; Ghebremeskel 2014; Hegarty 2012; Hendler 2017; Khandelwal 2012; Kodkhany 2017; Li 2013; Makrides 2013 (ORIP); Martini



2014 (CORDHA); Mbayiwa 2016; Murff 2017 (FORTUNE); Nishi 2015 (SYNCHRO); Wang 2018; Zielinsky 2015; Zimmermann 2018).

• 17 studies (20 reports) awaiting further classification (Farahani 2010; Gopalan 2004; Jamilian 2018; Kadiwala 2015; Laitinen

2013; Lazzarin 2009; Parisi 2013; Pavlovich 1999; Sajina-Stritar 1994; Sajina-Stritar 1998; Salvig 2009; Salzano 2001; Stoutjesdijk 2014; Vahedi 2018; Vakilian 2010; Valentine 2014; Valenzuela 2017).

See Figure 1 which outlines the study flow.



Figure 1. Study flow diagram.

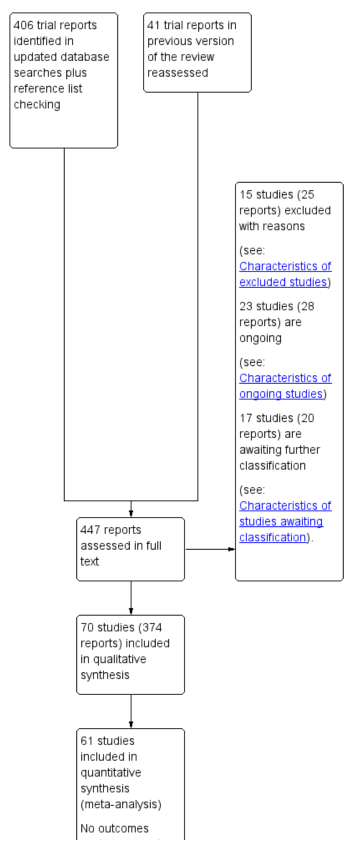




Figure 1. (Continued)

No outcomes could be reported for 9 studies

Included studies

Following application of eligibility criteria, we included 70 RCTs comparing an omega-3 fatty acid intervention (stand-alone or with a co-intervention), with placebo or no omega-3 fatty acids in this review (Ali 2017; Bergmann 2007; Bisgaard 2016; Boris 2004; Bosaeus 2015; Bulstra-Ramakers 1994; Carlson 2013; Chase 2015; D'Almedia 1992; de Groot 2004; Dilli 2018; Dunstan 2008; England 1989; Freeman 2008; Furuhjelm 2009; Giorlandino 2013; Gustafson 2013; Haghiac 2015; Harper 2010; Harris 2015; Hauner 2012; Helland 2001; Horvaticek 2017; Hurtado 2015; Ismail 2016; Jamilian 2016; Jamilian 2017; Judge 2007; Judge 2014; Kaviani 2014; Keenan 2014; Khalili 2016;. Knudsen 2006; Krauss-Etschmann 2007; Krummel 2016; Laivuori 1993; Makrides 2010; Malcolm 2003; Mardones 2008; Martin-Alvarez 2012; Miller 2016; Min 2014; Min 2016; Mozurkewich 2013; Mulder 2014; Noakes 2012; Ogundipe 2016; Oken 2013; Olsen 1992; Olsen 2000; Onwude 1995; Otto 2000; Pietrantoni 2014; Ramakrishnan 2010; Ranjkesh 2011; Razavi 2017; Rees 2008; Ribeiro 2012; Rivas-Echeverria 2000; Samimi 2015; Sanjurjo 2004; Smuts 2003a; Smuts 2003b; Su 2008; Taghizadeh 2016; Tofail 2006; Valenzuela 2015; Van Goor 2009; Van Winden 2017; Vaz 2017).

All the included trials were individually randomised. Ten were multi-arm trials (Bergmann 2007; Harris 2015; Jamilian 2017; Knudsen 2006; Krauss-Etschmann 2007; Laivuori 1993; Mozurkewich 2013; Oken 2013; Razavi 2017; Van Goor 2009).

A total of 19,927 women were involved in the included trials. Knudsen 2006 was the largest trial, randomising 3098 women, followed by Makrides 2010 and Olsen 2000, in which 2399 and 1647 women, respectively, were randomised. Ribeiro 2012 was the smallest trial, randomising 11 women, followed by Van Winden 2017 and Laivuori 1993 (14 and 18 women, respectively). For the majority of the included trials, fewer women were included in analyses than were randomised.

The included trials have been published over nearly three decades - from 1989 to 2018.

Review structure

The analyses in the review are structured as follows.

- **Overall:** omega-3 fatty acids versus placebo or no omega-3 fatty acids (Analysis 1)
- Type of intervention subgroups: omega-3 supplementation alone; combined with food and/or advice; omega-3 rich food; omega-3 plus another agent - all versus no omega-3 (Analysis 2)
- Dose subgroups (DHA/EPA): low (< 500 mg/day) versus mid (500 mg to 1 g/day) versus high (> 1 g/day) (Analysis 3)
- Timing subgroups: gestational age when omega-3 supplements commenced: ≤ 20 weeks' gestation versus > 20 weeks' gestation (Analysis 4)

- Type of omega-3: DHA/largely DHA; mixed EPA/DHA; mixed DHA/EPA/other (Analysis 5)
- Risk subgroups: increased/high risk versus low risk versus any/ mixed risk (Analysis 6)
- Direct comparisons of omega-3 doses (Analysis 7)
- **Direct comparisons of omega-3 types** (Analysis 8)
- Sensitivity analysis (Analysis 9)

Further details are given below and in the Characteristics of included studies tables.

Settings

The 70 trials were conducted in a wide range of countries, and most (but not all) in upper-middle or high-income countries:

- 16 trials were conducted in the USA (Carlson 2013; Chase 2015; Freeman 2008; Gustafson 2013; Haghiac 2015; Harper 2010; Harris 2015; Judge 2007; Judge 2014; Keenan 2014; Krummel 2016; Miller 2016; Mozurkewich 2013; Oken 2013; Smuts 2003a; Smuts 2003b);
- eight in Iran (Jamilian 2016; Jamilian 2017; Kaviani 2014; Khalili 2016; Ranjkesh 2011; Razavi 2017; Samimi 2015; Taghizadeh 2016);
- six in the UK (Malcolm 2003; Min 2014; Min 2016; Noakes 2012; Ogundipe 2016; Onwude 1995);
- four in the Netherlands (Bulstra-Ramakers 1994; de Groot 2004; Otto 2000; Van Goor 2009); and four in Demark (Bisgaard 2016; Boris 2004; Knudsen 2006; Olsen 1992);
- three in Australia (Dunstan 2008; Makrides 2010; Rees 2008); and three in Spain (Hurtado 2015; Martin-Alvarez 2012; Sanjurjo 2004);
- two each in Chile (Mardones 2008; Valenzuela 2015); Egypt (Ali 2017; Ismail 2016); Germany (Bergmann 2007; Hauner 2012); Italy (Giorlandino 2013; Pietrantoni 2014); Brazil (Ribeiro 2012; Vaz 2017); and Sweden (Bosaeus 2015; Furuhjelm 2009);
- and one each in Angola (D'Almedia 1992); Bangladesh (Tofail 2006); Canada (Mulder 2014); Croatia (Horvaticek 2017); Finland (Laivuori 1993); Mexico (Ramakrishnan 2010); Norway (Helland 2001); South Africa (England 1989); Taiwan (Su 2008); Turkey (Dilli 2018) and Venezuela (Rivas-Echeverria 2000).

Two of the 70 trials were performed in more than one country: Krauss-Etschmann 2007 (Germany, Spain and Hungary); and Olsen 2000 (Denmark, Scotland, Sweden, United Kingdom, Italy, the Netherlands, Norway, Belgium and Russia). Van Winden 2017 did not report where the study was conducted.

Participants

All participants were pregnant women (and their children). Most pregnancies were singletons, with some studies specifically excluding multiple births. Characteristics of the women are summarised below, including age, parity, eligibility criteria



relating to omega-3 consumption, socioeconomic status, ethnicity, smoking status and risk of adverse pregnancy outcomes. Further details are included in the Additional tables.

Age

Where reported, the mean age of the women ranged from 22 years in Smuts 2003a to 40 years in several studies. The mean age of the women in both groups was at least 30 years in 18 of the included trials (Bergmann 2007; Bisgaard 2016; Bosaeus 2015; Dilli 2018; Dunstan 2008; Furuhjelm 2009; Hauner 2012; Jamilian 2016; Jamilian 2017; Krauss-Etschmann 2007; Laivuori 1993; Miller 2016; Min 2014 [diabetic women]; Min 2016; Mulder 2014; Rees 2008; Su 2008; Van Goor 2009). Maternal age of women across the included trials is summarised further in Table 1.

Parity

Five trials specifically reported parity: Rivas-Echeverria 2000 excluded nulliparous women; Smuts 2003b excluded women with more than four prior pregnancies; Valenzuela 2015 included women with one to four prior births; Van Goor 2009 included women with a first or second pregnancy. Olsen 2000, for the prophylactic trials, included women who in an early pregnancy had experienced preterm birth (before 259 days gestation). Twenty-eight of the trials did not report baseline information related to parity clearly (Boris 2004; Bulstra-Ramakers 1994; Chase 2015; D'Almedia 1992; Dilli 2018; England 1989; Furuhjelm 2009; Giorlandino 2013; Gustafson 2013; Harper 2010; Harris 2015; Jamilian 2016; Jamilian 2017; Judge 2014; Kaviani 2014; Keenan 2014; Krummel 2016; Malcolm 2003; Martin-Alvarez 2012; Miller 2016; Noakes 2012; Ogundipe 2016; Ramakrishnan 2010; Razavi 2017; Ribeiro 2012; Samimi 2015; Taghizadeh 2016; Van Winden 2017). Both nulliparous and multiparous women were included in the remaining 38 trials (Ali 2017; Bergmann 2007; Bisgaard 2016; Bosaeus 2015; Carlson 2013; de Groot 2004; Dunstan 2008; Freeman 2008; Haghiac 2015; Hauner 2012; Helland 2001 (nulliparous and primiparous only); Horvaticek 2017; Hurtado 2015; Ismail 2016; Judge 2007; Khalili 2016; Knudsen 2006; Krauss-Etschmann 2007; Laivuori 1993; Makrides 2010; Mardones 2008; Min 2014; Min 2016; Mozurkewich 2013; Mulder 2014; Oken 2013; Olsen 1992; Olsen 2000 (therapeutic trials only); Onwude 1995; Otto 2000; Pietrantoni 2014; Ranjkesh 2011; Rees 2008; Sanjurjo 2004; Smuts 2003a; Su 2008; Tofail 2006; Vaz 2017). Detailed information relating to parity is reported in Table 2.

Eligibility criteria relating to omega-3 intake

Forty of the 70 trials reported eligibility criteria relating to omega-3 intake, such as excluding women with an allergy to fish or fish products and/or excluding women taking omega-3, fish oil or DHA supplements or regular/any intake of fish. However in one case, women were required to be consuming fish at least twice a week to be eligible for inclusion in the trial in addition to either omega-3 LCPUFA supplementation or placebo (Pietrantoni 2014). See Table 3 for further details for each relevant trial.

Socioeconomic status

The socioeconomic status of women at baseline was reported by a range of measures including, education, employment, household income, socioeconomic index, and welfare/benefit dependence. Education measures were reported by 21 trials (Bergmann 2007; Bosaeus 2015; Carlson 2013; de Groot 2004; Dunstan 2008; Freeman 2008; Gustafson 2013; Harper 2010; Hauner 2012; Helland 2001;

Judge 2007; Kaviani 2014; Khalili 2016; Krummel 2016; Makrides 2010; Mardones 2008; Pietrantoni 2014; Ramakrishnan 2010; Rees 2008; Tofail 2006; Vaz 2017), with all but seven of these trials suggesting that most women had at least 12 years education - see Table 4. Five trials reported other measures of socio-economic status - Bisgaard 2016 reported that 10% of participants had low incomes; D'Almedia 1992 reported that 69% of women were employed; Krauss-Etschmann 2007 stated that 40% of fathers had no training qualifications; Oken 2013 reported that 40% of women worked full-time and Smuts 2003a reported that "most subjects received government assistance for medical aid". The remaining 42 trials did not report socioeconomic status of participants.

Ethnicity or race

Most trials (46) reported no baseline information on ethnicity or race, though they did report the country where the study was conducted (with the exception of Van Winden 2017). Ten trials reported a mix of ethnicities, nine trials reported including only Caucasian women (understood to be white women) or women of similar ethnicities; two trials included African women, and one trial each reported including African-American women or Hispanic women - see Table 5.

Smoking

Thirteen trials reported excluding women who smoked. Twenty-three trials reported smoking rates in pregnancy ranged from several per cent to nearly 50% in one trial. The remaining 35 trials did not report maternal smoking status, see Table 6.

Women at risk

We defined increased/high risk as any factors which might increase the risk of adverse maternal and birth outcomes; these baseline risks included being at risk of pre-eclampsia, having a previous preterm birth, gestational diabetes mellitus (GDM), being overweight/obese or underweight, or being at risk of poor mental health - see Table 7.We classified trials into increased/high risk (34 trials); any or mixed risk (8 trials) and low risk (29 trials). One trial reported women with GDM and low risk women separately (Min 2014). We also performed a subgroup analysis based on risk (see Analysis 6 and results text).

Interventions

Overall analysis (Analysis 1)

Each of the 70 included trials compared an omega-3 fatty acid intervention (stand-alone or with a co-intervention); including 10 trials with a food or dietary advice component), with placebo or no omega-3 fatty acids, with 60 trials contributing data for this review.

Intervention type subgroup (Analysis 2)

As there was considerable variation between trials, we have subgrouped results by four main types of intervention (in addition to the overall analysis):

Intervention type 1: Omega-3 supplements only versus placebo or no omega-3 fatty acids (50 trials) (Bisgaard 2016; Boris 2004; Bulstra-Ramakers 1994; Carlson 2013; Chase 2015; Dilli 2018; Dunstan 2008; England 1989; Freeman 2008; Furuhjelm 2009; Giorlandino 2013; Gustafson 2013; Haghiac 2015; Harris 2015; Helland 2001; Horvaticek 2017; Ismail 2016; Jamilian 2016; Jamilian 2017*; Judge 2007; Judge 2014; Kaviani



2014; Keenan 2014; Khalili 2016; Knudsen 2006; Krummel 2016; Laivuori 1993; Makrides 2010; Malcolm 2003; Miller 2016; Min 2014; Min 2016; Mozurkewich 2013; Mulder 2014; Olsen 1992; Olsen 2000; Onwude 1995; Ramakrishnan 2010; Ranjkesh 2011; Razavi 2017*; Rees 2008; Ribeiro 2012; Samimi 2015; Sanjurjo 2004; Su 2008; Tofail 2006; Valenzuela 2015; Van Goor 2009; Van Winden 2017; Vaz 2017)

- * Most of these trials compared oral DHA and/or EPA (or mainly DHA/EPA) supplements with placebo or no omega-3 treatment. Four trials compared unspecified or other oral omega-3 fatty acid supplements with placebo or no omega-3 (Laivuori 1993; Ribeiro 2012; Samimi 2015; Valenzuela 2015), and one trial compared vaginal omega-3 supplementation with placebo (Giorlandino 2013). Some trials reported including small amounts of other agents in the intervention arm (e.g. vitamin E) but we judged these to have minimal effect on outcomes.
- Intervention type 2: Omega-3 supplements/enrichment plus food/dietary advice versus placebo or no omega-3 fatty acids (7 trials) (de Groot 2004; Hauner 2012; Hurtado 2015; Martin-Alvarez 2012; Pietrantoni 2014; Smuts 2003a; Smuts 2003b)
- Intervention type 3: Omega-3 food/dietary advice only versus placebo or no omega-3 fatty acids (3 trials): (Bosaeus 2015; Noakes 2012; Oken 2013)
- Intervention type 4: Omega-3 supplements plus other agents versus placebo or no omega-3 fatty acids (12 trials): the other agents used in these 12 trials were as follows.
 - arachidonic acid (AA) (Otto 2000; Van Goor 2009)
 - Aspirin (Ali 2017)
 - Aspirin + vitamins C + E (Rivas-Echeverria 2000)
 - Folate (Krauss-Etschmann 2007)
 - Gamma-linolenic acid (GLA) (D'Almedia 1992)
 - multiple micronutrients (Mardones 2008)
 - Prebiotics (Bergmann 2007)
 - Progesterone (Harper 2010)
 - Vitamin D (Jamilian 2017*; Razavi 2017*)
 - Vitamin E (high amounts) Taghizadeh 2016.

Jamilian 2017* and Razavi 2017* are multi-arm trials that span two of the above four intervention categories.

Multi-arm trials

Ten trials had multi-arm designs. We combined relevant groups in the multi-arm trials to create appropriate single pair-wise comparisons for inclusion in the main comparison, avoiding unit of analysis errors, specifically:

- Bergmann 2007: three arms (DHA/EPA + prebiotic versus prebiotic + vitamin/mineral versus vitamin/mineral); analysed as DHA/EPA + prebiotic versus the other two arms combined (prebiotic/vitamin/mineral and vitamin/mineral) in the overall comparison (Analysis 1).
- Harris 2015: three arms (300 g/day DHA versus 600 g/day DHA versus placebo); analysed as 300 g/day + 600 g/day combined versus placebo for the overall comparison (Analysis 1); we split doses in the dose subgroups (Analysis 3), and compared 300 g/day and 600 g/day directly in Analysis 8.
- Krauss-Etschmann 2007: four arms (DHA + EPA versus DHA + EPA + folate versus folate versus placebo, all using milk-based

- sachets); analysed as DHA + EPA and DHA + EPA + folate groups combined, compared with placebo and folate only combined.
- Jamilian 2017: four arms (DHA + EPA versus DHA + EPA + vitamin D versus vitamin D + placebo) - data from this trial were not included (no review outcomes reported).
- Knudsen 2006: seven arms (five different doses of DHA + EPA versus ALA versus no treatment/flax oil); analysed as six omega-3 groups combined versus no treatment/flax oil for the overall comparison (Analysis 1); omega-3 groups combined in two dose groups, < 1 g/day and ≥ 1 g/day in the direct dose comparison (Analysis 7); DHA + EPA versus ALA in the omega-3 supplement type direct comparison (Analysis 8).
- Laivuori 1993: three arms (DHA + EPA + other omega-3 versus linoleic acid (LA)/GLA versus placebo) - data from this trial were not included (no outcomes able to be used).
- Mozurkewich 2013: three arms (mainly DPA versus mainly EPA versus placebo); DPA + EPA groups pooled in analysis 1; DPA versus mainly EPA groups included in omega-3 supplement type comparison (Analysis 8).
- Oken 2013: three arms (voucher to purchase fish plus advice on which fish to consume versus advice on fish consumption only versus generic dietary advice only); analysed as intervention arms pooled versus generic dietary advice only.
- Razavi 2017 four arms (DHA + EPA versus DHA + EPA + vitamin D versus vitamin D versus placebo); two arms (DPA + EPA and DHA + EPA + vitamin D) pooled and compared with placebo in analysis.
- Van Goor 2009: three arms (DHA + AA versus DHA versus placebo); intervention arms pooled and compared with placebo for overall comparison (Analysis 1) and other analyses except for DHA + AA versus DHA for direct omega-3 supplement type comparison (Analysis 8).

For additional details on the omega-3 fatty acid interventions and how they varied across the trials see Characteristics of included studies.

Comparisons

Most comparisons were between omega-3 LCPUFA and placebo/ no omega-3. As well as contributing to the main omega-3 versus no omega-3 comparison in Analysis 1, five of the multi-arm trials contributed outcomes from direct comparisons of omega-3 supplement doses or omega-3 supplements for inclusion in the meta-analysis:

- Omega-3 supplement dose comparisons: two trials: Harris 2015 compared 600 mg versus 300 mg DHA/day; Knudsen 2006 compared six different doses which we collapsed into a comparison of < 1 g/day versus ≥ 1 g/day (see Analysis 7).
- Omega-3 supplements versus other omega-3 supplements: three trials: Knudsen 2006 compared EPA/DHA (five doses combined) versus ALA; Mozurkewich 2013 compared DHA versus EPA; and Van Goor 2009 compared DHA versus DHA/AA combined (see Analysis 8).

Subgroup analyses

Dose subgroup - DHA + EPA (Analysis 3)

 Low (< 500 mg/day): 24 trials (Ali 2017; Bergmann 2007; D'Almedia 1992; Harris 2015*; Hurtado 2015; Ismail 2016; Jamilian 2016; Judge 2007; Judge 2014; Khalili 2016; Knudsen



2006*; Malcolm 2003; Martin-Alvarez 2012; Miller 2016; Mulder 2014; Noakes 2012 (as fish); Ogundipe 2016; Pietrantoni 2014; Ramakrishnan 2010; Samimi 2015; Sanjurjo 2004; Smuts 2003a; Smuts 2003b; Van Goor 2009)

- Mid (500 mg/day to 1 g/day): 21 trials (Carlson 2013; Chase 2015; Dilli 2018; Giorlandino 2013; Gustafson 2013; Harris 2015*; Horvaticek 2017; Kaviani 2014#; Keenan 2014; Krauss-Etschmann 2007; Knudsen 2006*; Krummel 2016; Makrides 2010; Mardones 2008#; Min 2014; Min 2016; Otto 2000; Ranjkesh 2011; Razavi 2017; Rees 2008; Ribeiro 2012)
- * Of these 21 trials, 14 clearly reported doses of ≥ 500 mg DHA/day (Carlson 2013; Chase 2015; Giorlandino 2013; Gustafson 2013; Harris 2015 (one arm); Horvaticek 2017; Krauss-Etschmann 2007; Krummel 2016; Makrides 2010; Min 2014; Min 2016; Otto 2000; Rees 2008; Ribeiro 2012).
- High (> 1 g/day): 23 trials (Bisgaard 2016; Boris 2004; Bulstra-Ramakers 1994; Dunstan 2008; England 1989; Freeman 2008; Furuhjelm 2009; Haghiac 2015; Harper 2010; Hauner 2012; Helland 2001; Jamilian 2017; Knudsen 2006*; Laivuori 1993; Mozurkewich 2013; Olsen 1992; Olsen 2000; Onwude 1995; Rivas-Echeverria 2000; Su 2008; Tofail 2006; Van Winden 2017; Vaz 2017).
- Other: in five trials it was not possible to estimate DHA/EPA dose, or the omega-3 supplement was clearly not DHA or EPA (Bosaeus 2015; Oken 2013 (advice to consume fish); de Groot 2004 (margarine enriched to give 2.82 g/day ALA); Taghizadeh 2016 (ALA 400 mg/day (flax oil)); Valenzuela 2015 (10.1 g/day ALA).

*trials had more than one omega-3 group with different doses

#only specified as omega-3 and not DHA and/or EPA.

Timing subgroup - gestational age when omega-3 supplements commenced (Analysis 4)

- > 20 weeks' gestation: 33 trials (Ali 2017; Bergmann 2007; Bisgaard 2016; Boris 2004; Furuhjelm 2009; Giorlandino 2013; Hurtado 2015; Ismail 2016; Jamilian 2016; Jamilian 2017; Judge 2007; Judge 2014; Kaviani 2014; Knudsen 2006; Krummel 2016; Laivuori 1993; Martin-Alvarez 2012; Miller 2016; Min 2016; Olsen 1992; Onwude 1995; Ramakrishnan 2010; Razavi 2017; Rees 2008; Ribeiro 2012; Samimi 2015; Sanjurjo 2004; Smuts 2003a; Smuts 2003b; Taghizadeh 2016; Tofail 2006; Valenzuela 2015; Van Winden 2017)
- ≤ 20 weeks' gestation: 33 trials (Bosaeus 2015; Bulstra-Ramakers 1994; Carlson 2013; Chase 2015; D'Almedia 1992; de Groot 2004; Dilli 2018; Dunstan 2008; Gustafson 2013; Haghiac 2015; Harper 2010; Harris 2015; Hauner 2012; Helland 2001; Horvaticek 2017; Keenan 2014; Khalili 2016; Krauss-Etschmann 2007; Makrides 2010; Malcolm 2003; Mardones 2008; Min 2014; Mozurkewich 2013; Mulder 2014; Noakes 2012; Ogundipe 2016; Olsen 2000; Otto 2000; Pietrantoni 2014; Ranjkesh 2011; Su 2008; Van Goor 2009; Vaz 2017)
- Mixed: two trials (Freeman 2008; Oken 2013)
- Not reported: two trials (England 1989; Rivas-Echeverria 2000)

DHA/mixed subgroup (Analysis 5)

 DHA/largely DHA: 27 trials (Carlson 2013; Chase 2015; Dunstan 2008; Giorlandino 2013; Gustafson 2013; Harris 2015; Hauner 2012; Horvaticek 2017; Hurtado 2015; Judge 2007; Judge 2014; Keenan 2014; Krummel 2016; Makrides 2010; Malcolm 2003;

- Martin-Alvarez 2012; Miller 2016; Min 2014; Min 2014 [diabetic women]; Min 2016; Mulder 2014; Ogundipe 2016; Pietrantoni 2014; Ramakrishnan 2010; Rees 2008; Sanjurjo 2004; Smuts 2003a; Smuts 2003b)
- Mixed EPA + DHA: 25 trials (Bisgaard 2016; Boris 2004; Bulstra-Ramakers 1994; Dilli 2018; England 1989; Freeman 2008; Furuhjelm 2009; Haghiac 2015; Harper 2010; Helland 2001; Ismail 2016; Jamilian 2016; Jamilian 2017; Khalili 2016; Knudsen 2006; Mozurkewich 2013; Noakes 2012; Olsen 1992; Olsen 2000; Olsen 2000 [twins]; Onwude 1995; Ranjkesh 2011; Su 2008; Tofail 2006; Van Winden 2017; Vaz 2017)
- Mixed DHA + EPA + other: 18 trials: (Ali 2017; Bergmann 2007; Bosaeus 2015; D'Almedia 1992; de Groot 2004; Kaviani 2014; Krauss-Etschmann 2007; Laivuori 1993; Mardones 2008; Oken 2013; Otto 2000; Razavi 2017; Ribeiro 2012; Rivas-Echeverria 2000; Samimi 2015; Taghizadeh 2016; Valenzuela 2015; Van Goor 2009)

Risk subgroup: women at increased/high risk, any/mixed risk or low risk (Analysis 6)

Increased or high baseline risk of adverse maternal and birth outcomes included being at risk of pre-eclampsia, having a previous preterm birth, GDM, being overweight/obese or underweight, or being at risk of poor mental health - see Table 7.

- Increased or high risk: 34 trials (Ali 2017; Bulstra-Ramakers 1994; Chase 2015; Dilli 2018; England 1989; Freeman 2008; Giorlandino 2013; Haghiac 2015; Harper 2010; Horvaticek 2017; Ismail 2016; Jamilian 2016; Jamilian 2017; Kaviani 2014; Keenan 2014; Krummel 2016; Laivuori 1993; Mardones 2008; Min 2014 [diabetic women]*; Min 2016; Mozurkewich 2013; Ogundipe 2016; Olsen 2000; Onwude 1995; Ranjkesh 2011; Razavi 2017; Rees 2008; Rivas-Echeverria 2000; Samimi 2015; Su 2008; Taghizadeh 2016; Tofail 2006; Van Winden 2017; Vaz 2017)
- Any or mixed risk: eight trials (Bisgaard 2016; D'Almedia 1992; Knudsen 2006; Makrides 2010; Martin-Alvarez 2012; Miller 2016; Oken 2013; Ribeiro 2012)
- Low risk: 29 trials (Bergmann 2007; Boris 2004; Bosaeus 2015; Carlson 2013; de Groot 2004; Dunstan 2008; Furuhjelm 2009; Gustafson 2013; Harris 2015; Hauner 2012; Helland 2001; Hurtado 2015; Judge 2007; Judge 2014; Khalili 2016; Krauss-Etschmann 2007; Malcolm 2003; Min 2014; Mulder 2014; Noakes 2012; Olsen 1992; Otto 2000; Pietrantoni 2014; Ramakrishnan 2010; Sanjurjo 2004; Smuts 2003a; Smuts 2003b; Valenzuela 2015; Van Goor 2009)

*Min 2014 reported diabetic women separately.

Outcomes

Primary outcomes were reported in a format suitable for metaanalysis as follows:

- preterm birth < 37 weeks reported by 26 trials;
- preterm birth < 34 weeks reported by nine trials;
- prolonged gestation > 42 weeks reported by six trials.

Most of our secondary outcomes were reported in at least some of the trials.

We were unable to include any outcomes from 10 trials in our meta-analysis (Boris 2004; Bosaeus 2015; Chase 2015; Ismail 2016;



Jamilian 2017; Laivuori 1993; Martin-Alvarez 2012; Ogundipe 2016; Ribeiro 2012; Van Winden 2017). Of these, Boris 2004, Chase 2015, Ismail 2016, Jamilian 2017, Martin-Alvarez 2012 and Ribeiro 2012 did not report on any of our prespecified review outcomes. Bosaeus 2015 and Laivuori 1993 reported review outcomes, however, the data were not suitable for inclusion in the meta-analysis. Ogundipe 2016 only reported review outcomes overall, not separately by intervention and control group. Van Winden 2017 did report on maternal adverse effects, but narratively.

Sources of trial funding

Funding sources were reported by 56 of the 70 included trials (Bergmann 2007; Bisgaard 2016; Boris 2004; Bosaeus 2015; Carlson 2013; Chase 2015; D'Almedia 1992; de Groot 2004; Dilli 2018; Dunstan 2008; Freeman 2008; Furuhjelm 2009; Giorlandino 2013; Gustafson 2013; Haghiac 2015; Harper 2010; Harris 2015; Hauner 2012; Helland 2001; Hurtado 2015; Jamilian 2016; Jamilian 2017; Judge 2007; Judge 2014; Keenan 2014; Khalili 2016; Knudsen 2006; Krauss-Etschmann 2007; Krummel 2016; Laivuori 1993; Makrides 2010; Malcolm 2003; Mardones 2008; Min 2014; Min 2016; Mozurkewich 2013; Mulder 2014; Ogundipe 2016; Oken 2013; Olsen 1992; Olsen 2000; Otto 2000; Pietrantoni 2014; Ramakrishnan 2010; Razavi 2017; Rees 2008; Rivas-Echeverria 2000; Samimi 2015; Sanjurjo 2004; Smuts 2003a; Smuts 2003b; Su 2008; Taghizadeh 2016; Tofail 2006; Van Goor 2009; Vaz 2017). Funding bodies listed by the trials were mostly non-commercial $organisations \ (e.g.\ government\ funding\ bodies, universities, health$ services and other not-for-profit foundations, including the World Health Organization). However, commercial organisations - mainly pharmaceutical companies - were reported as the only or main funding sources in 11 trials (Bergmann 2007; de Groot 2004; Giorlandino 2013; Helland 2001; Laivuori 1993; Mardones 2008; Otto 2000; Sanjurjo 2004; Smuts 2003a; Smuts 2003b; Van Goor 2009). Thirteen trials did not report any funding (Ali 2017; Bulstra-Ramakers 1994; England 1989; Horvaticek 2017; Ismail 2016; Kaviani 2014; Martin-Alvarez 2012; Miller 2016; Ranjkesh 2011; Ribeiro 2012; Rivas-Echeverria 2000; Valenzuela 2015; Van Winden 2017).

Trial authors' declarations of interest

Eleven trials of the 70 trials reported information related to potential conflicts of interests for the trial authors, primarily related to income received from pharmaceutical and other

commercial organisations (Carlson 2013; Freeman 2008; Harper 2010; Hauner 2012; Helland 2001; Hurtado 2015; Krauss-Etschmann 2007; Makrides 2010; Mozurkewich 2013; Noakes 2012; Olsen 1992). A further 31 trials reported having no interests to declare (Ali 2017; Bergmann 2007; Bosaeus 2015; de Groot 2004; Dilli 2018; Dunstan 2008; Furuhjelm 2009; Haghiac 2015; Horvaticek 2017; Ismail 2016; Jamilian 2016; Jamilian 2017; Judge 2014; Kaviani 2014; Khalili 2016; Krummel 2016; Malcolm 2003; Mardones 2008; Min 2014; Min 2016; Mulder 2014; Oken 2013; Pietrantoni 2014; Ramakrishnan 2010; Razavi 2017; Ribeiro 2012; Taghizadeh 2016; Valenzuela 2015; Van Goor 2009; Van Winden 2017; Vaz 2017).

The remaining 28 trials did not report any information regarding declarations of interest (Bisgaard 2016; Boris 2004; Bulstra-Ramakers 1994; Chase 2015; D'Almedia 1992; England 1989; Giorlandino 2013; Gustafson 2013; Harris 2015; Judge 2007; Keenan 2014; Knudsen 2006; Laivuori 1993; Martin-Alvarez 2012; Miller 2016; Ogundipe 2016; Olsen 2000; Onwude 1995; Otto 2000; Ranjkesh 2011; Rees 2008; Rivas-Echeverria 2000; Samimi 2015; Sanjurjo 2004; Smuts 2003a; Smuts 2003b; Su 2008; Tofail 2006). For further details of the reported declarations, see Characteristics of included studies.

Excluded studies

We excluded 15 studies (Escobar 2008; Fievet 1985; Gholami 2017; Herrera 1993; Herrera 1998; Herrera 2004; Lauritzen 2004; Marangell 2004; Morrison 1984; Morrison 1986; Nishi 2016; Starling 1990; Valentine 2013; Velzing-Aarts 2001; Yelland 2016). Four trials assessed the effects of an omega-6 fatty acid intervention (linoleic acid) (Herrera 1993; Herrera 1998; Herrera 2004; Morrison 1984), and one trial assessed evening primrose oil (Fievet 1985). In Escobar 2008, participants were registered, but none were recruited. In five trials participants were not randomised (Gholami 2017; Marangell 2004; Nishi 2016; Starling 1990; Velzing-Aarts 2001), and in another it did not appear as if participants were randomised (Morrison 1986). In Lauritzen 2004 and Valentine 2013 women were supplemented with omega-3 during lactation only. The remaining trial was a methodological study that assessed aspects of several trials (Yelland 2016).

Risk of bias in included studies

For a summary of the risk of bias across the included trials, see Figure 2 and Figure 3.



Figure 2. Risk of bias summary: review authors' judgements about each risk of bias item for each included study.

	Random sequence generation (selection bias)	Allocation concealment (selection bias)	Blinding of participants and personnel (performance bias)	Blinding of outcome assessment (detection bias)	Incomplete outcome data (attrition bias)	Selective reporting (reporting bias)	Other bias
Ali 2017	•	•	?	?	?	?	•
Bergmann 2007	•	?	•	•	?	•	•
Bisgaard 2016	•	•	•	•	•	•	•
Boris 2004	?	?	?	?	•	?	•
Bosaeus 2015	•	?		?	•	?	?
Bulstra-Ramakers 1994	?	•	•	•	?	?	?
Carlson 2013	•	•	•	•	?	•	•
Chase 2015	?	?	?	?	?	?	?
D'Almedia 1992	•	•	•	•	?	?	?
de Groot 2004	?	?	?	?		?	?
Dilli 2018	?	?	•	?	•	?	?
Dunstan 2008	?	•	•	•	?	?	?
England 1989	?	?	?	?	?	?	•
Freeman 2008	?	?	•	•			?
Furuhjelm 2009	?	?	•	•	?	?	•
Giorlandino 2013	•	?	•	•	?	?	•
Gustafson 2013	•	•	•	?	?	•	•
Haghiac 2015	•	•	•	•		?	?
Harper 2010	•	•	•	•	•	•	•
Harris 2015						•	



Figure 2. (Continued)

Harris 2015								
Helland 2001 Horvaticek 2017 Hurtado 2015 Ismail 2016 Jamilian 2016 Jamilian 2017 Judge 2007 Judge 2014 Keenan 2014 Keenan 2014 Kawiani 2016 Knudsen 2006 Krauss-Etschmann 2007 Krummel 2016 Bakirides 2010 Makrides 2010 Makrides 2010 Miller 2016 Min 2014 [diabetic women] Min 2014 [diabetic women] Min 2014 Mozurkewich 2013 Mulder 2014 Mozurkewich 2013 Mulder 2014 Min 2014 Mozurkewich 2013 Mulder 2014 Min 2014 Min 2014 Min 2016 Mozurkewich 2013 Mulder 2014 Min 2014 Min 2016 Mozurkewich 2013 Mulder 2014 Min 2014 Min 2016 Mozurkewich 2013 Mulder 2016 Mozurkewich 2018 Mulder 2016 Mozurkewich 2018 Mulder 2016 Mozurkewich 2018 Mozurkewich 2018 Mozurkewich 2018 Mozurkewich 2018 Mozurkewich 2018	Harris 2015	•	•	•	•	•	?	•
Horvaticek 2017 7 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	Hauner 2012	•	?	•		•	?	?
Hurtado 2015 Ismail 2016 Jamilian 2017 Judge 2007 Judge 2007 Judge 2014 Kaviani 2014 Keenan 2014 Khalili 2016 Ninudsen 2006 Krauss-Etschmann 2007 Krummel 2016 Jaiviori 1993 Makrides 2010 Malcolm 2003 Mardones 2008 Martin-Alvarez 2012 Miller 2016 Min 2014 Min 2016 Mozurkewich 2013 Mulder 2014 M	Helland 2001	•	?	•	•	•	?	?
Ismail 2016	Horvaticek 2017	?	?	•	?	•	•	?
Jamilian 2016	Hurtado 2015	•	?	•	•	•	?	•
Jamilian 2017	Ismail 2016	•	•	•	•	?	?	•
Judge 2007	Jamilian 2016	•	•	•	•	?	•	•
Judge 2014 ? • • • • ? ? • • ? ? • ? <td< td=""><td>Jamilian 2017</td><td>•</td><td>?</td><td>•</td><td>?</td><td>•</td><td>?</td><td>•</td></td<>	Jamilian 2017	•	?	•	?	•	?	•
Kaviani 2014 ? <t< td=""><td>Judge 2007</td><td>?</td><td>?</td><td>•</td><td>?</td><td>•</td><td>?</td><td>?</td></t<>	Judge 2007	?	?	•	?	•	?	?
Keenan 2014 • • • • • • • • • • • • • • • • • • •	Judge 2014	?	•	•	•	•	•	?
Khalili 2016 Image: square	Kaviani 2014	?	?	•	?	?	•	?
Knudsen 2006 ? ? ? ?	Keenan 2014	•	•	•	•	?	?	?
Krauss-Etschmann 2007 ? ? . . ? ? .	Khalili 2016	•	•	•	•	?	•	•
Krummel 2016 + + + + + - <t< td=""><td>Knudsen 2006</td><td>?</td><td>?</td><td>?</td><td>?</td><td>•</td><td>•</td><td>•</td></t<>	Knudsen 2006	?	?	?	?	•	•	•
Laivuori 1993 ? ? • • • • ? • • • • Makrides 2010 • • • • • • • • • • • • • • • • • •	Krauss-Etschmann 2007	?	?	•	•	•	?	•
Makrides 2010 ⊕ ⊕ ⊕ ⊕ ⊕ ♀	Krummel 2016	•	•	•	•	•	?	?
Malcolm 2003 ? ? + ? + ? ? ? ? ? Mardones 2008	Laivuori 1993	?	?	•	•	•	?	•
Mardones 2008	Makrides 2010	•	•	•	•	•	•	•
Martin-Alvarez 2012 ? ? ? ? ? ? ? ? ? ? ? ? ? ?	Malcolm 2003	?	?	•	?	•	?	?
Miller 2016	Mardones 2008	•	•	?	?	•	?	?
Min 2014	Martin-Alvarez 2012	?	?	?	?	?	?	?
Min 2014 [diabetic women] Min 2016	Miller 2016	•	?	•	•	?	?	?
Min 2016	Min 2014	•	•	•	•	•	?	?
Mozurkewich 2013	Min 2014 [diabetic women]							
Mulder 2014	Min 2016	•	•	•	•	?	?	?
Noakes 2012	Mozurkewich 2013	•	•	•	?	•	•	•
Ogundipe 2016 ? ? ? ? ? ? ? ? Oken 2013 + + + + ? ? ?	Mulder 2014	•	?	•	•	?	?	?
Oken 2013 + + + + ? ? ?	Noakes 2012	•	?	?	•	?	?	•
	Ogundipe 2016	?	?	?	?	?	?	?
Olsen 1992 ? ? ? ? •	Oken 2013	•	•	•	•	?	?	?
	Olsen 1992	?	•	?	?	•	•	•

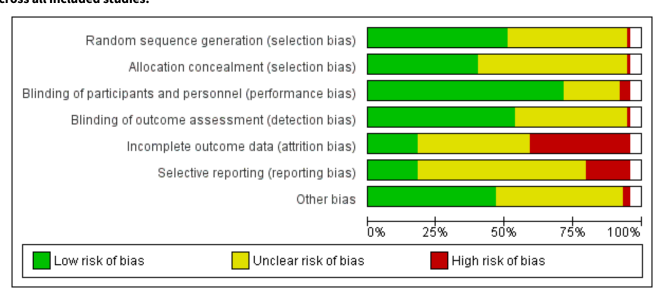


Figure 2. (Continued)

Olsen 1992	?	•	?	?	•	•	•
Olsen 2000	•	•	•	•	•	?	•
Olsen 2000 [twins]							
Onwude 1995	•	•	•	•	•	•	•
Otto 2000	?	?	•	?	•	•	•
Pietrantoni 2014	?	?	?	?	?	•	•
Ramakrishnan 2010	•	•	•	•	?	?	•
Ranjkesh 2011	?	?	•	?	•	?	•
Razavi 2017	•	•	•	•	•	?	•
Razavi 2017 [vit D]							
Rees 2008	•	•	•	•	•	•	•
Ribeiro 2012	?	?	?	?	?	?	?
Rivas-Echeverria 2000	?	?	•	?	?	•	?
Samimi 2015	•	•	•	•	•	?	•
Sanjurjo 2004	•	?	•	?	?	?	?
Smuts 2003a	•	?	•	•	?	•	•
Smuts 2003b	?	?	?	?	•	?	?
Su 2008	?	?	•	•	•	•	?
Taghizadeh 2016	•	•	•	•	?	•	•
Tofail 2006	?	?	•	•	•	•	?
Valenzuela 2015	?	?	?	?	•	?	•
Van Goor 2009	?	?	•	?	•	•	•
Van Winden 2017	?	?	•	?	•	?	?
Vaz 2017	?	?	•	?	•	?	?
	_						



Figure 3. Risk of bias graph: review authors' judgements about each risk of bias item presented as percentages across all included studies.



Allocation

Random sequence generation

We judged the methods used to generate the random sequence to be adequate in 37 of the 70 included trials (Ali 2017; Bergmann 2007; Bisgaard 2016; Bosaeus 2015; Carlson 2013; D'Almedia 1992; Giorlandino 2013; Gustafson 2013; Haghiac 2015; Harper 2010; Harris 2015; Hauner 2012; Helland 2001; Hurtado 2015; Ismail 2016; Jamilian 2016; Jamilian 2017; Keenan 2014; Khalili 2016; Krummel 2016; Makrides 2010; Miller 2016; Min 2014; Min 2016; Mozurkewich 2013; Mulder 2014; Noakes 2012; Oken 2013; Olsen 2000; Onwude 1995; Ramakrishnan 2010; Razavi 2017; Rees 2008; Samimi 2015; Sanjurjo 2004; Smuts 2003a; Taghizadeh 2016), with all trials using computer-generated methods or likely to have done so. We judged the risk of selection bias associated with sequence generation to be unclear in 32 trials, as many did not report how the random sequence was generated or provide sufficient information (Boris 2004; Bulstra-Ramakers 1994; Chase 2015; de Groot 2004; Dilli 2018; Dunstan 2008; England 1989; Freeman 2008; Furuhjelm 2009; Horvaticek 2017; Judge 2007; Judge 2014; Kaviani 2014; Knudsen 2006; Krauss-Etschmann 2007; Laivuori 1993; Malcolm 2003; Martin-Alvarez 2012; Ogundipe 2016; Olsen 1992; Otto 2000; Pietrantoni 2014; Ranjkesh 2011; Ribeiro 2012; Rivas-Echeverria 2000; Smuts 2003b; Su 2008; Tofail 2006; Valenzuela 2015; Van Goor 2009; Van Winden 2017; Vaz 2017). We judged one trial to be at high risk of selection bias, as alternation was used (odd and even numbers) (Mardones 2008).

Allocation concealment

We judged that 29 of the 70 trials had used adequate methods for allocation concealment (Ali 2017; Bisgaard 2016; Bulstra-Ramakers 1994; Carlson 2013; D'Almedia 1992; Dunstan 2008; Gustafson 2013; Haghiac 2015; Harper 2010; Harris 2015; Ismail 2016; Jamilian 2016; Judge 2014; Keenan 2014; Khalili 2016; Krummel 2016; Makrides 2010; Min 2014; Min 2016; Mozurkewich 2013; Oken 2013; Olsen 1992; Olsen 2000; Onwude 1995; Ramakrishnan 2010; Razavi 2017; Rees 2008; Samimi 2015; Taghizadeh 2016). Four of these reported using sequentially numbered, opaque sealed envelopes (Ali 2017;

Ismail 2016; Oken 2013; Olsen 1992). Three reported computer driven telephone or centre based randomisation (Harper 2010; Makrides 2010; Olsen 2000); 21 reported third party (pharmacy, health provider, supplement provider or external investigator) controlled randomisation (Bisgaard 2016; Bulstra-Ramakers 1994; Carlson 2013; D'Almedia 1992; Dunstan 2008; Gustafson 2013; Haghiac 2015; Harris 2015; Jamilian 2016; Judge 2014; Keenan 2014; Khalili 2016; Krummel 2016; Min 2014; Min 2016; Mozurkewich 2013; Ramakrishnan 2010; Razavi 2017; Rees 2008; Samimi 2015; Taghizadeh 2016), and one used sequentially numbered opaque sealed envelopes and third party (pharmacy) controlled randomisation (Onwude 1995).

Selection bias

We judged that the risk of selection bias associated with allocation concealment was unclear for 40 trials (Bergmann 2007; Boris 2004; Bosaeus 2015; Chase 2015; de Groot 2004; Dilli 2018; England 1989; Freeman 2008; Furuhjelm 2009; Giorlandino 2013; Hauner 2012; Helland 2001; Horvaticek 2017; Hurtado 2015; Jamilian 2017; Judge 2007; Kaviani 2014; Knudsen 2006; Krauss-Etschmann 2007; Laivuori 1993; Malcolm 2003; Martin-Alvarez 2012; Miller 2016; Mulder 2014; Noakes 2012; Ogundipe 2016; Otto 2000; Pietrantoni 2014; Ranjkesh 2011; Ribeiro 2012; Rivas-Echeverria 2000; Sanjurjo 2004; Smuts 2003a; Smuts 2003b; Su 2008; Tofail 2006; Valenzuela 2015; Van Goor 2009; Van Winden 2017; Vaz 2017), with either no methods of concealment detailed, or the methods described lacking sufficient detail. We judged one trial to be at high risk of selection bias, as alternation was used (odd and even numbers) (Mardones 2008).

Blinding

Blinding of participants and personnel

We judged blinding of participants and personnel to be adequate in 52 of the 70 included trials (Bergmann 2007; Bisgaard 2016; Bulstra-Ramakers 1994; Carlson 2013; D'Almedia 1992; Dilli 2018; Dunstan 2008; Freeman 2008; Furuhjelm 2009; Giorlandino 2013; Gustafson 2013; Haghiac 2015; Harper 2010; Harris 2015; Helland 2001; Horvaticek 2017; Hurtado 2015; Ismail 2016; Jamilian 2016;



Jamilian 2017; Judge 2007; Judge 2014; Kaviani 2014; Keenan 2014; Khalili 2016; Krauss-Etschmann 2007; Krummel 2016; Laivuori 1993; Makrides 2010; Malcolm 2003; Miller 2016; Min 2014; Min 2016; Mozurkewich 2013; Mulder 2014; Oken 2013; Olsen 2000; Onwude 1995; Ramakrishnan 2010; Ranjkesh 2011; Razavi 2017; Rees 2008; Rivas-Echeverria 2000; Samimi 2015; Sanjurjo 2004; Smuts 2003a; Su 2008; Taghizadeh 2016; Tofail 2006; Van Goor 2009; Van Winden 2017; Vaz 2017). We judged the risk of performance bias to be high in three trials, due to inadequate blinding of women and/or trial personnel (Bosaeus 2015; Hauner 2012; Otto 2000). For the remaining 15 trials, we judged the risk of performance bias to be unclear (Ali 2017; Boris 2004; Chase 2015; de Groot 2004; England 1989; Knudsen 2006; Mardones 2008; Martin-Alvarez 2012; Noakes 2012; Ogundipe 2016; Olsen 1992; Pietrantoni 2014; Ribeiro 2012; Smuts 2003b; Valenzuela 2015). Eleven of these trials did not provide sufficient information to allow confident assessment of blinding (Ali 2017; Chase 2015; de Groot 2004; England 1989; Knudsen 2006; Martin-Alvarez 2012; Noakes 2012; Ogundipe 2016; Pietrantoni 2014; Ribeiro 2012; Valenzuela 2015). Two trials reported that blinding of participants was partial (the notreatment groups were aware of this) (Boris 2004; Olsen 1992), and another two trials used food interventions which could not be fully blinded (Mardones 2008; Smuts 2003b).

Blinding of outcome assessors

Thirty-nine trials clearly indicated that blinded trial personnel were involved in outcome assessment or data collection, and we judged them to be at low risk of detection bias (Bergmann 2007; Bisgaard 2016; Bulstra-Ramakers 1994; Carlson 2013; D'Almedia 1992; Dunstan 2008; Freeman 2008; Furuhjelm 2009; Giorlandino 2013; Haghiac 2015; Harper 2010; Harris 2015; Helland 2001; Hurtado 2015; Ismail 2016; Jamilian 2016; Judge 2014; Keenan 2014; Khalili 2016; Krauss-Etschmann 2007; Krummel 2016; Laivuori 1993; Makrides 2010; Miller 2016; Min 2014; Min 2016; Mulder 2014; Noakes 2012; Oken 2013; Olsen 2000; Onwude 1995; Ramakrishnan 2010; Razavi 2017; Rees 2008; Samimi 2015; Smuts 2003a; Su 2008; Taghizadeh 2016; Tofail 2006). One trial reported that, except for ultrasound measurements (e.g. for fat mass measurements), assessors were not blinded and we judged to be at high risk of detection bias (Hauner 2012). For the remaining trials, we judged the risk of detection bias to be unclear, as most of them provided insufficient details about whether assessors and/or data collectors were blinded (Ali 2017; Boris 2004; Bosaeus 2015; Chase 2015; de Groot 2004; Dilli 2018; England 1989; Gustafson 2013; Horvaticek 2017; Jamilian 2017; Judge 2007; Kaviani 2014; Knudsen 2006; Malcolm 2003; Mardones 2008; Martin-Alvarez 2012; Mozurkewich 2013; Ogundipe 2016; Olsen 1992; Otto 2000; Pietrantoni 2014; Ranjkesh 2011; Ribeiro 2012; Rivas-Echeverria 2000; Sanjurjo 2004; Valenzuela 2015; Van Goor 2009; Van Winden 2017; Vaz 2017).

Incomplete outcome data

We judged 13 trials to be at low risk of attrition bias, with minimal losses to follow-up, and similar numbers/reasons for losses to follow-up in each group (Bisgaard 2016; Harper 2010; Jamilian 2017; Makrides 2010; Mozurkewich 2013; Olsen 1992; Olsen 2000; Onwude 1995; Otto 2000; Ranjkesh 2011; Razavi 2017; Samimi 2015; Valenzuela 2015).

We judged 27 trials to be at a high risk of attrition bias (Boris 2004; Bosaeus 2015; de Groot 2004; Dilli 2018; Freeman 2008; Haghiac 2015; Harris 2015; Hauner 2012; Helland 2001; Horvaticek 2017;

Hurtado 2015; Judge 2007; Judge 2014; Knudsen 2006; Krauss-Etschmann 2007; Krummel 2016; Laivuori 1993; Malcolm 2003; Mardones 2008; Min 2014; Rees 2008; Smuts 2003b; Su 2008; Tofail 2006; Van Goor 2009; Van Winden 2017; Vaz 2017). See Characteristics of included studies for further details.

We judged the remaining 30 trials to be at an unclear risk of attrition bias, often due to incomplete or unclear reporting, and complexity in some of the trials with several periods of childhood follow-up (Ali 2017; Bergmann 2007; Bulstra-Ramakers 1994; Carlson 2013; Chase 2015; D'Almedia 1992; Dunstan 2008; England 1989; Furuhjelm 2009; Giorlandino 2013; Gustafson 2013; Ismail 2016; Jamilian 2016; Judge 2007; Kaviani 2014; Keenan 2014; Khalili 2016; Martin-Alvarez 2012; Miller 2016; Min 2016; Mulder 2014; Noakes 2012; Ogundipe 2016; Oken 2013; Pietrantoni 2014; Ramakrishnan 2010; Rivas-Echeverria 2000; Smuts 2003a; Taghizadeh 2016; Van Goor 2009).

Selective reporting

We judged 13 trials to be at a low risk of reporting bias, as they provided data for the prespecified and/or expected outcomes (including from the published protocols) (Bergmann 2007; Bisgaard 2016; Carlson 2013; Harper 2010; Khalili 2016; Jamilian 2016; Makrides 2010; Mozurkewich 2013; Olsen 1992; Smuts 2003a; Taghizadeh 2016; Tofail 2006; Van Goor 2009). We judged 45 trials to be at an unclear risk of reporting bias (Ali 2017; Boris 2004; Bosaeus 2015; Bulstra-Ramakers 1994; Chase 2015; D'Almedia 1992; de Groot 2004; Dilli 2018; Dunstan 2008; England 1989; Furuhjelm 2009; Giorlandino 2013; Haghiac 2015; Harris 2015; Hauner 2012; Helland 2001; Hurtado 2015; Ismail 2016; Jamilian 2017; Judge 2007; Keenan 2014; Krauss-Etschmann 2007; Krummel 2016; Laivuori 1993; Malcolm 2003; Mardones 2008; Martin-Alvarez 2012; Miller 2016; Min 2014; Min 2016; Mulder 2014; Noakes 2012; Ogundipe 2016; Oken 2013; Olsen 2000; Ramakrishnan 2010; Ranjkesh 2011; Razavi 2017; Ribeiro 2012; Samimi 2015; Sanjurjo 2004; Smuts 2003b; Valenzuela 2015; Van Winden 2017; Vaz 2017). For the majority of these trials there was insufficient information to permit us to assess selective reporting confidently (i.e. no access to a published trial protocol). We judged the remaining 12 trials to be at a high risk of reporting bias (Freeman 2008; Gustafson 2013; Horvaticek 2017; Judge 2014; Kaviani 2014; Knudsen 2006; Onwude 1995; Otto 2000; Pietrantoni 2014; Rees 2008; Rivas-Echeverria 2000; Su 2008).

Kaviani 2014; Knudsen 2006; Pietrantoni 2014 and Rivas-Echeverria 2000 each reported only one of the expected or prespecified outcomes. Freeman 2008, Gustafson 2013; Judge 2014; Otto 2000; and Rees 2008 reported few of the prespecified or expected outcomes. Onwude 1995 reported a limited range of expected outcomes and incomplete data (no standard deviations) for two of the continuous outcomes (length of gestation and birthweight). Su 2008 reported few of the expected outcomes, and data were incomplete for birth outcomes.

Other potential sources of bias

We judged 34 trials to be at a low risk of other potential sources of bias (Ali 2017; Bergmann 2007; Bisgaard 2016; Boris 2004; Carlson 2013; England 1989; Furuhjelm 2009; Giorlandino 2013; Gustafson 2013; Harper 2010; Harris 2015; Hurtado 2015; Ismail 2016; Jamilian 2016; Jamilian 2017; Khalili 2016; Knudsen 2006; Krauss-Etschmann 2007; Makrides 2010; Mozurkewich 2013; Noakes 2012; Olsen 1992; Olsen 2000; Onwude 1995; Otto 2000; Pietrantoni 2014; Ramakrishnan 2010; Ranjkesh 2011; Razavi 2017; Samimi 2015;



Smuts 2003a; Taghizadeh 2016; Valenzuela 2015; Van Goor 2009). We judged another 34 trials to be at an unclear risk of other potential sources of bias (Bosaeus 2015; Bulstra-Ramakers 1994; Chase 2015; D'Almedia 1992; de Groot 2004; Dilli 2018; Dunstan 2008; Freeman 2008; Haghiac 2015; Hauner 2012; Helland 2001; Horvaticek 2017; Judge 2007; Judge 2014; Kaviani 2014; Keenan 2014; Krummel 2016; Malcolm 2003; Mardones 2008; Martin-Alvarez 2012; Miller 2016; Min 2014; Min 2016; Mulder 2014; Ogundipe 2016; Oken 2013; Ribeiro 2012; Rivas-Echeverria 2000; Sanjurjo 2004; Smuts 2003b; Su 2008; Tofail 2006; Van Winden 2017; Vaz 2017). We judged the remaining two trials, Laivuori 1993 and Rees 2008, to be at a high risk of other bias. In Laivuori 1993 there were substantial differences in the median length of supplementation between the three groups, which was a significant source of other bias, while in Rees 2008 women in the placebo group were more likely to have a co-morbid anxiety disorder (9/13 versus 3/13), which introduced significant baseline imbalance between groups that was relevant to all reported outcomes.

Effects of interventions

See: Summary of findings for the main comparison Birth/infant outcomes; Summary of findings 2 Maternal outcomes; Summary of findings 3 Child/adult outcomes; Summary of findings 4 Health service outcomes

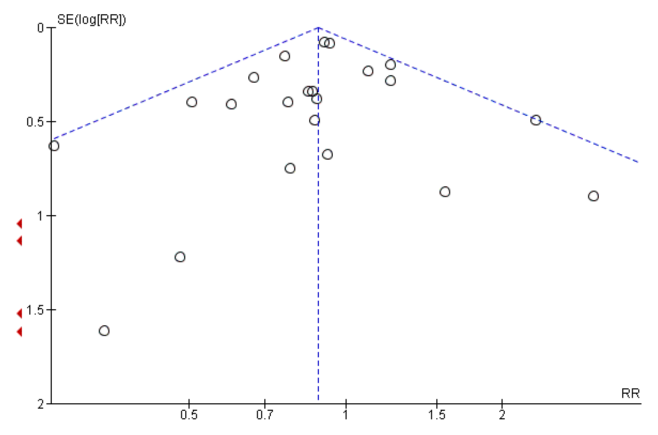
Omega-3 supplementation versus no omega-3

Primary outcomes

Preterm birth (< 37 weeks)

There was an 11% reduced risk of preterm birth (< 37 weeks) for omega-3 LCPUFA compared with no omega-3 (risk ratio (RR) 0.89, 95% confidence interval (CI) 0.81 to 0.97; 26 trials, 10,304 participants; high-quality evidence; Analysis 1.1). Some asymmetry was observed on visual assessment of a funnel plot for this outcome, suggesting an absence of some small negative studies (Figure 4), with little likely impact on the overall result.

Figure 4. Funnel plot of comparison: 1 Omega-3 versus placebo/no omega-3: OVERALL, outcome: 1.7 Preterm birth (< 37 weeks).



Early preterm birth (< 34 weeks)

There was a 42% lower risk of early preterm birth (< 34 weeks) for omega-3 LCPUFA compared with no omega-3 (RR 0.58, 95% CI 0.44 to 0.77; 9 trials, 5204 participants; high-quality evidence; Analysis 1.2).

Mother: secondary outcomes

Maternal death

Only four trials reported on maternal death (Bisgaard 2016; Makrides 2010; Oken 2013; Olsen 2000), with one maternal death reported in the omega-3 group in Oken 2013. There was no evidence of a difference in the risk of maternal death for omega-3 compared with no omega-3 (RR 1.69, 95% CI 0.07 to 39.30; 4 trials, 4830 participants; Analysis 1.4).

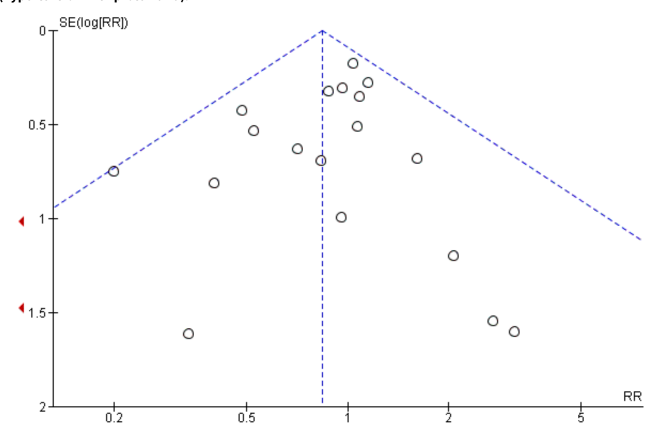


Pre-eclampsia (hypertension with proteinuria)

Pre-eclampsia (hypertension with proteinuria) may be reduced for omega-3 LCPUFA compared with no omega-3 group (RR 0.84, 95%

CI 0.69 to 1.01, 20 trials, 8306 participants; low-quality evidence; Analysis 1.5). No obvious asymmetry was observed on visual assessment of a funnel plot for this outcome (Figure 5).

Figure 5. Funnel plot of comparison: 1 Omega-3 versus placebo/no omega-3: OVERALL, outcome: 1.4 Pre-eclampsia (hypertension with proteinuria).



High blood pressure (without proteinuria)

There was no evidence of a difference in the risk of high blood pressure (without proteinuria) for omega-3 LCPUFA compared with no omega-3 (RR 1.03, 95% CI 0.89 to 1.20; 7 trials, 4531 participants; Analysis 1.6).

Eclampsia

Only one trial reported on eclampsia (D'Almedia 1992), and indicated no clear difference between omega-3 LCPUFA and no omega-3 (RR 0.14, 95% CI 0.01 to 2.70; 1 trial; 100 participants; Analysis 1.7).

Maternal antepartum hospitalisation

There was no evidence of a difference in risk of maternal antepartum hospitalisation between omega-3 LCPUFA and no omega-3 overall (RR 0.92, 95% CI 0.81 to 1.04; 5 trials, 2876 participants; Analysis 1.8).

Mother's length of stay in hospital (days)

Bisgaard 2016 and Olsen 2000 were the only trials to report data on the mother's length of stay in hospital, and showed no clear differences between omega-3 LCPUFA and no omega-3 (MD 0.18 days, 95% CI -0.20 to 0.57; 2 trials, 2290 participants; Analysis 1.9).

Maternal anaemia

Only Olsen 2000 reported on maternal anaemia and no difference was seen between omega-3 LCPUFA and no omega-3 (RR 1.16, 95% CI 0.91 to 1.48; 846 participants; Analysis 1.10).

Miscarriage (< 24 weeks)

There was no clear difference in miscarriage risk (< 24 weeks) for omega-3 LCPUFA compared with no omega-3 (RR 1.07, 95% CI 0.80 to 1.43; 9 trials, 4190 participants; Analysis 1.11).

Antepartum vaginal bleeding

There was no evidence of a difference in risk of antepartum vaginal bleeding for omega-3 LCPUFA compared with no omega-3 overall (RR 1.01, 95% CI 0.69 to 1.48; 2 trials, 2151 participants; Analysis 1.12).

Rupture of membranes

Carlson 2013, Harris 2015, Pietrantoni 2014 and Smuts 2003a reported on rupture of membranes (prelabour and preterm prelabour), and showed a lower risk overall with omega-3 LCPUFA compared with no omega-3 (RR 0.46, 95% CI 0.28 to 0.76; 4 trials, 1281 participants. The separate results for prelabour and preterm prelabour rupture are shown in Analysis 1.13.



Maternal admission to intensive care

Two trials reported on maternal admission to intensive care (Makrides 2010; Taghizadeh 2016), and saw no evidence of a difference in risk between omega-3 LCPUFA and no omega-3 (RR 0.56, 95% CI 0.12 to 2.63; 2 trials, 2458 participants; low-quality evidence; Analysis 1.14).

Maternal adverse events

Overall 16 trials reported on one or more maternal adverse effects. Using a random-effects model, there was no evidence of a difference in the risk of: severe adverse events (RR 1.04, 95% CI 0.40 to 2.72; 2 trials, 2690 participants; low-quality evidence), adverse events severe enough for cessation (RR 1.01, 95% CI 0.53 to 1.93; 6 trials, 1487 participants), any adverse effects (RR 1.38, 95% CI 1.16 to 1.65; $I^2 = 88\%$; 5 trials, 1480 participants), possibly due to

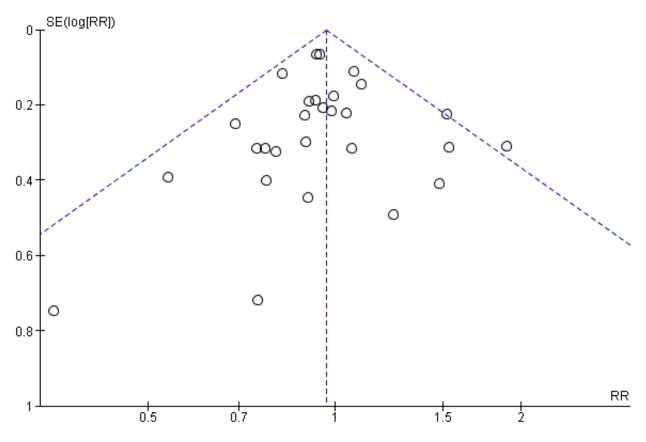
higher reports of belching/burping in the omega-3 LCPUFA group of Olsen 2000, and fewer reports of labour-related complications in the omega-3 LCPUFA group of Smuts 2003a (Analysis 1.15).

Very few differences were seen for individual adverse events, although unpleasant taste and belching/burping were more likely to be reported with omega-3 LCPUFA than with no omega-3 (Analysis 1.15).

Caesarean section

There was no evidence of a difference in the risk of caesarean section in omega-3 LCPUFA compared with no omega-3 (RR 0.97, 95% CI 0.91 to 1.03; 28 trials, 8481 participants; Analysis 1.16). No clear asymmetry was observed on visual assessment of a funnel plot for this outcome, although there was some indication that small negative trials may be missing (Figure 6). However this would be unlikely to affect the null findings.

Figure 6. Funnel plot of comparison: 1 Omega-3 versus placebo/no omega-3: OVERALL, outcome: 1.16 Caesarean section.



Induction (post-term)

Three trials reported on induction post-term (Harris 2015; Hauner 2012; Makrides 2010). The effect of omega-3 on post-term induction is uncertain due to the wide confidence intervals and variation between the results of the studies (average RR 0.82, CI 0.22 to 2.98; 2900 participants, 3 trials; $\text{Tau}^2 = 0.70$, P = 0.04, $I^2 = 77\%$; low-quality evidence; Analysis 1.17).

Blood loss at birth (mL)

There was no evidence of a difference in maternal blood loss at birth between omega-3 LCPUFA and no omega-3 (MD 11.50 mL, 95% CI -6.75 to 29.76; 6 trials, 2776 participants; Analysis 1.18).

Postpartum haemorrhage

Four trials reported on postpartum haemorrhage (Carlson 2013; Harper 2010; Makrides 2010; Olsen 1992), and found no evidence of a difference between omega-3 LCPUFA and no omega-3



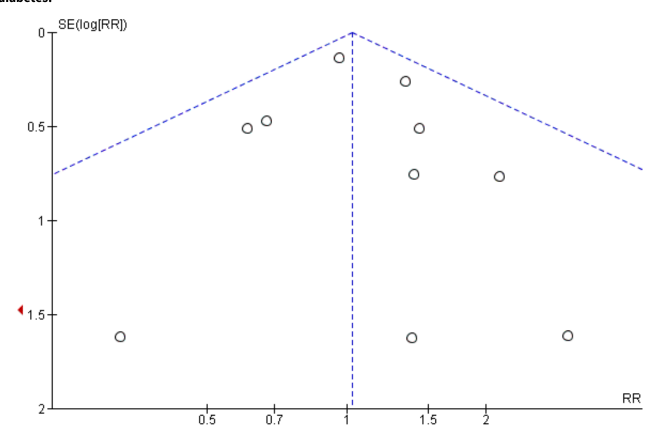
(RR 1.03, 95% CI 0.82 to 1.30; 4 trials, 4085 participants; Analysis 1.19).

Gestational diabetes

There was no evidence of a difference in the risk of GDM for omega-3 LCPUFA compared with no omega-3 (RR 1.02, 95% CI 0.83 to 1.26;

12 trials, 5235 participants; Analysis 1.20). No obvious asymmetry was observed on visual assessment of a funnel plot for this outcome (Figure 7).

Figure 7. Funnel plot of comparison: 1 Omega-3 versus placebo/no omega-3: OVERALL, outcome: 1.20 Gestational diabetes.



Maternal insulin resistance (HOMA-IR)

Only three trials reported on maternal insulin resistance (HOMA-IR) (Krummel 2016; Samimi 2015; Taghizadeh 2016), and showed no clear differences overall for omega-3 LCPUFA compared with no omega-3 (average MD -0.85, 95% CI -2.50 to 0.80; Tau 2 = 1.82; P = 0.0008; I 2 = 86%; 176 participants; Analysis 1.21). The high statistical heterogeneity may be due to different populations (overweight/obese women in Krummel 2016 and women with GDM in the other two trials).

Excessive gestational weight gain

Only Carlson 2013 reported on excessive gestational weight gain, and observed no evidence of a difference in the risk between

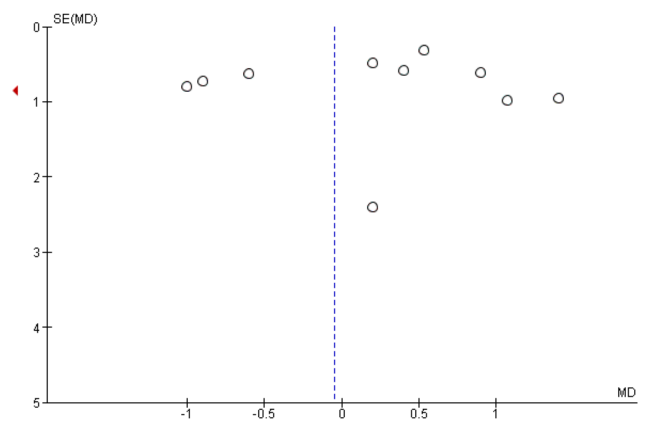
omega-3 LCPUFA and no omega-3 groups (RR 1.21, 95% CI 0.95 to 1.55; 350 participants; Analysis 1.22).

Gestational weight gain (kg)

There was no evidence of a difference in gestational weight gain for omega-3 LCPUFA compared with no omega-3 (MD -0.50 kg, 95% CI -0.68 to 0.59; 11 trials; random effects; $Tau^2 = 0.60$; P = 0.0006; $I^2 = 59\%$; 2297 participants; Analysis 1.23). The funnel plot was not markedly asymmetric (Figure 8). Dilli 2018 contributed to the high statistical heterogeneity, with a 3 kg lower gain in the omega-3 LCPUFA group compared with placebo.



Figure 8. Funnel plot of comparison: 1 Overall: omega-3 versus no omega-3, outcome: 1.23 Gestational weight gain (kg).



Depression during pregnancy: thresholds

Carlson 2013, Su 2008 and Vaz 2017 reported on different thresholds for depression during pregnancy (using the Hamilton Rating Scale for Depression (HAM-D), Edinburgh Postnatal Depression Scale (EPDS) and not specified), and showed no evidence of a difference between omega-3 LCPUFA and no omega-3 for each trial (Analysis 1.24).

Depression during pregnancy: scores

Depression scores during pregnancy were reported by five trials using four different methods (Beck Depression Inventory (BDI), HAM-D, EPDS and the Montgomery-Åsberg Depression Rating Scale (MADRS)). Only BDI showed a result favouring omega-3 LCPUFA over no omega-3 (MD -5.86 points 95% CI -8.32 to -3.39; 2 trials, 104 participants) with the other three comparisons showing no evidence of an effect (Analysis 1.25).

Anxiety during pregnancy

Only Carlson 2013 reported on anxiety during pregnancy, and observed no evidence of a difference between omega-3 LCPUFA and no omega-3 (RR 0.95, 95% CI 0.06 to 15.12; 301 participants; Analysis 1.26).

Difficult life circumstances (maternal)

Only Keenan 2014 reported on difficult life circumstances (maternal), indicating no evidence of a difference between omega-3

LCPUFA and no omega-3 (MD 0.32, 95% CI -0.15 to 0.79; 51 participants; Analysis 1.27).

Stress (maternal)

Keenan 2014 was also the only trial to report on maternal stress, showing no important difference between omega-3 LCPUFA and no omega-3 as measured by the perceived stress scale (MD -1.82 points, 95% CI -3.68 to 0.04; 51 participants; Analysis 1.28).

Depressive symptoms postpartum: thresholds

Postpartum depression scores were reported by four trials using three different methods (Postpartum Depression Screening Scale (PDSS) \geq 80, EPDS, and major depressive disorder), with none of the trials showing clear differences between omega-3 LCPUFA and no omega-3 (Analysis 1.29).

Depressive symptoms postpartum: scores

Only two trials reported on scores for postpartum depressive symptoms (Judge 2014; Mozurkewich 2013), and found no clear differences between omega-3 LCPUFA and no omega-3 for either BDI, PDSS overall or components of PDSS up to six months postpartum (Analysis 1.30).

Length of gestation (days)

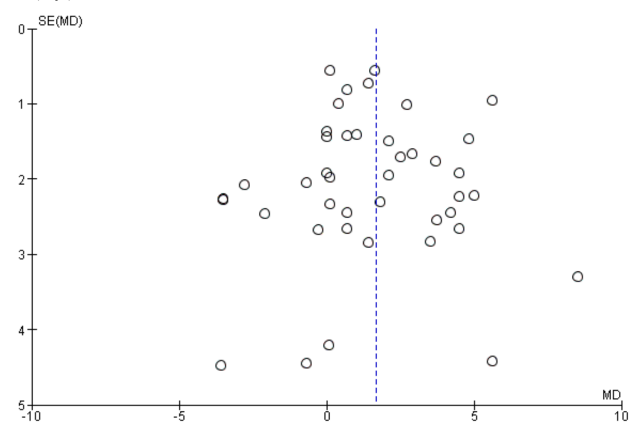
There was an increase in length of gestation with omega-3 LCPUFA compared with no omega-3 (average MD 1.67 days, 95% CI 0.95 to 2.39; Tau² = 2.33; P < 0.0001; I² = 52%; 41 trials, 12,517 participants;



moderate-quality evidence; Analysis 1.31). Reasons for the high statistical heterogeneity are not readily apparent, although there were wide variations in populations, inclusion criteria and doses of omega-3. Additionally, it was not always clear how length of

gestation was determined and this may have varied across studies. No obvious asymmetry was observed on visual assessment of a funnel plot for this outcome (Figure 9).

Figure 9. Funnel plot of comparison: 1 OVERALL omega-3 versus placebo/no omega-3, outcome: 1.31 Length of gestation (days).



Baby/infant/child

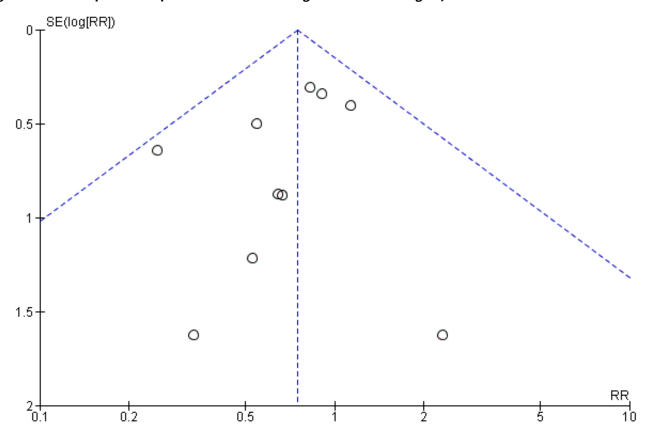
Perinatal death

There were fewer perinatal deaths in the omega-3 LCPUFA groups than the no omega-3 groups, though this did not reach

conventional statistical significance (RR 0.75, 95% CI 0.54 to 1.03; 10 trials, 7416 participants; moderate-quality evidence; Analysis 1.32). No obvious asymmetry was observed on visual assessment of a funnel plot for this outcome (Figure 10).



Figure 10. Funnel plot of comparison: 1 OVERALL omega-3 versus no omega-3, outcome: 1.32 Perinatal death.

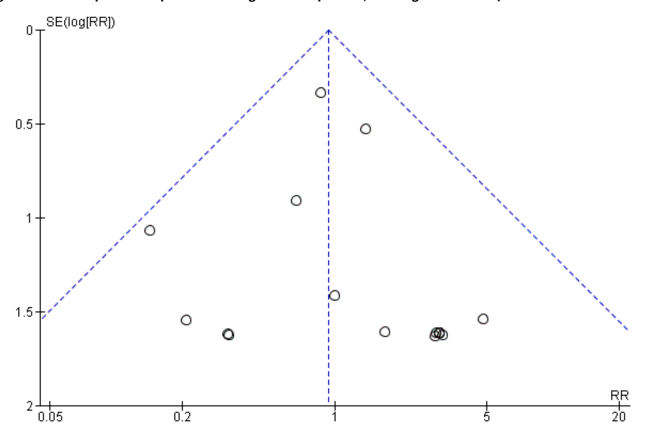


Stillbirth

No clear differences in still birth were seen between omega-3 LCPUFA and no omega-3 (RR 0.94,95% CI 0.62 to 1.42;16 trials, 7880 participants; Analysis 1.33). No obvious asymmetry was observed on visual assessment of a funnel plot for this outcome (Figure 11).



Figure 11. Funnel plot of comparison: 1 Omega-3 versus placebo/no omega-3: OVERALL, outcome: 1.32 Stillbirth.



Neonatal death

No clear difference between omega-3 LCPUFA and no omega-3 was seen for neonatal death (RR 0.61, 95% CI 0.34 to 1.11; 9 trials, 7448 participants; Analysis 1.34).

Infant death

Four trials reported on infant death (Carlson 2013; Makrides 2010; Mulder 2014; Tofail 2006), and observed no evidence of a difference in risk between the omega-3 LCPUFA and no omega-3 groups (RR 0.74, 95% CI 0.25 to 2.19; 3239 participants; Analysis 1.35).

Large-for-gestational age

Six trials reported on large-for-gestational age (generally defined as greater than the 90th percentile) (Dilli 2018; Harper 2010; Hauner 2012; Makrides 2010; Min 2014; Taghizadeh 2016), with a possible small increase in risk with omega-3 LCPUFA than no omega-3

(RR 1.15, 95% CI 0.97 to 1.03; 3722 participants; moderate-quality evidence; Analysis 1.36). This outcome was not prespecified in the protocol.

Macrosomia

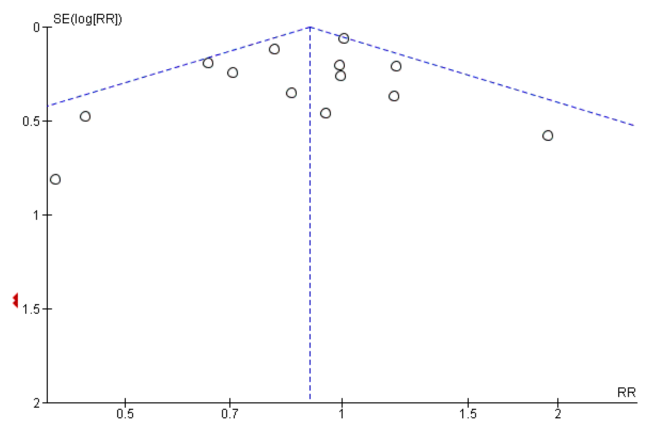
For macrosomia (generally defined as birthweight < 4000 g), no clear differences were seen between omega-3 LCPUFA and no omega-3 (RR 0.69, 95% CI 0.43 to 1.13; 6 trials, 2008 participants; Analysis 1.37). This outcome was not prespecified in the protocol.

Low birthweight (< 2500 g)

Rates of low birthweight (< 2500 g) showed a 10% relative risk reduction with omega-3 LCPUFA compared with no omega-3 (RR 0.90, 95% CI 0.82 to 0.99; 15 trials, 8449 participants; high-quality evidence Analysis 1.38). No obvious asymmetry was observed on visual assessment of a funnel plot for this outcome (Figure 12).



Figure 12. Funnel plot of comparison: 1 Omega-3 versus placebo/no omega-3: OVERALL, outcome: 1.38 Low birthweight (< 2500 g).



Small-for-gestational age or intrauterine growth restriction (IUGR)

There was little or no evidence of a difference in risk of small-forgestational age or IUGR between omega-3 LCPUFA and no omega-3 (RR 1.01, 95% CI 0.90 to 1.13; 8 trials, 6907 participants; moderate-quality evidence; Analysis 1.39).

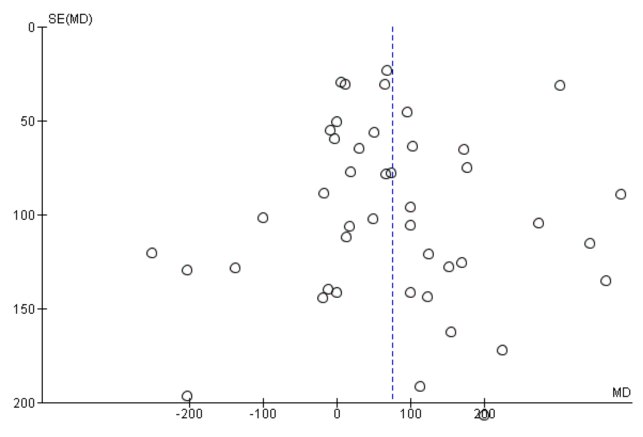
Birthweight (g)

Birthweight was higher in the omega-3 LCPUFA group than the no omega-3 group (average MD 75.74 g, 95% CI 38.05 to 113.43; Tau² =

7943.10; P < 0.00001; I² = 66%; 42 trials, 11,584 participants; Analysis 1.40). Reasons for the high statistical heterogeneity were not readily apparent, although there was a wide variation in birthweights between studies and inclusion criteria. No obvious asymmetry was observed on visual assessment of a funnel plot for this outcome (Figure 13).



Figure 13. Funnel plot of comparison: 1 Omega-3 versus placebo/no omega-3: OVERALL, outcome: 1.41 Birthweight (g).



Birthweight Z score

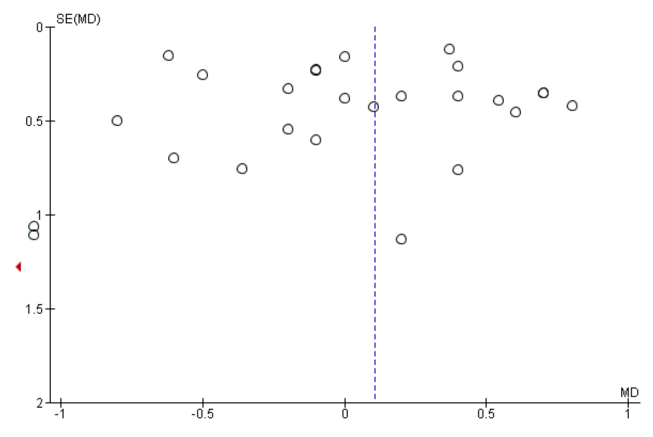
Four trials reported on birthweight Z score (Bergmann 2007; Krummel 2016; Makrides 2010; Mulder 2014), and there was no evidence of a difference between omega-3 LCPUFA and no omega-3 (MD 0.06, 95% CI -0.02 to 0.13; 2792 participants; Analysis 1.41).

Birth length (cm)

There was no evidence of a difference in birth length for omega-3 LCPUFA compared with no omega-3 (MD 0.11 cm, 95% CI -0.10 to 0.31; $Tau^2 = 0.13$; P = 0.0001, $I^2 = 57\%$; 28 trials, 8128 participants; Analysis 1.42). No clear asymmetry was observed on visual assessment of a funnel plot for this outcome (Figure 14).



Figure 14. Funnel plot of comparison: 1 Omega-3 versus placebo/no omega-3: OVERALL, outcome: 1.43 Birth length (cm).



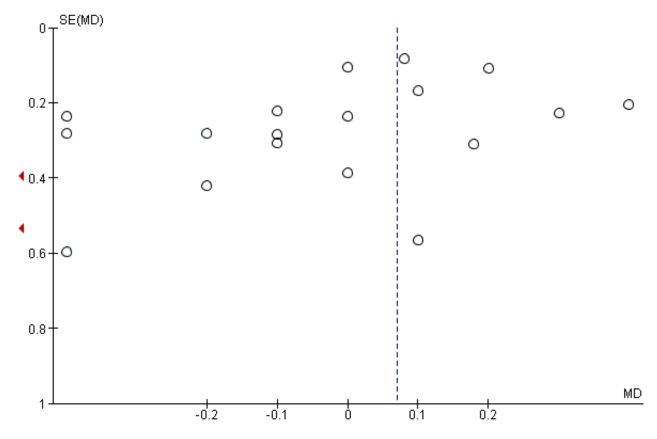
Head circumference at birth (cm)

There was no evidence of a difference in head circumference at birth for omega-3 LCPUFA compared with no omega-3 (average MD 0.07 $\,$

cm, 95% CI -0.05 to 0.19; 22 trials, 7161 participants; Tau^2 0.02, P = 0.06, $I^2 = 33\%$, Analysis 1.43). No clear asymmetry was observed on visual assessment of a funnel plot for this outcome (Figure 15).



Figure 15. Funnel plot of comparison: 1 Omega-3 versus placebo/no omega-3: OVERALL, outcome: 1.45 Head circumference at birth (cm).



Head circumference at birth Z score

Only two trials reported on head circumference at birth Z score (Krummel 2016; Makrides 2010), and there was no evidence of a difference between the omega-3 LCPUFA and no omega-3 groups (MD -0.03, 95% CI -0.14 to 0.07; 2462 participants; Analysis 1.44).

Length at birth Z score

Only two trials reported on length at birth Z score (Krummel 2016; Makrides 2010), and observed no clear difference between omega-3 LCPUFA and no omega-3 (average MD 0.18, 95% CI -0.18 to 0.54; Tau² = 0.05; P = 0.12; $I^2 = 59\%$; 2462 participants; Analysis 1.45).

Baby admitted to neonatal care

There was an 8% relative reduced risk of a baby being admitted to neonatal care with omega-3 LCPUFA compared with no omega-3, although this did not reach conventional statistical significance (RR 0.92, 95% CI 0.83 to 1.03; 9 trials, 6920 participants; Analysis 1.46).

Infant length of stay in hospital (days)

Only Olsen 2000 reported on infant length of stay in hospital and observed no evidence of a difference in length between the omega-3 LCPUFA and no omega-3 groups (MD 0.11 days, 95% CI -1.40 to 1.62; 2014 participants; Analysis 1.47).

Congenital anomalies

Three trials reported on congenital anomalies (Carlson 2013; Olsen 1992; Ramakrishnan 2010), and observed no clear difference

between the omega-3 LCPUFA and no omega-3 groups (RR 1.08, 95% CI 0.61 to 1.92; 1807 participants; Analysis 1.48).

Retinopathy of prematurity

Only one trial reported on retinopathy of prematurity (Harper 2010), and there was no clear difference between the omega-3 LCPUFA and no omega-3 groups (RR 1.20, 95% CI 0.32 to 4.44; 837 participants; Analysis 1.49).

Bronchopulmonary dysplasia

Only Harper 2010 and Makrides 2010 reported on bronchopulmonary dysplasia, and there was no clear difference between the omega-3 LCPUFA and no omega-3 groups (RR 1.06, 95% CI 0.45 to 2.48; 3191 participants; Analysis 1.50).

Respiratory distress syndrome

Two trials reported on respiratory distress syndrome (Carlson 2013; Harper 2010), and found no clear difference between the omega-3 LCPUFA and no omega-3 groups (average RR 1.17, 95% CI 0.54 to 2.52; Tau² = 0.21; P = 0.09; I² = 66%; 1129 participants; Analysis 1.51). Reasons for the statistical heterogeneity were not readily apparent although all the women in Harper 2010 had experienced a previous preterm birth and were treated with weekly intramuscular progesterone injections.



Necrotising enterocolitis (NEC)

Only Harper 2010 and Makrides 2010 reported on NEC, and found no clear difference between the omega-3 LCPUFA and no omega-3 groups (RR 0.97, 95% CI 0.26 to 3.55; 3198 participants; Analysis 1.52).

Neonatal sepsis (proven)

Harper 2010, Helland 2001 and Makrides 2010 reported on proven neonatal sepsis, and found no evidence of a difference in risk between the omega-3 LCPUFA and no omega-3 groups (RR 0.97, 95% CI 0.44 to 2.14; 3788 participants; Analysis 1.53).

Convulsion

Only Makrides 2010 reported on convulsion, and observed no clear difference between the omega-3 LCPUFA and no omega-3 groups (RR 0.09, 95% CI 0.01 to 1.63; 2361 participants; Analysis 1.54).

Intraventricular haemorrhage

Three trials reported on intraventricular haemorrhage (Harper 2010; Makrides 2010; Olsen 2000), and found no evidence of a difference in risk between the omega-3 LCPUFA and no omega-3 groups in any intraventricular haemorrhage (RR 1.00, 95% CI 0.29 to 3.49; random effects; Tau² = 0.63; P = 0.12, I² = 53%; 5423 participants). Although Makrides 2010 showed a marked reduction in intraventricular haemorrhage, reasons for the statistical heterogeneity were not clear. Harper 2010 also reported Grade 3 or 4 intraventricular haemorrhage, finding no clear differences between omega-3 LCPUFA and no omega-3 (RR 1.60, 95% CI 0.38 to 6.65; 837 participants; Analysis 1.55).

Neonatal/infant adverse events

There was possibly a small decrease for any adverse events in neonates/infants with omega-3 LCPUFA compared with no omega-3 (RR 0.92, 95% CI 0.82 to 1.02; 2 trials, 592 participants; Analysis 1.56). For serious neonatal/infant adverse events, there was a reduced risk with omega-3 LCPUFA compared with no omega-3 (RR 0.72, 95% CI 0.53 to 0.99; 2 trials, 2690 participants; low-quality evidence; Analysis 1.56).

Neonatal/infant morbidity

One trial of 291 infants reported on neonatal/infant cardiovascular, respiratory or morbidity caused by pregnancy/birth (Smuts 2003a), and found no evidence of difference between omega-3 LCPUFA and no omega-3 (Analysis 1.57; Analysis 1.58 and Analysis 1.59 respectively).

Another trial with 834 participants reported on neonatal/infant morbidity (Ramakrishnan 2010), and found no important differences in rates between omega-3 LCPUFA and no omega-3 for colds, fevers, rash, respiratory illnesses, vomiting, diarrhoea or other illnesses up to six months of age (Analysis 1.60).

Infant/child morbidity

Makrides 2010 reported on infant/child morbidity, and found a potentially reduced risk of ICU admissions for omega-3 LCPUFA compared with no omega-3 (RR 0.58, 95% CI 0.31 to 1.06; 1396 participants, but no clear differences for medical diagnosis of attention deficit hyperactivity disorder; autism spectrum disorder; other learning/behavioural disorders or other chronic health conditions; Analysis 1.61).

Ponderal index (g/m³ x 100)

There was no evidence of a difference in ponderal index between the omega-3 LCPUFA and no omega-3 groups (MD $0.05 \text{ g/m}^3 \times 100$, 95% CI -0.01 to 0.11; random effects, Tau² = 0.00, P = 0.07, I² = 50%, 6 trials, 887 participants; Analysis 1.62).

Infant/child weight (kg)

A total of 10 trials reported on infant/child weight at various time points, and there was no evidence of a difference between the omega-3 LCPUFA and no omega-3 groups at any of the time points from six weeks to seven years of age (Analysis 1.63).

Infant/child length/height (cm)

Ten trials reported on length/height at various time points, and there was no evidence of a difference between the omega-3 and no omega-3 groups from six weeks to five years. However there was evidence of child height being lower in the omega-3 LCPUFA compared with no omega-3 groups at seven years (MD-1.22 cm 95% CI-2.29 to -0.16; 2 trials, 393 participants; Analysis 1.64).

Infant/child head circumference (cm)

A total of 10 trials reported on infant/child head circumference at various time points, and there was no evidence of a difference between the omega-3 LCPUFA and no omega-3 groups at any of the time points from six weeks to six years of age (Analysis 1.65).

Infant/child length/height for age Z score (LAZ/HAZ)

Three trials reported on infant/child length/height for age Z score (LAZ/HAZ) at various time points (Mulder 2014; Ramakrishnan 2010; Tofail 2006), and observed no evidence of a difference between the omega-3 LCPUFA and no omega-3 groups at any of the time points from two months to five years (Analysis 1.66).

Infant/child waist circumference (cm)

Hauner 2012 and Makrides 2010 reported on infant/child waist circumference at various time points, and observed no evidence of a difference between the omega-3 LCPUFA and no omega-3 groups at any of the time points from two to five years (Analysis 1.67).

Infant/child weight-for-age Z score (WAZ)

Mulder 2014 and Ramakrishnan 2010 reported on infant/child weight-for-age Z score (WAZ) at various time points, and observed no evidence of a difference between the omega-3 LCPUFA and no omega-3 groups at any of the time points from one month to five years (Analysis 1.68).

Infant/child BMI Z score

Five trials reported on infant/child BMI Z score at various time points (Bergmann 2007; Carlson 2013; Ramakrishnan 2010; Krummel 2016; Makrides 2010), and found no evidence of a difference between the omega-3 LCPUFA and no omega-3 groups between any of the time points from 18 months to seven years of age (Analysis 1.69).

Infant/child weight for length/height Z score (WHZ)

Mulder 2014, Ramakrishnan 2010 and Tofail 2006 reported on infant/child weight for length/height Z score (WHZ) at various time points and there was no evidence of a difference between the omega-3 LCPUFA and no omega-3 groups at any of the time points from two months to 18 months old (Analysis 1.70).



Infant/child BMI percentile

Only Hauner 2012 reported on infant/child BMI percentile and there was no evidence of a difference between the omega-3 LCPUFA and no omega-3 groups at two, three and five years of age. However a higher infant/child BMI percentile was observed in the omega-3 LCPUFA compared with the no omega-3 group at 48 months (MD 13.00%; 95% CI 3.19 to 22.81; 107 participants; Analysis 1.71).

Child/adult BMI

Helland 2001, Makrides 2010; and Olsen 1992 reported on child/adult BMI at various time points and there was no evidence of a difference between the omega-3 LCPUFA and no omega-3 groups at any of the time points from three to 19 years of age (Analysis 1.72).

Infant/child body fat (%)

Carlson 2013, Hauner 2012 and Makrides 2010 reported on infant/child body fat at various time points, and found no evidence of differences at any points from one to seven years of age between the omega-3 LCPUFA and no omega-3 groups (Analysis 1.73).

Infant/child total fat mass (kg)

Hauner 2012 and Makrides 2010 reported on infant/child total fat mass at various time points, and found no evidence of differences at any points from one to seven years of age between the omega-3 LCPUFA and no omega-3 groups (Analysis 1.74).

Cognition: thresholds

Makrides 2010, Mulder 2014 and Ramakrishnan 2010 reported Bayley Scales of Infant Development (BSID) II or III cognition thresholds at 18 months, and found no evidence of differences between the omega-3 LCPUFA and no omega-3 groups, except for BSID III < 85, which favoured omega-3 LCPUFA, in Makrides 2010 (RR 0.49, 95% CI 0.24 to 0.98; 726 participants; Analysis 1.75).

Cognition: scores

Nine trials reported cognition scores at various time points. There was no evidence of a difference in cognition scores between the omega-3 LCPUFA and no omega-3 groups at any time point from nine months to 12 years of age, as measured by BSID II or III, Fagan novelty preference, Kaufman Assessment Battery for Children (K-ABC) mental processing composite, Griffith Mental Development Scale (GMDS) general quotient score, Differential Ability Scales (DAS) II, Wechsler Abbreviated Scale of Intelligence (WASI) full-scale intelligence quotient (IQ) or Wechsler Intelligence Scale for Children (WISC) IV full scale IQ (Analysis 1.76).

Attention: scores

Three trials reported on attention scores at various time points and used different assessment measures (Krauss-Etschmann 2007, Makrides 2010; Ramakrishnan 2010). There was no evidence of a difference between omega-3 LCPUFA and no omega-3 groups at any time point or with any measure from 27 months to 8.5 years except in Ramakrishnan 2010, where lower attention scores were seen at five years as measured by K-CPT omissions (MD -1.90, 95% CI -3.39 to -0.41; 797 participants; Analysis 1.77).

Motor: thresholds

Two trials reported thresholds for motor scores (Mulder 2014; Ramakrishnan 2010), and observed no differences between

omega-3 LCPUFA and no omega-3 groups at 18 months of age (Analysis 1.78).

Motor: scores

No difference was observed in motor scores between the omega-3 LCPUFA and no omega-3 groups, as measured by BSID III or II at 4 to 18 months of age across six trials (Analysis 1.79).

Language: thresholds

In one trial with 726 participants there was no evidence of a difference in BSID III language score thresholds between the omega-3 LCPUFA and no omega-3 groups at 18 months (Makrides 2010). Howevever in Mulder 2014 (154 participants), most Communicative Development Inventories (CDI) language thresholds were higher with omega-3 LCPUFA in children at 14 and 18 months (Analysis 1.80).

Language: scores

No differences between omega-3 LCPUFA and no omega-3 were seen in any communication or language scores in children from four months to seven years of age, across four trials (Analysis 1.81).

Behaviour: thresholds

In one trial with 730 participants no differences between omega-3 LCPUFA and no omega-3 were seen in behaviour thresholds for children at 18 months of age (Analysis 1.82) (Ramakrishnan 2010).

Behaviour: scores

There were few differences between omega-3 LCPUFA and no omega-3 in behaviour scores in children measured with different tools and over different time points from birth to 12 years. However, there was evidence of less difficult behaviour in the placebo compared with the omega-3 LCPUFA group as measured by the Strengths and Difficulties Questionnaire (SDQ) Total Difficulties at six to nine years in Makrides 2010 (MD 1.08, 95% CI 0.18 to 1.98; 543 participants; Analysis 1.83).

Vision: visual acuity (cycles/degree)

Only Mulder 2014 and Judge 2007 reported on visual acuity, observing no evidence of a difference between groups at two, four or six months (Analysis 1.84).

Vision: visual evoked potential

Three trials reported visual evoked potential at various time points from birth to six months, with no important differences seen between omega-3 and no omega-3 (Analysis 1.85).

Hearing: brainstem auditory-evoked responses

Only one trial reported on various ways of measuring brainstem auditory-evoked responses from one to three months (Ramakrishnan 2010), and found no evidence of differences between the omega-3 LCPUFA and no omega-3 groups (Analysis 1.87).

Neurodevelopment (overall): thresholds

Three trials reported various measures of overall neurodevelopment from six months to five years, and observed no clear differences between the omega-3 LCPUFA and no omega-3 groups (Analysis 1.88).



Neurodevelopment (overall): scores

One trial reported components of the Ages and Stages Questionnaire (ASQ) at four and six months (Khalili 2016), and found no clear differences except for improved communication with omega-3 LCPUFA at four months (MD 2.70 95% CI 0.41 to 4.99; 148 participants; Analysis 1.89).

Child Development Inventory

Only Hauner 2012 reported on the child development inventory (parent-reported), and observed no clear differences between the omega-3 LCPUFA and no omega-3 groups, across a range of measures when children were five years old (Analysis 1.90).

Infant sleep behaviour

Only Judge 2007 reported on infant sleep behaviour, and found fewer arousals in quiet and active sleep with omega-3 LCPUFA compared with no omega, but with no other differences between groups; 39 participants (Analysis 1.91).

Cerebral palsy

A single trial with 114 participants reported no cases of cerebral palsy in either the omega-3 LCPUFA or the placebo group (Van Goor 2009) (Analysis 1.92).

None of the trials reported caesarean section (post-term), jaundice, or use of community health services.

Subgroup analyses

By intervention type (Analysis 2)

Analyses were performed (where possible) based on type of omega-3 intervention (omega-3 LCPUFA supplements only; omega-3 LCPUFA supplements, omega-3 rich food and/or dietary advice; omega-3 LCPUFA supplements and other agents; and omega-3 supplements, omega-3 LCPUFA rich food and other agents). No clear or important subgroup differences were revealed for any outcome except for:

 birth length, where birth length was higher with omega-3 LCPUFA supplements alone, or with omega-3 rich food and/ or diet advice; and lower when the intervention was omega-3 LCPUFA combined with another non-omega-3 agent (Analysis 2.40).

However the small increase is not likely to be clinically meaningful.

By dose of DHA and EPA supplements (Analysis 3)

Subgroup analysis based on dose of omega-3 supplements for low dose (\leq 500 mg/day) versus mid dose (500 mg to 1 g/day) versus high dose (> 1 g/day) revealed no clear or important difference for any of the 12 prespecified outcomes except for:

- low birthweight, where the effect of low and mid doses of omega-3 LCPUFA (500 mg to 1 g/day) appeared more pronounced in reducing low birthweight than high dose (Analysis 3.10); Chi² 6.17, P = 0.05, I² 67.6%; and
- birthweight (Analysis 3.12); Chi² 8.34, P = 0.04, I² 64%, which
 appears to be driven by a single trial of flaxseed oil. When this
 trial is omitted, the subgroup analysis is no longer statistically
 significant.

Timing of intervention: gestational age when omega-3 supplements commenced (Analysis 4)

Subgroup analysis based on the time when omega-3 LCPUFA supplements started (≤ 20 weeks' gestation or > 20 weeks' gestation) revealed no clear or important differences for any of the 12 prespecified outcomes except for pre-eclampsia (Analysis 4.4). However as the single trial contributing to the heterogeneity did not report timing of intervention (Rivas-Echeverria 2000), this does not help elucidate the influence of timing start of supplement on this outcome.

Type of supplements (Analysis 5)

Subgroup analysis based on type of supplements (DHA/largely DHA versus mixed DHA/EPA versus mixed DHA/EPA/other) revealed no clear subgroup differences for the outcomes, except for:

- **pre-eclampsia** (likely due to the influence of a single study, Rivas-Echeverria 2000), in the mixed DHA/EPA/other subgroup (Analysis 5.4); Chi² 7.58, P = 0.02, I² = 73.6%; and
- caesarean section where incidence was higher in the mixed DHA/EPA subgroup (Analysis 5.5); Chi² 6.29, P = 0.04, I² = 68.2% than for the other DHA or EPA subgroups.

Risk (Analysis 6)

Analyses were performed based on risk of women - low risk (healthy women or health condition unlikely to affect birth outcomes, e.g. allergy) versus increased/high risk (e.g. women with hypertension, gestational diabetes mellitus, depression, a history of preterm birth) versus mixed risk (no inclusion criteria related to maternal health risk, or health risk not reported). No clear or important subgroup differences were seen for any of the outcomes except low birthweight where studies with women at low or any risk showed a reduction compared with the studies involving women at increased or higher risk (Analysis 6.10); Chi² 6.24, P = 0.04, I² 67.9%.

Omega-3 dose direct comparisons (Analysis 7)

One trial reported outcomes in 224 participants from a direct comparison of 600 mg and 300 mg DHA a day (Harris 2015); and another trial reported five different doses of DHA/EPA from 0.1 g/day to 2.8 g/day (which we collapsed into \leq 1 g/day and > 1 g/day DHA/EPA) (Knudsen 2006). Knudsen 2006 only reported gestational length.

Primary outcomes

In one trial, no difference between doses was seen for:

- early preterm birth (< 34 weeks) (RR 0.91, 95% CI 0.13 to 6.38; Analysis 7.1); or
- prolonged gestation (> 42 weeks) (RR 0.91, 95% CI 0.06 to 14.44; Analysis 7.2) (Harris 2015).

Mother: secondary outcomes

One trial (Harris 2015), observed no evidence of a difference between women who received 600 mg DHA/day compared with 300 mg DHA/day for:

- pre-eclampsia (RR 0.91, 95% CI 0.06 to 14.44; Analysis 7.3);
- induction post term (RR 0.10; 95% CI 0.01 to 1.87; Analysis 7.4);
- premature rupture of membranes (RR 0.30, 95% CI 0.03 to 2.89; Analysis 7.5); or



premature prelabour rupture of membranes (RR 1.22, 95% CI 0.28 to 5.32; Analysis 7.6).

Two trials (Harris 2015; Knudsen 2006), observed no difference in length of gestation between women who received higher and lower doses of omega-3 LCPUFA daily (MD 0.24 days, 95% CI -1.16 to 1.64; 1475 participants; Analysis 7.7).

Baby/infant/child

Harris 2015 observed no difference in women who received 600 mg DHA/day versus 300 mg DHA/day for:

- birthweight (-110.35 g, 95% CI -242.80 to 22.10; Analysis 7.8);
- length at birth (MD 0.05 cm, 95% CI -0.80 to 0.90; Analysis 7.9); or
- head circumference at birth (-0.24 cm, 95% CI 0.87 to 0.39) (Analysis 7.10).

Omega-3 type direct comparisons (Analysis 8)

Three trials reported direct comparisons of different types of omega-3 supplements (Knudsen 2006; Mozurkewich 2013; Van Goor 2009).

Primary outcomes

None of the three trials reported any of the review's primary outcomes.

Mother: secondary outcomes

Mozurkewich 2013 observed no evidence of a difference between women who received DHA compared with EPA for:

- caesarean section (RR 1.23, 95% CI 0.61 to 2.51; 77 participants; Analysis 8.2);
- cessation due to adverse events (RR 0.82, 95% CI 0.24 to 2.83; 77 participants; Analysis 8.3).
- blood loss at birth (MD 1.00 mL, 95% CI -181.94 to 183.94; 77 participants; Analysis 8.5);
- major depressive disorder at six to eight weeks postpartum (RR 0.68, 95% CI 0.12 to 3.87; Analysis 8.6); or in
- depressive symptoms postpartum as measured by the BDI at six to eight weeks (MD -1.40, 95% CI -3.75 to 0.95; Analysis 8.7).

In Mozurkewich 2013, there was a reduction in pre-eclampsia in women who received DHA compared with EPA (RR 0.26, 95% CI 0.06 to 1.13; 77 participants; Analysis 8.4), though this did not reach statistical significance.

Two trials reported on gestational diabetes (Mozurkewich 2013; Van Goor 2009); in Mozurkewich 2013 there was a reduction in gestational diabetes when women received DHA compared with EPA (RR 0.15, 95% CI 0.02 to 1.14; 77 participants), though this did not reach statistical significance. Van Goor 2009 observed no evidence of difference between women who received DHA versus DHA/AA (RR 0.33, 95% CI 0.01 to 7.96; 86 participants; Analysis 8.1).

Length of gestation was reported by three trials, two of which found no difference between different types of omega-3 supplements. Knudsen 2006 observed no evidence of a difference in length of gestation between women who received EPA/DHA compared with ALA (MD -0.29 days, 95% CI -2.33 to 1.75; 1250 participants). Van Goor 2009 found no evidence of a difference in this outcome between women who received DHA compared with DHA/AA (MD

0.00, 95% CI -3.31 to 3.31; 83 participants). However, Mozurkewich 2013 observed a greater length of gestation in women who received DHA compared with EPA (MD 9.10 days, 95% CI 5.24 to 12.96; 77 participants; Analysis 8.8).

Baby/infant/child

Mozurkewich 2013 observed no evidence of a difference between infants of women who received DHA and EPA in admission to neonatal care (RR 0.35, 95% CI 0.08 to 1.63; 78 participants; Analysis 8.9).

In Van Goor 2009, there was no evidence of a difference in birthweight between infants of women who received DHA compared with DHA/AA (MD -79.00 g, 95% CI -260.22 to 102.22; 83 participants). However, in Mozurkewich 2013 there was evidence of a higher birthweight in infants of mothers who received DHA compared with EPA (MD 372.00 g, 95% CI 151.90 to 592.10; 78 participants; Analysis 8.10).

Van Goor 2009 observed no evidence of a difference in infants of women who received DHA compared with infants of women who received DHA/AA for:

- weight (MD -0.20 kg, 95% CI -0.79 to 0.39; 80 participants; Analysis 8.11);
- height (MD -0.80 cm, 95% CI -2.50 to 0.90; 80 participants; Analysis 8.12);
- infant head circumference (MD 0.10, 95% CI -0.45 to 0.65; 80 participants; Analysis 8.13).
- cognition as measured by the BSID II (MD 0.90, 95% CI -4.71 to 6.51; 80 participants; Analysis 8.14); or
- motor development as measured by the BSID II (MD 3.40, 95% CI -1.07 to 7.87; 79 participants; Analysis 8.15).

One trial, Van Goor 2009, reported on neurodevelopment from a direct comparison of omega-3 (DHA versus DHA/AA). No differences between infants of mothers assigned to the two groups were observed for the neonatal neurological classification of mildly/ definitely abnormal at two weeks (RR 0.73 95% CI 0.28 to 1.87; 67 participants), or for general movement quality that was mildly/ definitely abnormal at two weeks (RR 1.08 95% CI 0.68 to 1.72; 67 participants), However, there was a higher risk of general movement quality being mildly or definitely abnormal at 12 weeks in infants of mothers in the DHA group (RR 1.81 95% CI 1.11 to 2.95; 83 participants; Analysis 8.16).

Van Goor 2009 observed no evidence of any difference between the infants of women who received DHA and infants of women who received DHA/AA in cerebral palsy (not estimable) (Analysis 8.17).

Sensitivity analyses (Analysis 9)

We included just over a third of the trials (24/70) that we considered to be at low risk of selection and performance bias in sensitivity analyses (Bisgaard 2016; Carlson 2013; D'Almedia 1992; Gustafson 2013; Haghiac 2015; Harper 2010; Harris 2015; Ismail 2016; Jamilian 2016; Keenan 2014; Khalili 2016; Krummel 2016; Makrides 2010; Min 2014; Min 2016; Mozurkewich 2013; Oken 2013; Olsen 2000; Onwude 1995; Ramakrishnan 2010; Razavi 2017; Rees 2008; Samimi 2015; Taghizadeh 2016). The 12 outcomes assessed in subgroup analyses 3 to 6 were included in these sensitivity analyses.



For preterm birth (< 37 weeks), the sensitivity analysis was similar to the overall analysis, although it lost conventional statistical significance (RR 0.92 95% CI 0.83 to 1.02; Analysis 9.1). Sensitivity analyses for early preterm birth (< 34 weeks) (RR 0.61 95% CI 0.46 to 0.82; Analysis 9.2) and prolonged pregnancy (> 42 weeks) (RR 2.32, 95% CI 1.26 to 4.28; Analysis 9.3) were very similar to the overall analyses.

The sensitivity analysis for pre-eclampsia indicated a null result (RR 1.00 95% CI 0.81 to 1.25; Analysis 9.4), in contrast to a possible benefit seen with omega-3 LCPUFA in the overall analysis.

For caesarean section, there was very little difference between the sensitivity analysis (RR 0.96 95% CI 0.89 to 1.04; Analysis 9.5) and the overall analysis. Length of gestation also showed similar results in the sensitivity analysis (1.42 more days with omega-3, 95% CI 0.73 to 2.11; Analysis 9.6, now as a fixed-effect model due to much lower statistical heterogeneity) and the overall findings.

The sensitivity analysis for perinatal death (RR 0.60, 95% CI 0.37 to 0.97; Analysis 9.7) now reached conventional statistical significance, but had a similar magnitude to the overall borderline analysis (Analysis 1.32).

In contrast, the sensitivity analysis for low birthweight lost statistical significance (RR 0.85 95% CI 0.68 to 1.06; Analysis 9.10) but was similar to the overall analysis which reached conventional statistical significance.

The sensitivity analyses for stillbirth (RR 0.72, 95% CI 0.35 to 1.52; Analysis 9.8), neonatal death (RR 0.56, 95% CI 0.25 to 1.27; Analysis 9.9), and small-for-gestational age (RR 0.94, 95% CI 0.78 to 1.12; Analysis 9.11) showed similar findings to their corresponding overall analyses (Analysis 1.33; Analysis 1.34; Analysis 1.39).

The sensitivity analysis for birthweight (MD 48.84 g; 95% CI 22.93 to 74.76; 17 trials, 7382 participants; Analysis 9.12) also showed a similar (but lower) result to the overall analysis (Analysis 1.40) (and had lower statistical heterogeneity).

DISCUSSION

Summary of main results

In this update we have included 70 trials with 19,927 women. Most of the trials evaluated omega-3 long-chain polyunsaturated fatty acids (LCPUFA) interventions compared with placebo or no omega-3. We grouped these as: omega-3 LCPUFA supplements (50 trials); omega-3 LCPUFA supplements combined with food or dietary additions (seven trials); food/dietary additions (three trials); and omega-3 fatty acid interventions combined with other agents (12 trials).

For our primary review outcomes, there was an 11% reduced risk (95% confidence interval (CI) 3% to 19%) of **preterm birth < 37 weeks** (high-quality evidence) and a 42% reduced risk (95% CI 23% to 56%) of **early preterm birth < 34 weeks** (high-quality evidence) for women receiving omega-3 LCPUFA compared with no omega-3. The number needed to treat for an additional beneficial outcome (NNTB) to prevent one preterm birth < 37 weeks is 68 (95% CI 39 to 238), and the NNTB to prevent a preterm birth < 34 weeks is 52 (95% CI 39 to 95). Conversely for **prolonged gestation** there was a probable 61% increase (95% CI 11% to 233%) with omega-3 LCPUFA (moderate-quality evidence) and the number

needed to treat for an additional harmful outcome (NNTH) for an additional pregnancy prolonged beyond 42 weeks is 102 (95% CI 47 to 568). In the sensitivity analysis for **preterm birth < 37 weeks**, conventional statistical significance was lost, although results were similar. Sensitivity analyses for **early preterm birth < 34 weeks** and **prolonged pregnancy > 42 weeks** were very close to the overall analyses.

Omega-3 LCPUFA probably reduces the risk of both perinatal death (moderate-quality evidence) and neonatal care admission (moderate-quality evidence); and reduces the risk of low birthweight babies (high-quality evidence). Little or no difference in small for gestational age (SGA) or intrauterine growth restriction (moderate-quality evidence) with a possible small increase in large for gestational age babies (moderate-quality evidence) was seen. For most maternal outcomes, we observed no differences between groups. Mean gestational length was greater in women who received omega-3 LCPUFA and pre-eclampsia may also have been reduced (low-quality evidence). For child/adult outcomes, very few differences between the antenatal omega-3 LCPUFA and no omega-3 groups were observed, indicating that there is uncertainty regarding the impact of omega-3 LCPUFA on child development and growth.

Subgroup analyses (based on type of intervention (e.g. supplementation, food or advice), dose of docosahexaenoic acid (DHA) or eicosapentaenoic acid (EPA), timing, type of omega-3 LCPUFA, and degree of risk for women) revealed few differences. In the subgroup analyses by intervention, omega-3 LCPUFA supplements and/or omega-3 rich food and dietary advice indicated a greater positive effect on birth length than for the other intervention types. In the dose subgroup analysis, there was a positive effect from the lower doses (< 1 g/day) for low birthweight. For the risk subgroup analysis, studies with low or any risk women showed a greater reduction in low birthweight compared with the studies involving women at increased or higher risk.

Direct comparisons of doses and types of omega-3 LCPUFA showed longer gestation and higher birthweight for DHA compared with EPA. There were few other differences apart from a possible reduction in pre-eclampsia and in gestational diabetes mellitus for DHA compared with EPA.

Sensitivity analysis (restricted to trials at low risk of selection and performance bias) largely supported the findings observed in the main analyses, except for pre-eclampsia (which no longer showed a reduction for omega-3 LCPUFA compared with no omega-3).

Overall completeness and applicability of evidence

Of the 70 trials included in this update, our three primary outcomes (preterm birth, early preterm birth and prolonged pregnancy) were reported by only 26, nine and six trials respectively, even though these could have been reported by most trials (e.g. as part of routine perinatal data collection). Birthweight was the most comprehensively reported outcome (41 trials). Longer-term childhood outcomes were sparsely reported and many different assessment measures were used, for example, in neurodevelopment.

Generally the larger trials reported most of the outcomes that we had prespecified as being important, and so the body of evidence in this review is reasonably complete. However, some



trials were conducted for specific reasons, such as assessing the effects of omega-3 LCPUFA supplementation on allergy outcomes, and these trials did not always report other perinatal outcomes extensively. (Allergy outcomes from these trials are included in a separate review (Gunaratne 2015).) Some trials even excluded preterm births altogether, which may have underestimated the differences seen in preterm birth in this review. For some of these trials we were able to deduce outcomes such as preterm birth from trial flow diagrams. Exclusion of preterm births may have also led to incomplete reporting of linked outcomes, such as gestational age.

In this update we broadened the scope of the review to ensure that we could track the evolution of the perceived benefits of omega-3 and present a single comprehensive review of the effects of omega-3 LCPUFA during pregnancy. For example, in the 2006 version of this review, the major benefit of omega-3 LCPUFA was thought to be in preventing pre-eclampsia and increasing the duration of gestation. In the next decade, there was an emphasis on assessing the role of omega-3 LCPUFA supplementation on child cognition and growth. More recently, there has been renewed interest in the role of omega-3 LCPUFA in preventing preterm birth.

We also included trials assessing omega-3 LCPUFA from food sources and omega-3 LCPUFA with co-interventions, although most of the 63 trials we have added in this update compare omega-3 supplementation (largely DHA and EPA) with placebo. We have presented overall analyses as well as analyses of the different comparisons. While trials were conducted in a broad range of countries, most were from high-income settings (although some of these studies recruited only disadvantaged women). Reporting of characteristics of women was both limited and variable, for example for baseline omega-3 LCPUFA concentrations, which may influence pregnancy and longer term outcomes.

Quality of the evidence

Overall study-level risk of bias was mixed, with selection and performance bias mostly at low risk, but there was high risk of attrition bias in some trials at some time points.

Most of the important perinatal outcomes assessed using GRADE had a rating of high-quality (e.g. preterm birth) or moderatequality evidence (e.g. perinatal death) (Summary of findings for the main comparison). For birth outcomes, we only downgraded for attrition bias if substantial losses happened around the time of birth, as later losses were not relevant here. For the other outcome domains GRADE ratings ranged from moderate to very low, with over half rated as low (maternal Summary of findings 2, child/adult Summary of findings 3, and health service Summary of findings 4, outcomes). Reasons for downgrading were mostly due to design limitations (largely due to high risk of attrition bias and selection bias; and unclear randomisation and blinding) and imprecision. Particularly for the longer-term child outcomes (Summary of findings 3), there were often low numbers of studies and thus imprecision. Due to the large number of studies following up participants or subsets of participants, often for quite lengthy periods, attrition bias was commonly evident for these longer-term outcomes.

Potential biases in the review process

Due to the rigorous methods we used (comprehensive searching, double screening and data extraction, and careful appraisal and

analysis), biases are likely to be low. We were able to include 12 funnel plots, most of which did not indicate evidence of publication bias.

While we applied wide inclusion criteria at the review level, some studies had quite restrictive criteria (e.g. excluding preterm births, as discussed above) and some reported small numbers of outcomes although potentially more (e.g. perinatal death) would have been readily available and could have been reported by the trial authors.

Agreements and disagreements with other studies or reviews

A large systematic review from the US Agency for Healthcare Research and Quality (AHRQ) on the effects of omega-3 fatty acids on child and maternal health concluded that, except for small beneficial effects on infant randomised controlled trials (RCTs) reporting preterm birth < 37 weeks and found no significant difference between omega-3 and no omega-3 (odds ratio (OR) 0.87 95% CI 0.66 to 1.15 for DHA and OR 0.86, 95% CI 0.65 to 1.15 for DHA/EPA, random-effects). We included 26 RCTs for this outcome and found a clear result in favour of omega-3 (risk ratio (RR) 0.89, 95% 0.81 to 0.97, fixed-effect). For comparison, our result converts to OR 0.85 95% CI 0.74 to 0.99, random-effects. The AHRQ review did not report on early preterm birth, which in our review showed a clear reduction with omega-3 LCPUFA (RR 0.58, 95% CI 0.44 to 0.77).

Imhoff-Kunsch 2012 reviewed 15 RCTs, finding like us, a clear reduction in early preterm birth, but only a suggestion of a reduction in preterm birth overall. In the Kar 2016 systematic review of nine included studies, a clear reduction was seen in both preterm and early preterm birth. Another systematic review of omega-3 LCPUFA concentrating on birth outcomes was also largely consistent with our findings (e.g. reduced risk of early preterm birth and preterm birth (though the authors used a broader definition of early preterm birth and used a fixed-effect model throughout the review) (Chen 2016). The same authors conducted a review of omega-3 LCPUFA addressing gestational diabetes, pregnancy hypertension and pre-eclampsia and found no evidence of differences (Chen 2015). Again their findings were largely consistent with ours, although we found a possible decrease in pre-eclampsia with omega-3 LCPUFA.

The Saccone 2016 systematic review included 34 studies, which comprised 17 omega-3 LCPUFA trials. They found no clear differences in preterm birth in seven trials including 3493 asymptomatic singleton pregnancies (RR 0.90, 95% CI 0.72 to 1.11) whereas we found an 11% decrease in preterm birth < 37 weeks (RR 0.89, 95% CI 0.81 to 0.97; 24 trials, 10,121 participants). (Saccone 2016 excluded trials of women who had experienced previous preterm birth. When we omitted these two trials, it made minimal difference to our results.) For perinatal death, Saccone 2016 found no differences overall between omega-3 LCPUFA and no omega-3 LCPUFA in six trials (RR 0.61, 95% CI 0.30 to 1.24) whereas we found a possible reduction (RR 0.75, 95% CI 0.54 to 1.03; 10 trials, 7416 women). Saccone 2016 also stated that when omega-3 LCPUFA supplementation started at 20 weeks' or less gestation this showed a large (73%) decrease in perinatal death with omega-3 LCPUFA compared with placebo in singleton pregnancies, but the authors did not use accepted subgroup interaction methods to test this (Higgins 2011). Our subgroup interaction test for perinatal death did not show a difference in timing of gestation - perinatal death



was reduced at any time that supplementation started (≤ 20 weeks' gestation or later) (Analysis 4.7).

Saccone and colleagues have also published three prior systematic reviews focusing on specific outcomes: Saccone 2015a: recurrent intrauterine growth (three trials); Saccone 2015b: prior preterm birth (two trials); and Saccone 2015c: preterm birth prevention (nine trials). The Saccone 2016 review includes all studies from Saccone 2015a and Saccone 2015c, but not Saccone 2015b, which included two trials in women with prior preterm birth. All relevant studies in the four Saccone reviews are included in our review (Saccone 2015a; Saccone 2015b; Saccone 2015c; Saccone 2016).

Gould and colleagues reviewed 11 RCTs involving 5272 participants and concluded that this body of evidence did not conclusively support or refute the role of omega-3 LCPUFA supplementation in pregnancy for improving cognitive or visual development (Gould 2013). Other recent reviews of child growth and development are consistent with our findings of little impact from omega-3 LCPUFA supplementation during pregnancy (Campoy 2012; Li 2017; Rangel-Huerta 2018).

AUTHORS' CONCLUSIONS

Implications for practice

Omega-3 long-chain polyunsaturated fatty acids (LCPUFA), particularly docosahexaenoic acid (DHA), supplementation during pregnancy is a simple and effective way to reduce preterm, early preterm birth and low birthweight, with low cost and little indication of harm. The effect of omega-3 LCPUFA on most child development and growth outcomes is minimal or remains uncertain. A universal strategy of supplementation may be reasonable, although ideally, with more knowledge, this would be targeted to women who would benefit the most. A further consideration is the present reliance on non-sustainable sources of fish to manufacture omega-3 LCPUFA supplements. Ideally, universal or targeted omega-3 LCPUFA supplementation would be accompanied by other ways of improving women's overall nutrition during pregnancy.

Implications for research

More studies comparing omega-3 LCPUFA and placebo are not needed at this stage. In addition to the 70 trials included in this review, there are 23 ongoing trials, including the large ORIP trial of over 5000 women which is due to report in 2019 (Makrides 2013)

(ORIP)). It is important for trials to assess longer-term outcomes for mother and child, in order to improve understanding of metabolic, growth, and neurodevelopment pathways, in particular.

Using data from completed trials and other studies, we also need to establish if, and how, outcomes vary by different types of omega-3 fatty acids, timing and doses; and by characteristics of women (such as baseline DHA status, body-mass index and previous pregnancy outcomes). Future priority research questions include establishing the minimum effective (and optimal) dose(s) of omega-3 LCPUFA, the optimal balance of DHA and eicosapentaenoic acid and effects of different forms of omega-3 LCPUFA. The ORIP trial will provide evidence on whether stopping supplementation at 34 weeks' gestation, instead of continuing supplementation until birth, helps prevent prolonged pregnancies. A planned individual participant meta-analysis will also address some of these questions. Further mechanistic studies are needed for a broader understanding of the anti-inflammatory actions of omega-3 LCPUFA and the circumstances under which these may prevent preterm birth and other adverse birth outcomes.

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CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Ali 2017

Methods	RCT: NCT02696577	
Participants	80 women randomised	
	Inclusion criteria: 20–35 years; 28-30 weeks' gestation; pregnancy complicated with asymmetrical IUGR (diagnosed by 2D trans-abdominal US when the abdominal circumference was reduced out of proportion to other fetal biometric parameters and was below the 10th percentile so there was an increased HC:AC ratio); with normal Doppler indices in uterine and umbilical arteries at time of recruitment (the normal value of S/D ratio was from 2.5 to 3.5; RI was from 0.60 to 0.75 and PI was from 0.96 to 1.270, respectively).	
	Exclusion criteria: ≤ 20 and ≥ 35 years; any hypertensive disorder; diabetes mellitus; smokers; multiple gestations, low amniotic fluid volume; premature prelabour rupture of membranes; antepartum haemorrhage and fetal congenital anomalies; women with abnormal Doppler indices, absent diastolic flow or reversed flow.	
	Setting: Assiut Woman's Health Hospital, Egypt	
Interventions	SUPPLEMENTATION + OTHER AGENT: DHA + EPA + wheat-germ oil + aspirin versus aspirin	
	Group 1: fish oil (1000 mg = DHA 9%, EPA 13%) plus 100 mg wheat-germ oil (LA 52%-59%) as a source of vitamin E, and aspirin 81 once daily: $n = 40$	
	Group 2: aspirin 81 once daily: n = 40	
	Timing of supplementation: 6 weeks (from ~28-30 weeks GA)	
	DHA + EPA dose/day: low: 90 mg DHA + 130 mg EPA	
Outcomes	Women/birth: gestational length; caesarean section; Doppler blood flow in uterine and umbilical arteries	
	Babies/infants/children: birthweight; perinatal mortality; admission to NICU	
Notes	Funding: not reported	
	Declarations of interest: none declared	

^{*} Indicates the major publication for the study



Ali 2017 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated random number table
Allocation concealment (selection bias)	Low risk	Quote: "Allocation concealment was done using serially numbered closed opaque envelopes. Each envelope was labelled with a serial number and had a card noting the intervention type. Allocation never changed after opening the closed envelopes"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Not reported
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	12/80 (15%) participants lost to follow-up (6/40 in both intervention (2 failure of treatment and 4 lost to follow-up) and control group (3 failure of treatment and 3 lost to follow-up).
Selective reporting (reporting bias)	Unclear risk	With no access to a trial protocol it was not possible to assess selective reporting confidently.
Other bias	Low risk	Baseline characteristics appeared similar, no obvious source of other bias identified.

Bergmann 2007

RCT (3 arms)
144 women randomised
Inclusion criteria: healthy pregnant Caucasian women at least 18 years of age and willing to breast feed for at least 3 months
Exclusion criteria: increased risk of preterm birth or multiple pregnancy, allergy to cow milk protein, lactose intolerance, smoking, diabetes, consumption of alcohol > 20 g/week, or participation in another study
Exclusions: infants born < 37 weeks GA, had major malformations or were hospitalised for > 1 week
Setting: Virchow-Klinikum of the Charité and other gynaecological practices in Berlin, Germany
SUPPLEMENTATION + OTHER AGENT: omega-3 + prebiotic versus vitamin/mineral + prebiotic versus vitamin/mineral
Group 1: 600 mg fish oil (with 200 mg DHA and low EPA) plus prebiotic (fructo-oligosaccharide (4.5 g)) daily; delivered in a tetrabox containing 200 mL milk-based supplement: n = 48 (40)
Group 2: control/comparison intervention: vitamin and mineral supplementation with or without additional prebiotic (fructo-oligosaccharide); delivered in a tetrabox containing 200 mL milk-based supplement: n = 96 (48 in each group) (74)



Bergmann 2007 (Continued)	Timing of supplementation: supplementation from 22 weeks GA to 37 weeks GA, resuming at 2 weeks postpartum until 3 months	
	DHA + EPA dose/day: low: 200 mg DHA; low EPA	
Outcomes	Women/birth: maternal weight gain (from 22 weeks GA to birth); GA; caesarean birth; breast milk composition; DHA RBC concentrations; preterm birth < 37 weeks;	
	Babies/infants/children: birthweight, birth length and head circumference at birth, 1, 3 and 21 months; Apgar score at 5 minutes; cord blood pH; chemokines; vaccine antibody responses	
	6-year follow-up: child weight, height, head circumference, skinfold thickness	
Notes	Funding: Nestec Ltd, Switzerland; Charité University Hospitals. Supplements were prepared and donated by Nestlé	
	Declarations of interest: none declared	
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Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "randomized by a computer program"
Allocation concealment (selection bias)	Unclear risk	Quote: "allocated to one of three groups"; no further detail reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The identity of supplements was blinded to the subjects, support staff and investigators"
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not reported but probably done
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	 27/144 (18.8%) lost to follow-up by birth higher rates of losses from birth to 6 years 6-year follow-up: 29/144 (20%) lost to follow-up: 7/48 (15%) in omega-3 group and 12/96 (12%) in control group
Selective reporting (reporting bias)	Low risk	No apparent selective outcome reporting (although reasons for exclusions differed between different follow-up periods)
Other bias	Low risk	Baseline characteristics similar between groups

Bisgaard 2016

Methods	RCT: NCT00798226
	Children of the mothers enrolled in this RCT formed the Copenhagen Prospective Studies on Asthma in Childhood (COPSAC).
Participants	736 women randomised
	Inclusion criteria: pregnant women, at least 18 years, between 22 and 26 weeks' gestation.



Bisgaard 2016 (Continued)

Exclusion criteria: women taking more than 600 IU of vitamin D per day, and women with any endocrine, heart or kidney disorder.

Setting: Copenhagen, Denmark (women recruited between November 2008 and November 2010).

Interventions

SUPPLEMENTATION: EPA + DHA versus placebo

Group 1: 2.4 g per day of omega-3 LCPUFA (55% EPA and 37% DHA) in triacylglycerol form (Incromega TG33/22, Croda Health Care). The omega-3 LCPUFA was administered in 4 identical 1 g capsules; n = 365

Group 2: placebo: olive oil, containing 72% omega-9 oleic acid and 12% omega-6 LA (Pharma-Tech A/S), administered in 4 identical 1 g capsules; n = 371

Timing of supplementation: intervention was given to women during the last 3 months (third trimester) of pregnancy and continued for 1 week after giving birth.

DHA + EPA dose/day: high: 890 mg DHA + 1320 mg EPA

Outcomes

Women/birth: preterm birth (< 37 weeks); caesarean section; PE; death/serious maternal morbidity/mortality; length of maternal hospital stay

Babies/infants/children: admission to NICU; neonatal death; growth, development (not yet reported)

Notes

Allergy outcomes from this trial will be reported in another Cochrane Review when it is updated (Gunaratne 2015).

Funding: "No funding agency played any role in the design or conduct of the trial, the collection, management, or interpretation of the data, the preparation, review, or approval of the manuscript for publication, or the decision to submit the manuscript for publication. In addition, no pharmaceutical company that produces n-3 [omega-3] LCPUFA was involved in the trial. The intervention was funded solely by COPSAC"

Declarations of interest: none reported

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The women were randomized using a computer-generated list of random numbers".
Allocation concealment (selection bias)	Low risk	Randomisation was "prepared by an external investigator with no other involvement in the trial".
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	For the outcomes reported and included in this review, the participants and personnel were blinded to intervention assignment. However, for the outcomes relating to the 5-year data collection time point, only the investigators were blind to group allocations.
		Quote: "Neither the investigators nor the participants were aware of group assignments during follow-up for the first 3 years of the children's lives, after which there was a 2-year follow-up period during which only the investigators were unaware of group assignments".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Probably done
Incomplete outcome data (attrition bias)	Low risk	Intervention: 365 women allocated to intervention, 21 withdrew during pregnancy due to intrauterine death (n = 2), disabling disease (n = 2), emigration (n



Bisgaard 2016	(Continued)
All outcomes	

= 1) and 16 were lost to follow-up; therefore, data from 344/365 intervention group women (94.2%) were available for inclusion in the analyses for maternal outcomes reported. There were 3 pairs of twins born to the intervention group women; therefore, data from 347 intervention group infants were available for inclusion in the analysis for infant outcomes.

Control: 371 women allocated to control, 22 withdrew during pregnancy due to intrauterine death (n = 2), disabling disease (n = 2), emigration (n = 2) and 16 were lost to follow-up; therefore, data from 349/371 control group women (94%) were available for inclusion in the analyses for maternal outcomes reported. There were 2 pairs of twins born to the control group women; therefore, data from 351 control group infants were available for inclusion in the analyses for infant outcomes.

Therefore, there were limited missing outcome data, and the missing data were balanced in numbers across intervention groups, with similar reasons for missing data across groups.

Selective reporting (re-
porting bias)	

Low risk

No evidence of selective reporting; data for prespecified outcomes (according to published protocol, made available as supplementary material with the online paper), have been reported. Further, the protocol provides a detailed description of the planned analysis which is reflected in the reporting of results.

Other bias Low risk

Baseline characteristics were similar between groups: Quotes: "The baseline characteristics of the pregnant women and their children showed that randomisation was not biased".

No indication of difference in intervention fidelity related to different levels of adherence between the 2 groups.

Boris 2004

Methods	Further randomisation (4 arms) of Olsen 1992 (supplementation until birth versus supplementation un-
	til 30 days after giving birth)

Participants

44 women randomised

Inclusion criteria: healthy pregnant women

Exclusion criteria: not reported

Setting: Aarhus, Denmark

Interventions

SUPPLEMENTATION: omega-3 (until birth) versus omega-3 (continuing for 30 days after giving birth) versus olive oil versus no supplementation

Group 1: omega-3 (1.3 g EPA and 0.9 g DHA per day) as 4×1 g gelatine capsules with Pikasol (Lube A/S, Hadsund, Denmark) fish oil (32% EPA (20:5n-3), 23% DHA, and 2 mg tocopherol/mL), stopping at birth: n = 11

Group 2: omega-3 (1.3 g EPA and 0.9 g DHA per day) as 4×1 g gelatine capsules with Pikasol (Lube A/S, Hadsund, Denmark) fish oil (32% EPA (20:5n-3), 23% DHA, and 2 mg tocopherol/mL), stopping 30 days after giving birth: n = 12

Group 3: 4 x 1 g capsules of olive oil per day (72% oleic acid (18:1n-9) and 12% LA (18:2n-6)), stopping at birth: n = 8

Group 4: no supplement, stopping 30 days after giving birth: n = 5

Timing of supplementation: from 30 weeks GA until birth or 30 days after giving birth



Boris 2004 (Continued)		
, , , , , , , , , , , , , , , , , , , ,	DHA + EPA dose/day: high: 900 mg DHA + 1300 mg EPA	
Outcomes	Omega-3 and lipid concentrations in breast milk	
Notes	Funding: The University of Aarhus (Aarhus, Denmark); Lube A/S ((Hadsund, Denmark), supplied Pikasol fish oil and olive oil capsules.	
	Declarations of interest: not reported	
	No outcomes able to be used in this review.	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomly allocated"
Allocation concealment (selection bias)	Unclear risk	Quote: "randomly allocated"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Partial (one group not supplemented; timing for omega-3 groups not able to be blinded)
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	3/26 in the intervention groups and 6/18 in the control groups were lost to follow-up (no reasons reported)
Selective reporting (reporting bias)	Unclear risk	Insufficient detail reported to determine confidently
Other bias	Low risk	Groups were similar for baseline characteristics with regard to maternal age at birth, and prepregnancy weight

Bosaeus 2015

	group) Setting: Sahlgrenska University Hospital, Gothenburg, Sweden			
	Exclusion after study entry: women having a miscarriage, abortion, intrauterine fetal death, sudden infant death, twin pregnancy or giving birth before 34 weeks' gestation (n = 1 but not reported which			
	Exclusion criteria: non-European descent, self-reported diabetes, use of neuroleptic drugs, and vegetarianism or veganism.			
	Inclusion criteria: pregnant women of normal weight (BMI 18.5 to 24.9), aged 20-45 years			
Participants	101 women randomised			
Methods	RCT: PONCH (Pregnancy Obesity Nutrition and Child Health Study)			



Bosaeus 2015 (Continued)

Group 1: dietary counselling (from registered dieticians): 3 sessions, 5 phone calls during pregnancy (from 8-12 weeks GA). Participants were advised to eat 3 meals of fish a week, with advice on types of fish to consume to avoid pollutants, to generally lower sugar intake to reach < 10% energy; to eat 500 g of vegetables and fruits a day; to increase daily energy intake by 350 kcal in the second trimester and by 500 kcal in the third trimester: n = 49

Group 2: control group: study visit each trimester (not further described): n = 52

Timing of counselling: from 8-12 weeks GA

DHA + EPA dose/day: other: unable to determine

Outcomes

Women/birth: fish intake; body composition; GWG; serum phospholipid fatty acids (in all 3 trimesters); fat mass (air-displacement plethysmography); size, number and lipolytic activity of adipocytes; and adipokine release and density of immune cells and blood vessels in adipose tissue

Babies/infants/children: birthweight (numerical results not reported)

Notes

Funding: "Supported by grants from Novo Nordisk Foundation, the Swedish Research Council (No. 12206), the Swedish Research Council (Project No. 2013-28632-103061-41), the Swedish Diabetes Association Research Foundation, the Swedish Federal Government under the LUA/ALF agreement, IngaBritt and Arne Lundbergs Foundation, Freemasonry Barnhus Board in Gothenburg, Olle Engkvist Building contractor Foundation (210/56) and Queen Silvia's Jubilee Fund".

Declarations of interest: none declared

Women in the intervention group did not use supplements containing fish oil or omega-3 fatty acids during pregnancy but in the control group, 1 woman in the first trimester, 2 in the second trimester and 4 women in the third trimester used these supplements.

No outcomes could be used in this review.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Randomization was done by a computerized programmatched for age, BMI and parity"
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not feasible to blind
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	66/101 (65%) attrition (complete measurements from all trimesters): 31/49 (63%) in the intervention group and 35/52 (67%) in the control group lost to follow-up
Selective reporting (reporting bias)	Unclear risk	'Exclusions' not always reported by intervention or control group; preterm < 37 weeks not reported; birthweight not fully reported



Bosaeus 2015 (Continued)

Other bias Unclear risk Baseline characteristics comparable except for women in the intervention re-

porting lower fish consumption and being shorter than women in the control

Identical placebos (not reported whether women could guess their treatment)

group

Bulstra-Ramakers 1994

Methods	RCT			
Participants	68 women randomised			
		men 12-14 weeks GA with a history of IUGR (birthweight < 10th centile), \pm PIH* in y; or chronic renal disease or placental abnormalities of an impaired uteropla-		
	Exclusion criteria: women with diabetes, systemic lupus erythematosus or other connective tissue disease, or women who had already agreed to be treated with low dose aspirin because of their obstetric history			
	Setting: University Ho	spital and regional hospitals in the north of the Netherlands		
	*PIH defined as an increase in diastolic pressure of at least 25 mmHg during the course of pregnancy with a final diastolic pressure > 90 mmHg.			
Interventions	SUPPLEMENTATION:	EPA + DHA versus placebo		
	Group 1: EPA (n = 34 randomised; 32 analysed): 3 g/day, given as 12 capsules/day (each capsule contained 250 mg EPA); no information about the DHA content of the capsules Group 2: 12 capsules coconut oil/day (n = 34; 31 analysed)			
	Timing of supplementation: from 12-14 weeks GA "onwards" - until birth			
	DHA + EPA content/day: high: DHA not stated + 3 g EPA			
Outcomes	Women/birth: PIH; preterm birth < 37 weeks; preterm birth < 34 weeks; antenatal hospitalisation; miscarriage; mode of birth; high uric acid; low platelets; 2nd trimester Hb decrease < 5%; duration of pregnancy; adverse effects			
	Babies/infants/children: SGA (birthweight < 10th percentile); LGA (> 10th percentile); stillbirth; neonatal death; perinatal death			
Notes	Funding: not reported			
	Declarations of interest: not reported			
	Sample size estimate v	vas based on the first randomised study of aspirin in high-risk pregnancies.		
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Unclear risk	Not reported		
Allocation concealment (selection bias)	Low risk	Quote: "Randomisation was performed by the hospital pharmacy"; placebo capsules were identical to treatment capsules		

Blinding of participants

and personnel (perfor-

mance bias)

Low risk



Bulstra-Ramakers 1994 (Continued)

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Blinding of outcome assessment (detection bias) All outcomes	Low risk	Outcome assessments were blinded
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	5/68 (7.3%) post-randomisation exclusions: 2/34 in the EPA group due to non-adherence and 3/34 in the placebo group (1 miscarriage and 2 due to non-adherence). Non-adherence was due to the perceived effects of nausea and vomiting.
Selective reporting (reporting bias)	Unclear risk	Most expected outcomes were reported, but GA was not reported as mean and SD.
Other bias	Unclear risk	Some baseline imbalance between groups: previous PIH in 24/32 women in the EPA group and 15/31 in the control group.

Carlson 2013

Methods	RCT: NCT00266825 (KUDOS)				
Participants	350 women randomised				
	Inclusion criteria: women who were English speaking, between 8 and 20 weeks of gestation, between 16 and 35.99 years of age, and planning to give birth at a hospital in the Kansas City metropolitan area				
	Exclusion criteria: carrying more than 1 fetus, had pre-existing diabetes mellitus or SBP ≥ 140 mmHg at enrolment, or had any serious health condition likely to affect the prenatal or postnatal growth and development of their offspring, including cancer, lupus, hepatitis, HIV/AIDS, or a diagnosed alcohol or chemical dependency, BMI ≥ 40 (self-reported); taking DHA supplement 300 mg or more/day				
	Characteristics: baseline DHA status mean 4.3 [1] g/100 g total fatty acids; 42% women enrolled in KUDOS were African-American which is higher than the national average of 16%				
	Setting: Kansas City metropolitan area, KS, USA. Study conducted from January 2006 and October 2011.				
Interventions	SUPPLEMENTATION: omega-3 (DHA) versus placebo				
	Group 1: 600 mg DHA/day: 3 capsules/day of a marine algae-oil source of DHA (200 mg DHA/capsule) DHASCO;				
	n = 178				
	Group 2: placebo: 3 capsules containing half soybean and half corn oil. The placebo capsules did not contain DHA but did contain a-linolenic acid; n = 172.				
	Timing of supplementation: < 20 weeks (~14 weeks GA) until birth				
	DHA + EPA dose/day: mid: 600 mg DHA + EPA negligible				
Outcomes	Women/birth: adherence; DHA concentrations (maternal and cord blood); DHA concentrations (by FADS genotypes in a subset of 250 women); length of gestation; miscarriage; severe PE; gestational diabetes; caesarean section; maternal adverse effects; PPH; placental abruption; preterm birth < 37 weeks; early preterm birth < 34 weeks; low birthweight; very low birthweight; antenatal hospital admission; PPROM; GWG, costs.				



Carlson 2013 (Continued)

Babies/infants/children: birthweight; birth length; head circumference at birth; ponderal index; NICU admissions, length of hospital stay; mortality; congenital anomalies; visual habituation at 4, 6 and 9 months; and at 5 years, fat mass; fat-free mass; body fat; weight; height, BMI Z-score

Notes

Adherence: "Capsule compliance was similar for the 2 groups: placebo (76% consumed) and DHA (78% consumed)".

Funding: NIH (R01 HD047315) and the Office of Dietary Supplements; Kansas Intellectual and Developmental Disabilities Research Center (P30 HD02528). DSM Nutritional Products (formerly Martek Biosciences) donated the placebo and DHA capsules.

Declarations of interest: "SEC has given talks for several companies, including Martek, Mead Johnson Nutrition, and Nestle on results from our studies and the results of others who study the effects of DHA on infant and child outcomes. She is the President of the International Society for the Study of Fatty Acids and Lipids, which has corporate members who produce sources of DHA. JC consults with several companies on developmental measures to assess cognitive development of infants and children. None of the other authors declared a potential conflict of interest."

Bias	Authors' judgement	Support for judgement Quote: "The study biostatistician generated randomization schedules for 2 maternal age groups (16–25.99 and 26–35.99 y), and each sequence of 8 random numbers included 4 assignments per group to stratify by age and treatment"		
Random sequence generation (selection bias)	Low risk			
Allocation concealment (selection bias)	Low risk	Quote: "The Investigational Pharmacy personnel assigned women to placebo or DHA based on the age shared by the study personnel"		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Participants and data collectors were blinded to allocation, as were a investigators until children were 18 mo of age and had completed early cognitive and visual acuity development testing"		
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Participants and data collectors were blinded to allocation, as were all investigators until children were 18 mo of age and had completed early cognitive and visual acuity development testing"		
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	24/178 (13.5%) lost from DHA group: 9 changed hospitals/clinics 4 moved from city 4 miscarried 1 voluntary abortion 1 incarcerated 2 no longer interested 2 illness 1 wanted to take DHA 25/172 (14.5%) lost from placebo group: 7 changed hospitals/clinics 4 moved from city		
		 4 moved from city 3 miscarried 5 no longer interested 2 illness 1 wanted to take DHA 		



Carlson 2013 (Continued)		3 primary caregivers said no
		At 5-year follow-up, data were available for 88 children in the omega-3 group and 83 children in the placebo group, equating to 179/350 (51%) lost to follow-up.
Selective reporting (reporting bias)	Low risk	Most expected outcomes were reported.
Other bias	Low risk	No apparent source of other bias.

Chase 2015

Methods	RCT (pilot): NCT00333554				
Participants	41 infants (randomised during pregnancy). A further 21 mothers (~33%) received either DHA or placebo during their last trimester, but discontinued post birth.				
	Inclusion criteria: women 18 years of age or older, from 24 weeks GA whose babies may be at higher risk for T1D based on family history (had T1D, or child's father or a full or half sibling of the child had T1D)				
	Exclusion criteria: any condition investigators believed would put the mother or her fetus at an unacceptable medical risk; known complication of pregnancy causing an increased risk for the mother of fetus prior to entry into the study; have previously had 2 or more preterm births (< 36 weeks); were diabetic and had a known HbA1c > 9% at any time during the pregnancy, plan to take DHA during the pregnancy				
	Setting: 9 clinical sites across USA				
Interventions	SUPPLEMENTATION: DHA versus placebo				
	Group 1: algal DHA daily while pregnant and lactating (if choosing to breastfeed); 800 mg DHA per day (4 capsules); infant received ~ 150 mg/day from mother or from formula; then 400 mg/day as toddlers (1-2 years of age): total number randomised: n = unclear (21 reported)				
	Group 2: corn/soy oil (placebo): total number randomised: n = unclear (16 reported)				
	Timing of supplementation: supplementation began immediately after randomisation (start of third trimester of pregnancy) and continued at least until the HLA type of the infant was known; if an infant entered the study antenatally, duration of supplementation would be a minimum of 36 months				
	DHA + EPA dose/day: mid: 800 mg DHA + EPA negligible				
Outcomes	Women/birth: breastmilk DHA				
	Babies/infants/children: RBC DHA, IL 1-betaC, CRP and other inflammatory mediators; infant vitamin D				
Notes	Funding: NIDDK branch of the NIH, the ADA, and the Juvenile Diabetes Research Foundation (JDRF)				
	Supplements from DHASCO-S oil, Martek Biosciences Corporation, Columbia, MD				
	Declarations of interest: not reported				
	No outcomes could be used in this review				



Chase 2015 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
Allocation concealment (selection bias)	Unclear risk	Quote: "randomly assigned"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Quote: "double blinded"
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Not fully reported
Selective reporting (reporting bias)	Unclear risk	Limited number of outcomes reported
Other bias	Unclear risk	Insufficient information to determine

D'Almedia 1992

Methods	RCT (3 arms)				
Participants	100 women (from 2 of the 3 arms)				
	Inclusion criteria: primiparous and multiparous women, aged 14-40 years and ≤ 16 weeks' gestation.				
	Exclusion criteria: none reported				
	Characteristics: 76% had a recent history of malaria or fever of unknown origin, 34% had a history of sickle cell trait or disease, 37% had a history of anaemia, 21% had a history of pregnancy hypertension or other hypertension and 4% had a previous preterm birth.				
	Setting: Luanda, Angola				
Interventions	SUPPLEMENTATION + OTHER AGENT: GLA + EPA + DHA (omega 6/omega 3) versus placebo				
	Group 1: 8 capsules/day evening primrose oil + fish oil, providing a total of 296 mg GLA, 144 mg EPA and 80 mg DHA/day: total number randomised = 50				
	Group 2: 8 capsules olive oil/day (without vitamin E): total number randomised = 50				
	(The third arm (magnesium oxide: $n = 50$) was not considered for this review):				
	Timing of supplementation: 6 months				
	DHA + EPA dose/day: low: 80 mg DHA + 144 mg EPA				
Outcomes	Women: PIH, PE (hypertension (rise in SBP > 30 mmHg and/or a rise in DBP > 15 mmHg); oedema (visible fluid accumulation in the ankles and feet), and proteinuria (protein > 1 determined by test tape) any time during the pregnancy), eclampsia.				



D	Αl	med	lia	1992	(Continued)
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Babies/infants/children: birthweight (< 2000 g and > 3000 g (not used for LGA outcome)).

Notes

Funding: GLA, EPA, DHA tablets and placebo tablets were prepared by Efamol Research Institute and

Efamol Ltd

Declarations of interest: not reported

Reported dietary intake of women in all groups at study entry was poor.

Risk of bias

Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Women "randomly assigned using a random number table"	
Allocation concealment (selection bias)	Low risk	Quote: "the code of the capsules was not made known by the manufacturer, until the end of the treatment period"	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Olive oil and evening primrose oil + fish oil capsules identical (but both different to magnesium oxide), so fully blinded with regard to the fish oil/evening primrose and placebo comparison.	
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Outcome assessments partially blinded - olive oil and evening primrose oil + fish oil capsules identical (but both different to magnesium oxide), so fully blinded with regard to the fish oil/evening primrose and placebo comparison.	
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Not specifically reported	
Selective reporting (reporting bias)	Unclear risk	Outcomes such as preterm birth and perinatal mortality were not reported	
Other bias	Unclear risk	Baseline nutritional profiles (determined by dietary recall) differed (placebo group higher caloric intake; higher animal protein; higher total fat; higher "fish fat"; higher cholesterol; higher fibre; higher potassium)	

de Groot 2004

Methods	RCT: parallel	
Participants	79 women randomised	
	Inclusion criteria : white origin, GA < 14 weeks, normal health (not suffering from any hypertensive, metabolic, cardiovascular, renal, psychiatric, or neurologic disorder), fish consumption < 2 times per week	
	Exclusion criteria : DBP > 90 mmHg, multiple pregnancy, use of medications, use of (LC)PUFA rich supplements, origin other than Caucasian	
	Setting: region around Maastricht, Heerlen and Sittard in the Netherlands	
Interventions	SUPPLEMENTATION/ENRICHMENT: ALA + LA versus LA (in margarine)	



de Groot 2004 (Continued)

Group 1: ALA: daily ≥ 25 g ALA-enriched high-LA margarine from week 14 of pregnancy until birth (with the requested intake of 25 g margarine/day women consumed 2.82 g ALA + 9.02 g LA per day) - 40 women randomised; 29 analysed

Group 2: No ALA: daily ≥ 25 g of high-LA margarine without ALA from week 14 of pregnancy until birth (with the requested intake of 25 g of margarine/day women consumed 10.94 g LA and 0.03 g ALA per day) - 39 women randomised; 29 analysed

All women: every 3 weeks the volunteers received 3 tubs each containing 250 g margarine. Women were instructed to consume the margarine primarily on bread (if consumption was lower than required, they were advised to put it on top of potatoes or pasta; they were not allowed to use it for baking because of possible adverse effects on the polyunsaturated fatty acid content of the margarine). They were allowed to maintain their usual diets during the course of the study, with the exception of the use of butter/their usual margarine.

Timing of supplementation: from 14 weeks GA to birth

DHA + EPA dose/day: other (2.82 g ALA)

Outcomes

Women/birth: maternal cognitive functioning; caesarean section; gestational diabetes; depression (postnatal); antenatal admission to hospital (long-term hospitalisation); gestational length; fatty acid concentrations; breastfeeding

Babies/infants/children: preterm birth < 36 weeks; stillbirth; birthweight; Apgar score

Notes

Funding: grant from Unilever Research and Development (Vlaardingen, Netherlands), which also donated the margarines used in the study.

Declarations of interest: none declared by authors of main reference, not reported in other references

Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Quote: "randomly allocated"; no further details reported	
Allocation concealment (selection bias)	Unclear risk	Quote: "randomly allocated"; no further details reported	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Described as "double blind"; no further details reported	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported	
Incomplete outcome data (attrition bias) All outcomes	High risk	21/79 (27%) women were lost to follow-up: 11 from the ALA group: 2 preterm birth < 36 weeks 3 not motivated 2 non-adherence 1 severe morning sickness 1 long-term hospitalisation 1 abroad 1 insufficient blood samples	



de Groot 2004 (Continued)		10 in the no ALA group:
 1 preterm birth < 36 weeks 1 not motivated 1 non-adherence 2 disliked intervention 1 severe morning sickness 1 stillbirth 1 gestational diabetes 1 long-term hospitalisation 1 insufficient blood samples 		 1 preterm birth < 36 weeks 1 not motivated 1 non-adherence 2 disliked intervention 1 severe morning sickness 1 stillbirth 1 gestational diabetes 1 long-term hospitalisation
		After giving birth, a further 2 women were lost to follow-up, both in the ALA group (1 moved away; 1 postnatal depression).
Selective reporting (reporting bias)	Unclear risk	Not all expected outcomes were reported; some outcomes treated as exclusions and therefore may be incompletely reported
Other bias	Unclear risk	More breastfeeding mothers in the ALA group

Dilli 2018

Methods	RCT: NCT02371343 (MaFOS-GDM)			
Participants	140 women randomised			
	Inclusion criteria: pregnant women, 18-40 years old, between 24-28 weeks GA, residents of one of the study centres, planning to remain in the area for the next year, subsequently diagnosed with gestational diabetes mellitus			
	Setting: three tertiary maternity and children's hospitals from different regions in Turkey (trial conducted from January 2015 to January 2017)			
Interventions	SUPPLEMENTATION: EPA + DHA versus placebo			
	Group 1: omega-3 LCPUFA 1200 mg/day: 384 mg EPA; 252 mg DHA (Ocean Plus): n = 70			
	Group 2: placebo (sunflower oil - similar in appearance and taste to the fish oil capsules): n = 70			
	Timing of supplementation: from 26-27 weeks till birth (~ 9 weeks)			
	DHA + EPA dose/day: mid: 252 mg DHA + 384 mg EPA			
Outcomes	Women/birth: GWG; caesarean; preterm birth < 37 weeks			
	Babies/infants/children: cord Insulin-like growth factor 1 (IGF)-1 DNA methylation; birth weight; macrosomia (> 90th percentile for GA); head circumference at birth; hospitalisation (not further specified)			
Notes	Funding: Republic of Turkey Ministry of Health Central Directorate for Health Research			
	Declaration of interest: authors declared no conflict of interest			
Risk of bias				
Bias	Authors' judgement Support for judgement			



Dilli 2018 (Continued)		
Random sequence generation (selection bias)	Unclear risk	Quote: "balanced blocks"
Allocation concealment (selection bias)	Unclear risk	Quote: "sealed envelopes"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: placebo-controlled
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data	High risk	Omega-3 LCPUFA: 18/70 lost to follow-up or refused to continue
(attrition bias) All outcomes		Placebo: 2/70 refused to continue
		Judged to be at high risk due to differential rates of losses between omega-3 and placebo groups (also see other bias text).
Selective reporting (reporting bias)	Unclear risk	Limited number of pregnancy outcomes reported
Other bias	Unclear risk	No apparent evidence of other bias, though GDM was more often managed by diet only in the omega-3 group than in the -placebo group

Dunstan 2008

Methods	RCT: ACTRN12611000041954		
Participants	98 women randomised		
	Inclusion criteria: all women had a history of physician-diagnosed allergic rhinitis and/or asthma and 1 or more positive skin prick test to common allergens, but who were otherwise healthy, with healthy full-term infants; and recruited < 20 weeks GA.		
	Exclusion criteria: normal consumption of fish meals exceeding 2 per week, women who smoked, had other medical problems, complicated pregnancies, seafood allergy		
	Setting: antenatal clinic, St John of God Hospital, Perth, Western Australia		
Interventions	SUPPLEMENTATION: DHA + EPA versus olive oil		
	Group 1: omega-3 LCPUFA: 3300 mg/day (DHA 2200 mg/day): $4 \times 1 g$) omega-3 LCPUFA capsules comprising 2.07 g DHA and 1.03 g EPA per day (total number randomised = 52)		
	Group 2: control: $4 (x 1 g)$ capsules of olive oil per day containing 66.6% omega-9 oleic acid and $< 1\%$ omega-3 LCPUFAs (total number randomised = 46)		
	Timing of supplementation: 20 weeks GA to birth		
	DHA + EPA dose/day: high: 2.07 g DHA + 1.03 g EPA		
Outcomes	Women/birth: food frequency questionnaire (20 and 30 weeks GA); adherence; allergen-specific T-cell responses in cord blood, neonatal cord blood CD4+ T-cell DNA methylation; fatty acid composition (including in breast milk), length of gestation, elective caesarean, spontaneous labour, induction; ma-		



Dunstan 2008 (Continued)

ternal BP; pregnancy weight gain (subsample in Keelan 2015); Beck Depression Inventory (depressed mood = score ≥ 10) (not reported by randomised groups)

Babies/infants/children:

- birthweight, birth length, head circumference, 5 minute Apgar score; neonatal T-cell protein kinase
 C; T-cell cytokines; medically-diagnosed allergies including incidence of asthma, atopic eczema, and food allergy at 1 year of age
- 2.5 years: infant growth and development quotients (Griffiths Mental Development Scales); receptive language (Peabody Picture Vocabulary Test) IIIA; Child Behaviour Checklist (1.5-5 years);
- 12 years: full-scale IQ (Weschler Intelligence Scale for Children-IV); Child Behaviour Checklist (both parent and child forms); Beery-Buktenica Developmental Test of Visual-Motor Integration; Children's Communication Checklist; telomere length; oxidative stress; specialised pro-resolving mediators.

Notes

Funding: National Health and Medical Research Council of Australia (APP139025 and APP1010495); National Heart Foundation of Australia (G 09P 4280); Raine Medical Research Foundation; Ada Bartholomew Trust; and McMaster University. Dr Janet Dunstan was supported by the Child Health Research Foundation of Western Australia Women and Infants Research Foundation.

Declarations of interest: none declared

Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Not reported	
Allocation concealment (selection bias)	Low risk	Quote: "Randomization and allocation of capsules occurred at a different centre separate from the recruitment of participants. Capsules were administered to the participants by someone separate from those doing the allocation"; and, "staff dispensing the capsules were blinded to the allocation".	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Participants, research scientists and paediatrician remained blinded to group allocations for the duration of the study.	
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not reported but probably done	
Incomplete outcome data	Unclear risk	Birth: 15/98 (15%) excluded or lost to follow-up:	
(attrition bias) All outcomes		From the omega-3 group 12/52 (23%):	
		• 7 withdrew	
		3 births < 36 weeks GA2 sick infants	
		From the placebo group 3/46 (7%):	
		2 withdrew1 birth < 36 weeks GA	
		Child follow-up at 2.5 years: 26/98 (27%) lost to follow-up:	
		From the omega-3 group 12 (plus a further 7 = 19/52 (37%)):	
		• 4 moved	
		• 3 withdrew	



Dunstan 2008 (Continued)		From the placebo group 3 (plus a further 4 = 7/52 (13%)):
		1 moved3 withdrew
Selective reporting (reporting bias)	Unclear risk	Some outcomes (e.g. preterm births) treated as exclusions.
Other bias	Unclear risk	Possible baseline imbalance with 52 and 46 randomised.

England 1989

Methods	RCT: parallel		
Participants	40 women randomised		
	Inclusion criteria: women with severe gestational proteinuric hypertension (BP > $140/90$; proteinuria > $0.3 \text{ g}/24 \text{ hour}$)		
	Exclusion criteria: not reported		
	Setting: University of Witwatersrand and the South African Institute for Medical Research, Johannesburg, South Africa		
Interventions	SUPPLEMENTATION: EPA versus placebo		
	Group 1: EPA 3 g/day: total number randomised = 20*		
	Group 2: placebo (not further described): total number randomised = 20*		
	Timing of supplementation: not reported		
	DHA + EPA dose/day: high: 3 g EPA; DHA not stated		
	*assumed, not specifically stated.		
Outcomes	Women/birth: requirement for pregnancy to be terminated; mean time to termination of pregnancy; amount of proteinuria; platelet and serum membrane EPA in first 2 weeks of treatment; amount hypertensive therapy required;		
	Babies/infants/children: birthweight		
Notes	Funding: not reported		
	Declarations of interest: not reported		
Risk of bias			

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
Allocation concealment (selection bias)	Unclear risk	Quote: "prospective randomized double blind trial"
Blinding of participants and personnel (perfor- mance bias)	Unclear risk	Quote: "prospective randomized double blind trial"; no details about the placebo were reported.



Eng	land	1989	(Continued)

All outcomes

Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	3/20 women in the EPA arm and 2/20 in the placebo arm required termination of pregnancy due to fulminating hypertension in the first week and were subsequently excluded from analysis.
Selective reporting (reporting bias)	Unclear risk	Only a few outcomes reported fully.
Other bias	Low risk	The 2 groups were similar in terms of maternal age, GA, level of hypertension and amount of proteinuria at baseline.

Freeman 2008

Methods	RCT: parallel				
Participants	59 women randomised (25 were pregnant and thus only these women were eligible for this review)				
	Inclusion criteria: perinatal women (pregnant (n = 25) and postpartum (n = 34) with major depressive disorder, 12-32 weeks GA or postpartum (within 6 months of childbirth); 18-45 years of age; scored ≥ 9 on EPDS, outpatient status				
	Exclusion criteria: previous intolerance to omega-3 fatty acids, current use of antidepressants or anticoagulants, psychosis, diagnosis of bipolar disorder, active substance use, or active suicidal ideation.				
	Setting: University of Arizona, USA				
Interventions	SUPPLEMENTATION: EPA + DHA versus placebo				
	Group 1: EPA + DHA: $1.9~g$ /day ($1.1~g$ EPA and $0.8~g$ DHA, total 4 capsules/day) for 8 weeks: total 12 pregnant women randomised				
	Group 2: placebo: corn oil with a small amount of fish oil for 8 weeks: total 13 pregnant women randomised (9 completed study)				
	Timing of supplementation: pregnant women: from 12-32 weeks GA				
	All women: were provided with manualised supportive psychotherapy				
	DHA + EPA dose/day: high: 0.8 g DHA + 1.1 g EPA				
Outcomes	Women/birth: Hamilton Rating Scale Depression (HAM-D) biweekly; Edinburgh Postnatal Depression Scale (EPDS) biweekly; Clinical Global Impression; tolerability of omega-3				
Notes	Funding: NIMH K23MH066265; Pronova/EPAX provided study drug and placebo at no cost.				
	Declarations of interest:				
	Dr Marlene Freeman: Research support: NIMH, U.S. FDA, Institute for Mental Health Research (Arizona), Forest, Reliant, Lilly; honorarium from AstraZeneca;				
	Dr Katherine Wisner: Research support: NIMH, Stanley Medical Research Foundation, New York-Mid Atlantic Consortium for Genetics and Newborn Screening Services (NYMAC), State of Pennsylvania, American Society for Bariatric Surgery, Pfizer, Wyeth (pending)				



Freeman 2008 (Continued)

Dr Alan Gelenberg: Consultantships: Eli Lilly, Pfizer, Best Practice, Astra/Zeneca, Wyeth, Cyberonics, Novartis, Forest, GlaxoSmithKline; Stock Options: Vela Pharmaceuticals; Speakers Bureau: Pfizer Pharmaceuticals, GlaxoSmithKline

Dr Joseph Hibbeln, Dr. Priti Sinha, Dr Melinda Davis: nothing to disclose

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Described as randomised; no further details reported
Allocation concealment (selection bias)	Unclear risk	Described as randomised; no further details reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Placebo-controlled
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not reported, but probably done.
Incomplete outcome data (attrition bias) All outcomes	High risk	7/59 women dropped out after the baseline visit; a further woman was diagnosed with hyperthyroidism after randomisation and was then excluded as being ineligible (leaving 51 women who completed at least 2 assessments) – 21 of these women were pregnant meaning 4/25 (16%) pregnant women were lost to follow-up (all 4 were from the placebo group).
Selective reporting (reporting bias)	High risk	Few outcomes were reported
Other bias	Unclear risk	Some baseline differences – omega-3 group more likely to be Caucasian; and to enrol earlier in their pregnancy and more likely to take antidepressants

Furuhjelm 2009

urunjeun 2009	
Methods	RCT: NCT00892684
Participants	145 women randomised
	Inclusion criteria: women affected by allergies themselves, or having a husband or an older child with current or previous allergic symptoms, i.e. bronchial asthma diagnosed by a doctor, atopic eczema, allergic food reactions, itching and running eyes and nose on exposure to pollen, pets or other known allergens.
	Exclusion criteria: mothers with an allergy to soy or fish or undergoing treatment with anticoagulants or commercial omega-3 fatty acid supplements
	Characteristics: 73% of women in the omega-3 group and 63% in the placebo group had allergic symptoms; average registered dietary intake of DHA and EPA at inclusion was 0.2 g/day and 0.1 g/day, respectively, thus the daily dose of omega-3 LCPUFA was increased 8–10 times by the supplementation (this corresponds to a meal of approximately 100 g salmon daily).
	Setting: Linköping and Jönköping, Sweden



Furuhjelm 2009 (Continued)

		ve			

SUPPLEMENTATION: EPA + DHA versus placebo (soy oil)

Group 1: EPA/DHA: 9 x 500 mg capsules a day containing 35% EPA, and 25% DHA, to provide 1.6 g of EPA and 1.1 g of DHA; plus 28 mg alphatocopherol: total number randomised = 70

Group 2: placebo: 9 soy oil capsules a day, containing 58% LA to provide 2.5 g LA/day and 6% ALA to provide 0.28 g ALA/day, plus 36 mg alphatocopherol: total number randomised = 75

Timing of supplementation: 25 weeks GA to birth, and encouraged to continue during lactation (average 3-4 months).

DHA + EPA dose/day: high: 1.1 g DHA + 1.6 g EPA

Outcomes

Women/birth: GA at birth; maternal BMI at end of gestation; breastfeeding duration; diet at 6 months postpartum

Babies/infants/children: birthweight; Apgar scores at 10 minutes; medically diagnosed allergy outcomes at 3, 6, 12 months and 2 years of age including: IgE antibody analysis, food allergy and eczema

Notes

Funding: Medical Research Council of Southeast Sweden (FORSS), The Östergötland County Council, The Ekhaga Foundation, Swedish Asthma and Allergy Association, The Swedish Research Council for Environment, Agricultural Sciences and Spatial Planning (FORMAS), The Swedish Society of Medicine and Glaxo Smith Kline, Sweden;

Bio Marin capsules were supplied at a reduced price from Pharma Nord, Denmark.

Declarations of interest: none declared

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "block randomization"
Allocation concealment (selection bias)	Unclear risk	Quote: "randomly allocated"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Placebo-controlled; however women may have been able to detect if they were in the omega-3 group through fishy burps.
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Outcome assessors were blinded.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Total 28/145 (19%) women not included in analysis: 25 women did not complete the requested 15 week intervention period (16, 23% omega-3 and 9, 12% placebo) and were excluded from the analysis, 1 withdrew postpartum, 2 not followed as moved (group not stated). 2 year follow-up: 17/70 omega-3 group (24%) and 12/75 (16%) placebo group were lost to follow-up
Selective reporting (reporting bias)	Unclear risk	Mostly allergy outcomes; some child development outcomes not fully reported



Furuhjelm 2009 (Continued)

Other bias Low risk Maternal LA and AA levels at study entry inclusion were not equal in the 2 groups

Giorlandino 2013

Methods	RCT: ISRCTN39268609			
Participants	43 women randomised			
		men at high risk of preterm birth (history of previous IUGR, fetal demise or PE) s preterm birth and/or ultrasonographic findings of cervical incompetence		
	bleeding episode in the to fish, regular intake o	on-viable fetus (before or after randomisation), a history of placental abruption, e present pregnancy, use of (or used) PG inhibitors, multiple pregnancy, allergy of fish oil, a positive cervical swab for chlamydia, mycoplasma/ureaplasma and ections, major fetal abnormalities.		
	Setting: Artemisia Medical Centre, Rome, Italy			
Interventions	VAGINAL APPLICATIO	N: DHA versus placebo		
	Group 1: DHA (1 g/day) vaginally: n = 22			
	Group 2: placebo vaginally: n = 21			
	Timing: from 21 weeks to 37 weeks 0 days			
	DHA + EPA dose/day: high: 1 g DHA			
Outcomes	Women/birth: GA at b	irth		
	Babies/infants/childr	en: birthweight		
Notes	Funding: Pharmarte Srl (Italy) and sponsors Italian Society of Prenatal Diagnosis and Fetal Maternal Medicine (S.I.Di.P.) (Italy) and the Artemisia Foundation in Fetal-Maternal Medical Research. The authors report that the funders had no role in data collection, data analysis, data interpretation or writing of the report. Declarations of interest: not reported			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Low risk	Quote: "customised randomisation programme that generated a random number for each participant, with equal ratio of selection"		
Allocation concealment	Unclear risk	Not reported		

Physicians and women were blinded to treatment.

Not reported, but probably done.

(selection bias)

mance bias) All outcomes

All outcomes

Blinding of participants

and personnel (perfor-

Blinding of outcome as-

sessment (detection bias)

Low risk

Low risk



Giorlandino 2013 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	1 woman in the omega-3 group was lost to follow-up (1/22); and women whose condition worsened were taken off treatment (1/22 in the omega-3 group and 7/21 in the placebo group).
Selective reporting (reporting bias)	Unclear risk	Birthweight was only reported for women who gave birth at 37 weeks' gestation or later (and was therefore not included in the meta-analysis).
Other bias	Low risk	No apparent risk of other bias.

Gustafson 2013

RCT: NCT01007110 (HOPE)				
67 women randomised				
Inclusion criteria: women 16 to < 40 years old with a singleton pregnancy, 12-20 weeks GA				
Exclusion criteria: any serious health condition likely to affect the growth and development of the fetus or the health of the mother including cancer, lupus, hepatitis, diabetes mellitus (type 1, 2 or gestational) or HIV/AIDS at baseline. Women who self-reported illicit drug use or alcohol use during pregnancy and those with hypertension or BMI ≥ 40 were excluded. Women who were taking more than 200 mg/day DHA in prenatal vitamins or over-the-counter supplements were excluded from participation.				
Setting: Kansas City, Kansas, USA				
SUPPLEMENTATION: DHA versus placebo				
Group 1: DHA 600 mg/day: contained 500 mg of oil: algal oil as a source of DHA (200 mg of DHA per capsule; 3 capsules a day): total number randomised = 35				
Group 2: placebo (3 placebo capsules a day containing 50% soy and 50% corn oil): total number randomised = 32				
Timing of supplementation: 14.4 weeks GA \pm 4 weeks; women were advised to stop taking capsules once they had given birth				
DHA + EPA dose/day: mid: 600 mg DHA + EPA negligible				
Women/birth: DHA RBC concentrations; GA at birth				
Babies/infants/children: fetal heart rate, heart rate variability (at 24, 32 and 36 weeks GA); birthweight; birth length; DHA RBC concentrations; NBAS at 1-14 days postpartum				
Funding: Eunice Kennedy Shriver National Institute of Child Health and Development, Kansas Intellectual Development and Disabilities Research Center; study product donated by DSM Nutritional Products (P30NICHDHD002528).				
Declarations of interest: not reported				

Bias	Authors' judgement Support for judgement		
Random sequence generation (selection bias)	Low risk	Computer-generated random sequence	
Allocation concealment (selection bias)	Low risk	Quote: "only members of the investigational pharmacy knew the subject allocation"	



Gustafson 2013 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Participants and all members of the investigational team were blinded to the intervention assignment.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported - insufficient information to make any judgement.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	33% loss to follow-up overall: to birth (23/69): In the control group, 8/32 (25%): 3 withdrawals 1 did not meet inclusion criteria 1 miscarriage 1 high-risk pregnancy 1 congenital anomalies 1 preterm birth In the DHA group: 13/35 (37%): 1 fetal demise 1 miscarriage 6 lost contact/transferred 1 withdrawal 1 excessive morning sickness 1 cholelithiasis 1 miscalculated due date 1 preterm birth NBAS: a further 12 from the control group and a further 7 from the DHA group did not have NBAS assessments
Selective reporting (reporting bias)	High risk	Few maternal and birth outcomes reported
Other bias	Low risk	Maternal characteristics at trial entry were similar, no other sources of bias were apparent.

Haghiac 2015

Methods	RCT: NCT00957476
Participants	72 women randomised
	Inclusion criteria: overweight/obese pregnant women (BMI ≥ 25 at first antenatal visit); singleton pregnancy and GA between 8 weeks and 16 weeks
	Exclusion criteria : known fetal anomaly, regular intake of fish oil supplements (> 500 mg per week in the previous 4 weeks), daily use of NSAIDs; pre-existing metabolic disorder such as hypertension, diabetes or hyperthyroidism; allergy to fish or fish products; gluten intolerance; women who are vegetarians and do not eat any fish; planned termination of pregnancy or birth at another hospital; known HIV-positive, illicit drug or alcohol use during current pregnancy



Hagh	iac 2015 ((Continued)
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Setting: MetroHealth Medical Center, Ohio, USA (participants recruited September 2009 to August 2011)

Interventions

SUPPLEMENTATION: DHA + EPA versus placebo

Group 1: DHA plus EPA (total 2 g/day): 800 mg DHA (22:6n-3) and 1200 mg EPA (20:5n-3): 4 capsules (2 x twice a day). Total number randomised: n = 36 (25)

Group 2: placebo (2 capsules twice a day); contains wheat germ oil. Total number randomised: n = 36 (25)

Timing of supplementation: weeks 10-16 to term

DHA + EPA dose/day: high: 800 mg DHA + 1200 mg EPA

Outcomes

Women/birth: length of gestation; maternal plasma omega-3 and omega-6 concentrations; CRP; TLR4, IL6, IL8 (in adipose and placental tissue); glucose concentrations; insulin sensitivity (narrative report only); adiponectin; leptin; spontaneous abortions; stillbirth; gestational diabetes; placental gene expression; placental triglycerides

Babies/infants/children: birthweight; neonatal lean mass; fat mass; body fat; pea pod lean mass; pea pod fat mass; pea pod body fat

Notes

Notes relating to intervention: adherence run-in: consenting eligible women were given 1 week's supply of placebo capsules; they were not allowed to participate in the trial if they did not return or if they had taken < 50% of the placebo capsules.

Funding: NIH RHD057236. Emiment supplied the study supplements.

Declarations of interest: "The authors declared that they have no conflicts of interest".

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computerised random number generation
Allocation concealment (selection bias)	Low risk	Quote: "Randomization and treatment assignment were carried out by the research coordinators"
		Probably done
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Study group assignment was not known by study participants, their health care providers, or the research staff"
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not reported but probably done
Incomplete outcome data (attrition bias) All outcomes	High risk	21/72 (29%) attrition: Omega-3 group, lost 10: 7 missed second visit 1 spontaneous abortion 1 unable to contact 1 moved away



Haghiac 2015 (Continued)		
		Control group, lost 11:
		 7 missed second visit
		 1 spontaneous abortion
		• 2 unable to contact
		 1 moved away (slightly different numbers reported in different parts of the papers)
Selective reporting (reporting bias)	Unclear risk	Few maternal, birth and neonatal outcomes reported
Other bias	Unclear risk	Baseline characteristics were similar except for higher average weight (but not BMI) in the omega-3 group.

Harper 2010

Methods RCT: NCT00135902

Participants

852 women randomised

Inclusion criteria: women with at least 1 prior spontaneous preterm birth; singleton pregnancies; GA at randomisation between 16 and 22 (21 6/7) weeks. An US examination was required between 14 weeks GA and enrolment to screen for major anomalies.

Exclusion criteria: major fetal anomaly or demise; regular intake of fish oil supplements (> 500 mg per week at any time during the preceding month); daily use of NSAIDs; allergy to fish or fish products; gluten intolerant; heparin use or known thrombophilia; haemophilia; planned termination; current hypertension or current use of antihypertensive medications; uncontrolled thyroid disease; type D, F or R diabetes; maternal medical complications; current or planned cerclage; illicit drug or alcohol abuse during current pregnancy; plan to, or give birth at a non-network hospital; participation in another pregnancy intervention study; participation in this trial in a previous pregnancy; seizure disorder.

Characteristics: 30% women never consumed fish or consumed fish < once per month; 9% consumed fish > 3 times a week.

Setting: antenatal clinics in 13 network centres, USA: recruitment between January 2005 and October 2006

Interventions

SUPPLEMENTATION: DHA + EPA + PG versus placebo + PG

Group 1: 1200 mg EPA; 800 mg DHA for a total of 2000 mg of omega-3 long-chain polyunsaturated acids, divided into 4 capsules per day: total number randomised: n = 434

(Source of omega-3 LCPUFA was deep ocean fish; each capsule contained 10 IU of vitamin E as a preservative.)

Group 2: matching placebo (4 capsules containing a minute amount of inert mineral oil per day). Total number randomised: n = 418

Timing of supplementation: 16-22 weeks GA (mean 19.6 weeks) to 36 weeks GA

All women: received 17α -hydroxyprogesterone caproate (weekly intramuscular injections: 250 mg); participants received no dietary advice as part of the study and otherwise received usual clinical care.

DHA + EPA dose/day: high: 800 mg DHA + 1200 mg EPA

Outcomes

Women/birth: fatty acid status; diet (fish intake); DNA; PE; PPH; adverse events; gestational diabetes; preterm birth < 37 weeks; spontaneous preterm birth < 37 weeks; preterm birth < 34 weeks; birth > 40 weeks; low birthweight < 2500 g; SGA; LGA; GA reported as IQR



Harper 2010 (Continued)

Babies/infants/children: birthweight (median); NEC; RDS (clinical diagnosis of RDS and oxygen therapy ($FiO_2 > 0.40$); neonatal sepsis; BPD; ROP, IVH, perinatal mortality (pregnancy loss and neonatal death); NICU/intermediate care nursery admission; neonatal morbidity composite (ROP, grade III or IV IVH, Patent ductus arteriosus, NEC, culture-proven sepsis, respiratory morbidity, and perinatal death)

Notes

Compliance run-in: consenting women received an injection of 250 mg 17α -hydroxyprogesterone caproate (17P) and a 7-day supply of placebo capsules. Those who did not return after 5 days and before 21 6/7 weeks GA or had taken < half of the placebo capsules were not allowed to participate. Women passing the compliance run-in were randomly assigned to EPA/DHA or placebo.

Funding: The Eunice Kennedy Shriver National Institute of Child Health and Human Development Grants (HD27860, HD27917, HD40560, HD34208, HD40485, HD21410, HD27915, HD40500, HD40512, HD40544; MO1-RR-000080; HD34136; HD27869; HD40545; HD36801 and HD19897).

Declarations of interest: "Dr Esplin serves on the scientific advisory board and holds stock in Sera Prognostics, a private company that was established to create a commercial test to predict preterm birth and other obstetric complications. Dr Manuck is also on the scientific advisory board for Sera Prognostics". None of the other authors reported any conflicts of interest.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "randomly assigned"; "simple urn method of randomization with stratification according to clinical center to create a randomization sequence for each center"
Allocation concealment (selection bias)	Low risk	Quote: "randomly assigned"; "simple urn method of randomization with stratification according to clinical center to create a randomization sequence for each center"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "double-masked"; "Study group assignment was not known by study participants, their health care providers or the research personnel"; placebo control
Blinding of outcome assessment (detection bias) All outcomes	Low risk	After giving birth, "records of the participants and their newborns were reviewed by study personnel, unaware of treatment assignments, who abstracted delivery date, birth weight, occurrence of maternal or neonatal complications and interventions"
Incomplete outcome data	Low risk	No losses to follow-up (for primary outcome)
(attrition bias) All outcomes		Denominators for liveborns = 427 and 410 (2 neonates in the omega-3 group and 7 in the placebo group died before admission to NICU and were not included in liveborn neonate outcomes).
Selective reporting (reporting bias)	Low risk	Large number of relevant outcomes reported (some as medians).
Other bias	Low risk	Baseline demographics, risk factors for recurrent preterm birth and dietary fish intake were similar between omega-3 and placebo groups.

Harris 2015

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Harris 2015 (Continued)

Participants

843 pregnant women randomised (634 to the 3 supplement arms and 209 to the single nutrition arm)

Inclusion criteria: women recruited at 16-20 weeks of gestation or at WIC intake visits with singleton pregnancies; 18 to 40 years of age, able to sign informed consent and Health Insurance Portability and Accountability Act forms in English or Spanish

Exclusion criteria: women presenting with known medical or obstetrical complications associated with increased risk for preterm birth including cervical incompetence, presence of cervical cerclage, placenta praevia, intrauterine infection, known substance abuse, multiple fetuses, current PE, pre-existing diabetes, or a history of gestational diabetes in a prior pregnancy. Women were also excluded if they were taking NSAIDS or if they consumed salmon, mackerel, rainbow trout, or sardines at least once weekly or if they had known allergies to fish or any constituent of the nutritional supplement.

Characteristics: low-income population, but at lower risk of preterm birth with predominantly Hispanic women included

Setting: antenatal clinics, Denver Health Hospitals (Denver, Colorado, USA)

Interventions

SUPPLEMENTATION: DHA, 300 mg versus 600 mg, as bars) versus placebo

Group 1: supplementation: 300 mg algae-derived DHA (200 women randomised) – provided in the form of 300 Kcal supplement bars containing DHASCO-S oil (DHA single-cell oil)

Group 2: supplementation: 600 mg of algae-derived DHA (221 women randomised) – provided in the form of 300 Kcal supplement bars containing DHASCO-S oil (DHA single-cell oil)

Group 3: placebo: olive oil (213 women randomised) – provided in the form of 300 Kcal supplement bars

All women: gel capsules containing the test oil or olive oil were available for those who refused the bars (51 women opted for gel capsules - groups not reported).

Timing of supplementation: from 20 weeks GA to birth

DHA + EPA dose/day: low and mid: 300 mg and 600 mg DHA/day + negligible EPA

Outcomes

Women/birth: adherence; DHA concentrations at birth; dietary intake of DHA-rich foods; adverse events (including vaginal infection, vaginal bleeding during pregnancy and preterm labour); GA at birth; preterm birth < 34 weeks (preterm birth < 280 days was reported but not by group); post-term birth; mode of birth; type of rupture of membranes; type of labour onset; estimated blood loss; birth complications.

Babies/infants/children: birthweight; birth length; head circumference

Notes

- Women at risk for preterm birth were excluded
- A 4th non-randomised arm offered nutrition education, canned fish and egg coupons to 209 women attending a WIC clinic from 18 to 20 weeks' gestation (191 women completed). Results for this arm were similar to the DHA supplement arms (apart from a higher rate of induced labour > 40 weeks in the nutrition education arm).
- Same NCT # allocated to Miller 2016

Funding: United States Department of Agriculture. Bars and gel capsules were supplied by Martek Biosciences, Columbia, MD, USA.

Declarations of interest: none reported

Risk of bias

Bias Authors' judgement Support for judgement



Harris 2015 (Continued)		
Random sequence generation (selection bias)	Low risk	Quote: "stratified block randomization schedule, generated using a randomization table by staff at Martek Biosciences, to insure equal group assignment from each of three clinics participating in the supplement trial"
Allocation concealment (selection bias)	Low risk	As above; probably adequate allocation concealment (third party)
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Both participants and all study personnel were blinded.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not reported, but probably done.
Incomplete outcome data (attrition bias) All outcomes	High risk	53.5% and 52.9% were lost to follow-up from the 2 intervention arms (leaving 107/200 in the 300 mg DHA arm and 117/221 in the 600 mg DHA arm: 56.8% (121/213) were lost to follow-up from control arm. Reasons were not given.
Selective reporting (reporting bias)	Unclear risk	Insufficient information to make judgement; preterm birth not fully reported
Other bias	Low risk	Comparable baseline characteristics for total fatty acids, maternal BMI and maternal age, parity and previous preterm birth (and for completers and withdrawals)

Hauner 2012	
Methods	RCT: NCT00362089 (INFAT - The Impact of Nutritional Fatty acids during pregnancy and lactation for early human Adipose Tissue development)
Participants	208 women randomised
	Inclusion criteria: healthy pregnant women < 15 weeks GA, aged 18-43 years, BMI at conception between 18 and 30, sufficient German language; for infants at follow-up, GA at birth between 37-42 weeks, appropriate size for GA, and Apgar score > 7 at 5 minutes
	Exclusion criteria: high-risk pregnancy (multiple pregnancy, hepatitis B or C infection, parity > 4), hypertension, chronic diseases such as diabetes or gastrointestinal disorders, psychiatric disorders, supplementation with omega-3 fatty acids before randomisation, alcohol abuse, hyperemesis gravidarum, smoking; known metabolic defects
	Characteristics: mean baseline BMI of 22; women were relatively well educated
	Setting: University Hospital Klinikum rechts der Isar, Technische Universität München, Germany (women recruited between 2006 and 2009)
Interventions	SUPPLEMENTATION + DIET ADVICE: DHA + EPA + diet advice versus diet advice
	Group 1: omega-3 LCPUFA (180 mg EPA and 1020 mg DHA (= 1200 mg omega-3) and 9 mg vitamin E), taken as 3 capsules per day and requested to restrict consumption of AA-rich foods (e.g. meat (500 g a week = $2-3$ portions), meat products and eggs); n = 104
	Group 2: brief semi-structured counselling on a healthy diet according to the guidelines of the German

frain from taking fish oil or DHA supplements: n = 104

Nutrition Society for a healthy balanced diet; women in the control group were specifically asked to re-



Hauner 2012 (Continued)

All women: participants of both groups were also offered individual nutrition counselling based on the 7-day dietary record.

Timing of supplementation: women were randomised and began the study at 15 weeks GA and continued supplementation until 4 months lactating (or time when ceased breastfeeding if earlier)

DHA + EPA dose/day: high: 1020 mg DHA + 180 mg EPA

Outcomes

Women: adherence; preterm birth; post-term birth; induction (all at term); GWG; blood loss at birth; gestational diabetes; pathological cardiotocography; cessation of labour; retained placenta; mode of birth (spontaneous birth, caesarean section, vacuum extraction); breastfeeding; blood lipid concentrations (triglycerides, total cholesterol, high- and low-density lipoproteins) at baseline, 32 weeks GA, birth, 6 weeks and 4 months postpartum in pregnant and lactating women; leptin; fatty acid pattern in erythrocytes and plasma in maternal blood as well as umbilical cord blood samples and adipokines in maternal plasma, as well as umbilical cord plasma samples and breastmilk samples at 6 weeks and 4 months; maternal 7-day dietary questionnaire (energy, protein, carbohydrates, lipids, AA); maternal plasma levels of DHA, EPA and AA reported as per cent weight of total fatty acids; maternal RBC fatty acid baseline (16-21 weeks GA) – before study drug was dispensed (reported in Hauner 2009 only by fish consumption), insulin resistance; maternal leptin; cord blood insulin concentrations

Babies/infant/children: Apgar score; LGA > 90th percentile; adipose tissue mass (skinfold thickness) 3-5 days after birth, 6 weeks, 4 and 12 months postpartum; subgroup: subcutaneous and visceral fat mass ultrasonography at 6 weeks, 4 and 12 months postpartum and MRI at 6 weeks and 4 months postpartum; birthweight; birth length; head circumference; upper arm circumference); body weight and length, head circumference and upper arm circumference, BMI (kg/m²) - all at birth/3-5 days after birth, 6 weeks, 4 and 12 months postpartum; weight/length (g/cm) at birth; ponderal index (kg/m³) at birth; fetal leptin; annual body-composition measurements including skinfold thickness measurements (primary outcome) - up to 5 years, a sonographic assessment of abdominal subcutaneous and preperitoneal fat, and child growth at 2, 3, 4 and 5 years (weight, height, head circumference, BMI percentile, waist circumference); abdominal MRI was performed in a subgroup of 5-year-old children; dietary intake at 3, 4 and 5 years; physical activity at 3, 4 and 5 years; Child Development Inventory at 4 and 5 years; hand movement test at 5 years (mirror movements reported as medians).

Notes

Funding: Else Kröner-Fresenius Foundation, Bad Hamburg, the International Unilever Fund, EU-funded EARNEST Consortium (subcontractor Numico, Frankfurt), and the German Ministry of Education and Research via the Competence Network on Adiposity; Danone Research-Centre for Specialised Nutrition, Friedrichsdorf, Germany. The analysis of fatty acids was performed by the laboratory of Lipid Research, Danone Research-Centre for Specialised Nutrition by using coded samples. "There was no intervention from any sponsor with any of the research aspects of the study including study design, intervention, data collection, data analysis and interpretation as well as writing of the manuscript." "Danone as a Funding source and cooperation partner in fatty acid analysis was recruited after the study design was fully established."

Declarations of interest: "HH has received grants from Riemser and Weight Watchers for clinical trials and payment for lectures from Novartis, Roche Germany, and Sanofi-Aventis". The other authors reported no conflicts of interest related to the study.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computerised randomisation
Allocation concealment (selection bias)	Unclear risk	Quote: "randomly assigned"
Blinding of participants and personnel (perfor- mance bias)	High risk	Not blinded; "open label"



Haun	er	2012	(Continued)
A 1.1			

ΛII	loutcomes
ΑU	outcomes

Blinding of outcome as-
sessment (detection bias)
All outcomes

High risk

No (except for US measurements, e.g. for fat mass measurements)

Incomplete outcome data (attrition bias)
All outcomes

High risk

Omega 3: 17/104 (16%) lost at 12 months postpartum:

- 8 personal reasons
- 3 lost to follow-up
- · 1 with group allocation
- 2 intolerance to supplements
- 1 moved away
- 1 repeated non-attendance
- 1 genetic disorder in child diagnosed

Control: 21/104 (20%) lost at 12 months:

- 3 personal reasons
- 2 lost to follow-up
- · 6 unhappy with group allocation
- · 4 moved away
- 3 repeated non-attendance
- 3 preterm birth

2 years and longer:

- 118 children remained at 2 years of age (56.7%)
- 120 children at 3 years of age (57.7%)
- 107 children at 4 years of age (51.4%), and
- 114 children at 5 years of age (54.8%), with similar numbers between study groups (at 5 years: intervention group, n = 58; control group, n = 56). The most common reasons for dropout were a lack of time or relocation.

Unclear why 3/4 preterm births in the control group were treated as exclusions, whereas none of the 3 preterm births in the omega-3 group were.

Selective reporting (re-
porting bias)

Unclear risk

Focus on biochemical and skinfold measurements rather than clinical outcomes

Other bias

Unclear risk

Baseline demographics and characteristics were comparable, except for higher rates of smoking and alcohol use during pregnancy in the control group.

Helland 2001

Methods

RCT

Participants

590 women

Inclusion criteria: healthy women 19-35 years of age with singleton pregnancies, nulliparous or primiparous, intending to breastfeed, no omega-3 supplementation earlier in the pregnancy, 17-19 weeks' gestation

Exclusion criteria: already taking DHA, preterm births, birth asphyxia, general infections, anomalies in infants requiring special attention



Helland 2001 (Continued)

Setting: attendance at routine US scans, Rikshospitalet University Hospital and Baerum Central Hospital, Oslo, Norway; women recruited between December 1994 to October 1996.

Interventions

SUPPLEMENTATION: DHA + EPA (cod-liver oil) versus placebo (corn oil)

Group 1: cod-liver oil (10 mL = 1183 mg DHA, 803 mg EPA (total omega-3 LCPUFA 2494 mg); total number randomised: n = 301

Group 2: liquid oil (corn oil); (10 mL = 4747 mg LnA, 92 mg ALA); total number randomised: n = 289

Fat-soluble vitamin content was identical for both groups (117 μ g/mL vitamin A; 1 μ g/mL vitamin D; 1.4 mg/mL dl- α); cod-liver oil added 42 mg cholesterol per 10 mL.

Timing of supplementation: 18 weeks GA to 3 months infant age

DHA + EPA dose/day: high: 1183 mg DHA + 803 mg DHA

Outcomes

Women: length of gestation; maternal and infant diet (food frequency questionnaires); placental weight; fatty acids in breast milk; breastfeeding; BMI at birth

Babies/infants/children: birthweight; birth length; head circumference at birth; fatty acids (cord blood and infants at 4 weeks and 3 months); electroencephalography (2 days and 3 months); Fagan (infant intelligence) (6 and 9 months); K-ABC (IQ) (4 and 7 years)

Notes

Funding: Peter Möller (Oslo, Norway) provided both oils; Orkla ASA; Eckbos Legater; Aktieselskabet Freia Chocolade-fabriks Medicinske Fond

Declarations of interest: Dr Helland's scholarship during the study period was funded by the Peter Möller Department of Orkla ASA, and Dr Saarem was working at the Peter Möller Department of Orkla ASA during the study period and Dr Drevon has been a consultant for the Peter Möller Department of Orkla ASA; Drs Smith, Saugstad, and Ms Blomén indicated they had no financial relationships relevant to disclose.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "randomization was performed by a computer program"
Allocation concealment (selection bias)	Unclear risk	Quote: "randomization was performed by a computer program"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "double-blinded"
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not specifically stated, but probably done.
Incomplete outcome data (attrition bias) All outcomes	High risk	249/590 (42%) lost to follow-up by time of birth: 126/301 (42%) in the cod-liver oil group and 123/289 (43%) in the corn oil group. There were 27 exclusions and 222 withdrawals (mostly due to "feeling discomfort taking the oil").
		K-ABC at 4 years of age: 51 of the 135 children invited (38%) were lost to follow-up: 90 came for assessment (84 children completed the assessment).
		K-ABC at 7 years: 119 of the 262 children tested on Fagan intelligence scale during their first year of life were invited were lost to follow-up (45%).



Helland 2001 (Continued)		
Selective reporting (reporting bias)	Unclear risk	Individual K-ABC scales not fully reported; did not have SDs reported at 4 years; no SDs reported at 7 years.
Other bias	Unclear risk	Women in the omega-3 group were on average 1 year older than women in the corn oil group.

Horvaticek 2017

Methods	RCT		
Participants	109 women		
	Inclusion criteria: pre	gnant women with T1D mellitus (not further specified)	
	Exclusion criteria: not	t reported (post-randomisation exclusions: fetal loss, preterm birth)	
	Setting: Referral Center for Diabetes in Pregnancy Ministry of Health Republic of Croatia, Department of Obstetrics and Gynecology, Zagreb University of Hospital Center, Zagreb, Croatia, conducted 1 January 2014 to 30 September 2016		
Interventions	SUPPLEMENTATION: betic diet	DHA + EPA + standard diabetic diet versus placebo (corn oil) + standard dia-	
		capsules twice daily (each capsule contained EPA 60 mg and DHA 308 mg, there- 16 mg DHA daily); total number randomised: n = 56 (47 included in main analysis)	
	Group 2: placebo caps	sules (corn oil; total number randomised: n = 53 (43 included in main analysis)	
	Timing of supplementation: from 9 weeks GA		
	DHA + EPA dose/day: mid: 616 mg DHA + 120 mg EPA		
Outcomes	Women/birth: adherence; preterm birth (< 37 weeks); early preterm birth (< 34 weeks); length of tion; PE (and hypertension); GWG Babies/infants/children: miscarriage; stillbirth; perinatal mortality; birthweight > 4.5 kg (macros mia); birthweight; birth length; head circumference		
Notes	Funding: not reported Declarations of interest: none declared		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Method used to generate a random sequence not reported.	
Allocation concealment (selection bias)	Unclear risk	Quote: "It was randomly decided which pregnancy women with type-1 diabetes would take EPA and DHA or which placebo".	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Placebo-controlled	



sessment (detection bias) All outcomes Incomplete outcome data (attrition bias) All outcomes High risk 90/10t = 47 ir rando flow composition possible for the second possibl	
(attrition bias) = 47 in rando flow c 56 wo spectition p Interv • 4s • 1 fe • 3 p Control • 3 s • 3 g • 4 p Selective reporting (reporting bias) High risk Outcomes	ported
• 4s • 1 fe • 3 p Control • 3s • 3 g • 4 p Selective reporting (re-porting bias) High risk ber of from fe	(81%) women included in the study remained at the birth time point (n intervention group and n = 43 in control group). The number of women mised to the intervention and control groups was not reported. From nart (Figure 1) we might assume (though not with 100% certainty) that men were assigned to the intervention, and 43 to the control groups revely. Before the end of pregnancy/birth pregnancy (main) data collectiont:
• 1 fe • 3 p Control • 3 s • 3 g • 4 p Selective reporting (re-porting bias) High risk Outcoporting bias) Outcoporting bias	ention group lost 9/56 (16%):
• 3 p Control • 3 s • 3 g • 4 p Selective reporting (re- porting bias) High risk ber of from f	pontaneous abortions
Selective reporting (reporting bias) Control 3 s 4 p Selective reporting (refrom from from from from from from from	tal demise
Selective reporting (reporting bias) High risk Outco ber of from f	reterm births
Selective reporting (reporting bias) High risk Outcon ber of from f	ol group lost 10/53 (19%):
Selective reporting (reporting bias) High risk Outcon ber of from f	pontaneous abortions
Selective reporting (re- porting bias) High risk ber of from f	ave birth in other clinics
porting bias) ber of from f	reterm births
Other bias Unclear risk Data p	mes of trial poorly defined and no protocol available. Additionally, num- participants randomised to each group not reported in text (inferred owchart of participants in the trial).
•	rovided on participant characteristics at baseline insufficient to confiassess similarity of baseline characteristics.

Hurtado 2015

Methods	RCT: NCT01947426 (NUGELA)
Participants	110 women randomised
	Inclusion criteria: healthy term infants with no presence of diseases that may affect the normal development of pregnancy or lactation, singleton gestation, normal course of pregnancy, BMI of 18 to 30 kg/m² at the start of pregnancy, weight gain of 8 kg to 12 kg since pregnancy onset, no intake of DHA supplements during pregnancy, term birth, spontaneous vaginal birth, appropriate weight for GA, Apgar index ≥ 7 at 1st and fifth minute of life, normal monitoring results, and breast-feeding of the neonate
	Exclusion criteria: see above
	Setting: 2 hospitals, Hospital Materno-Infantil (Granada, Spain) and Hospital Universitario Materno-Infantil (Las Palmas de Gran Canaria, Spain), between June 2009 and August 2010
Interventions	SUPPLEMENTATION + FOOD: DHA + EPA versus placebo (in the form of a dairy product)
	Group 1: Fish oil enriched dairy drink (400 mL enriched with omega-3 (total 392 mg – 72 mg EPA and 320 mg DHA/day; fish oil from tuna)): total number randomised = 56
	Group 2: dairy drink with no fish oil: 400 mL: total number randomised = 54
	Timing of supplementation: from 28 weeks GA to 4th month of lactation
	Both groups of women received dietary advice



Hurtac	o 2015	(Continued)
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DHA + EPA dose/day: low: 320 mg DHA + 72 mg EPA

Outcomes

Women/birth: fatty acid profiles were determined in the mother's (at enrolment, at birth, and at 2.5 and 4 months) and newborn (at birth, and at 2.5 months) placenta and breast milk (colostrum and at 1, 2, and 4 months); maternal diet (enrolment, 1 month after enrolment and first month of lactation); GWG; GA; placenta DMT1, FPN1, TfR1 and Hamp1 mRNA and protein expression; hepcidin expression; oxidative damage biomarkers (enrolment, birth, 2.5 and 4 months postpartum); inflammatory markers (birth and 2.5 months); cytokines

Babies/infants/children: hepcidin expression; oxidative damage biomarkers (birth; 2.5 months); Apgar at 1 and 5 minutes; birthweight; birth length; head circumference at birth; pattern reversal visual evoked potentials (VEPs) (at 2.5 and 7.5 months) and Bayley test (at 12 months) – BSID II MDI; PDI

Notes

Funding: Excellence grant (mP-BS-9) from the Campus de Excelencia Internacional GREIB (Granada Research of Excellence Initiative on BioHealth)

Declarations of interest: "F.L.-V is an employee of Lactalis Puleva". No other conflicts declared.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "unpredictable sequence computer-generated"
Allocation concealment (selection bias)	Unclear risk	Quote: "randomly assigned"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Identical white packaging used; trial investigators and participants were unaware of the treatment allocation.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not reported, but probably done.
Incomplete outcome data (attrition bias) All outcomes	High risk	34/110 (31%) lost to follow-up by the end of the intervention. In the omega-3 group 18/56: 1 lactose intolerant 4 did not attend 1 gestational diabetes 3 no reason given 1 DHA supplemented 4 did not like the milkshakes 4 did not breastfeed
		In the control group 16/54: 1 lactose intolerant 2 did not attend 3 no reason given 4 DHA supplemented 1 congenital heart disease 1 moved 4 did not breastfeed



Hurtado 2015 (Continued)		Likely to have been post randomisation exclusions (e.g. preterm) but none were mentioned.
		At 12 months: 24/56 (43%) in the DHA group and 25/54 (46%) in the placebo group were lost to follow-up.
Selective reporting (reporting bias)	Unclear risk	Inadequate information to assess confidently.
Other bias	Low risk	Similar baseline characteristics

Ismail 2016			
Methods	RCT: NCT01990690		
Participants	140 women randomised		
	Inclusion criteria: pregnant women with singleton pregnancy (30-34 weeks' gestation) with an ultrasonographic diagnosis of oligohydramnios (amniotic fluid index ≤ 5 cm); women aged 20-35 years with normal Doppler indices in uterine and umbilical arteries at the time of recruitment (the normal value of S/D ratio is from 2.5 to 3.5; RI is from 0.60 to 0.75; and PI is from 0.96 to 1.27)		
	paired liver or kidney for non-reactive non-stres	men with evidence of IUGR, history of premature rupture of membranes, imunction, PE or long-term diabetes and any placental abnormalities; women with s test or women using NSAIDs or who had any congenital fetal malformation; bnormal Doppler indices at the time of recruitment	
	Setting: Department of Obstetric and Gynecology, Woman's Health Hospital, Assiut University, Assiut, Egypt (conducted between 1 January 2015 and 1 August 2015)		
Interventions	SUPPLEMENTATION: EPA + DHA versus placebo		
	Group 1: omega-3 capsules: 1000 mg fish oil (containing 13% EPA and 9% DHA plus 100 mg wheat-germ oil (LA 52% to 59%) as a natural source of vitamin E once daily for 4 weeks: n = 70		
	Group 2: placebo: once daily empty soft gelatin capsules of the same shape, size, colour and weight for the same duration: Total number randomised: n = 70		
	Timing of supplementation: from 30-34 weeks' gestation for 4 weeks		
	DHA + EPA dose/day: low: 90 mg DHA + 130 mg EPA		
Outcomes	Women: improvement in amniotic fluid volume 4 weeks after start of treatment; Doppler blood flow changes in the uterine artery after 4 weeks of treatment; uterine artery Doppler indices		
Notes	No outcomes could be used in this review.		
	Funding: not reported		
	Declarations of interest: none declared		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Computer-generated random table	



Ismail 2016 (Continued)		
Allocation concealment (selection bias)	Low risk	Allocation concealment was carried out using serially numbered closed opaque envelopes. Each envelope was labelled with a serial number and had a card noting the intervention type inside. Allocation was never changed after opening the envelopes. Preparation and sorting of the serially numbered envelopes was carried out by an investigator who did not participate in the evaluation of patients.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Placebo controlled
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not specifically reported, but probably done.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	14.3% (20/140) lost to follow-up: 10/70 in omega-3 group and 10/70 in the placebo group.
Selective reporting (reporting bias)	Unclear risk	No perinatal health outcomes reported.
Other bias	Low risk	Similar baseline characteristics

Jamilian 2016

Methods	RCT: IRCT201406305623N20		
Participants	54 women randomised		
	Inclusion criteria: women with GDM not on oral hypoglycaemic agents (diagnosed by 1-step 2-hour 75 g OGTT at 24-28 weeks GA - ADA criteria) and singleton pregnancy		
	Exclusion criteria: pre-existing diabetes, required complex diets, chronic medical conditions (e.g. valvular heart disease), significant psychiatric disease, smokers, kidney or liver diseases, chronic hypertension or hypothyroidism, those needing to commence insulin therapy during the intervention		
	Setting: Arak, Iran		
Interventions	SUPPLEMENTATION: omega-3 (EPA + DHA) versus placebo		
	Group 1: omega-3 (1000 mg omega-3 pearl (containing 180 mg EPA and 120 mg DHA) per day): total number randomised: $n=27$		
	Group 2: placebo 1 per day (appearance, colour, shape size, and packaging identical to omega-3 capsules): total number randomised: n = 27		
	Timing of supplementation: from 24-28 weeks GA for 6 weeks		
	All women were asked to maintain their usual diet and physical activity levels throughout the study period.		
	All women were also consuming both 400 mg/day folic acid from the beginning of pregnancy and 60 mg/day ferrous sulphate from the second trimester.		
	DHA + EPA dose/day: low: 120 mg DHA + 180 mg DHA		



Jamilian 2016 (Continued)

Outcomes

Women/birth: diet and physical activity at weeks 2, 4 and 6 of intervention; weight at end of intervention; BMI at end of intervention; maternal polyhydramnios; PE; GA; caesarean section; birthweight; birth length; birth head circumference; inflammatory factors, biomarkers of oxidative stress and metabolic biomarkers; need for insulin therapy after intervention; antenatal hospitalisation

Babies/infants/children: Apgar score; hyperbilirubinaemia; intrauterine fetal death; macrosomia (> 4000 g); ponderal index; 1 and 5 minute Apgar; newborn hyperbilirubinaemia; newborn hospitalisation

Notes

Adherence was 100%

Funding: Arak University of Medical Sciences (grant no. 93-165-20)

Declarations of interest: none declared

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "computer-generated random numbers"
Allocation concealment (selection bias)	Low risk	Quote: "Random assignment was done by the use of computer-generated random numbers randomization and allocation were concealed from the researchers and participantsA trained midwife at the maternity clinic did the randomized allocation sequence, enrolled participants, and assigned participants to intervention"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Placebo-controlled
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not reported, but probably done.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	3/27 women in the omega-3 group withdrew, and 2/27 women in the place- bo group withdrew, all 5 for personal reasons. Data for all 54 women were analysed, using Last Observation Carried Forward for missing data.
Selective reporting (reporting bias)	Low risk	Most expected perinatal outcomes were reported.
Other bias	Low risk	Baseline characteristics were similar

Jamilian 2017

Methods	RCT (4 arms, see below)
Participants	140 women randomised

Inclusion criteria: women 18-40 years; without prior diabetes; diagnosed with GDM by 1-step 2-hour 75-g OGTT at 24-28 weeks' gestation referred to Kosar Clinic in Arak, Iran. GDM was diagnosed according to the ADA guidelines: those whose plasma glucose met 1 of the following criteria were considered as having GDM: FPG ≥ 92 mg/dL, 1-hour OGTT ≥ 180 mg/dL, and 2-hour OGTT ≥ 153 mg/dL.



Jamilian 2017 (Continued)

Exclusion criteria: taking vitamin D and/or omega-3 fatty acid supplements; taking insulin; placental abruption; PE; hypothryroidism and hyperthyroidism; smokers; those with kidney or liver disease

Setting: Kosar Clinic, Arak, Iran; women were recruited from March 2016 to July 2016

Interventions

SUPPLEMENTATION + OTHER AGENT: omega-3 versus omega-3 + vitamin D versus vitamin D versus placebo

Group 1: omega-3 fatty acids (2000 mg; 720 mg EPA and 480 mg DHA per day) as 2 capsules (produced by Zahravi Pharmaceutical Company, Tabriz, Iran): n = 35 randomised (n = 32 completed study)

Group 2: omega-3 fatty acids (2000 mg; 720 mg EPA and 480 mg DHA per day) as 2 capsules plus vitamin D (50,000 IU every 2 weeks): n = 35 randomised (and completed study)

Group 3: vitamin D (50,000 IU every 2 weeks) plus omega-3 placebo capsules: n = 35 randomised (and completed study)

Group 4: no supplement (placebo) as 2 capsules that were indistinguishable in colour, shape, size, and packaging, smell, and taste from the vitamin D and omega-3 fatty acids capsules: n = 35 randomised (32 completed study)

Timing of supplementation: 6 weeks (start 24-28 weeks' gestation)

DHA + EPA dose/day: high: 480 mg DHA + 720 mg EPA

Outcomes

Women/birth: insulin metabolism (primary); lipid concentrations (secondary)

Notes

No outcomes could be included in this review.

Funding: grant from the Vice-Chancellor for Research, AUMS (no.1394.373)

Declarations of interest: none declared

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Random sequence generation was ensured using a computer-generated random numbers table.
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "appearance of placebo capsule was indistinguishable in color, shape, size, and packaging, smell, and taste from Vitamin D and omega-3 fatty acids capsules"; considering the nature of intervention and placebo, participants and personnel could have been effectively blind to group assignments throughout the trial.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	140 women were randomised, 35 each to: placebo; vitamin D; omega-3; and vitamin D + omega-3. A total of 6 women (3 placebo group, 3 omega-3), were lost to follow-up, leaving 134/140 (97%) women randomised in the data collection. The difference in the proportions of randomised women included in the data collection was marginal: 100% of women in the vitamin D and vitamin D + omega 3 groups; and 91% of the women in the remaining 2 groups.



Jamilian 2017 (Continued)			
Selective reporting (reporting bias)	Unclear risk	Insufficient information for confident assessment	
Other bias	Low risk	Baseline characteristics similar; no obvious other bias identified	
Judge 2007			
Methods	RCT		
Participants	ticipants 73 women randomised		
	Inclusion criteria: women aged 18-35 years, primiparous or not been pregnant for the past 2 years, with no pregnancy complications		
	Exclusion criteria: women with a history of drug or alcohol addiction; hypertension, smoking, hyperlipidaemia, renal disease, liver disease, diabetes, or psychiatric disorder, parity > 5, history of chronic hypertension, heart disease, thyroid disorder, multiple gestations or pregnancy-induced complications including hypertension, PE or preterm labour; treated during labour with analgesics such as Stadol (butorphanol tartrate) that may cause infant respiratory distress. Infants born preterm and infants with less than 4 hours of crib time in the first and second days postpartum were excluded from the analyses.		
	Setting: Hartford	Hospital, Connecticut, USA	
Interventions	SUPPLEMENTATI	ON: DHA versus placebo (cereal bars)	
	Group 1: DHA-containing cereal-based bars (300 mg DHA/92 kcal bar, with 8:1 ratio of DHA to EPA); average consumption of 5 bars a week = average 214 mg/day (1.7 g of micro-encapsulated fish oil in each 23 g bar). Total number randomised: n = 37		
	Group 2: cereal-based placebo bars: 1.7 g of corn oil in each 23 g bar (same total macronutrient content as intervention with respect to carbohydrate, protein and fat). Total number randomised: n = 36		
	Timing of supplementation: 24 weeks GA to birth (38-40 weeks GA)		
	DHA + EPA dose/o	day: low: 191 mg DHA + 23 mg EPA	

Outcomes

Women/birth: GWG; GA; birthweight; head circumference; length; sexually transmitted infections; maternal depression; mode of birth; maternal dietary intake; DHA status

Infant/child: Apgar score at 1 and 5 minutes; Infant Planning test (total intention scores over 5 trials and intentional solutions – retrieving a toy) at age 9 months; Fagan Test of Infant Intelligence (recognition memory) at age 9 months; mode of feeding; infant sleep; ponderal index; visual acuity. (*Developmental assessments were only conducted on healthy infants*.)

Notes

Funding: US Department of Agriculture Initiative for Future Agriculture and Food Systems; NESTEC; US Department of Agriculture Agricultural Research Service; University of Connecticut Research Foundation; National Fisheries Institute; American Dietetic Association Foundation

Declarations of interest: not reported

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomly assigned"; no further details reported



Judge 2007 (Continued)		
Allocation concealment (selection bias)	Unclear risk	Quote: "randomly assigned"; no further details reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "double-blind, placebo-controlled"
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Problem solving trials were administered and scored by a single tester who was blinded to test groups (one-third of videos were scored by an additional rater).
Incomplete outcome data (attrition bias) All outcomes	High risk	Apparently large and differential losses to follow-up.
Selective reporting (reporting bias)	Unclear risk	Some outcomes not fully reported.
Other bias	Unclear risk	Baseline characteristics were mostly similar between groups; more women in the placebo group had higher BMI and received WIC support compared with the omega-3 group.

Judge 2014

Methods	RCT
Participants	73 women randomised
	Inclusion criteria: no other births in previous 2 years; ≤ 20 weeks' gestation; aged 18-35 years
	Exclusion criteria: women with a self-reported significant medical history (e.g. currently being treated for depression/psychiatric illness, addiction problems, hyperlipidaemia, hypertension, renal disease, liver disease or diabetes)
	Setting: several WIC's offices and hospitals in New England, USA
Interventions	SUPPLEMENTATION: DHA versus placebo
	Group 1: DHA (1 fish oil capsule 300 mg DHA 5 days per week): total number randomised: n = 37 (20)
	Group 2: placebo (1 corn oil capsule, no DHA 5 days per week): total number randomised: n = 36 (22)
	Timing of supplementation: 24 weeks GA to 40 weeks GA (or to birth)
	DHA + EPA dose/day: low: 215 mg DHA; EPA not stated
Outcomes	Women/birth: maternal postpartum depressive symptomatology at 2 and 6 weeks; and 3 and 6 months postpartum; repeated measures over 6 months (assessed with the PDSS and CES-D; RBC DHA (weight%)
Notes	Funding: Patrick and Catherine Weldon Donaghue Medical Research Foundation, Hartford CT; LodersCroklaan (supplied capsules), University of Connecticut School of Nursing and Agricultural Center and Pennington Biomedical Research Center, Louisana State University
	Declarations of interest: none declared



Judge 2014 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomized utilizing a coded marble system"
Allocation concealment (selection bias)	Low risk	Quote: "randomized utilizing a coded marble system and assigned to groups by a trained individual who was not a research team member. Packages containing capsules were labeled identically and listed only sequential study iden tification numbers."
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "A record linking participant names with group assignments was maintained in a secure location away from the researchers to ensure adequate blinding throughout the investigation from recruitment to the completion of data analysis. Identical numbered packages were assembled in advance for use by the research team in enrolling participants. Participants were blinded to group allocation through identical dose and packaging."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not specifically stated, but probably done.
Incomplete outcome data (attrition bias) All outcomes	High risk	Intervention group lost 17/37 (46%) to follow-up: 7 unable to contact/noncompliant 1 transportation issues 1 desire to take fish oil supplements 4 unrelated medical conditions 1 unrelated fetal demise 3 missing data Control group lost 14/36 (39%) to follow-up: 4 unable to contact/noncompliant 1 too busy 1 social issues 1 unrelated medical complication 1 "suspect" 6 missing data
Selective reporting (reporting bias)	High risk	Only 2 outcomes reported (depression and DHA concentrations)
Other bias	Unclear risk	Baseline depressive symptoms (CES-D) were similar; however baseline RBC DHA concentrations were higher in the intervention group.

Kaviani 2014

Methods	RCT: IRCT201212101011717	
Participants	80 women randomised	
	Inclusion criteria: primiparous women > 20 weeks' gestation, with mild depression BDI score between 14 to 19, > 18 years of age, not consuming fish more than twice a week#, not suffering from schizophrenia, bipolar disorders, blood disorders, such as Von Willebrand, hypertension, hyperlipidaemia, renal	



Kaviani 2014	(Continued)
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and thyroid diseases, not taking anticoagulants or antidepressants, not smoking or using narcotics, or not participating in activities such as yoga, relaxation, and psychological consultations.

#those consuming fish more than twice a week were replaced by the next individual

Exclusion criteria: allergic or digestive reactions to study medications

Setting: 2 randomly selected health centres in Shiraz, Iran

Interventions

SUPPLEMENTATION: omega-3 versus placebo

Group 1: omega-3 LCPUFA: 1 g/day*: total number randomised = 40

Group 2: placebo (olive oil): total number randomised = 40

Timing of supplementation: women in the omega-3 group were supplemented for 6 weeks**

DHA + EPA dose/day: unclear

*not further specified

**gestational ages not specified apart from being > 20 weeks' gestation

Outcomes

Women/birth: depression during pregnancy (Beck Depression Inventory)

Notes

Funding: not reported

Declarations of interest: none declared

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "permuted block randomisation"
Allocation concealment (selection bias)	Unclear risk	Quote: "permuted block randomisation"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Both mothers and researchers were blinded to drug and placebo"
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	No losses to follow-up reported; however women consuming fish more than twice a week were replaced by the next individual (number of instances not reported).
Selective reporting (reporting bias)	High risk	Only 1 outcome was reported.
Other bias	Unclear risk	Similar baseline characteristics except that all participants in placebo group were employed, compared with only 5% of participants in the intervention group.



Keenan 2014			
Methods	RCT: NCT01158976: Nu	trition and Pregnancy Study (NAPS)	
Participants	64 women randomised (2:1 ratio)		
	Inclusion criteria: pregnant women living in urban low-income environments, enrolled at 16-21 weeks' gestation and 'demographically eligible' (Medicaid insured or eligible, African American, and aged 20-30 years)		
	Exclusion criteria: 2 or more servings of sea fish per week, known medical complications (gestational diabetes, PE, subchorionic haematoma), regular use of steroid medications, regular alcohol use, cigarettes or use of illegal substances (by maternal report), use of blood thinners or anticoagulants, use of psychotropic medications, BMI > 40, allergy to iodine or soy		
		SA: University of Pittsburgh Medical Center obstetric clinics (from 2010-2012). in urban areas, and provide health services to women living in low-income	
Interventions	SUPPLEMENTATION:	DHA + EPA + DPA + ETA versus placebo	
	Group 1: omega-3 LCPUFA (2 gel capsules providing 450 mg DHA, 40 mg DPA and ETA, 90 mg EPA and 10 mg vitamin E): n = 43		
	Group 2: placebo capsules - matched in size, colour and smell to the study drug (900 mg soybean oil, 16.5 mg vitamin E, 10 mg EPA/DHA): n = 21		
	Timing of supplementation: 16-21 weeks GA to birth		
	To support adherence with the intervention, research assistants contacted participants by phone 3 times per week to ask the time of day that the supplement was taken, and gathered data on perception of taste and possible gastrointestinal side effects.		
	DHA + EPA dose/day: mid: 450 mg DHA + 90 mg EPA		
Outcomes	Women/birth: Perceived Stress Scale (self report) at 24 and 30 weeks' gestation; Trier Social Stress Test (cortisol response - saliva samples before and after test completion; baseline, 24 and 30 weeks' gestation); symptoms of depression (Edinburgh Postnatal Depression Scale (EPNS)) at 24 and 30 weeks; Difficult Life Circumstances Scale; length of gestation		
	Babies/infants/children: birthweight; 1-minute Apgar score; cortisol concentrations; BSID-III at 4 months; Face-to-Face Still Face at 4 months		
Notes	Funding: capsules provided by Nordic Naturals; supported by NIH grant. "This study was supported by NIH grants R21 HD058269 and RO1HD084586 to Dr Keenan, with additional support from the University of Chicago Institute for Translational Medicine (UL1TR000430). DHA supplement and placebo were supplied by Nordic Naturals".		
	Declarations of interest: none reported		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera-	Low risk	Computer-generated random assignment	
tion (selection bias)		Quote: "The pharmacist at the University of Pittsburgh used computer-generated random assignment of identification numbers to active supplement or placebo in blocks of nine"	
Allocation concealment (selection bias)	Low risk	Pharmacist (third party)	



Keenan 2014 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Placebo controlled
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Placebo capsules matched in size, colour and smell to the study drug; probably done
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	13/64 (20%) women were lost to follow-up: 21% in the omega-3 group and 19% in the placebo group. 2 participants in the placebo group withdrew due to miscarriage and mood changes; 2 also withdrew in the omega-3 group - 1 with headaches and 1 with an upset stomach.
		Data for 15/64 (23%) infants were not available at the 4 months postpartum assessment.
		End of pregnancy/birth mother and infant outcomes, including length of gestation and birthweight: at the 36-week gestation data collection point (last before birth), data were collected from 36/43 (84%) and 17/21 (81%) of mothers randomised to the intervention and control groups respectively. Number of mothers and infants from whom data were collected at the end of pregnancy/birth time point was not reported.
		3-month postpartum outcomes, including infant neurological/neurosensory and developmental outcomes and maternal stress: data were collected from 34/43 (79%) and 15/21 (71%) of infants born to mothers randomised to the intervention and control groups respectively.
Selective reporting (reporting bias)	Unclear risk	Not all expected outcomes were reported (e.g. preterm birth).
Other bias	Unclear risk	Only baseline comparisons for maternal mental health characteristics were reported (baseline groups wrong way round in Table 1 of 2014 paper).

Khalili 2016

Khalili 2016	
Methods	RCT: IRCT2013100914957N1
Participants	150 women randomised
	Inclusion criteria: women aged 18-35 years; 1st to 5th pregnancy, with a household health record in the participating health centres, ability to read and write, singleton pregnancy, stable phone access
	Exclusion criteria: bleeding during pregnancy, placenta praevia, abruption or cerclage in the present pregnancy, history of allergy to fish oil or other fish products, allergy to gelatin, history of previous underlying diseases such as heart disease, kidney disease, hyperlipidaemia, taking medication for 1 of these, consuming fish more than twice a week, smoking or drug addiction, bleeding disorders or taking anticoagulants, BMI > 30, participating in another interventional study
	Characteristics: very low baseline omega-3 concentrations
	Setting: 27 health care centres, Tabriz, Iran
Interventions	SUPPLEMENTATION: DHA + EPA versus placebo
	Group 1: 1000 mg fish oil supplements (120 mg DHA and 180 mg EPA). Total number randomised: n = 75



Khalili 2016 (Continued)	Group 2: placebo (similar shape, size and weight); liquid paraffin. Total number randomised: n = 75				
	Timing of supplementation: from 20 weeks GA to 30 days after birth (women were recruited between 16-20 weeks GA)				
	DHA + EPA dose/day: low: 120 mg DHA + 180 mg EPA				
Outcomes	Women/birth: maternal serum fatty acid profiles at 35-37 weeks GA; adherence; adverse events (nausea, unpleasant taste, vomiting diarrhoea, stomach pain); caesarean section				
	Babies/infants/children: neonatal death; perinatal death; birthweight; low birthweight; birth length; head circumference at hirth; growth at 4 and 6 months; ASO (2nd edition) at 4 and 6 months (scores				

read circumference at birth; growth at 4 and 6 months; ASQ (2nd edition) at 4 and 6 months (scores and thresholds)

Funding: Tabriz University of Medical Sciences. follow-up study supported by Tabriz Health Services

Management Research Centre (Grant No. 5/77/5241)

Declarations of interest: none declared

Risk of bias

Notes

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computerised random number table generator
Allocation concealment (selection bias)	Low risk	Quote: "randomly assigned by a staff member not involved in the research"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Placebo controlled (although smell or taste of paraffin may have been able to be detected)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not reported, but probably done.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	8/75 women in the omega-3 group and 7/75 did not have blood samples at 35-37 weeks GA (omega-3 group: 5 not interested in sampling; 2 preterm labour: placebo group: 6 not interested in sampling; 2 preterm labour); 1 women in the placebo group stopped capsules due to nausea. For other health outcomes, 0/75 in the omega-3 group and 4/75 in the placebo
Salastive reporting (re	Low risk	group were lost to follow-up.
Selective reporting (reporting bias)	LOW HSK	No apparent evidence of selective reporting (though stillbirth not explicitly reported).
Other bias	Low risk	Baseline characteristics appeared similar.

Knudsen 2006

Methods	RCT: parallel with 7 groups (6 treatment and 1 control group in 1:1:1:1:1:2 ratio)
Participants	3098 women randomised; a letter was mailed to women in the 6 treatment groups, with a 56% take-up rate overall (1291/2324)



Knudsen 2006	(Continued)
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Inclusion criteria: women at gestation week 17 to 27, with limited fish intake, no use of fish oil capsules in pregnancy; only liveborn singleton pregnancies were analysed

Exclusion criteria: none stated

Characteristics: about 86% women consumed some fish at baseline.

Setting: subgroup of the National Danish Birth Cohort, Denmark

Interventions

SUPPLEMENTATION: omega-3 (5 different doses DHA/EPA; ALA (flax oil)) versus no treatment

Group 1: 0.1 g/day EPA + DHA: total number randomised = 389 (234 participated: 229 analysed)

Group 2: 0.3 g/day EPA + DHA: n = 385 (231 participated: 224 analysed)

Group 3: 0.7 g/day EPA + DHA: n = 385 (228 participated: 222 analysed)

Group 4: 1.4 g/day EPA + DHA: n = 383 (223 participated: 212 analysed)

Group 5: 2.8 g/day EPA + DHA: n = 393 (195 participated: 187 analysed)

Group 6: 2.2 g/day ALA: n = 389 (180 participated: 176 analysed) – 4 x 1 g flax oil

Group 7: no treatment: (774 randomised:748 analysed) (not contacted at all)

Timing of supplementation: women were asked to stop taking capsules on the date of expected birth. On average, women stopped taking the capsules 12 days before giving birth; with a mean baseline supplementation commencement of 22-23 weeks' gestation (approximate estimate of 16 weeks mean supplementation length).

DHA + EPA dose/day: different doses - see above

Outcomes

Women/birth: GA at birth

Notes

Unclear how recruitment by mail may have influenced results.

Funding: Danish National Research Foundation, Pharmacy Foundation, Egmont Foundation, Augustinus Foundation, Health Foundation, March of Dimes Birth Defects Foundation, EU, Heart Foundation. Dansk Droge and Biooriginal Food Science Corp donated the fish oil and flax oil capsules respectively

Declarations of interest: not reported

Bias	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Quote: "block-wise stratified randomisation"; no further details reported.
Allocation concealment (selection bias)	Unclear risk	Quote: "block-wise stratified randomisation"; no further details reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Not clearly stated, but we assume that women in the 6 treatment groups did not know the dosage (or type) of omega-3 they were taking.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not explicitly reported; "date of birth was extracted from the Danish Civil Registration System".



Knudsen 2006 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	High risk	Between 40% and 54% of women failed to participate in the 6 treatment groups.
Selective reporting (reporting bias)	High risk	Only gestational length reported.
Other bias	Low risk	Similar baseline characteristics

Methods	RCT: NCT01180933 (NUHEAL Nutraceuticals for a Healthier Life); 2 x 2 factorial				
Participants	315 women randomised (4 groups - see below)				
	Inclusion criteria: apparently healthy women < 20 weeks GA; singleton pregnancy, intention to give birth in 1 of the obstetric centres listed below; body weight from > 50 kg to 92 kg and > 18–41 years old				
	Exclusion criteria: women with serious chronic illness (e.g. diabetes, hepatitis, or chronic enteric disease) or who used fish oil supplements since the beginning of pregnancy or folate or vitamin B-12 supplements after gestation week 16				
	Setting: Departments of Obstetrics at Ludwig Maximilians University, Munich, Germany; the University of Granada, Granada, Spain; and the University of Pecs, Pecs, Hungary				
Interventions	SUPPLEMENTATION + OTHER AGENTS: DHA + EPA versus DHA + EPA + folate versus folate versus placebo - all in a milk base				
	Group 1: DHA/EPA: 15 g milk-based supplement with 500 mg DHA and 150 mg EPA daily: total number randomised: $n = 77$				
	Group 2: DHA/EPA/folate:15 g milk-based supplement with 500 mg DHA and 150 mg EPA, 400 μ g 5-MTHF daily: total number randomised: n = 77				
	Group 3: folate:15 g of a milk-based supplement with 400 μ g 5-MTHF daily: total number randomised: = 80				
	Group 4: control: 15 g of a milk-based supplement placebo: total number randomised: n = 81				
	Timing of supplementation: 20 weeks GA to birth (infant formula (see below) until 6 months of age if child not breastfed)				
	All women: all sachets contained vitamins and minerals in amounts that met the recommended intakes during the second half of pregnancy for European women (minerals: 300 mg Ca, 240 mg P, 93 mg Mg, 3 mg Zn, 66 μg I; vitamins: 330 μg vitamin A, 1.5 μg vitamin D, 3 mg vitamin E, 0.36 mg thiamine, 1.5 mg riboflavin, 4.5 mg vitamin B-3, 1.9 mg vitamin B-6, 3.5 μg vitamin B-12, 270 mg vitamin C).				
	All women were encouraged to breastfeed. For infants who were not fully breastfed: if the newborns were born to fish oil-supplemented women, they received an infant formula containing 0.5% of total fatty acids as DHA and 0.4% as AA, whereas children born to mothers in the placebo or 5-MTHF groups received a formula virtually free of DHA and AA during the first 6 months of postnatal life.				
	DHA + EPA dose/day: mid: 500 mg DHA + 150 mg EPA				

weeks 20 and 30 and at birth; indicators of pregnancy outcome, and fetal development; placental samples; proliferation cell nuclear antigen; mRNA expression of placental proteins; TLR2, TLR4 and CD14 mRNA (maternal blood, placenta, cord blood); GWG, mode of birth; preterm birth < 35 weeks; postnatal depression 8 weeks after birth (EPDS); proteinuria, BP, and eclampsia for gestation weeks 22 and 30



Krauss-Etschmann 2007 (Continued)

and at birth; blood loss at birth; birthweight, birth length; head circumference at birth (the previous 3 measurements reported for only a sample of babies); MTHFR C677T polymorphism; fatty acids.

Babies/infants/children: Apgar score; umbilical pH; visually evoked potentials at 8 weeks (Germany and Spain); Bayley Mental Development Test at 6 months old (Spain); skin-prick test at 6 months old (Spain); paediatric symptoms and illness questionnaire at birth, 8 and 24 weeks; Hempel examination at 4 years; Touwen examination at 5.5 years; minor neurological dysfunction, NOS; fluency score; cognitive development (Kaufman Assessment Battery) at 6.5 years; response conflict-resolution ability; alerting, and spatial orienting of attention (Attention Network Test), ERPs, and sLORETA at 8.5 years; MTHFR C677T polymorphism

Notes

DHA and EPA provided by Pronova Biocare, Lysaker, Norway: Folate from BASF, Ludwigshafen, Germany

Adherence: 89.5% of the women in the second trimester gestation and 87.4% in the 3rd trimester missed < 5 days of supplementation

Funding: Commission of the European Research and Technological Development Programme "Quality of Life and Management of Living Resources" within the 5th Framework Programme (contract QLK1-CT-1999-00888 (NUHEAL "Nutraceuticals for a Healthier Life")) and the European Community's 7th Framework Programme (FP7/2008-2013) under grant agreement 212652 (NUTRIMENTHE Project "The Effect of Diet on the Mental Performance of Children"); the University Science Program of Ludwig Maximilians University; a Freedom to Discover Award from the Bristol Myers Squibb Foundation; Spanish Ministry of Economy and Competitiveness grant (State Secretariat for Research, Development, and Innovation; PSI2012-39292); European Research Council advanced grant ERC-2012-AdG (no.322605 META-GROWTH).

Declarations of interest: "Disclosure of potential conflict of interest: The authors have received grant support from the Commission of the European Communities and the Danone Institute of Nutrition". No other potential or actual conflicts of interest declared.

Bias	Authors' judgement	Support for judgement			
Random sequence generation (selection bias)	Unclear risk	Quote: "blockwise randomization"			
Allocation concealment (selection bias)	Unclear risk	Quote: "envelopes containing cards with 1 of 4 numbers (1, 2, 3, or 4) according to the 4 intervention groups were mixed and put into a closed box. By drawing envelopes, intervention group numbers were consecutively assigned to subject identity numbers."			
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Neither the participating women nor the study personnel knew the content of the sachets"			
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not specifically reported, but probably done.			
Incomplete outcome data (attrition bias) All outcomes	High risk	45/315 (14.3%) did not complete the study at first follow-up:			
		Group 1: DHA/EPA 8/77 (10.4%)			
		Group 2: DHA/EPA + folate 13/77 (16.9%)			
		Group 3: folate 15/80 (18.8%)			
		Group 4: placebo 9/81 (11.1%)			



Krauss-Etschmann 2007 (Continued)

There were 4 post randomisation exclusions: 2 women weighed > 92 kg, 1 of whom used commercial fish oil preparations; and 2 women regularly consumed fish oil preparations. Reasons for dropping out (n = 41) were noncompliance (n = 2), relocation (n = 1), aversion to or bad taste of the supplement (n = 9), and loss of contact (n = 2). In the remaining 17^* cases, the reasons for dropping out were not known. Reasons were not reported by group and there were differential rates of loss between groups.

*could be 27

Later follow-up (4 and 5.5 years)

270 mother-infant pairs were invited for neurological assessment; 175 complied with the request at the age of 4 years and 157 complied at 5.5 years of age.

Dropout rates were a further 35.18% at the age of 4 years and 41.9% at the age of 5.5 years, with no differences in the dropout rates between groups.

Main reasons for dropping out were relocation (n = 3), loss of contact (n = 65, n = 76), and unwillingness to continue in the study (n = 27, n = 34). 4 of the children examined at the age of 4 years and 5 of those examined at 5.5 years were born prematurely before week 35 of pregnancy and were therefore excluded from the analyses. Except for 1 child who was born with a congenital left side anophthalmus, no other serious congenital disorder was observed. In the health screening questionnaire at 4 years of age, 1 child was reported to have left side deafness, another had developed craniosynostosis and had surgery at the age of 6 months, and 1 child suffered from a developmental retardation of unknown etiology. These children were also excluded from the analyses, which left 167 and 148 children at the age of 4 and 5.5 years, respectively.

6.5 year follow-up

161 children participated (exclusions: 4 children born < 35 weeks, 1 child was born with a congenital left-side anophthalmus, 1 child developed craniosynostosis, and another was reported to have left-side deafness).

Dropout rates were similar between groups. Main reasons for dropping out were relocation (n = 3), loss of contact (n = 74), and unwillingness to continue (n = 30).

There was a higher attrition of children whose fathers had a high educational level in the placebo and FO + 5-MTHF groups, as well as differences in several outcomes (e.g. length of gestation, birth length) between children followed up and those lost to follow-up.

8.5 year follow-up

130 children participated; 37 FO, 27 folate; 32 placebo and 34 FO/folate (32 from Germany, 96 from Spain and 8 from Hungary; (exclusions: 4 children born < 35 weeks, 1 child was born with a congenital left-side anophthalmus, 1 child developed craniosynostosis, and another was reported to have left-side deafness). Dropout rates were similar between groups and had similar sociodemographic profiles. Main reasons for dropping out were relocation (n = 7), loss of contact (n = 75), and unwillingness to continue (n = 45).

Selective reporting (reporting bias)	Unclear risk	Preterm births treated as exclusions and not reported by group.
Other bias	Low risk	Similar baseline characteristics



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Methods	RCT: NCT00865683: The Omega-3 pregnancy study				
Participants	91 women randomised				
	Inclusion criteria: women with prepregnancy BMI \geq 25 kg/m ² , \sim 26 weeks' gestation, 18-40 years of age, and a singleton pregnancy (stated as BMI \geq 30 kg/m ² in Foster 2017).				
	Exclusion criteria: diseases affecting study outcomes (e.g. gestational or other diabetes mellitus, hypertension, or concurrent inflammatory, vascular or metabolic disease); high unusual intake of DHA (more than 1 fish meal per week, use of DHA-fortified foods or supplements); current or previous use of tobacco, illicit drugs, or medications such as corticosteroids that affect inflammatory markers; inability to travel to the research centre for study visits.				
	Setting: General Clinical Research Center, Cincinnati, Ohio, USA (December 2009 to June 2013)				
Interventions	SUPPLEMENTATION: DHA versus placebo				
	Group 1: DHA (800 mg/day) for 10 weeks, in capsule form; number randomised not reported				
	Group 2: placebo: corn/soy oil daily for 10 weeks, in capsule form: number randomised not reported				
	Timing of supplementation: from 26 to 36 weeks' gestation				
	DHA + EPA dose/day: mid: 800 mg DHA; EPA not stated				
Outcomes	Women/birth: maternal adverse effects; gestational diabetes; birthweight, length and adiposity (BMI Z score); length of gestation; DHA concentrations (erythrocyte and placenta); fasting blood glucose; cytokines; metabolic hormones; lipids				
	Babies/infants/children: child growth (including adiposity, weight, height, arm circumference and arm skinfold); child neurological/neurosensory and developmental outcomes (as measured by the Bayley Scales of Infant and Toddler Development)				
Notes	Funding: "Research reported in this publication was supported by National Institutes of Health grant R21HL093532, 8UL1TR000149 and the Mike Hogg Fund (T.L.P). The two-year follow-up was supported by the Rita and William Head endowment for studies on environmental influences on Prematurity (R.R.). B.A.F.was supported by a grant from the National Institute of Diabetes and Digestive and Kidney Diseases of the National Institutes of Health under Award Number K23DK109199."				
	Declarations of interest: none declared				
Risk of bias					

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The randomization scheme was prepared prior to start of the trial by the study pharmacist using a tested system utilizing a random-number generator. Each study subject was assigned to study group on a consecutive basis"
Allocation concealment (selection bias)	Low risk	Pharmacy-controlled randomisation
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quotes: "Study staff, subjects, and investigators were blinded to the group assignment. At the end of the study, pharmacy staff mailed the study group assignment to the subject and investigators"; and "the gelcaps were identical in size, appearance, taste, and smell (orange flavoured)"
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quotes: "Study staff, subjects, and investigators were blinded to the group assignment. At the end of the study, pharmacy staff mailed the study group as-



Krummel 2016 (Continued)		signment to the subject and investigators"; and "the gelcaps were identical in size, appearance, taste, and smell (orange flavoured)"
Incomplete outcome data (attrition bias) All outcomes	High risk	The 2 papers reported different numbers of participants remaining in the trial at the 36-week and birth data collection points (end of main study). Of the 91 women, initially randomised, 31 (34%) or 28 (30%) were not included the analysis. It was unclear which groups the excluded women had been randomised to, and therefore it was not possible to assess potential systematic differences between groups in withdrawals from the study confidently. No further losses to follow-up (in the follow-up study) were reported.
Selective reporting (reporting bias)	Unclear risk	Some mismatches with the outcomes reported in the trial registration entry; outcomes such as GDM and BSID III incompletely reported in results; unclear whether some standard errors were SDs
Other bias	Unclear risk	Higher BMIs and body weight in the placebo group

Laivuori 1993

Methods	RCT (3-arms)		
Participants	18 women randomised		
	Inclusion criteria: women with PE (admitted to hospital between 26 and 37 weeks' gestation because of BP consistently exceeding 140/100 mmHg; 8 women also had proteinuria)		
	Exclusion criteria: none reported		
	Setting: Helsinki, Finland		
Interventions	SUPPLEMENTATION: omega-3 versus LA + GLA (Primrose oil) versus placebo		
	Group 1: fish oil: 10 MaxEPA capsules (180 mg EPA, 120 mg DHA per capsule): total number randomised: n = 5 (3)		
	Group 2: primrose oil: 10 Preglandin capsules (375 mg linolenic acid and 45 mg gammalinolenic acid per capsule): total number randomised: n = 7 (4)		
	Group 3: placebo (10 capsules each containing 500 mg of maize oil and 500 mg olive oil); total number randomised: $n = 6$ (5)		
	All women: intervention between 31-36 weeks' gestation; bed rest in hospital for 2 days before randomisation		
	Median duration and range of supplementation was:		
	• fish oil: 32 days (13-54)		
	 evening primrose oil: 17 days (7-28) 		
	• placebo: 24 days (14-39)		
	Women were advised to follow their normal diets.		
	DHA + EPA dose/day: daily dose unclear		
Outcomes	Women/birth: prostanoids (urine); clinical signs of PE; birthweight (reported as median and range); BF (reported as % of pretreatment levels)		
Notes	2 women used betablockers (metoprolol 100 mg/day) and 1 used dihydrazaline 50 mg/day. No aspirin-like drugs were allowed.		



Laivuori 1993 (Continued)

No outcomes could be meta-analysed

Funding: Preglandin capsules (Suomen Rohdos, Turku, Finland); MaxEpa and placebo capsules (Orion OY, Kuopio, Finland)

Declarations of interest: not reported

Risk of bid	as
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Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Reported as "in randomized order"
Allocation concealment (selection bias)	Unclear risk	Sealed envelopes
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Capsules of identical appearance
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not reported, but probably done.
Incomplete outcome data (attrition bias) All outcomes	High risk	6/18 (33%) women excluded: 3 women gave birth before samples could be collected and 3 additional women failed to collect adequate urine samples.
Selective reporting (reporting bias)	Unclear risk	Limited number of outcomes reported.
Other bias	High risk	Median length of supplementation differed substantially between the 3 groups.

Makrides 2010

Methods

RCT: ACTRN012605000569606 (DOMInO main trial); allergy follow-up: ACTRN12610000735055; 3- and 5-year follow-up: ACTRN12611001127998; 4-year follow-up: ACTRN12611001125910; 7-year follow-up: ACTRN12614000770662

Participants

2399 women randomised

Inclusion criteria: singleton pregnancy < 21 weeks GA, no known fetal abnormality, and not taking medication where tuna oil was contraindicated

Exclusion criteria: women already taking a DHA supplement, with a bleeding disorder in which tuna oil was contraindicated, taking anticoagulant therapy, had a documented history of drug or alcohol abuse, were participating in another fatty acid trial, fetus had a known major abnormality, or were unable to give written informed consent or if English was not the main language spoken at home

Setting: 5 Australian perinatal centres, recruiting from October 2005 to January 2008

Pregnant women were approached to enter the allergy follow-up, Palmer 2012, after randomisation into the DOMInO trial. Only Adelaide-based women were eligible for the allergy follow-up. Women were eligible if the unborn baby had a mother, father, or sibling with a history of any medically diagnosed allergic disease (asthma, allergic rhinitis, eczema) (n = 706).



Makrides 2010 (Continued)

Interventions

SUPPLEMENTATION: DHA + EPA versus placebo

Group 1: DHA-rich fish oil capsules: 3 x 500 mg/day (providing 800 mg DHA + 100 mg EPA per day): n = 1197

Group 2: matching placebo vegetable oil blend of 3 non-genetically modified oils (rapeseed, sunflower, palm) in equal proportions: n = 1202

Timing of supplementation: from trial entry (~20 weeks) to birth

DHA + EPA dose/day: mid: 800 mg DHA + 100 mg EPA

Outcomes

Women/birth: length of gestation; adherence (28 weeks); PPH; log blood loss at birth; preterm birth < 37 weeks; early preterm birth < 34 weeks; PIH; PE; PE (clinical diagnosis in medical records); caesarean section; post-term induction or post-term prelabour caesarean; post-term induction; serious morbidity composite; renal failure; liver failure; death; eructations (28 and 36 weeks); diarrhoea; gestational diabetes (based on glucose tolerance test); gestational diabetes (based on clinical diagnosis in medical record); postnatal depression (all women: EPDS > 12 at 6 weeks, 6 months postpartum); postnatal depression (women with previous or current depression at trial entry: EPDS > 12 at 6 weeks, 6 months postpartum); new diagnosis of depression in study period; new or existing diagnosis of depression during study period; antenatal admission to hospital; maternal admission to intensive care unit (level III antenatal hospitalisation); vitamin D concentrations (cord blood)

Babies/infants/children: perinatal death; birthweight; birth length; head circumference at birth; birthweight z score; birth length z score; head circumference at birth z score; low birthweight < 2.5 kg); SGA (weight < 10th percentile), length, head circumference); LGA (weight (> 90th percentile), length, head circumference); birthweight > 4 kg; any serious adverse event; IVH (and grade); NEC; sepsis; convulsion; BPD (oxygen required for treatment of chronic lung disease); neonatal hypoglycaemia; resuscitation at birth; bone fracture; NICU admission; breastfeeding; morbidities up to 5 years; BSID at 18 months in 600 randomly selected infants; visual development outcomes at 4 months; attention and working memory and inhibitory control at 27 months; general cognitive function (DAS II), executive function, language, behaviour at 4 years; BMI z-scores; body fat; BP; and insulin sensitivity at 3 and 5 years (HOMA-IR); BMI z-score and percentage body fat at 3 and 5 years of age

(Allergy outcomes are reported in another Cochrane review (Gunaratne 2015)).

Notes

Funding: supported by grants from the Australian National Health and Medical Research Council and Australian Egg Corporation Limited. Treatment and placebo capsules were donated by Efamol, UK and Croda Chemicals.

Declarations of interest: Professor Makrides reported serving on scientific advisory boards for Nestle, Fonterra, True Origins and Nutricia. Professor Gibson reported serving on scientific advisory boards for Nestle and Fonterra. Associated honoraria for Professors Makrides and Gibson were paid to their institutions to support conference travel and continuing education for postgraduate students and early career researchers. Dr Makrides reported receiving non financial support from Clover Corporation and Nestle Nutrition. Dr Makrides, through the Women's & Children's Health Research Institute, has a patent pending: "Methods and compositions for promoting the neurological development of an infant". Dr Gould reports honoraria paid to her institution from the Nestle Nutrition Institute. Dr John Colombo served on the advisory boards for Nestle, Fonterra and Nutricia. Dr Muhlhausler had given lectures on maternal nutrition for Aspen Nutrition and Danone Nutritia. Linda Tapsell served on the Science Advisory Committees of the California Walnut Commission and the McCormicks Science Institute. Dr Palmer had consulted for and received payment for lectures from Nestle Nutrition. Dr Prescott reported receiving honorariums from the Nestle Nutrition Institute and Fonterra and being paid for lectures by the World Allergy Association and the American Academy of Asthma, Allergy and Immunology. Dr Heddle reported being paid for expert testimony from Analysis Plus and Rodika Research Services, and receiving grants from Commonwealth Serum Laboratories, Vaxine, GLaxoSmithKline, and Healthed. Dr Beverley Muhlhausler had given lectures on maternal nutrition for Aspen Nutrition and Danone Nutricia. No other author declarations.



Makrides 2010 (Continued)

Pine	Authoral independent	Cumpant for independent
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Women were randomly assigned a unique study number and treat- ment group allocation through a computer-driven telephone randomization service according to an independently generated randomization schedule"
Allocation concealment (selection bias)	Low risk	As above
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "All capsules were similar in size, shape and color", and, "trial investigators, staff and participants were unaware of the treatment allocation"; however more women in the treatment group guessed that they were taking fish oil capsules (67% in the treatment group compared with 13% in the control group)
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Described as "double blind"; Zhou 2012* describes a "blinded audit of medical records"; Smithers 2011* describes "a blinded assessment of a subset of healthy full-term infants; Makrides 2014* specifies "with a psychologist who was blinded to group allocation"
		*These references are listed under Makrides 2010.
Incomplete outcome data (attrition bias) All outcomes	Low risk	1179/1197 (98%) women in the omega-3 group and 1166/1202 (97%) in the placebo group completed 6-month postpartum follow-up (though all women were included in the primary analyses): "adequate data for the analysis of the primary outcome were available for 2320 women (97.3% in the DHA group and 96.1% in the control group)".
		• 18 month follow-up: 726 infants; 333/351 (95%) of infants in the omega-3 group and 361/375 (96%) in the placebo group completed the 18-month BSID III (though all infants were included in analyses: "694 children (95.6% of those selected for follow-up) were assessed at 18 months".
		• 4-month visual acuity tests: 185 children were enrolled (91 omega-3 and 94 control); 89/91 (98%) and 93/94 (99%) were included in analyses.
		• 27-month follow-up: 184 children were eligible (1 had died): rates of loss to follow-up were 8.8% in the omega-3 group and 18.1% in the placebo group: "The follow-up sample was comparable to that in the overall DOMInO trial, with the exception of the proportion of mothers in the follow-up group who had completed tertiary education and compliance was slightly higher in the follow-up group".
		• 3-year follow-up: of the 1531 children (95% of those eligible) participating, there were BMI z-scores for 96% and body fat measurements for 83%.
		• 4-year follow-up: of the 726 children selected for 18 month follow-up, 703 were eligible for 4 year follow-up and 646 (92%) were included in the analyses.
		• 5-year follow-up: of the 1531 children (95% of those eligible) participating, there were BMI z-scores for 88% and body fat measurements for 73%.
		 7-year follow-up: 232 children (113 omega-3; 119 control completed 7-year follow-up for the outcomes of body fat (Wood 2017).
Selective reporting (reporting bias)	Low risk	No indication of selective reporting bias.
Other bias	Low risk	The demographic and clinical characteristics of the women at randomisation were comparable between the 2 groups, though some divergence was seen over time in the follow-ups, particularly with regard to sociodemographic characteristics.



Malcolm 2003

Methods	RCT			
Participants	100 women randomised			
	Inclusion criteria: women who were expected to deliver their infants at term and planned to feed them on breast and/or formula milk			
	Exclusion criteria: women with diabetes, twin pregnancies, PE/toxaemia, a past history of abruption or PPH, allergy to fish products, a thrombophilic tendency, or who were receiving drugs that affect thrombocyte function; pregnancies concluded prematurely before 36 weeks, in which the neonate had an Apgar score < 7 at 5 minutes, had weight below the 3rd centile for GA, or had medical or developmental problems were not included in the final analysis of results.			
	Setting: antenatal clin	Setting: antenatal clinic (elective); Yorkhill NHS Trust, Stirling, Scotland		
Interventions	SUPPLEMENTATION:	DHA versus placebo		
	Group 1: fish oil; blend 50 randomised	led, Marinol D40 (200 mg DHA/day, 100 mg per capsule); 2 capsules per day; n =		
	Group 2: placebo capsules containing high-oleic acid sunflower oil, with 810 mg oleic acid/g (200 mg per capsule), devoid of any omega-3 fatty acid; 2 capsules per day; n = 50 randomised			
	Timing of supplementation: 15 weeks GA to birth			
	All women: advised to follow their normal diet during pregnancy			
	DHA + EPA dose/day: low: 200 mg DHA			
Outcomes	Women/birth: DHA and other fatty acid status (RBC and plasma); length of gestation; birthweight; length at birth; head circumference at birth; preterm birth			
	Babies/infants/children: longer-term growth at 50 and 66 weeks post-conceptional age; visual development; DHA and other fatty acid status (cord blood)			
Notes	Funding: University of Glasgow and the Chief Scientist Office, Scotland. RP Scherer Ltd UK donate capsules.			
	Declarations of interest: none declared			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Unclear risk	Quotes: "double blind, prospective, randomised, and controlled in design"; "The women were then randomised"		
Allocation concealment (selection bias)	Unclear risk	As above, no further detail provided.		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The study supplements were identical in appearance and could not be identified on the basis of scent or taste" and "The capsules were identical in appearance, taste and odour."		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not specifically detailed		



Malcolm 2003 (Continued)

Incomplete outcome data
(attrition bias)
All outcomes

High risk

Of 100 women recruited to the study, 29 (29%) (15 in DHA group; 14 in placebo group) withdrew before 28 weeks, and a further 7 (4 in DHA group; 3 in placebo group) withdrew before birth (total n=36). Common reasons for withdrawing were poor adherence (n=16); frequent nausea/vomiting (n=13); loss of contact (n=3); anxiousness (n=2); unspecified (n=3).

A further 3 infants (all from the placebo group) were excluded from the postnatal portion of the study after birth: born before 36 weeks (n = 1); Apgar score < 7 at 5 minutes (n = 2).

A further infant was discharged before testing. Therefore birth outcomes were available for 60/100 women; birth vision tests were performed on 59 infants. ERG studies (retina) were performed on 56 infants.

(Other paper reports 2 excluded from placebo group due to prematurity, and SGA) $\,$

Selective reporting (re- U porting bias)
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Unclear risk

Unclear; no access to trial protocol

Other bias Unclear risk

Insufficient detail to determine risk of other bias

Mardones 2008

Methods	RCT			
Participants	1173 women			
	Inclusion criteria: women age 18 years and over, parity 0-5, up to 20 weeks' gestation (confirmed by US), non-consumers of drugs and alcohol, and underweight (BMI ≤ 21.2 at 10 weeks' gestation, as defined by Chilean charts for pregnant women)			
	Exclusion criteria: multiple pregnancies; suffering from chronic diseases that could affect fetal growth; smokers; and disease diagnosed during pregnancy			
	Setting: 19 urban health clinics belonging to the Servico de Salud Metropolitano Sur-Oriente (Southeast Metropolitan Public Health Services), Santiago, Chile (study dates not reported).			
	Mainly low-income, ethnically diverse families (Ameri-Indian and Hispanic).			
Interventions	SUPPLEMENTATION + OTHER AGENT: omega-3 + omega-6 + multiple micronutrients + milk versus milk only			
	Group 1: milk product fortified with omega-3 LCPUFA (0.6g/day) omega- 6LCPUFA (3g/day), multiple micronutrients: total number randomised: n = $589 $			
	Group 2: regular powdered milk; total number randomised: n = 552			
	Timing of supplementation: < 20 weeks to birth (presumed)			
	DHA + EPA dose/day: mid: 600 mg DHA; EPA not stated			
Outcomes	Women/birth: preterm birth (< 37 weeks); preterm birth (< 34 weeks); miscarriage; PE; length of gestation; caesarean section; GWG; stillbirth; neonatal death; perinatal death			
	Babies/infants/children: birthweight; infant growth (crown-heel length, head circumference)			
Notes	Funding: Parmalat SpA, Italy, the company that provided the Maman (intervention) product.			



Mardones 2008 (Continued)

Declarations of interest: none declared

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Quote: "Midwives in charge assigned the women in their initial pregnancy visit using the order of arrival: odd numbers to the experimental group and even numbers to the control group".
Allocation concealment (selection bias)	High risk	Quote: "Midwives in charge assigned the women in their initial pregnancy visit using the order of arrival: odd numbers to the experimental group and even numbers to the control group".
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Authors reported that: "the study could not be blinded because Chilean regulations do not allow delivery of food without information on its composition". However, lack of blinding of participants and personnel is unlikely to have introduced bias due to the objective nature of the outcomes measured and reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "For those co-authors who performed the calculations, groups were simply labelled 1 or 2, and the coding was not known by them either. These measurements should suffice to minimise the possible effects of non-blinding". No information was provided on blinding of the individuals who performed the assessments.
Incomplete outcome data (attrition bias) All outcomes	High risk	There were 32/1173 post randomisation exclusions. Data were available for 333/552 (60%) and 365/589 (62%) of women assigned to the control and intervention group respectively.
Selective reporting (reporting bias)	Unclear risk	insufficient information to permit confident assessment.
Other bias	Unclear risk	Among women who discontinued the trial, GA at recruitment was lower in the intervention group.

Martin-Alvarez 2012

Methods	RCT		
Participants	60 women		
	Inclusion criteria: not reported		
	Exclusion criteria: not reported		
	Setting: Virgen de las Nieves Hospital, Granada, Spain		
Interventions	SUPPLEMENTATION + FOOD: DHA versus placebo (in the form of a dairy product)		
	Group 1: dairy product (2 glasses a day) supplemented with DHA (400 mg/day): total number randomised: $n=30$		
	Group 2: dairy product (2 glasses a day) with no DHA supplement: total number randomised: n = 30		
	Timing of supplementation: from 28 weeks' gestation to when breastfeeding stopped		
	DHA + EPA dose/day: low: 400 mg DHA		



Martin-Alvarez 2012 (Continued)

Outcomes Women: plasma antioxidant capacity; adipokines

Babies/infants/children: antioxidant capacity (umbilical cord; newborn); peroxides; superoxide dis-

mutase activity; adipokines (birth; 2.5 months)

Notes Abstracts only available

No outcomes could be included in this review

Funding: not reported

Declarations of interest: not reported

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Quote: "double-blind"; not further described
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Not reported
Selective reporting (reporting bias)	Unclear risk	Insufficient detail
Other bias	Unclear risk	Insufficient detail

Miller 2016

Methods	RCT: NCT02219399
Participants	115 women randomised
	Inclusion criteria: women aged 18-42 years, singleton pregnancies and willingness to breastfeed exclusively for first 3 months
	Exclusion criteria: maternal age < 18 years, multiple pregnancies, diabetes, HIV-positive, chronic illnesses or other conditions which could preclude breastfeeding, and any known allergies to seafood or fish oils
	Setting: private practice gynaecology and obstetrics clinics, Fort Collins, Colorado, USA
Interventions	SUPPLEMENTATION: omega-3 (DHA + EPA) versus placebo



Miller 2016 (Continued)	Group 1: tupa fish oil <i>t</i>	300 mg DHA and 67 mg EPA); hard capsule: n = 60		
		ntical Sunola hard capsule (high oleic acid sunflower oil placebo): n = 55		
		tation: from the last trimester of pregnancy through to the first 3 months of		
	breastfeeding			
	DHA + EPA dose/day: \	low: 300 mg DHA + 67 mg EPA		
Outcomes	Women/birth: maternal FFQ; lipids (blood, breastmilk); Home Screening Questionnaire at 9 months; abbreviated Wechsler Adult Intelligence Scale (at baseline); gestational length (last menstrual period method); preterm birth < 37 weeks; later term birth > 40 weeks; caesarean birth; breastfeeding			
	Babies/infants/children: BSID-III, MDI at 4 months and 12 months (cognitive, language, social-emotional and general adaptive behaviour scales)			
Notes	Same NCT # allocated	to Harris 2015 (but author has confirmed this is a separate trial)		
	Funding: not reported			
	Declarations of intere	est: not reported		
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Low risk	Computer-generated block randomisation		
Allocation concealment (selection bias)	Unclear risk	Method of allocation not reported		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Participating women, all data collectors and investigators were blinded to supplement allocation until all study children were 12 months of age and had completed the cognitive testing. After all study data was collected, the study was un-blinded only to study investigators for analysis".		
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Participating women, all data collectors and investigators were blinded to supplement allocation until all study children were 12 months of age and had completed the cognitive testing. After all study data was collected, the study was un-blinded only to study investigators for analysis".		
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	In omega-3 group 12/60 (20%) loss to follow-up for BSID: • 1 lost contact at birth • 3 discontinued breastfeeding • 2 withdrew • 2 lost contact at 2 months postpartum • 1 discontinued breastfeeding at 4 months postpartum • 1 withdrew • 2 lost contact at 12 months postpartum		
		In placebo group 20/55 (36%) lost to follow-up:		
		 2 lost contact at birth 10 discontinued breastfeeding 3 withdrew 3 lost contact at 2 months postpartum 2 lost contact at 4 months postpartum 		



Miller 2016 (Continued)		
Selective reporting (reporting bias)	Unclear risk	Secondary outcomes (including infant birth length, infant growth velocity, infant birthweight or infant head circumference at birth) prespecified in the trial registration were not reported.
Other bias	Unclear risk	Baseline characteristics similar; however all women were allowed to take voluntary fish oil supplements (63% in the DHA group and 71% in the placebo group did so). Additionally women in the placebo group took higher amounts than the DHA group.

Min 2014

Methods	RCT: ISRCTN68997518: FOSIP				
Participants	173 women randomised (88 healthy women and 85 women with pre-existing type 2 diabetes)				
	Inclusion criteria: women 17–45 years old with singleton pregnancies with either pre-existing type 2 diabetes; or without any known medical condition (uncomplicated pregnancy group).				
	Exclusion criteria: multiple pregnancy; known major fetal anomaly; current or planned corticosteroid therapy; asthma requiring medication; current or planned beta-adrenergic therapy; chronic medical conditions such as HIV/AIDS, kidney disease, or congenital heart disease; haematologic or autoimmune disease such as sickle cell disease, other haemoglobinopathies, lupus, or antiphospholipid syndrome; previous or planned tocolytic therapy to induce labour or increase contraction strength				
	Characteristics: high proportion of South Asian and African/Caribbean women				
	Setting: antenatal clinic, Newham University Hospital, London, UK (women were recruited during thei first visit to the antenatal clinic between January 2008 and December 2011)				
Interventions	SUPPLEMENTATION: DHA + EPA versus placebo				
	Group 1: fish oil: 2 capsules per day (600 mg DHA): n = 86 (41 women with type 2 diabetes; 45 healthy women)				
	Group 2: placebo: 2 capsules per day; 82.6% oleic acid (sunflower oil): n = 87 (47 women with type 2 diabetes; 40 healthy women)				
	Timing of supplementation: recruited between 10-12 weeks' gestation, with supplementation continuing until birth				
	All women: both supplements contained vitamin E				
	DHA + EPA dose/day: mid: 600 mg DHA; EPA not stated				
Outcomes	Women/birth: caesarean section; miscarriage (< 24 weeks); GA (reported only as median and range) Babies/infants/children: preterm birth < 37 weeks; preterm birth < 34 weeks; shoulder dystocia; still-birth; birthweight; low birthweight (< 2500 g); anthropometric measures at birth: head circumference; length; femur length; humerus length; biparietal diameter; occipito-frontal diameter; arm and thigh lean and fat mass; abdominal circumference; abdominal fat mass; shoulder circumference, mid-arm circumference				
Notes	Funding: FP6 Marie Curie Actions-Transfer of Knowledge, The Foyle Foundation, Newham University Hospital NHS Trust, Diabetes Research Network (North East London Diabetes Local Research Network) Equazen/Vifor Pharma Ltd., London Metropolitan University, The Letten Foundation, The Mother and Child Foundation, Sir Hally Stewart Trust, Emeritus Professor Clara Lowy				
	Declarations of interest; none declared				



Min 2014 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was carried out using a random code generated by the supplement provider.
Allocation concealment (selection bias)	Low risk	Randomisation was carried out using a random code generated by the supplement provider (third party).
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Participants, midwives and all investigators were blinded to allocation until all the analysis was completed and the data recorded.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Participants, midwives and all investigators were blinded to allocation until all the analysis was completed and the data recorded; probably done.
Incomplete outcome data (attrition bias)	High risk	30% (26/86) in the fish oil arm and 35% (30/87) in the placebo arm were excluded or lost to follow-up at birth.
All outcomes		Diabetic women:
		Fish oil: 13/41 (32%); 4 dropouts, 7 miscarriages, 2 stillbirths
		Placebo: 17/47 (36%); 11 dropouts, 6 miscarriages
		Healthy women
		Fish oil: 13/45 (29%); 6 dropouts, 2 moved, 1 termination, 4 miscarriages, 3 developed GDM
		Placebo: 13/40 (33%); 8 dropouts, 2 moved, 3 miscarriages
Selective reporting (re-	Unclear risk	Some protocol changes after trial registration
porting bias)		Under the same registry number as FOSIP ((Min 2014; Min 2016); not entirely clear whether these trials were conducted jointly or in tandem).
Other bias	Unclear risk	Possibly some baseline imbalance: more smokers and more planned pregnancies in the omega-3 group.

Min 2014 [diabetic women]

Methods	See Min 2014 above
Participants	Subset of women with type 2 diabetes (41 from fish oil group: 47 from placebo group)
Interventions	
Outcomes	
Notes	



lin 2016					
Methods	RCT: ISRCTN68997518: FOSIP				
Participants	138 women randomised				
	Inclusion criteria: women 17–45 years old with singleton pregnancies diagnosed with GDM (some were tested early in pregnancy)				
	Exclusion criteria: planning to receive tocolytic or corticosteroid therapy or taking fish oil supplements				
	Characteristics: high proportion of South Asian and African/Caribbean women				
	Setting: antenatal clinic, Newham University Hospital, inner-city London, UK				
Interventions	SUPPLEMENTATION: omega-3 + AA versus placebo (oleic acid)				
	Group 1: omega-3: 2 capsules of fish oil = 600 mg of DHA; 84 mg EPA; 16.8 mg AA (plus vitamin E): tota number randomised = 67				
	Group 2: placebo: 1442 mg high oleic acid sunflower oil/day: total number randomised = 71				
	Timing of supplementation: GA at recruitment: median 28 weeks (range 17 to 33)				
	Duration of supplementation: median 10 weeks (range 4 to 20)				
	All women: both supplements contained vitamin E				
	DHA + EPA dose/day: mid: 600 mg DHA + 84 mg EPA				
Outcomes	Women/birth: RBC membrane phospholipid DHA levels in the women and their neonates at birth; GA (median and range); caesarean; miscarriage; stillbirth; congenital anomaly; preterm birth < 37 weeks; preterm birth < 34 weeks; late preterm birth, low birthweight; birthweight > 4 kg; birthweight, birth length; birth head circumference; shoulder, mid arm and abdominal circumferences at birth				
	Babies/infants/children: neonatal hyperglycaemia				
Notes	Funding: grants from FP6 Marie Curie Actions-Transfer of Knowledge (MTKD-CT-2005-029914), Foyle Foundation, Newham University Hospital NHS Trust, Diabetes Research Network (North East London Diabetes Local Research Network), Equazen/Vifor Pharma Ltd., London Metropolitan University, Sir Halley Stewart Trust, The Mother and Child Foundation, Letten Foundation and personal donation from Emeritus Professor Clara Lowy. The supplements (Mumomega and placebo) used in this study were prepared and provided by Equazen/Vifor Pharma Ltd free of charge.				
	Declarations of interest: none declared				
	Trial registration is the same as for Min 2014.				

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was carried out using a random code generated by the supplement provider.
Allocation concealment (selection bias)	Low risk	Randomisation was carried out using a random code generated by the supplement provider (third party).
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Participants, midwives and all investigators were blinded to allocation until all the analysis was done.



Min 2016 (Continued)		
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Participants, midwives and all investigators were blinded to allocation until all the analysis was done.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The omega-3 + AA arm lost 9/67 (13%) to follow-up: • 5 withdrew • 4 moved The placebo arm lost 13/71 (18%) to follow-up: • 8 withdrew • 5 moved • 1 miscarriage = 56 live births
Selective reporting (reporting bias) Other bias	Unclear risk Unclear risk	Some protocol changes after trial registration. Under the same registry number as FOSIP ((Min 2014; Min 2016); not clear whether these trials were conducted jointly or in tandem). Possible imbalance: most women with a preterm birth in the omega-3 group had a history of preterm birth in contrast to 1 woman in the placebo group.

Methods	RCT: NCT00711971: Mothers, Omega-3, and Mental Health Study
	3 arms
Participants	126 women randomised (2 omega-3 groups and 1 control group)
	Inclusion criteria: a past history of depression (EPDS score 9-19), singleton gestation, a maternal age of ≥18 years, and a GA of 12-20 weeks
	Exclusion criteria: women with history of a bleeding disorder, thrombophilia requiring anticoagulation, multiple gestation, bipolar disorder, current major depressive disorder, current substance abuse, lifetime substance dependence or schizophrenia. Women were also ineligible if they were taking omega-3 fatty acid supplements or antidepressant medications or eating more than 2 fish meals per week. The Mini-International Neuropsychiatric Interview was also used to exclude current major depressive disorder, bipolar disorder, current substance abuse or dependence, suicidal ideation or schizophrenia.
	Setting: The University of Michigan Health System and St Joseph's Mercy Hospital Health System, in southeastern Michigan, USA from October 2008 to May 2011
Interventions	SUPPLEMENTATION: EPA versus DHA versus placebo (soy oil)
	Group 1: EPA-rich fish oil supplementation (1060 mg EPA plus 274 mg DHA): 2 large EPA-rich fish oil capsules and 4 small placebo capsules (double-dummy design); total number randomised: n = 42 (39)
	Group 2: DHA-rich fish oil supplementation (900 mg DHA plus 180 mg EPA): 2 large placebo capsules and 4 small DHA-rich fish oil capsules; total number randomised: n = 42 (38)
	Group 3: soy oil placebo (98% soybean oil and 1% each of lemon and fish oil): 2 large and 4 small place bo capsules; total number randomised: $n = 42$ (41)
	Timing of supplementation: from 12-20 weeks GA to 6-8 weeks postpartum
	DHA + EPA dose/day:



Mozurkewich 2013 (Continued)

Group 1: high: 274 mg DHA + 1060 mg EPA

Group 2: high: 900 mg DHA + 80 mg EPA

Outcomes

Women/birth: PE/hypertension; caesarean section; induction of labour; PPH (blood loss); adverse effects; cessation of the intervention; gestational diabetes; postnatal depression; length of gestation; spontaneous vaginal birth; operative vaginal birth; birthweight; cytokines

Babies/infants/children: admission to NICU; cytokines (cord blood)

(Allergy (childhood eczema at 36 months) was reported, but is covered by another Cochrane review, Gunaratne 2015)

Notes

Funding: NIH grant R21 AT004166-03S1 (National Center for Complementary and Alternative Medicine) and a University of Michigan Clinical Research Initiatives grant and by the University of Michigan General Clinical Research Center, now the Michigan Clinical Research Unit. This study was also supported (in part) by the NIH through the University of Michigan's Cancer Center Support Grant (P30 CA046592), and by the National Center for Research Resources and the National Center for Advancing Translational Sciences, NIH, through grant 8UL1TR000041, for the Clinical and Translational Science Center, University of New Mexico. The Nordic Naturals Corporation donated both active supplements and placebos to the trial.

Declarations of interest: "E.L.M. was an invited speaker at the Nutracon 2012 Conference, sponsored by the Global Organization for EPA and DHA Omega-3s (GOED), Anaheim, CA, March 7–8, 2012, and received reimbursement for travel expenses. The remaining authors report no conflict of interest"

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Randomization was carried out using a random number table maintained in the University of Michigan Investigational Drug Service."
Allocation concealment (selection bias)	Low risk	Quote: "Randomization was carried out using a random number table maintained in the University of Michigan Investigational Drug Service." (central randomisation)
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The placebos were formulated to be identical in appearance to both the EPA- and DHA-rich supplements Because the EPA and DHA capsules were not identical in appearance, we used a double-dummy design to maintain blinding."
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not specifically detailed
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up (and exclusion from analyses): • EPA group: 3/42 • DHA group: 4/42 • Placebo group 1/42 (7/8 women were lost to follow-up; 1 woman in the DHA group had a sec-
Selective reporting /re	Low risk	ond-trimester pregnancy loss attributed to cervical insufficiency).
Selective reporting (reporting bias)		Trial protocol available; no evidence of selective reporting (though data on adverse effects reported in text only).
Other bias	Low risk	Baseline characteristics reported were well balanced across the 3 groups.



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Methods	RCT: NCT00620672					
Participants	270 women randomised					
	Inclusion criteria: women at 16 weeks' gestation or less, not taking any lipid or fatty acid supplement, who were expected to deliver 1 infant at full-term gestation, with no maternal or fetal complications Exclusion criteria: diabetes, cardiac disease, renal disease, tuberculosis, HIV/AIDS, hepatitis, previous pregnancy complications, substance abuse					
	Setting: Vancouver, Ca	anada: 2004 to 2008				
Interventions	SUPPLEMENTATION: DHA versus placebo					
	Group 1: DHA: 400 mg/day (given as algal oil triglycerides). The supplements were provided in identical capsules in bottles with more than sufficient supplements to cover the study interval: total number randomised = 132					
	Group 2: placebo: equivalent amount of corn and soybean oil blended to reflect the dietary 18:2 omega-6 and 18:3 omega-3 ratio (but in amounts quantitatively insignificant compared to usual intakes): total number randomised n = 138					
	Timing of supplementation: from enrolment (~16 weeks) to 36 weeks GA					
	DHA + EPA dose/day: low: 400 mg DHA + EPA not stated					
Outcomes	Women/birth: RBC DHA; dietary intake at 16 and 36 weeks GA; GWG; breast milk DHA; breastfeeding monthly					
	Babies/infants/children: birthweight; infant mortality; weight at 2, 6, 9, 12 and 18 months; length at 2, 6, 9, 12 and 18 months; weight for length at 18 months; weight for age at 18 months; visual acuity at 2 months, 12 months and 5.75 years; language at 14-18 months and 5.75 years; problem-solving; MacArthur Communicative Development Inventory; BSID-III; all at 18 months					
	at 5.75 years: PPVT, K-ABC scores, lipids, attention (TOVA)					
Notes	Funding: Canadian Institutes for Health Research					
	Declarations of interest: none declared					
Risk of bias						
Bias	Authors' judgement	Support for judgement				
Random sequence generation (selection bias)	Low risk	Computer-generated, random codes				
Allocation concealment (selection bias)	Unclear risk	Use of sealed, opaque envelopes				
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The supplements were identical in appearance, and contained an orange flavour mask.				
Blinding of outcome assessment (detection bias) All outcomes	Low risk	As above; plus "testers blinded to the infant group"				



Mulder 2014 (Continued)

Incomplete outcome data (attrition bias)
All outcomes

Unclear risk

Lost to follow-up at 36 weeks from:

DHA group: 27/132 (20%):

- 16 self-withdrawal from 16-36 weeks
- 6 protocol non-adherence
- 5 preterm birth, miscarriage, elective termination, other pregnancy complications

Placebo group: 23/138 (17%):

- 9 self-withdrawal from 16-36 weeks
- 7 protocol non-adherence
- 5 preterm birth, miscarriage, elective termination, other pregnancy complications
- 1 twins
- 1 lost blood sample

At 18 months losses to follow-up were:

- DHA group: 36/132 (25%)
- Placebo group: 34/138 (25%)

(with higher losses for some of the anthropometric assessments)

At 5.75 years losses to follow-up were:

- DHA group: 86/132 (65%)
- Placebo group: 86/138 (62%)

Selective reporting (reporting bias)	Unclear risk	Adverse birth outcomes treated as exclusions; not clearly reported.
Other bias	Unclear risk	There were more boys born in the placebo group than the DHA group; no other obvious sources of bias identified.

Noakes 2012

Methods	RCT: NCT00801502 (SiPs)		
Participants	123 women randomised		
	Inclusion criteria: women with a diet low in oily fish (excluding canned tuna) (intake ≤ twice per month); with a family history of atopy, allergy or asthma (1 or more of the first-degree relatives of the infant affected by atopy, allergy or asthma by self-report); age 18-40 years; < 19 weeks GA; healthy uncomplicated singleton pregnancy; not using fish oil supplements currently or in the previous 3 months		
	Exclusion criteria: participation in another research study; known diabetic; presence of any autoimmune disease, learning disability, terminal illness or mental health problems		
	Setting: catchment area of the Princess Anne Hospital, Southampton University Hospitals NHS Trust, Southampton, UK (recruitment and study dates not reported)		
Interventions	FISH DIET versus USUAL DIET (low in oily fish)		
	Group 1: farmed salmon: women were provided with 2 x 150 g portions of farmed salmon a week and a cookbook of salmon recipes). The salmon (see below) was delivered to the homes of these women in individual frozen and vacuum-packed portions (150 g) on a monthly basis; sufficient portions were pro-		



Noakes 2012 (Continued)

vided for each woman and her partner. Salmon for use in the SiPS were raised at Skretting Aquaculture Research Centre, Stavanger, Norway. Each 150-g salmon portion contained (on average) 30.5 g protein, 16.4 g fat, 0.57 g EPA, 0.35 g DPA (22:5n23), 1.16 g DHA, 3.56 g total omega-3 PUFA, 4.1 mg α-tocopherol, 1.6 mg γ-tocopherol, 6 μg vitamin A (sum of all retinols), 14 μg vitamin D3, and 43 μg selenium; variance in content of all nutrients among several analysed portions was: 5% for protein and fat; 10% for individual fatty acids, α-tocopherol, and γ-tocopherol; and 20% for vitamin A, vitamin D3, and selenium. Thus, 2 portions of salmon/wk would typically provide 3.45 g EPA +DHA, 28 μg vitamin D3, and 86 μg selenium. Total number randomised: n = 62

Group 2: usual diet: women were asked to continue their habitual diets; these women also received the information sheet that described the possible health benefits of consuming oily fish during pregnancy and the government recommendation that pregnant women consume 1 or 2 oily fish meals/week. In addition, they received a cookbook providing recipes for healthy eating during pregnancy.

All women: received a diary in which to record any seafood consumed during the course of the study and the nature of its preparation and cooking. Total number randomised: n = 61

Timing of supplementation: 20 weeks GA to birth

DHA + EPA dose/day: low: 330 mg DHA + 160 mg EPA

Outcomes

Women/birth: adherence, caesarean section, adverse effects; length of gestation

Babies/infants/children: birthweight; birth length; head circumference; ILs (cord blood); IgE (birth and at 6 months); incidence and severity of atopic eczema at 6 months, skin prick test positivity at 6 months

Notes

Funding: supported by the European Commission under Framework 6: Sustainable aqua feeds to maximize the health benefits of farmed fish for consumers (Aquamax; FOOD-CT-2006-16249). 2 researchers were supported by the Southampton NIHR Biomedical Research Unit in Nutrition, Diet & Lifestyle. Salmon was donated by the University of Bergen, Norway.

Declarations of interest: Grethe Roselunf is employed by a company that produces feed for farmed salmon. None of the other authors had any personal or financial conflicts of interest.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The women were allocated to 1 of 2 groups according to a previously generated random number table".
Allocation concealment (selection bias)	Unclear risk	No details reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Quote: "single blind"
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Researchers responsible for assessing outcomes measures (both laboratory and clinical) remained blinded to the groups.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	 123 enrolled at 19-20 weeks (62 salmon, 61 control): withdrew before week 34 of pregnancy: salmon: 7, control: 5 withdrew between 34-38 weeks: salmon: 1, control: 2 withdrew between 38 weeks and birth: salmon: 1, control: 0 withdrew before 6-month visit: salmon: 1, control: 7 unable to contact: salmon: 4, control: 9



Noakes 2012 (Continued)		Therefore 53/62 (86%) remained in salmon group and 54/61 (89%) in control group at birth.
Selective reporting (reporting bias)	Unclear risk	Very few clinical outcomes have been reported; trial registration brief and no access to trial protocol.
Other bias	Low risk	Baseline characteristics were comparable between groups.

Ogundipe 2016

<u> </u>			
Methods	RCT: FOSS: ISRCTN24068733		
Participants	300 women randomised (normal healthy controls (n = 50), women identified at risk of having a low birthweight baby either spontaneously (n = 100) or because of developing PE (n = 100), and women at risk of gestational diabetes (n = 50).		
	Inclusion criteria: healthy women and women at risk of developing pregnancy-related complications PE, fetal growth restriction, gestational diabetes		
	Exclusion criteria: women with known allergy to fish and fish oil, non-English speakers who decline the use of an interpreter and those unable or unwilling to attend follow-up appointments; women with chronic disease such as HIV, cirrhosis or other chronic liver disease, hepatitis B and C carriers; women previously on regular pre-conceptual fish oil supplement or who, for different reasons, were not able to give competent, written consent		
	Setting: Chelsea and Westminster Hospital, London, UK		
Interventions	SUPPLEMENTATION + OTHER AGENT: DHA + EPA + AA versus placebo		
	Group 1: 2 capsules daily of DHA-enriched formula (each capsule contained 300 mg of DHA, 42 mg of EPA and 8.4 mg of AA): total number randomised unclear		
	Group 2: placebo: 2 capsules daily (high oleic acid sunflower seed oil – 721 mg oleic acid); total number randomised unclear		
	Timing of supplementation: from 8-12 weeks' gestation		
	DHA + EPA dose/day: mid: 600 mg DHA + 84 g EPA		
Outcomes	Women/birth: maternal neurobehavioural outcomes (listed in trial registration entry), maternal lipid profile		
	Babies/infants/children: MRI brain scan findings; infant developmental outcomes (no outcomes yet reported by intervention and control group - Ogundipe 2016 reports overall GA, birthweight, birth length, head circumference at birth, low birthweight)		
Notes	No outcomes could be used in this review to date.		
	Funding: The Mother and Child Foundation, Letten Foundation, Waterloo Foundation and Vifor Pharma, Switzerland.		
	Declarations of interest: not reported		
Risk of bias			
Bias	Authors' judgement Support for judgement		
Random sequence generation (selection bias)	Unclear risk Not reported		



Ogundipe 2016 (Continued)		
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Not reported
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Not reported
Selective reporting (reporting bias)	Unclear risk	Not reported
Other bias	Unclear risk	Not reported

Oken 2013

Methods	3-arm trial; NCT01126762: 'Food for thought' (pilot study)		
Participants	61 women randomised		
	Inclusion criteria: women 12-22 weeks GA, consuming ≤ 2 fish servings/month, ≥18 years of age, singleton pregnancy, planning to remain in Boston till the birth, women with no contraindications to fish consumption such as allergy, or self-restrictions such as vegetarian diet		
	Exclusion criteria: as indicated above		
	Setting: Harvard Medical School or participant's homes, greater Boston, USA (recruited April to October 2010)		
Interventions	DIETARY ADVICE: Advice to consume fish versus advice + vouchers versus generic advice		
	Group 1: advice to consume low-mercury/high-DHA fish; 8-page booklet and resources on safe fish consumption and health benefits; weekly emails; total number randomised: n = 20 (17)		
	Group 2: advice and grocery store gift cards to purchase fish; 8-page booklet on safe fish consumption and health benefits; weekly emails; USD 120 value in gift cards (USD 40 at baseline, first month and second month = USD 10 a week); total number randomised: n = 20 (18)		
	Group 3: generic advice (control): 8-page generic advice on pregnancy nutrition; weekly emails; total number randomised: n = 21 (20)		
	All women: USD 25 gift card at baseline and completion		
	DHA + EPA dose/day: unclear		
Outcomes	Women/birth: fish intake (using 1 month fish intake FFQ) and fruit, vegetables, dairy, nuts and meat; use of DHA supplements; women's opinions and attitudes about fish consumption; DHA (plasma); mercury (blood and hair); preterm birth; maternal mortality; stillbirth; gestational diabetes; PE; gestational hypertension; induction of labour; caesarean birth; postpartum depressive symptoms		



Oken 2013 (Continued)

Notes

Funding: NIH; HSPH-NIEHS Center for Environmental Health, Harvard Clinical Nutrition Research Center; Harvard Pilgrim Health Care Institute

Declarations of interest: none declared

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Random number table
Allocation concealment (selection bias)	Low risk	Sealed opaque envelopes, sequentially opened
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "All study staff were blinded to group assignment before baseline measures were collected. To minimize bias introduced by non-blinding of the single research assistant who both delivered the intervention and collected follow-up data, self-reported data were collected by self-administered question-naire rather than by interview. Laboratory staff, statistical analysts, and study investigators remained blinded to group assignment throughout data collection and analysis."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Laboratory staff, statistical analysts, and study investigators remained blinded to group assignment throughout data collection and analysis."
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	6/61 (10%) women were lost to follow-up (2 in the advice group discontinued intervention (1 died)); and 1 and 3 in the control and advice + gift card groups respectively
		Post birth outcomes were not available for 13/61 (21%) women – 9/40 in the 2 intervention arms and 4/21 in the control arm.
Selective reporting (reporting bias)	Unclear risk	Specific figures for most birth outcomes not reported (only direction of effect).
Other bias	Unclear risk	Baseline characteristics similar, except for a higher proportion of women working full time in the advice + gift card group.

Olsen 1992

Interventions	SUPPLEMENTATION: omega-3 versus olive oil versus no supplement
	Setting: main midwifery clinic, Aarhus, Denmark
	Exclusion criteria: history of placental abruption, a serious bleed in the current pregnancy, use of PG inhibitors, multiple pregnancy, fish allergy or regular intake of fish oil
	Inclusion criteria: healthy women, at approximately 30 weeks' gestation, aged 18-44 years
Participants	533 women randomised
	3-arm RCT in a ratio of 2:1:1: for fish oil, olive oil, and no supplement
Methods	NCT01353807



Olsen 1992 (Continued)

Group 1: fish oil (2.7 g omega-3 fatty acids/day) given as 4 x 1 g capsules/day containing fish oil (Pikasol 32% EPA (20:5n-3), 23% DHA (22:6n-3)) and 2 mg tocopherol/mL: total number randomised = 266 **Group 2:** 4 x 1 g capsules olive oil/day: total number randomised = 136

Group 3: no supplement: total number randomised = 131

Timing of supplementation: from ~30 weeks GA to birth

DHA + EPA dose/day: high: 864 mg DHA + 621 mg EPA

Outcomes

Women/birth: SBP and DBP, PIH, PE, food frequency questionnaire*, preterm birth < 37 weeks, caesarean, congenital anomalies, blood loss, maternal adverse effects

Babies/infants/children/adults: stillbirth, duration of gestation, birthweight, birth length, BMI

At 16-year follow-up: asthma, atopic dermatitis, allergic rhinitis

At 19-year follow-up: insulin, glucose, glycated Hb, leptin, adiponectin, insulin-like growth factor 1, high sensitivity CRP, height, weight, BMI, waist circumference

*enabled women to be classified into low, medium and high habitual intake of fish

Notes

Sample size estimates were done but not reported in the papers because they were regarded as posthoc by authors (personal communication).

Women completed baseline information regarding fish intake.

Outcome assessment was blinded, but 85% of women in the fish oil group correctly identified their group allocation, whereas for olive oil 50% identified the correct oil.

Funding: "This study was supported by the Danish Medical Research council (J No 12-9052) and 12-9144), Sygekassernes Helsefond, Weirnan's Legat and Michaelsen Fonden". Capsules were provided by Lube Ltd, Hadsund, Denmark.

Follow-up was supported by the EU FP5 consortium, Early Nutrition Programming Project, NIH, The Danish Strategic Research Council, The Danish Heart Foundation, The Novo Nordisk Foundation, The Danish Diabetes Foundation, The Aase and Ejnar Danielsens Foundation, and the National Center for Complementary and Alternative Medicine.

Declarations of interest: JE Chavarro and Sjurdur F Olsen received research support from the NIH. The rest of the authors declared that they had no relevant conflicts of interest.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Stratified and block randomisation
Allocation concealment (selection bias)	Low risk	A sealed, opaque envelope containing a randomisation number which identified either a particular package of oil capsules or no treatment.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Capsules and their boxes looked identical for fish oil and olive oil; however the no treatment group was unblinded.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported, apart from that at 19-year follow-up outcome assessors were blinded to group allocation.
Incomplete outcome data (attrition bias)	Low risk	No post-randomisation exclusions and no losses to follow-up; at 19-year follow-up: n = 243 completed physical examination (out of 517 mother/child



Olsen 1992 (Continued) All outcomes		dyads alive and still living in Denmark); 41% were from the fish oil group and 53% were from the olive oil/no oil group.
Selective reporting (reporting bias)	Low risk	Most expected outcomes were reported.
Other bias	Low risk	Similar baseline characteristics

Olsen 2000

Methods

Multicentre RCT: NCT02229526 (FOTIP) with 6 different subsets (A to F) of eligibility criteria. The 6 subsets had a standard protocol, and were mutually exclusive.

Trials A-D were prophylactic trials in women after 16 weeks' gestation, with uncomplicated pregnancies:

A: previous preterm birth (before 259 days gestation); n = 232

B: IUGR (< 5th centile); n = 280

C: PIH (DBP > 100 mmHg); n = 386

D: current twin pregnancies: n = 579.

Trials E and F were therapeutic trials, enrolling women around 33 weeks' gestation:

E: signs or symptoms of PE in the current pregnancy (± IUGR): n = 79; or

F: suspected IUGR (< 10th centile by ultrasonography) in current pregnancy: n = 63.

Participants

1647 women randomised (fish oil: 818, olive oil: 829)

Inclusion criteria

Prophylactic trials: women after 16 weeks' gestation with an uncomplicated pregnancy, who in an earlier pregnancy had experienced preterm birth (before 259 days gestation); IUGR (< 5th centile); PIH (DBP > 100 mmHg); or women with current twin pregnancies

Therapeutic trials: women with threatening PE (women with signs or symptoms or PE) or suspected IUGR (< 10th centile).

Exclusion criteria: diabetes mellitus in or before pregnancy, diagnosed severe fetal malformation or hydrops in current pregnancy, suspicion in current pregnancy or occurrence in an earlier pregnancy of placental abruption, drug or alcohol abuse, regular intake of fish oil or of NSAIDs or other drugs with an effect on thrombocyte function or eicosanoid metabolism, allergy to fish products. In the therapeutic trials, an additional exclusion criterion was: high probability of delivering soon after randomisation (within 1 week).

Setting: 19 hospital centres in Denmark, Scotland, Sweden, England, Italy, the Netherlands, Norway, Belgium and Russia

Interventions

SUPPLEMENTATION: omega-3 (EPA/DHA) versus control (olive oil)

Group 1: omega-3 (2.7 g/day (1.3 g EPA and 0.9 g DHA)), given as 4 capsules/day in the prophylactic trials and 9 capsules per day in the therapeutic trials (6.1 g/day: 2.9 g EPA and 2.1 DHA): total n = 818 **Group 2:** matching olive oil capsules: total n = 829

Timing of supplementation: prophylactic trials: ~20 weeks' gestation to birth; therapeutic trials ~33 weeks' gestation to birth

DHA + EPA dose/day: prophylactic trials: high: 900 mg DHA + 1300 mg EPA



Olsen 2000 (Continued)

DHA + EPA dose/day: therapeutic trials: high: 2100 mg DHA + 2900 EPA

Outcomes

Subset A: preterm birth (< 37 weeks), early preterm birth (< 34 completed weeks), low birthweight, length of gestation, birthweight

Subset B: SGA, low birthweight, birthweight.

Subset C: PIH (DBP > 90 mmHg at rest, PE (combination of PIH and proteinuria), antihypertensive therapy. BP

Subset D: preterm birth (< 37 weeks), early preterm birth (< 34 completed weeks), PE (combination of PIH and proteinuria), antihypertensive therapy, BP, SGA, low birthweight, length of gestation, birthweight

Threat-PE trial: duration until birth

Susp-IUGR trial: weight for GA

For the combined subsets: length of gestation, preterm birth (minus elective births), early preterm birth (minus elective births), prolonged gestation (42 completed weeks), maternal morbidity and mortality, infant mortality and morbidity

Elective births were defined as induced vaginal births or prelabour caesareans.

Notes

Sample size estimates were modified during the course of the study: sample size determinations were undertaken twice in the study; after 4 years of enrolment, it was realised the original estimated sample sizes were unrealistically large, therefore a stopping strategy was developed (based on a number of primary hypotheses, defined a priori for the prophylactic trials).

Funding: Concerted Action (ERB-BMH1-CT92-1906) and PECO (ERB-CIPD-CT94-0235) programmes of the European Commission, and the Danish National Research Foundation. Lube Ltd provided Pikasol fish oil and olive oil capsules.

Declarations of interest: not reported

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Restricted blockwise computer-generated randomisation was used
Allocation concealment (selection bias)	Low risk	Quote: "Randomisation identified a package number at the relevant centre, where packages were ordered in a random way as to oil type". Placebo capsules were identical looking but contained olive oil.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Both oils were provided in 1 g identical looking gelatine capsules, which were not identical in taste; packages with capsules were identified by a cryptographed number, the code of which was known only by the data manager"; however a questionnaire completed by a subgroup of women indicated that 80% of women in the fish oil group could guess their allocation.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessments were blinded (correspondence with Olsen).
Incomplete outcome data (attrition bias) All outcomes	Low risk	Overall, 1647 women randomised (fish oil: 818, olive oil: 829), trial entry forms received for: 98% in both groups, follow-up forms received for 97% in both groups (and compliance forms for 68% in both groups).
Selective reporting (reporting bias)	Unclear risk	Not all outcomes were measured in all groups.



Olsen 2000 (Continued)

Other bias Low risk Baseline characteristics were similar between groups except for women with suspected IUGR where GA at enrolment was higher in the olive oil group.

Olsen 2000 [twins]

Methods	Twins (see Olsen 2000)
Participants	
Interventions	
Outcomes	
Notes	

Onwude 1995

Methods	RCT			
Participants	232 women randomised (161 multigravida and 72 primigravida)			
	Inclusion criteria: mean 24 weeks' gestation with high-risk singleton pregnancy: history of 1 or more small babies (birthweight < 3rd percentile), history of pregnancy hypertension, history of unexplained stillbirth, or primigravida with abnormal uterine Doppler at 24 weeks' gestation			
	Exclusion criteria: history of diabetes mellitus, chronic hypertension, asthma, use of anticoagulants, multiple pregnancy			
	Setting: antenatal clinic, St James's University Hospital, Leeds, UK			
Interventions	SUPPLEMENTATION: EPA + DHA versus placebo			
	Group 1: 2.7 g of MaxEPA/day (n = 113): 9 capsules/day provided 1.62 g EPA and 1.08 g DHA Group 2: control: matching air-filled capsules (n = 119)			
	Timing of supplementation: from a mean of 24 weeks to 38 weeks' gestation			
	EPA + DHA dose/day: high: 1080 mg DHA + 1620 mg EPA			
Outcomes	Women/birth: length of gestation; preterm birth < 37 weeks; preterm birth < 32 weeks; duration and mode of onset of labour; mode of birth; PIH, PE, adverse events (75 women only) Babies/infants/children: stillbirth; neonatal mortality; birthweight; birthweight < 3rd percentile			
Notes	Sample size estimate was given for proteinuric hypertension			
	All women were asked to avoid NSAIDs			
	Adherence: 50% of women in the fish oil group and 57% of women in the placebo group took < 70% of capsules.			
	Protocol variations: 1 woman in the omega-3 arm took aspirin and 1 women in the placebo arm purchased fish oil privately.			
	Funding: Yorkshire Region Locally Organised Research; GLAXO (Leeds); Sevens Seas (Hull)			
	Declarations of interest: not reported			



Onwude 1995 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Random numbers were generated by computer.	
Allocation concealment (selection bias)	Low risk	Random numbers were kept in sealed, opaque, numbered envelopes in the hospital pharmacy and pharmacy staff allocated the trial treatments.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Placebo capsules were identical to treatment capsules (44% of a subgroup of women identified that they were in the fish oil group).	
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Outcome assessments were blinded.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	1/233 (0.4%) post-randomisation exclusions (one woman with a multiple pregnancy was randomised in error) Adverse events were reported only for a small subsample; 76/233 (33%)	
		women returned the questionnaires.	
Selective reporting (reporting bias)	High risk	Limited range of outcomes reported; no SDs reported for continuous outcomes (length of gestation; birthweight).	
Other bias	Low risk	Baseline characteristics were similar in supplement and placebo groups.	

Otto 2000

Jtto 2000	
Methods	RCT
Participants	24 women randomised
	Inclusion criteria: healthy pregnant women in the second trimester, aged 20-38 years, with singleton pregnancies
	Setting: Department of Obstetrics and Gynecology, University Hospital Maastricht; and midwifery practices in the Maastricht area, the Netherlands
Interventions	SUPPLEMENTATION + OTHER AGENT: DHA + AA versus no treatment
	Group 1: DHA capsules (algal oil: 0.57 g DHA/day) and AA (fungal-derived oil: 0.26 g/day) for 4 weeks: total number randomised = 12
	Group 2: no supplements: total number randomised = 12
	Timing of supplementation: from 17-20 weeks' gestation
	DHA + EPA dose/day: mid: 570 mg DHA; negligible EPA
Outcomes	Women: fish and seafood consumption at the end of the study; plasma phospholipid fatty acids after 4 weeks; adverse events
Notes	3 women in each group reported consuming fish once a week during the 4 week study period.



Otto 2000 (Continued)

Funding: Martek Corporation (donation of capsules); Numico BV, the Netherlands

Declarations of interest: not reported

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomly assigned"
Allocation concealment (selection bias)	Unclear risk	Quote: "randomly assigned"
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not blinded
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	All 12 women completed the study.
Selective reporting (reporting bias)	High risk	Limited number of outcomes reported.

Baseline characteristics were similar, apart from more primigravida in the sup-

plement group (8/12 versus 5/12 in the control group).

Pietrantoni 2014

Other bias

Pietrantoni 2014	
Methods	RCT
Participants	300 women
	Inclusion criteria: < 8 weeks' gestation; Caucasian women aged 22-35 years; singleton pregnancy; BMI ≥ 18.5 and ≤ 25; habitual fish consumption (at least twice a week); high school or university degree; average socioeconomic status; absence of uterine abnormalities (fibroids, cervical incompetence; uterine malformations etc.)
	Exclusion criteria: smoking, substance abuse including alcohol; allergy to fish or derivatives; diabetes, hypertension, metabolic, cardiovascular, renal, psychiatric, neurological, thrombophilic, thyroid or autoimmune disease; previous pregnancy complications (miscarriage, preterm or operative birth); previous uterine surgery (myomectomy, cervical conisation, trachelorraphy, caesarean etc.); recurrent genito-urinary infections
	Setting: Department of Obstetrics and Gynaecology, San Camillo Forlanini Hospital, Rome, Italy
Interventions	SUPPLEMENTATION + DIET: DHA + fish versus placebo + fish
	Group 1: omega-3 (2 capsules of DHA (100 mg each) administered daily); total number randomised: n = unclear (129)
	Group 2: placebo (2 capsules of olive oil); total number randomised: n = unclear (126)

Low risk



Pietranton	2014	(Continued)
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All women: controlled diet (high protein ~17%; low in carbohydrates ~54%; fat ~2%; omega-3 (600 g fish/week); increased calorie requirements (+200 Kcal); food diary for a week a month

Timing of supplementation: from 8 weeks GA until birth

DHA + EPA dose/day: low: 200 mg DHA; EPA not stated

Outcomes Women: lipids; food diary; rupture of membranes

Notes Funding: Ministry of Health, Rome, Italy

Declarations of interest: none declared

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Not reported
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Not clear, but if 300 women were randomised, then 45/300 (15%) were lost to follow-up
Selective reporting (reporting bias)	High risk	Only rupture of membranes reported
Other bias	Low risk	Similar baseline characteristics

Ramakrishnan 2010

Methods	RCT: parallel; NCT00646360: POSGRAD (Prenatal Omega-3 fatty acid Supplements, Growth, and Development)
Participants	1094 women randomised
	Inclusion criteria: women 18-35 years old, at 18-22 weeks' gestation, who planned to give birth at the Mexican Institute of Social Security General Hospital (IMSS) in Cuernavaca, exclusively or predominately breastfeeding for at least 3 months, and planned to live in the area for at least 2 years after giving birth
	Exclusion criteria: high-risk pregnancy (history or prevalence of pregnancy complications: abruptio placenta, PE, PIH, any serious bleeding episode in current pregnancy, physician referral), lipid metabolism or abruption disorders, regular intake of fish oil or DHA supplements, or chronic use of certain medications (e.g. for epilepsy)



Ramakrishnan 2010 (Continued)

Characteristics: medium-low socioeconomic status; low baseline omega-3; high omega 6:omega 3 ratio

Primiparae and multipara results presented separately.

Setting: Mexico: hospital (routine antenatal); IMMS, Cuernavaca, Mexico, and 3 small health clinics (study conducted between February 2005 and February 2007

Interventions

SUPPLEMENTATION: DHA versus placebo

Group 1: DHA: 400 mg daily; capsules contained 200 mg DHA derived from an algal source. Women were instructed to take 2 capsules, together, at the same time each day: total number randomised: n = 547

Group 2: placebo: daily; placebo capsules contained olive oil or soy/corn mix and were similar in appearance and taste to the DHA capsules: total number randomised: n = 547

Timing of supplementation: 18-22 weeks GA (mean 20.6 weeks) to birth

DHA + EPA dose/day: low: 400 mg DHA; EPA not stated

Outcomes

Women: nausea; vomiting; vaginal bleeding; GA; caesarean section; cessation of supplements; adherence

Birth/infant/child:

- **Birth to 18 months:** preterm birth < 37 weeks; birthweight; birth length; head circumference at birth; SGA (IUGR < 10th centile); low birthweight; congenital anomalies; serious adverse events; fetal loss; stillbirth (28 weeks); neonatal deaths; infant deaths; cord blood (DNA analysis); child health at 3 and 6 months; BSID-II (Spanish version) MDI and PDI at 18 months, HOME inventory at 12 months; Behaviour Rating Scale at 18 months; brainstem auditory-evoked responses (singletons only) at 1 and 3 months; visual-evoked potentials (singletons only) at 3 and 6 months; anthropometric measures at 1, 3, 6, 9, 12 and 18 months;
- 4-year follow-up: weight, height, BMI, z-scores, overweight/obese; glucose and lipid concentrations; insulin
- 5-year follow-up: child weight, length, BMI, McCarthy Test for Global Development (Spanish Language version), K-CPT, BASC-PRS

Notes

Adherence: 88%; did not differ by treatment group

Funding: NIH (HD-043099) and the March of Dimes Foundation

Declarations of interest: Y Gutierrez-Gomez, AD Stein, U Ramakrishnan, A Barraza-Villarreal, H Moreno-Macias, C Aguilar-Salinas, I Romieu, and JA Rivera declared no conflicts of interest

Bias Authors' judgement Support for judgement		Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "We used block randomization to randomly create balanced replication of four treatments (two colors for DHA and two for control) using a block size of eight. The list was generated for a sample size of 1,104" - probably done
Allocation concealment (selection bias)	Low risk	Quote: "The assignment codes were placed in sealed envelopes at the beginning of the study, and these envelopes were held in a sealed location by a faculty member who was not involved in the study."
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "All the study participants and members of the study team remained blinded to the treatment scheme throughout the intervention period of the study. Data were unblinded for the analytical study team after the last baby in the study was born and had reached 6 months of age, at which time the partic-



Ramakrishnan 2010 (Continue	d)	ipants were no longer taking supplements. Since the study is ongoing for follow-up of child development, the participants and fieldworkers in Mexico remain blinded to the treatment allocation"
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "All the study participants and members of the study team remained blinded to the treatment scheme throughout the intervention period of the study. Data were unblinded for the analytical study team after the last baby in the study was born and had reached 6 months of age, at which time the participants were no longer taking supplements. Since the study is ongoing for follow-up of child development, the participants and fieldworkers in Mexico remain blinded to the treatment allocation"
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	1094 randomised (547 to each group); birthweight analysed for 487/547 in the DHA group and 486/547 in the placebo group (and GA analysed for 486/547 and 484/547) = 11% loss to follow-up in the DHA group and 11% in the placebo group.
		Reasons for loss to follow-up similar among groups; though the women in the final sample with birth outcomes (973) were of higher socioeconomic status than randomised women for whom birth outcomes were not available (121).
		18-month follow-up (growth): 739 children at 18 months (76.0% of birth cohort), loss to follow-up did not differ by treatment assignment; children lost to follow-up were lighter at birth, had smaller head circumferences at birth, and were more likely to have a low birthweight compared with those assessed at 18 months (and in the 18-month sample, women who received DHA were shorter than those who received placebo).
		18-month follow-up (development: 730 included in analyses ("Comparison of the final sample with outcome data (n = 730) to those randomised but lost to follow-up (n = 364) showed that the offspring in the final sample were similar in terms of selected maternal and infant characteristics including treatment group").
		4-year follow-up: data available for DHA: 51% (276) children and placebo: 45% (248) children
		5-year follow-up: 802 children (DHA 403: placebo 399) = 73% of those randomised.
Selective reporting (reporting bias)	Unclear risk	Trial registered retrospectively
Other bias	Low risk	No obvious sources of other bias; similar baseline characteristics

Ranjkesh 2011

Methods	RCT: IRCT138706061113N1
Participants	100 women
	Inclusion criteria: women with risk of PE (primiparous women, aged < 20 years and > 40 years, previous history of PE or a positive family history, twin pregnancy, BMI > 29, history of renal disease and hypertension), not using any anticoagulant or antihypertension drugs at the time of entering the study
	Exclusion criteria: none specified
	Setting: Qazvin city, Iran



Ran	kesh 2	011	(Continued)
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Interventions SUPPLEMENTATION: omega-3 (EPA + DHA) versus placebo

Group 1: omega-3 (EPA + DHA - individual doses not specified), 1 g daily: n = 50

Group 2: placebo (starch): n = 50

Timing of supplementation: from 14-18 weeks GA to end of pregnancy

DHA + EPA dose/day: unclear

Outcomes Women/birth: BP (mmHg); PE, hypertension, caesarean birth; birthweight

Babies/infants/children: Apgar score at 5 minutes

Notes Funding: not reported

Declarations of interest: not reported

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomly divided"
Allocation concealment (selection bias)	Unclear risk	Quote: "randomly divided"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Described both as single- and double-blind; placebo-controlled so probably done.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	No losses to follow-up reported.
Selective reporting (reporting bias)	Unclear risk	Some outcomes not fully reported.
Other bias	Low risk	Baseline characteristics were similar.

Razavi 2017

Methods	RCT (4 arms, see below)
Participants	120 women randomised
	Inclusion criteria: 18-40 years; without prior diabetes; diagnosed with GDM at 24-28 weeks' gestation
	Exclusion criteria: taking omega-3 fatty acid supplements; insulin therapy; placental abruption; PE; eclampsia: hypo- and hyperthyroidism: smokers



Razav	i 201	7 (Continued)
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Setting: Ardabil, Iran (conducted from September 2016 to March 2017)

Interventions

SUPPLEMENTATION: omega-3 versus omega-3 + vitamin D versus vitamin D versus placebo

Group 1: omega-3 fatty acids (2000 mg size of capsules containing 360 mg EPA and 240 mg DHA per day) as 2 capsules plus placebo vitamin D capsules; n = 30 randomised (all completed)

Group 2: omega-3 fatty acids (2000 mg size of capsules containing 360 mg EPA and 240 mg DHA per day) as 2 capsules, plus vitamin D (50 000 IU every 2 weeks): n = 30 randomised (all completed)

Group 3: vitamin D (50 000 IU every 2 weeks) plus placebo omega-3 capsules: n = 30 randomised (all completed)

Group 4: no supplement (2 placebo capsules, each containing 500 mg of liquid paraffin): n = 30 randomised (all completed)

Timing of supplementation: from 24-28 weeks' gestation (for 6 weeks)

All women: requested not to change their routine physical activity or usual dietary intakes throughout study and not to consume any supplements other than the one provided to them by the investigators, as well as not to take any medications that might affect findings during the 6-week intervention

DHA + EPA dose/day: mid: 240 mg DHA + 360 mg EPA

Outcomes

Women/birth: inflammatory biomarkers; biomarkers of oxidative stress; caesarean section; preterm birth < 37 weeks; polyhydramnios; PE; length of gestation

Babies/infants/children: macrosomia (> 4000 g); birthweight; length at birth; head circumference at birth; Apgar score; newborn hyperbilirubinaemia; newborn hospitalisation; newborn hypoglycaemia

Notes

Funding: research grant from Research Deputy of Ardabil University of Medical Sciences (AUMS)

Declarations of interest: none declared

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Assignment was conducted using computer-generated random numbers.
Allocation concealment (selection bias)	Low risk	Computer based randomisation; and "a person who was not involved in the trial and not aware of random sequences, assigned the subjects to the numbered bottles of capsules"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Randomisation and allocation were concealed from the researchers and participants until the final analyses were completed"
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not reported; however outcomes probably assessed by the researchers (who were blind to group assignments), and all outcomes included for analysis objective.
Incomplete outcome data (attrition bias) All outcomes	Low risk	All 120 participants randomised included for analysis.
Selective reporting (reporting bias)	Unclear risk	Outcomes specified in published protocol reported; no obvious sign of selective reporting, however limited range of outcomes for babies/infants/children reported.



Razavi 2017 (Continued)

Other bias Low risk No clear baseline differences between the 4 groups of participants.

Razavi 2017 [vit D]

Methods	With Vitamin D (see Razavi 2017)
Participants	
Interventions	
Outcomes	
Notes	

Rees 2008

Methods	RCT		
5			

Participants 26 women randomised

Inclusion criteria: current episode of major depression or dysthymia, according to DSM-IV criteria, confirmed by both CIDI-structured interview criteria and clinical assessment by a psychiatrist; at least 21 years of age; from the third trimester of pregnancy to 6 months postnatal; baseline score ≥ 13 on the EPDS and either > 14 on the HDRS or > 25 on MADRS.

Exclusion criteria: bipolar disorder, psychosis, drug and alcohol abuse, obsessive-compulsive disorder, eating disorder or personality disorder, an unstable medical condition, diabetes, receipt of anticoagulants or having a fish allergy; those already receiving an antidepressant or any psychological therapy, as well as those taking fish oil supplements or eating more than 3 oily fish portions per week. Those with comorbid anxiety disorders were not excluded.

Setting: perinatal depression clinic and midwives' antenatal clinic at Royal Hospital for Women, Sydney, Australia

Interventions

SUPPLEMENTATION: DHA + EPA versus placebo

Group 1: fish oil (soft gelatin capsules, DHA 0.77 g/day, EPA 0.23 g/day); vitamin E (80 mg) was added to prevent oxidation of the oil: n = 13 randomised

Group 2: placebo (sunola oil (85% monounsaturated fats, 7% saturated fats)): n = 13 randomised

All women: peppermint oil was added to all capsules to disguise any fish taste and may have minimised any gastrointestinal effects.

Timing of supplementation: either from 28-33 weeks' gestation; or from 11-19 weeks postnatal for 6 weeks intervention

DHA + EPA dose/day: mid: 770 mg DHA + 230 mg EPA

Outcomes

Women/birth: depression scores at 6 weeks (EPDS, HDRS, MADRS); adherence; omega-3 plasma concentrations; adverse events

Notes

Funding: NSW Institute of Psychiatry; Pfizer Neuroscience Research Grant, NHMRC Program Grant 222708, NSW Department of Health Infrastructure Grant; Numega Ingredients and Clover Corporation for supply of fish oil and placebo capsules

Low risk



Rees 2008 (Continued)

Blinding of outcome as-

All outcomes

sessment (detection bias)

Declarations of interest: none reported

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "computer-based random number generation"
Allocation concealment (selection bias)	Low risk	Quote: "dispensed by the hospital pharmacy"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "dispensed by the hospital pharmacy to ensure blinding of the investigators and raters"; in "identical plastic containers"; "peppermint used to mask fish oil taste"

Incomplete outcome data (attrition bias) All outcomes	High risk	Fish oil: 1/13 (8%) discontinued (possibly hypomanic)
		Placebo: 4/13 (31%) discontinued (nausea (1); suicidal (1); non-adherent (1); other treatment (1))
		Results for all 26 women were analysed; with depression scores extrapolated using the last-observation-carried-forward method.
Selective reporting (re-	High risk	The only clinical outcomes reported were depression and adverse events.

until the study had been completed"

Quote: "Subjects were interviewed by the first author, who remained blind to

treatment assignment and assessed weekly by her. The blind was not broken

porting bias)		
Other bias	High risk	Women in the placebo group were more likely to have a co-morbid anxiety disorder (9/13 vs $3/13$).

Ribeiro 2012

RIDEII 0 2012	
Methods	RCT
Participants	11 women randomised
	Inclusion criteria: age 20-30 years; in 30th week of pregnancy; not using any form of medication; not showing any signs of intolerance or allergy to fish; not using any dietary supplements containing omega-3 and omega-6 PUFA; and not feeding the baby exclusively on their own milk
	Exclusion criteria: unable to attend all the scheduled study sessions; non-authorisation by the gynae-cologist responsible; embarking on a medication treatment after the study had begun; or deciding not to participate after consenting for the study
	Setting: Sao Francisco University Hospital, Brazil (participants were selected from May 2009 to February 2010)
Interventions	SUPPLEMENTATION: omega-3 (fish oil) versus primrose oil
	Group 1: fish oil capsules containing 0.72 g omega-3 PUFA/day; total number randomised = 7
	Group 2: primrose oil capsules containing 1.46 g omega-6 PUFA/day; total number randomised = 4



Ribeiro 201	2 (Continued)
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Timing of supplementation: for 15 days (no further details)

All women: instructed to maintain their usual diet for 7 days, and then to include the capsules during the period of the study intervention (15 days)

DHA + EPA dose/day: mid? - (unclear)

Outcomes Women/birth: dietary intake; fatty acid composition of the erythrocyte membranes and breastmilk

Notes No outcomes could be used in this review.

Funding: not reported

Declarations of interest: none declared

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Method used to generate a random sequence not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment method not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Method used to blind participants and personnel not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Post randomisation dropouts and exclusions not reported.
Selective reporting (reporting bias)	Unclear risk	Few outcomes reported, and without access to a published protocol it was not possible to assess risk of reporting bias confidently.
Other bias	Unclear risk	Inadequate reporting of methods made confident assessment of risk of other bias impossible.

Rivas-Echeverria 2000

Methods	RCT
Participants	127 women randomised
	Inclusion criteria: pregnant women < 29 weeks' gestation at high risk of PE, nulliparity, previous PE, obesity, hypertension, < 20 years old, diabetes, nephropathy, mean arterial pressure < 85 mmHg, positive roll-over test, "black race", family history of hypertension or PE, twin pregnancy, poor socioeconomic conditions
	Exclusion criteria: none reported
	Setting: Mérida, Venezuela



Rivas-Echeverria 2000 (Continued)

Interventions SUPPLEMENTATION + OTHER AGENTS: omega-3 (+ aspirin, vitamins C and E) versus	placebo
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Group 1: fish oil capsules 3 times a day (omega-3 content not reported); aspirin 100 mg 3 times a week,

vitamin C 500 mg 3 times a day, vitamin E 400 IU a day; total number randomised = 63

Group 2: placebo (not further described); total number randomised = 64

Timing of supplementation: not reported

DHA + EPA dose/day: unclear

Outcomes Women/birth: PE; "no serious maternal or neonatal side effects of treatment occurred in either group"

Notes Funding: none reported

Conflicts of interest: none reported

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomly divided"
Allocation concealment (selection bias)	Unclear risk	Quote: "randomly divided"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "triple-blind"; probably done
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Not reported
Selective reporting (reporting bias)	High risk	Only 1 outcome fully reported.
Other bias	Unclear risk	Not sufficient reporting to determine risk of other bias.

Samimi 2015

Methods	RCT: IRCT20131226562N16
Participants	56 women randomised
	Inclusion criteria: pregnant women 18-40 years, diagnosed with GDM (1 step 2-hour 75 g OGTT at 24-28 weeks GA using ADA 2014 criteria, i.e. fasting glucose ≥ 92 mg/dL,1-hour ≥ 180 mg/dL and 2-hour ≥ 153 mg/dL)
	Exclusion criteria: women requiring insulin therapy, women with PPROM, placental abruption, PE, eclampsia, chronic hypertension, hypothyroidism, urinary tract infection, smokers and those with kidney or liver diseases or those taking oestrogen therapy



Samim	i 2015	(Continued)
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Characteristics: women had high omega-6 concentrations

Setting: Kashan, Iran (January-March 2014)

Interventions

SUPPLEMENTATION: EPA + DHA + other omega-3 versus placebo

Group 1: 1000 mg omega-3 fatty acid (1 'pearl' per day for 6 weeks; the pearl contained 70% LCPUFA (180 mg EPA, 120 mg DHA and 400 mg of other omega-3 fatty acids)); total number randomised: n = 28

Group 2: placebo (1 'pearl' per day for 6 weeks; the pearl contained 500 mg liquid paraffin); total number randomised: n = 28

Timing of supplementation: 6 weeks from 24-28 weeks GA

All women: asked not to alter their routine physical activity or usual dietary intakes throughout the study and not to consume any supplements other than the 1 provided to them by the investigators

DHA + EPA dose/day: low: 120 mg DHA + 120 mg EPA

Outcomes

Women: insulin resistance (HOMA-IR); HOMA-B; plasma glucose; QUICKI; lipid profile; dietary records; adherence, CRP (ng/mL)

Babies/infants/children: nil

Notes

Funding: Kashan University of Medical Sciences

Declarations of interest: none reported

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated random numbers
Allocation concealment (selection bias)	Low risk	Quote: "Randomization and allocation were concealed from the researchers and participants until the main analyses were completed. A trained midwife at maternity clinic did the randomized allocation sequence, enrolled participants and assigned participants to intervention"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The appearance of placebo color, shape, size, and packaging, were identical to omega-3 fatty acid capsule"
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not reported, but probably done.
Incomplete outcome data (attrition bias) All outcomes	Low risk	3/28 women in each group were lost to follow-up (omega-3: 1 insulin therapy and 2 hospitalised); placebo (2 insulin therapy and 1 placental abruption). However results for all 56 women were analysed.
Selective reporting (reporting bias)	Unclear risk	No baby or child outcomes reported.
Other bias	Low risk	No clear baseline differences.



Methods	RCT		
Participants	20 women randomised	İ	
	Inclusion criteria: nor nutrition	rmal pregnant women without risk of systemic diseases or the existence of mal-	
	Exclusion criteria: multiple pregnancies; IUGR; diabetes or hypertension		
	Setting: Baracaldo, Spain		
Interventions	SUPPLEMENTATION:	DHA + EPA versus placebo	
	Group 1: DHA: 200 mg DHA/day and 40 mg EPA in a supplement containing 2 g fat; n = 10 randomised, 8 analysed		
	Group 2: placebo: n = 10 randomised, 8 analysed		
	Timing of supplementation: 26-27 weeks GA to birth		
	DHA + EPA dose/day: low: 200 mg DHA + 40 mg EPA		
Outcomes	Women/birth: plasma AA; plasma DHA; GA		
	Babies/infants/children: DHA; birthweight		
Notes	Funding: supported in part by grants from Novartis op		
	Declarations of intere	est: not reported	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Computer-generated random number table	
Allocation concealment (selection bias)	Unclear risk	Not reported	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "the women were given either 200 mg/day of DHA (group A) on a b basis (through a dietary formula for pregnant women containing 2 g of fat which 200 mg are DHA and 40 mg EPA) or placebo"	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported	
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Quote: "the number of biochemical samples obtained and maternal-fetal studies completed was 16 (eight from group A and eight from group B)"	
Selective reporting (reporting bias)	Unclear risk	Without access to a published protocol it was not possible to assess selective reporting confidently.	
Other bias	Unclear risk	Information on methods insufficient to assess this domain confidently.	



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Methods	RCT			
Participants	350 women randomised			
	Inclusion criteria: singleton pregnancies, women aged 16-36 years; 24-28 weeks' gestation at enrolment; able and willing to consume eggs; access to refrigeration			
		ight > 109 kg at baseline; serious illness such as cancer, lupus, hepatitis; known serious infectious disease; diabetes or gestational diabetes at baseline; elevated nuse		
	Characteristics: most	women were socially disadvantaged, and most were African-American (73%).		
	Setting: Department of Dietetics and Nutrition, University of Kansas Medical Center, Kansas City, Kansas, USA			
Interventions	OMEGA-3-ENRICHED I	FOOD: DHA-enriched eggs versus CONTROL (ordinary eggs)		
	but reported eating 5.5 Group 2: ordinary eggs	d eggs: each egg had 133 mg DHA. Women were asked to eat 12 eggs per week (731.5 mg DHA): n = 176 randomised (142 could be analysed) s: each egg had 33 mg DHA. Women were asked to eat 12 eggs per week but reveek (178.2 mg DHA): n = 174 randomised (149 could be randomised)		
	Timing of supplementation: 24-28 weeks GA to birth			
	DHA + EPA dose/day: low: 100 mg DHA/day; EPA not stated			
Outcomes	Women/birth: gestational diabetes; PE/eclampsia; duration of gestation, preterm birth (< 37 weeks); caesarean; maternal RBC phospholipid DHA concentration at enrolment and at birth			
	Babies/infants/children: birthweight; birth length, head circumference; low birthweight; meconium staining; admissions to neonatal care; neonatal RBC phospholipid DHA concentration at birth; serious adverse events (life-threatening event, inpatient hospitalisation, or prolonging of an existing hospitalisation, a persistent or significant disability/incapacity, or a congenital anomaly/birth defect			
Notes	tion on which to base a	285. Because there were no published data for low-level DHA supplementa- a power analysis, a blinded review of the data was undertaken after the first 100 analysis. Sample size was increased to 350 after the blinded analysis.		
	Funding: OmegaTech Inc, Colorado, USA			
	Declarations of interest: not reported			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Low risk	Computer-generated randomisation schedule		
Allocation concealment (selection bias)	Unclear risk	Not reported		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "double blinded"		
Blinding of outcome assessment (detection bias)	Low risk	Outcome assessments were blinded		



Smuts 2003a (Continued)

All outcomes

Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Overall 59/350 (16.9%) lost to follow-up:
		DHA-enriched eggs group lost 34/176 (19.3%):
		• 6 moved
		7 withdrew consent
		2 never received eggs
		6 birthed elsewhere
		 1 second pregnancy
		• 1 low age
		• 11 unknown reasons
		Ordinary eggs group lost 25/174 (14.4%):
		• 4 moved
		• 5 withdrew consent
		 1 never received eggs
		• 1 birthed elsewhere
		 1 second pregnancy
		• 13 unknown reasons
Selective reporting (reporting bias)	Low risk	Most expected outcomes were reported.
Other bias	Low risk	Baseline characteristics similar in each group.

Smuts 2003b

Smuts 2003b	
Methods	RCT: parallel (feasibility study)
Participants	52 women randomised (52 randomised to the 2-egg groups: 25 to the regular-egg group and 27 to the high-DHA egg group. (Another 21 women consented to the study but were not randomised and were not given eggs (low-egg intake group).)
	Inclusion criteria: women 24-28 weeks pregnant by obstetric assessments (either date of last menstrual period or US), aged 16-35 years, were accessible by telephone, and planned to give birth at the Regional Medical Center (Memphis, TN)
	If women said they ate eggs, they were asked for informed consent to be randomised to ordinary or high-DHA eggs.
	Exclusion criteria: any chronic illness, PIH, PE, or pregnancy-induced diabetes at the time of enrolment. Women were excluded if they had > 4 prior pregnancies.
	Characteristics: women were mainly African-African and rarely consumed fish; however they commonly consumed eggs.
	Setting: Regional Medical Center (Memphis, TN), USA (trial recruitment dates not reported)
Interventions	OMEGA-3-ENRICHED FOOD: high-DHA eggs versus ordinary eggs
	Group 1: high-DHA eggs (135 mg DHA/egg): total number = 27 randomised (18 could be analysed)
	Group 2: ordinary eggs (18 mg DHA/egg): total number = 25 randomised (19 could be analysed)



Smuts 2003b (Continued)

Timing of supplementation: egg consumption started at ~27 weeks' gestation and continued for ~13 weeks

During the course of the study, women were sent 2 dozen eggs (i.e. 24 eggs) every 2 weeks by courier. After the first delivery, they were interviewed before each subsequent delivery and asked how many eggs they had consumed. In addition, the unused eggs were returned by the courier, counted, the number recorded, and the eggs destroyed. Women were asked to keep a written record of their egg intake on forms supplied to them and to return these with the uneaten eggs; however, few were compliant with this request.

DHA + EPA dose/day: low: up to 135 mg DHA; EPA not stated

Outcomes

Women/birth: GA, GWG, caesarean section, gestational diabetes, maternal plasma and RBC lipids just prior to birth, maternal antibiotics, preterm birth, low birthweight, placental weight, PE, birthweight, length at birth; head circumference at birth, birthweight < 37 weeks

Babies/infants/children: newborn plasma and RBC lipids; admission to NICU/Special Care Unit; not routine hospital care, meconium

Notes

Funding: OmegaTech, Inc (Boulder, CO) supplied both ordinary and high-DHA eggs (Gold Circle Farms).

Declarations of interest: not reported

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Method of sequence generation not reported - "using a randomization in blocks of 6 to ensure that the groups remained relatively balanced".
Allocation concealment (selection bias)	Unclear risk	Quote: "women were randomized to the two egg groups"; no further information provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Quote: "The ordinary and high-DHA eggs had white shells but came in cartons of different colors. Carton color remained the same throughout the study" indicating that study personnel and possibly women could deduce which group they were in.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	9/27 in the high DHA and 7/25 in the control group lost to follow-up; reasons for losses not clearly reported.
Selective reporting (reporting bias)	Unclear risk	No protocol or trial registration entry; unable to determine
Other bias	Unclear risk	Women assigned to consume ordinary eggs were significantly older than those in the high DHA egg group.

Su 2008

Methods	RCT: parallel: NCT00618865
Participants	36 women randomised



Su 2008 (Continued)

Inclusion criteria: pregnant women aged 18-40 years, with major depressive disorder (DSM-IV) with onset between 16 weeks and 32 weeks GA; and to not have taken psychotropic agents for at least 1 month, to have a score of at least 18 on the 21-item HAM-D at screening; and to have good physical health as determined by medical history, physical examination, blood laboratory results, electrocardiogram, chest radiography and urinalysis. No psychotropic agents were given during the study period.

Exclusion criteria: DSM-IV diagnosis of bipolar disorder, psychotic disorder, or substance abuse/dependence or any Axis II diagnosis of borderline or antisocial personality disorder.

Setting: China Medical University Hospital, Taiwan (June 2004 to June 2006).

Interventions

SUPPLEMENTATION: DHA + EPA versus placebo

Group 1: omega-3 LCPUFA (3.4 g/day; 2.2 g EPA and 1.2 g DHA) (produced from Menhaden fish body oil concentrate). Total number randomised: n = 18 (13 completed)

Group 2: placebo: 5 identical gelatin capsules per day (olive-oil ethyl-esters). Total number randomised: n = 18 (11 completed)

Timing of supplementation: 8 weeks to ~30-32 weeks GA

Before random assignment, all consenting participants took part in a placebo trial for 1 week – those showing a decrease of 20% or more in HAM-D scores (placebo responders) were not permitted to proceed to the randomisation stage.

All capsules (fish oil and placebo) were vacuum deodorised, amended by blending with orange flavour and supplemented with tertiary butylhydroquinone (0.2 mg/g) and tocopherols (2 mg/g) as antioxidants

DHA + EPA dose/day: high: 1.2 g DHA + 2.2 g EPA

Outcomes

Women/birth: HAM-D (every other week); EPDS (Taiwanese version); Beck Depression Inventory and brief adverse effect checklist at week -1 (lead-in period), week 0 (baseline) and weeks 2, 4, 6 and 8; blood samples taken at week 1 and week 8 for omega-3 PUFA analysis (EPA; DHA)

"No obstetric complication was noted in any participant"

Babies/infants/children: "all the newborns were normal in general physical and neurobehavioral examination at birth"

Notes

Funding: National Science Council, Department of Health, and China Medical University and Hospital,

Declarations of interest: none reported

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Placebo used
Blinding of outcome assessment (detection bias)	Low risk	Not reported, but probably done.



Su 2008 (Continued) All outcomes

Incomplete outcome data (attrition bias) All outcomes High risk The intention-to-treat population included all women who had a baseline and at least 1 post baseline observation. The per protocol population included all women who completed 8 weeks of treatment. 12/36 (33%) lost to follow-up: Omega-3: 5/18 (28%) lost to follow-up: • 2 did not return • 3 'unsatisfactory response' Placebo: 7/18 (39%) lost to follow-up: • 2 did not return • 3 'unsatisfactory response' • 1 severe nausea Selective reporting (reporting (reporting bias) Only depressive symptoms, adverse events and omega-3 status assessed, e.g. no birth outcomes reported numerically. Other bias Unclear risk Baseline characteristics similar between study groups. Not clear what constituted "unsatisfactory response" (3 women in each group).			
The per protocol population included all women who completed 8 weeks of treatment. 12/36 (33%) lost to follow-up: Omega-3: 5/18 (28%) lost to follow-up: 2 did not return 3 'unsatisfactory response' Placebo: 7/18 (39%) lost to follow-up: 2 did not return 3 'unsatisfactory response' 1 severe nausea Selective reporting (reporting (reporting bias) Only depressive symptoms, adverse events and omega-3 status assessed, e.g. no birth outcomes reported numerically. Other bias Unclear risk Baseline characteristics similar between study groups. Not clear what constituted "unsatisfactory response" (3 women in each	(attrition bias)	High risk	
Omega-3: 5/18 (28%) lost to follow-up:			
2 did not return 3 'unsatisfactory response' Placebo: 7/18 (39%) lost to follow-up: 2 did not return 3 'unsatisfactory response' 1 severe nausea Selective reporting (reporting (reporting bias) Only depressive symptoms, adverse events and omega-3 status assessed, e.g., no birth outcomes reported numerically. Other bias Unclear risk Baseline characteristics similar between study groups. Not clear what constituted "unsatisfactory response" (3 women in each			12/36 (33%) lost to follow-up:
Placebo: 7/18 (39%) lost to follow-up: 2 did not return 3 'unsatisfactory response' 1 severe nausea Selective reporting (reporting bias) Only depressive symptoms, adverse events and omega-3 status assessed, e.g. no birth outcomes reported numerically. Other bias Unclear risk Baseline characteristics similar between study groups. Not clear what constituted "unsatisfactory response" (3 women in each			Omega-3: 5/18 (28%) lost to follow-up:
Selective reporting (reporting bias) High risk Only depressive symptoms, adverse events and omega-3 status assessed, e.g. no birth outcomes reported numerically. Other bias Unclear risk Baseline characteristics similar between study groups. Not clear what constituted "unsatisfactory response" (3 women in each			Placebo: 7/18 (39%) lost to follow-up:
porting bias) no birth outcomes reported numerically. Other bias Unclear risk Baseline characteristics similar between study groups. Not clear what constituted "unsatisfactory response" (3 women in each			3 'unsatisfactory response'
Not clear what constituted "unsatisfactory response" (3 women in each		High risk	
	Other bias	Unclear risk	Baseline characteristics similar between study groups.

Taghizadeh 2016

Methods	RCT: IRCT201506085623N43		
	(also IRCT201507035623N47 (Asemi 2015; Jamilian 2016)		
Participants	60 women randomised		
	Inclusion criteria: women with GDM (diagnosed with 'one-step' 2-hour 75 g OGTT at 24-28 weeks' gestation using ADA 2014 criteria, i.e. plasma glucose meeting any of the following criteria: fasting plasma glucose ≥ 92 mg/dL, 1-hour OGTT ≥ 180 mg/dL and 2-hour OGTT ≥ 153 mg/dL), who were not taking orahypoglycaemics, aged 18-40 years, without prior diabetes.		
	Exclusion criteria: premature preterm rupture of membranes, placental abruption, PE, eclampsia, hypo- and hyperthyroidism, urinary-tract infection, smokers, women with kidney or liver diseases, and women commencing insulin therapy during intervention.		
	Setting: Kosar Clinic, Kashan, Iran (study conducted May 2015 to July 2015)		
Interventions	SUPPLEMENTATION: omega-3 (ALA) + vitamin E versus placebo		
	Group 1: omega-3 and vitamin E co-supplementation (1000 mg omega-3 fatty acids from flaxseed oil; 400 mg α -linoleic acid, plus 400 IU vitamin E): total number randomised: n = 30		
	Group 2: placebo: colour, shape, size, and packaging identical to combined fatty acids and vitamin E pearl; total number randomised: n = 30		
	Timing of supplementation: 6 weeks from trial entry (24-28 weeks GA)		



Taghizadeh 2016 (Continued)

All women: 400 μ g/day vitamin B9 from the beginning of pregnancy and 60 mg/day ferrous sulphate from the second trimester. Although the intervention was for 6 weeks only, all women were followed up to the time of the birth (mean follow-up ~13 weeks)

DHA + EPA dose/day: not applicable

Outcomes

Women/birth: fasting plasma glucose; serum insulin concentrations; homeostasis models of assessment-estimated insulin resistance and beta cell function; quantitative insulin sensitivity checklist; serum triglycerides; VLDL-cholesterol; low-density lipoprotein; HDL-cholesterol; total antioxidant capacity, nitric oxide, plasma malondialdehyde concentrations, plasma glutathione and serum high-sensitivity CRP concentrations; maternal BMI; GWG, DBP and SBP, need of insulin therapy after intervention, maternal hospitalisation, GA, polyhydramnios, preterm birth, PE; 3 day dietary intakes (baseline, week 1, 3, 5 and end of trial); maternal adherence; birthweight; birth length and head circumference at birth

Babies/infants/children: hyperbilirubinaemia; 1- and 5-minute Apgar scores, newborn hospitalisation rates; hypoglycaemia

Notes

Funding: supplements and placebos were supplied by Barij Essence Pharmaceutical Company, Kashan, Iran; grant from KUMS, Kashan, Iran.

Declaration of interests: none declared

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated random number sequence
Allocation concealment (selection bias)	Low risk	Quote: "randomly allocated into two groups"; a trained midwife assigned participants and "randomization and allocation were hidden from the researchers and patients until the main analyses were completed"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "double-blind placebo-controlled"
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Not specifically reported, but probably done.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	3/30 women in each group withdrew due to personal reasons.
Selective reporting (reporting bias)	Low risk	No apparent selective outcome reporting bias.
Other bias	Low risk	Baseline characteristics similar

Tofail 2006

Methods	RCT (parallel)
Participants	400 women randomised



Гofail 2	.006 (Continued)
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Inclusion criteria: pregnant women at 25 weeks' gestation

Exclusion criteria: not reported

Characteristics: 28% of mothers were under-nourished; 82% of babies predominantly breastfed

Setting: Dhaka, Bangladesh (January to March 2000): participants from low-income households (as-

sessed via a household survey)

Interventions

SUPPLEMENTATION: DHA + EPA versus soy oil

Group 1: fish oil: 4 capsules (1 g in each) as a single daily dose. Total daily fish oil supplement contained 1.2 g of DHA and 1.8 g of EPA. Total number randomised: n = 200

Group 2: soy oil: 4 capsules (1 g in each) as a single daily dose. Soy oil supplement contained 2.25 g of LA, and 0.27 g of ALA. Total number randomised: n = 200

Timing of supplementation: 25 weeks GA to birth

DHA + EPA dose/day: high: 1.2 g DHA + 1.8 g EPA

Outcomes

Women/birth: length of gestation; birthweight; birth length; head circumference at birth; ponderal index; preterm birth; stillbirth; neonatal death; perinatal mortality; infant death; low birthweight

Babies/infants/children: duration of exclusive breastfeeding; at 10 months: Bayley Mental developmental index; Bayley psychomotor developmental index; Behaviour (Wolke) (Approach; Activity; Cooperation; Emotional tone; Vocalisation) using 0-9 scales where higher scores are better; head circumference; weight-for-height z-score; weight-for-age z-score; height-for-age z-score; HOME (modified for Bangladesh)

Notes

Tofail 2012 was a follow-up study but results were not reported by supplementation/no supplementa-

Funding: World Bank

Declarations of interest: not reported

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Capsules were identical in appearance
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Two female psychologists, unaware of the infant's group assignment, tested all infants"
Incomplete outcome data (attrition bias)	High risk	76/400 women were lost before birth due to out-migration or refusal to take capsules (fish oil: 41/200, soy oil: 35/200).
All outcomes		Of the mothers included at birth (159 fish oil; 165 soy oil), further losses occurred after birth (stillbirth: 8 and 6; early neonatal death: 4 and 5; outmigration: 19 and 26; infant death: 3 and 4);



Tofail 2006 (Continued)		Therefore 249 infants (62%) were included in the 10-month follow-up (125/200 in the fish oil, and 124/200 in the soy oil group).
Selective reporting (reporting bias)	Low risk	No apparent risk of selective reporting.
Other bias	Unclear risk	Baseline characteristics: mothers in the fish oil group were younger and had fewer children.

Valenzuela 2015

Methods	RCT
Participants	40 women randomised
	Inclusion criteria: women aged 22–35 years; GA of at least 22-25 weeks according to the date of the last menstrual period and confirmed by US; 1–4 children
	Exclusion criteria: women with a history of drug or alcohol consumption; a diet including polyunsaturated fatty acids (PUFA, ALA supplements) or LCPUFA (EPA and or DHA supplements); underweight as defined by the Chilean chart for pregnant women; history of twins or of suffering from chronic diseases such as diabetes, arterial hypertension, obesity, or other illness that could affect fetal growth
	Characteristics: recruited women were mostly of low- and middle-socioeconomic status (according to the ESOMAR); all were of Hispanic origin.
	Setting: Obstetrical and Gynecology Health Service of the University of Chile Hospital, Chile (study conducted January 2012-December 2013)
Interventions	SUPPLEMENTATION: ALA versus soy oil
	Group 1: chia oil (provided in 240 mL bottles, 4500 mL in total). Women were requested to consume 16 mL/day (10.1 g ALA/day). Total number randomised: n = 19
	Group 2: sunflower/soy oil (provided in 240 mL bottles, 4500 mL in total). Women were requested to consume 16 mL/day. Total number randomised: n = 21
	Timing of supplementation: from the 6th month of pregnancy until the 6th month of nursing (total of 9 months)
	All women: received a complete nutritional interview including nutritional diagnosis and counselling according to the dietary guidelines for pregnant women, were given plastic teaspoons (4 mL) for measuring consumption of intervention/control oil; received a dietary record to register the daily consumption of oil; visited weekly to assess oil consumption
	DHA + EPA dose/day: not applicable
Outcomes	Women/birth: maternal diet (energy and composition of diet ingested by mothers during pregnancy and nursing); fatty acid composition of erythrocyte phospholipids of mothers during pregnancy and nursing; fatty acid profile of breast milk; length of gestation
	Babies/infants/children: birthweight; head circumference at birth
Notes	Funding: not reported
	Declarations of interest: none declared
Risk of bias	



Valenzuela 2015 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "the pregnant women were randomly assigned to either the control group (n=21) or to the experimental group that received the dietary supplementation with chia oil"
Allocation concealment (selection bias)	Unclear risk	Quote: "the pregnant women were randomly assigned to either the control group (n=21) or to the experimental group that received the dietary supplementation with chia oil"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Not reported
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing outcome data: all women randomised (to intervention and control), and their infants, were included for analyses.
Selective reporting (reporting bias)	Unclear risk	Few outcomes reported; additionally, without access to a published protocol it is not possible to assess selective outcome reporting confidently.
Other bias	Low risk	Data reported on the characteristics of women (age; weight; BMI; socioeconomic status) suggests groups were similar at baseline.

Van Goor 2009

Methods	RCT: ISRCTN58176213	
Participants	183 women randomised (3 groups - see below)	
	Inclusion criteria: apparently healthy women with a low-risk first or second pregnancy	
	Exclusion criteria: preterm births (< 37 weeks) or GA > 42 weeks and any maternal or neonatal complications; vegetarians or vegans	
	Characteristics: low background DHA intake (fish intake once a week)	
	Setting: Groningen, the Netherlands: assumed to be during antenatal care (recruited by midwives and obstetricians)	
Interventions	SUPPLEMENTATION: DHA + AA versus DHA versus placebo	
interventions	SUPPLEMENTATION: DHA + AA versus DHA versus placebo	
interventions	Group 1: DHA + AA (220 mg each/day): total number randomised: 58 (41)	
merventions	·	
merventions	Group 1: DHA + AA (220 mg each/day): total number randomised: 58 (41)	
merventions	Group 1: DHA + AA (220 mg each/day): total number randomised: 58 (41) Group 2: DHA (220 mg/day) + 1 soy capsule/day: 63 (41) Group 3: placebo (2 soy capsules/day); equivalent to 535 mg LA/day: total number randomised: n = 62	



Van Goor 2009 (Continued)

Outcomes

Women/birth: preterm birth; GA; GDM; fatty acids in plasma (at 16 and 36 weeks GA); EPDS at 16 and 36 weeks GA and 6 weeks pp; blues questionnaire within 1 week pp; DHA and AA in breastmilk (2 and 12 weeks pp); sleep quality (36 weeks GA and 4 weeks pp); Obstetric Optimality Score (74 items covering socioeconomic status, non-obstetric conditions during pregnancy, obstetric history, diagnostic and therapeutic measures, parturition and neonatal condition immediately after birth; food frequency questionnaires (16 and 36 weeks GA, 12 weeks pp); fatty acids (in umbilical cord); birthweight; duration of breastfeeding

Babies/infants/children: general movement quality; NOS; Hempel neurological examination; BSID II (Dutch version); weight at 18 months; height at 18 months; head circumference at 18 months; cerebral palsy

(EPDS and Hempel reported as median and ranges only)

Notes

Funding: Friesland Foods, the Netherlands

Declarations of interest: none declared

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Block randomisation; not further reported
Allocation concealment (selection bias)	Unclear risk	Quote: "randomized into three groups"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Double-blind placebo-controlled
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	58/183 (32%) "lost interest" (23 placebo; 20 DHA; 15 DHA/AA); 6 pregnancy complications (3 placebo, 1 DHA, 2 DHA/AA), leaving 119 for analysis
Selective reporting (reporting bias)	Low risk	Most expected outcomes were reported (although numbers per group were not reported for EPDS > 12 and blues score > 12)
Other bias	Low risk	Baseline characteristics similar

Van Winden 2017

Methods	RCT (pilot study)	
Participants	14 women randomised (2 groups, see below)	
	Inclusion criteria: women with diet-controlled GDM, between 24-30 weeks' gestation	
	Exclusion criteria: not reported in conference abstract	
	Setting: not reported in conference abstract	



Van Winden 2017 (Continued)

Interventions

SUPPLEMENTATION: DHA + EPA versus placebo

Group 1: commercial fish oil (2200 mg/day; 1300 mg EPA and 900 mg DHA). Total number randomised: n = 7 (4 completed)

Group 2: placebo, identical (no further details in conference abstract). Total number randomised: n = 7 (5 completed)

Timing of supplementation: 6 weeks (start 24-30 weeks' gestation)

DHA + EPA dose/day: high: 900 mg DHA + 1300 mg EPA

Outcomes Women/birth: maternal adverse effects (narrative report only)

Notes No outcomes included that could be used in meta-analysis

Funding: not reported.

Declaration of interests: none declared

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "randomised in a prospective, double-blind fashion to receive either 2200 mg daily of commercial fish oil or an identical placebo."
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	14 GDM patients randomised to treatment with fish oil ($n = 7$) or placebo ($n = 7$); 36% (3 in the treatment group, 2 in the placebo group) did not complete the treatment course due to intolerance.
Selective reporting (reporting bias)	Unclear risk	Confident assessment was not possible with conference abstract report only.
Other bias	Unclear risk	Information on methods insufficient to assess this domain confidently.

Vaz 2017

Methods	RCT: NCT01660165	
Participants	60 women randomised	
	Inclusion criteria: women 5-13 weeks' gestation; aged 20-40 years at the time of enrolment; free from any known chronic diseases; residing in the study catchment area; intending to continue prenatal care in the public health centre; classified as at risk for postpartum depression (reported a past history of depression or presented a depressive symptom score at baseline ≥ 9 based on the EPDS).	



Vaz 2017 (Continued)

Exclusion criteria: depressed or presented with psychotic symptoms; past history of mania; or at suicidal risk; or taking any psychiatric medication (as anti-depressives and anxiolytics); or being seen by a psychologist or psychiatrist. Women taking any oil supplementation (such as fish oil, flaxseed oil or cod liver oil), were also excluded.

Setting: Rio de Janeiro, Brazil (enrolled November 2009-October 2011 and the last follow-up visit occurred in July 2012).

Interventions

SUPPLEMENTATION: omega-3 versus placebo

Group 1: omega-3: 6 capsules/day, 1 g each, for a total dose of 1.8 EPA and 0.72 g DHA (The capsules were deodorised, and contained 0.2 mg/g of vitamin E as antioxidant. Women were advised to take 3 capsules at lunch and 3 capsules at dinner): n = 32

Group 2: placebo: n = 28

Timimg of supplementation: second trimester to 16 weeks postpartum

All women: 400 μ g/day of folic acid from the beginning of pregnancy, and 60 mg/day ferrous sulphate from the 2nd trimester until birth, as provided during standard prenatal care in Brazil. Participants were asked to not alter their usual dietary habits and not consume any supplements other than the ones provided by the research team and antenatal care service.

DHA + EPA dose/day: high: 720 mg DHA + 1.8 g EPA

Outcomes

Women/birth: length of gestation; depression (at 30-32 weeks' gestation and 4-6 weeks postpartum); adverse effects (gastrointestinal and non-gastrointestinal); EPA, DHA and omega-6/omega-3 concentrations

Babies/infants/children: birthweight

The EPDS was used to measure depression, and scored at 5-13 weeks' gestation, 22-24 weeks' gestation (baseline for RCT), 30-32 weeks' gestation, and 4-6 weeks postpartum. The Portuguese version of the scale was validated in a sample of mothers from Pelotas, southern Brazil. EPDS score of \geq 11 was the cut off for depressive symptoms.

Notes

The fish oil supplement was manufactured by Galena Nutrition Química Industrial, São Paulo, Brazil.

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Declarations of interest: "The authors declare they have no competing interests".

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "The randomization was performed by a researcher not involved in the data collection using the participant identification (subject ID) after stratification for EPDS score and previous history of depression".
Allocation concealment (selection bias)	Unclear risk	Quote: "The randomization was performed by a researcher not involved in the data collection using the participant identification (subject ID) after stratification for EPDS score and previous history of depression".
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Participants and all research assistants and technicians responsible for running both the cohort study and the RCT were blinded to the study allocation"



Vaz 2017 (Continued)		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information provided by the authors on whether assessors were blinded, for any of the outcomes reported.
Incomplete outcome data (attrition bias) All outcomes	High risk	Of the 32 women randomised to the intervention, 6 did not receive intervention (2 transferred antenatal care to another health centre; 3 exceeded 13 weeks of gestation at the baseline), and 7 dropped out before the end of pregnancy time-point, leaving 17/32 women (53%) contributing data to the analysis for end of pregnancy/birth outcomes. A further 2 women were lost to follow-up before the 4-6-week postpartum data collection time point (did not visit); therefore only 15/32 women (47%) of the women randomised to the intervention were included in the analysis for the postpartum outcomes. Of the 28 women randomised to the control, 4 did not receive intervention (1 transferred prenatal care to another health centre; 1 miscarriage; 2 missed 2nd trimester visit), and 5 dropped out before the end of pregnancy time-point, leaving 17/28 women (60%) contributing data to the analysis for end of pregnancy/birth outcomes.
Selective reporting (reporting bias)	Unclear risk	Reporting of birthweight and GA incomplete (the total number of participants in the groups is not reported).
Other bias	Unclear risk	Baseline characteristics similar between except for ethnicity.

Abbreviations.

2D: 2-dimensional

AA: arachidonic acid; **ALA:** alpha-linolenic acid; **AIDS:** acquired immune deficiency syndrome; **ADA:** American Diabetes Association; **ASQ:** Ages and Stages Questionnaire

BASC-PRS: Behaviour Assessment for Children: Parent Rating Scale; **BDI:** Beck Depression Inventory; **BMI:** body-mass index; **BP:** blood pressure; **BPD:** bronchopulmonary dysplasia; **BSID:** Bayley Scales of Infant Development

CES-D: Center for Epidemiologic Studies Depression Scale; **CRP:** C-reactive protein

DBP: diastolic blood pressure; **DHA:** docosahexaenoic acid; **DNA:** deoxyribonucleic acid; **DPA:** docosapentaenoic acid;**DSM:** Diagnostic and Statistical Manual;

EPA: eicosapentaenoic acid; **EPDS:** Edinburgh Postnatal Depression Scale; **ERG:** electroretinography; **ERP:** electroencephalography/event-related potentials; **ESOMAR:** European Society for Opinion and Marketing Research; **ETA:** eicosatetranoic acid; **EU:** European Union **FADS:** fatty acids desaturase; **FDA:** Food and Drug Administration; **FFQ:** food frequency questionnaire; **FiO2:** fraction of inspired oxygen;

FPG: fasting plasma glucose

GA: gestational age; GDM: gestational diabetes mellitus; GLA: gamma linolenic acid; GWG: gestational weight gain

HAM-D: Hamilton Rating Scale Depression; **Hb:** haemoglobin; **HbA1c:** glycated haemoglobin; **HC:AC:** head circumference/abdominal circumference; **HDL-cholesterol:** high density lipoprotein-cholesterol; **HDRS:** Hamilton Rating Scale for Depression; **HLA:** human leukocyte antigen; **HIV:** human immunodeficiency virus; **HOMA-B:** Homeostatic Model Assessment of Beta cell function; **HOMA-IR:** Homeostatic Model Assessment of Insulin Resistance; **HOME:** home observation measurement of the environment

IgE: immunoglobulin E; **IGF:** Insulin-like growth factor; **IL:** interleukin;**IQ:** intelligence quotient; **IQR:** interquartile range;**IU:** international units; **IUGR:** intrauterine growth restriction; **IVH:** intraventricular haemorrhage

K-ABC: Kaufman Assessment Battery for Children; K-CPT: Conners Kiddie Continuous Performance Test

LA: linoleic acid; LC: long-chain; LCPUFA: long-chain polyunsaturated fatty acid; LGA: large-for-gestational age

MADRS: Montgomery-Åsberg Depression Rating Scale; MD: Maryland; MDI: mental development index (BSID); MRI: magnetic resonance imaging; mRNA: messenger ribonucleic acid; MTHF: methyltetrahydrofolic acid

n-3: omega-3; **n-6:** omega-6; **n-9:** omega-9; **NBAS:** Neonatal Behavioral Assessment Scale; **NCT:** Clinical Trials.gov registry; **NEC:** necrotising enterocolitis; **NICU:** neonatal intensive care unit; **NIDDK:** National Institute of Diabetes and Digestive and Kidney Diseases; **NIH:** National Institutes of Health; **NIMH:** National Institute of Mental Health; **NOS:** neurological optimality score; **NSAIDS:** nonsteroidal anti-inflammatory drugs

OGTT: oral glucose tolerance test

PDI: psychomotor development index (BSID); **PDI:** postnatal depression inventory; **PDSS:** Postpartum Depression Screening Scale;**PE:** pre-eclampsia; **PG:** prostaglandin; **PI:** pulsatility index; **PIH:** pregnancy-induced hypertension; **PONCH:** Pregnancy Obesity Nutrition and Child Health Study; **PPH:** postpartum haemorrhage; **PPROM:** preterm prelabour rupture of membranes

QUICKI: quantitative insulin sensitivity check index

RBC: red blood cell; **RCT:** randomised controlled trial; **RDS:** respiratory distress syndrome;**RI:** resistance index; **ROP:** retinopathy of prematurity



S/D: systolic/diastolic; **SBP:** systolic blood pressure; **SD:** standard deviation; **SGA:** small-for-gestational age; **SiPS:** Salmon in Pregnancy

Study;**sLORETA:** standardised low-resolution brain electromagnetic tomography **T1D:** type 1 diabetes; **TLR:** toll-like receptor; **TOVA:** Test of Variables of Attention

US: ultrasound; **U.S. FDA:** United States Food and Drug Administration

VLDL-cholesterol: very low density lipoprotein-cholesterol

WIC: Women, Infants and Children

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Escobar 2008	Trial registered but no participants were recruited
Fievet 1985	Intervention (evening primrose oil) is not an omega-3 fatty acid
Gholami 2017	Not randomised
Herrera 1993	Intervention (linoleic acid) is an omega-6 fatty acid
Herrera 1998	Intervention (linoleic acid) is an omega-6 fatty acid
Herrera 2004	Intervention (linoleic acid) is an omega-6 fatty acid
Lauritzen 2004	Women supplemented with omega-3 fatty acids only during lactation
Marangell 2004	Non-randomised pilot study
Morrison 1984	Intervention (linoleic acid) is an omega-6 fatty acid
Morrison 1986	Trial does not appear to be randomised.
Nishi 2016	Non-randomised pilot study
Starling 1990	Not randomised: "divided into two groups"
Valentine 2013	Lactating women only (human milk donor pilot study)
Velzing-Aarts 2001	Not randomised
Yelland 2016	Methodological study across several trials

Characteristics of studies awaiting assessment [ordered by study ID]

Farahani 2010

Methods	RCT	
Participants	120 women randomised	
Interventions	1) salmon fish oil capsule (1000 mg/day) from 16 weeks GA to birth 2) standard prenatal care	
Outcomes	Systolic and diastolic blood pressure	



Farahani 2010 (Continued)

Notes In Arabic

Gopalan 2004

Methods	Comparison of 3 groups - not clear if this was a randomised study	
Participants	900 pregnant women; low socioeconomic status, attending government antenatal centres in India	
Interventions	1) 900 mg alpha linolenic acid daily from 22 weeks GA + iron/folate supplementation from 20 weeks' gestation	
	2) iron/folate (100 mg/500 μ g) daily from 20 weeks GA	
	3) control	
Outcomes	Birthweight; low birthweight; preterm birth < 37 weeks	
Notes	Abstract only - no details of how groups were allocated	

Jamilian 2018

Methods	RCT: IRCT201610015623N90	
Participants	40 women aged 18-40 diagnosed with GDM, based on the American Diabetes Association guide- lines, without prior history of diabetes	
Interventions	 1) 1000 mg fish oil capsules, containing 180 mg EPA and 120 mg (DHA) (n = 20) 2) placebo (n = 20) twice a day for 6 weeks from ~25 weeks gestation 	
Outcomes	PPAR-v gene expression, low-density lipoprotein receptor (LDLR), interleukin-1 (IL-1), interleukin-8 (IL-8), and TNF-a; birth outcomes; infant outcomes	
Notes	need to clarify if this is a separate study or an additional report	

Kadiwala 2015

Methods	Abstract
Participants	
Interventions	
Outcomes	
Notes	Abstract only; no further information available



aitinen 2013	
Methods	RCT (4 arms): NCT01922791
Participants	Recruitment target: 440 women.
	Inclusion criteria: less than 17 gestational weeks; overweight; healthy.
	Exclusion criteria : diabetes (Type 1 or 2); coeliac disease; increased bleeding tendency.
	Setting: Turku University Hospital, Finland.
Interventions	Intervention groups
	1) probiotic dietary supplements; 2) women will receive fish oil; and 3) probiotics and fish oil (from early pregnancy until 6 months after birth, no further information provided).
	Control group
	4) placebo (from early pregnancy until 6 months after birth).
Outcomes	Primary outcomes : GDM (at 24-28 weeks' gestation; fasting glucose levels (assessed at third trimester of pregnancy); prevalence of allergy in child (at 12 and 24 months of age).
	Other outcomes : need for medication for management of GDM (insulin or metformin); body composition of mother (during and after pregnancy); immunologic and metabolic markers (during and after pregnancy); and faecal microbiota (during and after intervention); body composition, growth, development and metabolic markers of child 9 (0-12 months).
Notes	awaiting to see if there are further reports of other outcomes

Lazzarin 2009

Methods	RCT
Participants	60 women with unexplained recurrent miscarriage
Interventions	1) aspirin (20 women)
	2) omega-3 LCPUFA (20 women)
	3) aspirin and omega-3 LCPUFA (20 women)
Outcomes	Doppler uterine artery pulsatility index
Notes	awaiting to see if there are further reports of other outcomes

Parisi 2013

Methods	RCT
Participants	35 healthy singleton nulliparous pregnant women (20-22 weeks' gestation)
Interventions	Not reported
Outcomes	Not reported



Parisi 2013 (Continued)

Notes Insufficient detail in abstract	Notes	Insufficient detail in abstract
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Pavlovich 1999

Methods	RCT
Participants	60 pregnant women at high risk of developing placental insufficiency
Interventions	Picasol (omega-3 fatty acid preparation) placebo
Outcomes	Triglyceride and cholesterol concentrations
Notes	In Russian

Sajina-Stritar 1994

Methods	"comparative trial"; "randomly allocated"
Participants	20 women at high risk of gestational hypertension
Interventions	Omega-3 fatty acids versus aspirin
Outcomes	Hypertension and other pregnancy outcomes
Notes	Results not reported numerically

Sajina-Stritar 1998

Methods	"randomly allocated"
Participants	48 women at high risk of gestational hypertension
Interventions	1) aspirin (12 women)
	2) fish oil (11 women)
	3) no treatment (25 women)
Outcomes	Gestational hypertension
Notes	Not clear that this study is a RCT

Salvig 2009

Methods RCT



Salvi	g 2009	(Continued)
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Participants	190 women with a previous preterm birth < 34 weeks GA
Interventions	Omega-3 (2.7 g/day) versus olive oil
Outcomes	Preterm birth; GA
Notes	Abstract; states only that no differences were found and no numeric results were reported

Salzano 2001

Methods	"divided into 2 unequal groups"
Participants	65 primigravid women at risk of hypertension
Interventions	calcium, linoleic acid, mono and polyunsaturated fatty acids (40 women) no treatment (25 women)
Outcomes	Gestational hypertension
Notes	not clear if this is a RCT

Stoutjesdijk 2014

Methods	RCT (4-arm trial)
Participants	43 healthy women in first trimester of pregnancy, intending to breastfeed
Interventions	4 different doses of DHA/EPA
Outcomes	Maternal RBC and milk DHA/EPA concentrations at 4 weeks postpartum
Notes	no maternal or birth outcomes yet reported

Vahedi 2018

Methods	RCT
Participants	pregnant women
Interventions	1) fish oil supplementation
	2) control
Outcomes	maternal glucose concentrations, haemoglobin, haematocrit
Notes	not enough detail yet available



Vakilian 2010	
Methods	RCT
Participants	100 women with a singleton pregnancy, aged between 18-40, any acute or chronic diseases, antenatal care before 16 weeks' gestation, non-smoking
Interventions	1) omega 3 (1000 mg) from 16 weeks to 40 weeks' gestation
	2) no treatment
Outcomes	Lipid peroxide, thiol group, and ferric reducing antioxidant plasma
Notes	Completed but no English translation is available

Valentine 2014

Methods	RCT
Participants	90 pregnant women (19-20 weeks GA); > 18 years; diagnosed with hypertension; without bleeding disorders, lupus, autoimmune diseases, or presence of infant congenital (trisomy 13, 18, 21, urethral, gastrointestinal and cardiac defects)
Interventions	1) 200 mg DHA daily, from 18-20 weeks GA to 6 weeks postpartum
	2) 1000 mg DHA daily, from 18-20 weeks GA to 6 weeks postpartum
Outcomes	Primary: endothelial health (measures not reported); secondary: maternal homoeostasis (blood and cord blood concentrations of pro-inflammatory cytokines IL-6, I L-8, TNF a, and receptor sRAGE
Notes	May have been withdrawn/never commenced

Valenzuela 2017

Methods	abstract
Participants	
Interventions	
Outcomes	
Notes	not enough detail provided in the abstract

GA: gestational age

RCT: randomised controlled trial

Characteristics of ongoing studies [ordered by study ID]

Albert 2017

Trial name or title	Fish oil in pregnancy for a healthy start to life for the children of overweight mothers
Methods	RCT: ACTRN12617001078347p



Albert 2017 (Continued)

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Recruitment target: 160 women.

Inclusion criteria: age > 18 < 40 years; pregnant; BMI 30-45; singleton pregnancy; between 12-16 weeks of gestation.

Exclusion criteria: current use of tobacco or nicotine; illicit drugs or medications that influence blood pressure; lipid metabolism or insulin sensitivity. Women were also excluded if they had diabetes mellitus or chronic illnesses such as autoimmune disease or malignancy.

Setting: Auckland, New Zealand.

Interventions

Intervention group

3 g of omega-3 PUFA rich fish oil taken in capsules on each day of pregnancy and for 3 months during the breastfeeding period. Note that if the mother chooses to stop breastfeeding then supplementation will stop early, this will be at birth if the mother decides not to breast feed at all. Compliance will be assessed by return of unused capsules, and secondarily by measurement of omega-3 PUFA levels in maternal red blood cells. Note that the expected concentration of omega-3 PUFAs in this oil is 33% EPA/22% DHA, but this will be independently verified.

Control group

3 g of olive oil taken in capsules on each day of pregnancy and for 3 months during the breastfeeding period (if the mother chooses to breast feed)

Outcomes

Primary outcome: whole body percentage body fat, as measured by DXA scan (in the offspring).

Other outcomes: peripheral quantitative computed tomography derived measures of bone strength from the lower Tibia (in the offspring, at 2 weeks of age); birthweight; whole body percentage body fat, measured by DXA scan (in the offspring, at 3 months of age); HOMA-IR (in the offspring, at 3 months of age); weight in the offspring (at 12 months of age); HOMA-IR (in the mother, at 30 weeks' gestation); Ponderal Index (in the offspring, at 12 months of age); Insulin sensitivity as determined by a modified IVGTT with minimal modelling (offspring, 4-7 years of age).

Starting date

2 October 2017

Contact information

Dr Benjamin Albert, Liggins Institute, University of Auckland. E-mail: b.albert@auckland.ac.za

Notes

<u>Funding and collaborators</u>: A Better Start National Science Challenge, Funded by the Ministry of Business, Innovation and Employment (government body, New Zealand); Cure Kids (charity, New Zealand); Health Research Council (charity, New Zealand); Liggins Institute, University of Auckland.

Carlson 2017 ADORE

Trial name or title	ADORE
Methods	RCT (multi-centre, adaptive design, 3 arms): NCT02626299
Participants	Recruitment target: 1200 women.
	Inclusion criteria : women ≥ 18 years; 12–20 weeks of gestation; agree to consume study capsules and a typical prenatal supplement of 200 mg CHA; and available by telephone.
	Exclusion criteria : expecting multiple infants; gestational age at baseline < 12 weeks or > 20 weeks; unable or unwilling to agree to consume capsules until birth; unwilling to discontinue use of another prenatal supplement with DHA; and with allergy to any component of DHA product (including algae), soybean oil or corn oil.



Carlson 2017 ADORE (Continued)	
	Setting: Kansas, USA.
Interventions	Intervention group
	DHA supplements (800 mg/day), administered in 2 400 mg capsules, plus 1 200 mg/capsule per day of DHA that is a common amount in prenatal vitamins.
	Control group
	Standard care (200 mg/day of DHA) administered in 2 capsules (masked) containing half soybean oil and half corn oil equalling 800 mg. the soybean and corn oil combination does not contain DHA.
Outcomes	Primary outcome: early preterm birth < 34 weeks (baseline to 34 weeks).
	Other outcomes : change in plasmal soluble(s) RAGE concentration (baseline to 34 weeks); and adverse events (34 weeks).
Starting date	8 June 2016
Contact information	Dr Susan Carlson, University of Kansas Medical Center, USA, E-mail: scarlson@kumc.edu
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of Cincinnati, Ohia State University, Nationwide Children's Hospital.

Funding and collaborators: NICHD R01 HD083292, University of Kansas Medical Center, University

Carvajal 2014

Notes

Trial name or title	Docosahexaenoic acid (DHA) supplementation during pregnancy to prevent deep placentation disorders: a randomized clinical trial and a study of the molecular pathways of abnormal placentation prevention
Methods	RCT
Participants	2400 women 18 years or older; GA < 16 weeks
	Setting: Chile
Interventions	DHA (600 mg/day) from early pregnancy until the ed of pregnancy versus placebo
Outcomes	Composite of preterm birth < 34 weeks or pre-eclampsia before 34 weeks or severe fetal growth restriction < 34 weeks GA
Starting date	May 2015
Contact information	Jorge Carvajal, email: jcarva@med.puc.cl
Notes	

de Carvalho 2017

Trial name or title	Gestational obesity and interventions with probiotics or fish oil trial: GOPROFIT
Methods	RCT (4 arms)
Participants	80 women at 13 weeks GA, aged between 19 to 40, pre-pregnancy BMI > 29.9



de Carvalho 2017 (Continued)	Setting: Brazil
Interventions	1) DHA (100 mg) + EPA (137 mg) a day
	2) probiotic 1
	3) probiotic 2
	4) placebo
Outcomes	Inflammation
Starting date	March 2015
Contact information	Fátima Lúcia C Sardinha, email: sardinhaflc@nutricao.ufrj.br
Notes	

Dos Santos 2018

Trial name or title	Omega-3 supplementation during pregnancy to prevent postpartum depressive symptoms and possible effect on breastfeeding, child growth and development
Methods	RCT
Participants	80 women with postpartum depression score greater than or equal to 10 and need for medical treatment and/or medical follow-up
Interventions	1) fish oil (1000mg DHA and 400mg EPA per day)
	2) placebo capsules (olive oil)
	Both groups will be instructed to ingest 2 x 1000mg capsules per day for 16 weeks.
Outcomes	Prevention of postpartum depressive symptoms (Edinburgh Postnatal Depression Scale)
Starting date	August 2018
Contact information	Luana Caroline dos Santos, email: luanacstos@gmail.com
Notes	Universidade Federal de Minas Gerais

Dragan 2013

Trial name or title	The impact of EPA and DHA supplementation on the content of lipids in the pregnant women and the fetus
Methods	RCT
Participants	Recruitment target: 87 women.
	Inclusion criteria : healthy; with a singleton pregnancy; BMI < 25 kg/m ² ; willing to provide informed consent.



Dragan 2013 (Continued)	Exclusion criteria: pregnancy terminated as preterm birth; with chronic illness; gestational diabetes mellitus or pre-eclampsia. Setting: Bosnia, Herzegovina.
Interventions	Intervention group
	360 mg EPA (eicosapentanoic fatty acid) and 240 mg DHA (docosahexanoic fatty acid) per day during pregnancy, from baseline (14th week of gestation) until birth.
	Control group
	No supplementation of omega-3 fatty acids during pregnancy.
Outcomes	Primary outcomes : concentration of omega-3 fatty acids in total serum lipids, measured using gas chromatography at the end of pregnancy; concentration of omega-3 fatty acids in umbilical vein serum, measured using gas chromatography at time of birth; concentration of monounsaturated fatty acids in serum total lipids of umbilical vein serum, measured by gas chromatography at time of birth; concentration of monounsaturated fatty acids in serum total lipids of the mother's serum, measured by gas chromatography at the end of pregnancy; weight of mother, measured by weighing scale at recruitment, 20th week of gestation, 30th week of gestation and before birth. Secondary outcomes : weight of mother (measured at recruitment, 20 weeks of gestation, 30 weeks of gestation and before birth).
Starting date	May 2013
Contact information	Dr Soldo Dragon, Department of Obstetrics and Gynecology School of Medicine Mostar, Kralja Tvrt-ka, b.b. Mostar 88000, Bosnia, Herzegovina. E-mail:dragan.soldo3@tel.net.ba
Notes	Funding and collaborators: investigator initiated and funded.

FOPCHIN	
Trial name or title	FOPCHIN
Methods	RCT (3 arms): NCT02770456
Participants	Recruitment target: 5531 women
	Inclusion criteria: women 20 to 44 years and pregnant without known complications.
	Exclusion criteria : regular user of fish oil; regular user of NSAIDs; known twin pregnancy.
	Setting: China*
Interventions	Intervention group
	1) High-dose intervention group: 4 0.72-g gelatine capsules daily with fish oil providing 2.0 g/d Ic-n3FA, from 16-24 weeks' gestation until they have completed the preterm period (i.e. at 37 full gestation weeks) or until they deliver.
	2) Low-dose intervention group will receive 4 0.72-g gelatine capsules daily with a mixture of fish oil and olive oil providing 0.5 g/d Ic-n3FA, from 16-24 weeks' gestation until they have completed the preterm period (i.e. at 37 full gestation weeks) or until they deliver.
	Control group



FOPCHIN (Continued)	Placebo (4×0.72 -g gelatine capsules daily with olive oil providing 0 g/d Ic-n3FA), from 16-24 weeks' gestation until they have completed the preterm period (i.e. at 37 full gestation weeks) or until they deliver.
Outcomes	Primary outcome: preterm birth.
	No other outcomes specified.
Starting date	March 2008
Contact information	Dr Sjurdur F Olsen, Statens Serum Institut. E-mail: SFO@ssi.dk
Notes	Funding and collaborators: Centre for Fetal Programming, Denmark; Shanghai Institute of Planned Parenthood Research.
	* assumed, not specifically stated

Garg 2017	
Trial name or title	Omega-3 fish oil for the prevention of gestational diabetes
Methods	RCT: ACTRN12617000177358
Participants	Recruitment target: 74 women
	Inclusion criteria: < 14 weeks pregnant; aged 18-40; with any 1 of the following: a) PAPP-A between 0.3 and 0.6 MoM in their Nuchal Translucency Scan b) previous history of gestational diabetes c) at risk of developing gestational diabetes
	Exclusion criteria : BMI greater than 45 kg/m ² ; any incidence of ongoing bleeding beyond 8 weeks' gestation in the current pregnancy; on anti-coagulant therapy or known to have clotting disorders; known to be pregnant with multiples; lactating; established diabetes prior to pregnancy or currently taking anti-diabetic medications; being diagnosed with gestational diabetes in this pregnancy prior to enrolment in the study; known allergies to seafood or corn; currently on medication with aspirin and warfarin; has significant current gastrointestinal disease; incapable of giving informed consent; history of new investigational drug 3 months prior to this trial; currently consuming more than 200 g oily fish per week or taking supplements delivering 150 mg or more of DHA/day; unable to fast for 10 hr before obtaining blood sample
	Setting: John Hunter Hospital, Newcastle, Australia
Interventions	Intervention group
	2 x 1 g fish oil capsules each day (each capsule containing 60mg eicosapentaenoic acid and 430 mg DHA) from $^{\sim}$ 14 weeks' gestation until 34 weeks' gestation
	Control group
	Placebo: 1 x 1 g corn oil capsules/day from ~ 14 weeks' gestation until 34 weeks' gestation
Outcomes	Primary outcome : insulin resistance, as measured by HOMA-IR fasting glucose X fasting insulin/22.5 (at 14, 20, 28 & 34 weeks' gestation).
	Other outcomes : plasma inflammatory markers (IL-6, TNF-alpha, CRP, IL-1beta) and adipokines (adiponectin, leptin), measured using ELISA assays (at 14, 20 & 34 weeks' gestation); blood pressure (at 14, 20 & 34 weeks' gestation); newborn whole-blood fatty acid composition (48-72 hours after birth); Matsuda Index (calculated from 2 hour oral glucose tolerance test at 28 weeks' gesta-



Garg 2017 (Continued)	
San Continued	tion); erythrocyte fatty acid composition, measured by gas chromatography from a fasting blood sample (at 14, 20 & 34 weeks' gestation)
Starting date	1 March 2017
Contact information	Prof Manohar Garg, University of Newcastle. E-mail: manohar.garg@newcastle.edu.au
Notes	Funding and collaborators: University of Newcastle, Australia and EPAX, Norway.
Garmendia 2015	
Trial name or title	Diet and physical activity counselling and n3-long chain (PUFA) supplementation in obese pregnant women (MIGHT)
Methods	RCT (cluster, with 4 arms): NCT02574767
Participants	Recruitment target: 1000 women.
	Inclusion criteria : ≤ 14 weeks' gestational age at first prenatal visit; BMI > 30 kg/m² at first prenata visit; singleton pregnancy; plan to deliver at Sotero del Rio Hospital.
	Exclusion criteria : pre-existing diabetes (known or diagnosed at first control (fasting plasma glucose > 126 mg/dl or 2 h plasma glucose > 200 mg/dl during an OGTT; insulin or metformin use; known medical or obstetric complications which restrict physical activity; history of eating disorders; high risk of haemorrhagic bleeding; high-risk pregnancy according to national guidelines.
	Setting : 12 primary healthcare centres (PHCC) and Sotero del Rio Hospital, Chile.
Interventions	Intervention groups
	1) 'Lifestyle counselling + PUFA supplement group': home-based diet & physical activity counselling (2 home visits of 1 hour duration consisting of individually tailored dietary educational and behaviour education plus PUFA supplementation (n3LC-PUFAs oral supplementation based on Schizochytrium oil (S-oil) containing 800 mg DHA acid/day, administered as capsular preparations containing 200 mg DHA/capsule (4 capsules/day).
	2) 'PUFA supplementation" group': routine dietary and physical activity counselling plus n3LC-PU-FAs oral supplementation based on Schizochytrium oil (S-oil) containing 800 mg DHA acid/day, administered as capsular preparations containing 200 mg DHA/capsule (4 capsules/day).
	Control groups
	3) 'Llifestyle counselling + PUFA placebo group': home-based diet & physical activity counselling (2 home visits of 1 hour duration consisting of individually tailored dietary educational and behaviou education plus capsular preparations containing 50 mg DHA capsule (4 capsules/day), delivered in the same way as the n3LC-PUFA supplementation.
	4) 'Routine diet & PA + PUFA placebo group': routine dietary and physical activity counselling plus capsulare preparations containing 50 mg DHA/capsule (4 capsules/day), delivered in the same way as the n3LC-PUFA supplementation.
Outcomes	Primary outcomes : GDM (at 24-28 weeks of gestation according to ADA 2011 guidelines); macroso mia (birthweight > 4000 g); prevalence of insulin resistance at birth.
	Other outcomes : low birthweight (below 2500 g); excess weight gain during pregnancy; preeclampsia (at 24-28 weeks of gestation); preterm birth (< 37 weeks); caesarean.
Starting date	August 2015



Garmendia 2015 (Continued)	
Contact information	Dr Maria Luisa Garmendia. E-mail: mgarmendia@inta.uchile.ch
Notes	<u>Funding and collaborators</u> : University of Chile; Fondo Nacional de Desarrolo Cientificao y Technológico, Chile; DSM Nutritional Products, Inc; Corporación de Apoyo de la Investigatión Científic en Nutrición.
ihebremeskel 2014	
Trial name or title	DHA4PREG: DHA for PREGnant women: is the current recommendation appropriate for women with very low intake and status?
Methods	RCT (with 3 arms)
Participants	Recruitment target: 180 women
	Inclusion criteria : healthy pregnant women with a singleton pregnancy (with very low DHA intak and status)
	Exclusion criteria : pre-existing chronic medical conditions such as diabetes, high blood pressure, congenital heart disease, kidney disease, very preterm birth; sickle cell disease or haemoglo binopathies; history of pre-eclampsia, stillbirth, fetal death, major fetal abnormality; smoking or legal substance use.
	Setting: Sudan
Interventions	Intervention group
	1) Low-dose intervention group: 575 mg omega-3 (322.5 mg DHA and 47.2 mg EPA)
	2) High-does intervention group: 1725 mg omega-3 (967.7 mg DHA and 141.5 mg EPA)
	Control group
	3) placebo
Outcomes	Maternal DHA concentrations (blood and breast milk); fetal growth; preterm birth; gestational age at birth; birthweight; head circumference; length
Starting date	September 2014
Contact information	Professor Kebreab Ghebremeskel: k.ghebremeskel@londonmet.ac.uk
Notes	<u>Funding and collaborators</u> : Lipidomics and Nutrition Research Centre, London Metropolitan University, London (UK).
legarty 2012 Trial name or title	STABIL: Fish oil for mood stabilization during pregnancy in women with bipolar disorder
Methods	RCT: ACTRN12612000405819
Participants	Recruitment target: 200 women Inclusion criteria: pregnant women (up to 10 weeks of pregnancy, clinical diagnosis of bipolar di order I or II, using MS medication with an intention to either continue or discontinue MS medication throughout pregnancy, no experience of a mood episode reaching DSM IV-TR criteria within 4
Omega-3 fatty acid addition du	ring pregnancy (Review)



Hegarty 2012 (Continued)

weeks of recruitment, prepared to continue regular visits to personal treating medical professional/s throughout study period for ongoing psychiatric and antenatal care, able to give informed consent.

Exclusion criteria: under 18 years of age, poor written and/or spoken English, diagnosed with schizophrenia or schizoaffective disorder, taking any daily supplement containing more than 120 mg EPA or more than 500 mg EPA + DHA, at high risk of suicide, participating in another clinical trial, current drug or alcohol problems, unstable medical condition or unstable thyroid dysfunction or lipid metabolism disorder, current use of anticoagulant therapy, have a bleeding disorder, fish/seafood allergy.

Setting: Australia

Interventions

Intervention group

5 g concentrated omega-3 triglycerides from fish, containing 1000 mg DHA and 1500 mg EPA daily, taken as an oral capsule once daily, from enrolment (up to 10 weeks of pregnancy) till 12 weeks postpartum

Control group

1 g concentrated omega-3 triglycerides from fish, containing 430 mg DHA and 90 mg EPA, plus 4 g medium chain fatty acids from coconut, taken as an oral capsule once daily, from enrolment (up to 10 weeks of pregnancy) till 12 weeks postpartum

Outcomes

Primary outcomes: number of mood episode recurrences (defined by DSM-IV criteria for major depression, hypomania or mania, or the need for rescue medication to be provided by the treating clinician due to deteriorating mood)

Other outcomes: time to onset of first mood episode recurrence; severity of depressive and manic symptoms (Montgomery Asberg Depression Rating Scale (MADRS) and Young Mania Rating Scale (YMRS) will be used to assess the severity of mood symptoms)

Starting date	1 May 2012
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Contact information B Hegarty. E-mail: b.hegarty@unsw.edu.au

Notes

<u>Funding and collaborators</u>: Department of Health and Aging, Canberra; University of New South Wales, Black Dog Institute (Australia)

Hendler 2017

Trial name or title	The effect of alpha linolenic acid (ALA) supplementation during pregnancy
Methods	RCT (3 arm): NCT03040856
Participants	150 pregnant women aged 18 to 45 years attending the high-risk clinic (GA 12-16 weeks)
Interventions	ALA (1260 mg/day) versus DHA + EPA (480 mg DHA and 720 mg EPA/day) versus placebo (olive oil)
Outcomes	Omega-3 fatty acid concentrations, expression of messenger RNAs
Starting date	May 2017
Contact information	Aya Mohr Sasson, email: mohraya@gmail.com
Notes	Sheba Medical Center, Israel



Trial name or title	Effect of Docosa-Hexanoic Acid (DHA) Supplements During Pregnancy on Newborn Outcomes in India: (DHANI)
Methods	RCT: NCT 01580345.
Participants	Recruitment target: 600 women
	Inclusion criteria : 18 to 35 years or older; ≤ 20 weeks of gestation; willing to participate in the study and perform all measurements for self, husband and offspring; willing to provide signed and dated informed consent.
	Exclusion criteria : allergic (if aware) to any of the test products; at high risk for hemorrhagic bleed ing, clotting (if aware); high-risk pregnancy; consuming omega-3 supplements or having used these in 3 months preceding the intervention period; reported participation in another biomedical trial 3 months before the start of the study or during the study.
	Setting : KLEUs Jawaharlal Nehru Medical College, Prabhakar Kore Charitable Hospital, Belgaum, Karnataka, India.
Interventions	Intervention group
	400 mg of DHA daily, from ≤ 20 weeks' gestation until birth.
	Control group
	400 mg/day of placebo (corn/soy oil), from ≤ 20 weeks' gestation until birth.
Outcomes	Primary outcomes: newborn anthropometry (birthweight, length, and head circumference).
	Other outcomes : gestational age; new born APGAR score (at 1 minute and 5 minutes); unfavourable pregnancy outcomes (stillbirth, low birthweight babies, and preterm babies)
Starting date	December 2015
Contact information	Dr Shweta Khandelwal, Senior Public Health Nutritionist, Centre for Chronic Disease Control, India. E-mail: shweta.khandelwal@phfi.org
Notes	<u>Funding and collaborators</u> : Centre for Chronic Disease Control, India; Department of Science and Technology, Government of India; Jawaharlal Nehru Medical College.

Kodkhany 2017

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Trial name or title	Maternal DHA Supplementation and Offspring Neurodevelopement in India (DHANI-2)
Methods	RCT: NCT03072277
Participants	957 participants
	Inclusion criteria : 18 to 35 year old pregnant women (singleton) at ≤ 20 weeks GA, willing to participate in study and perform all measurements for self, husband and offspring including anthropometry, dietary assessment, questionnaires and biological samples (blood and breast milk), willing to provide signed and dated informed consent
	Exclusion criteria : allergic (if aware) to any of the test products, at high risk for haemorrhagic bleeding or clotting (if aware), high-risk pregnancies (history and prevalence of pregnancy complications, including abruptio placentae, pre-eclampsia, pregnancy-induced hypertension, any seri-



Kodkhany 2017 (Continued)	ous bleeding episode in the current pregnancy, and/or physician referral); and/or diagnosed chronic degenerative disease(s) such as diagnosed heart disease, cancer, stroke or diabetes (as omega-3 could raise blood sugar and lower insulin promotion) Setting: KLEUs Jawaharlal Nehru Medical College, Prabhakar Kore Charitable Hospital, Belgaum, Karnataka, India.
Interventions	Intervention group
	DHA (400 mg/day) from ≤ 20 weeks of gestation through 6 months postpartum
	Control group
	Placebo (400 mg/day corn/soy oil) from ≤ 20 weeks of gestation through 6 months postpartum
Outcomes	Primary outcomes : infant neurodevelopment defined as measured by the mean difference in the average Developmental Quotient (DQ) scores of the 2 groups
	Other outcomes: DHA levels (fatty acid levels in maternal and infant blood)
Starting date	December 2015
Contact information	Public Health Foundation of India cited as responsible party, no further details.
Notes	Funding and collaborators : Public Health Foundation of India, Jawaharlal Nehru Medical College, India Alliance
	Estimated study completion date: December 2020 (primary completion August 2019)

Li 2013

Trial name or title	Effect of omega-3 fatty acids on insulin sensitivity in Chinese gestational diabetes patients		
Methods	RCT: NCT01912170		
Participants	Recruitment target: 75 women.		
	Inclusion criteria: 18-40 years; with gestational diabetes; willing to participate in the trial.		
	Exclusion criteria : type 1 diabetes mellitus or type 2 diabetes mellitus before pregnancy; not willing to provide informed consent; with a family history of hypertriglyceridaemia or fasting triglyceride > 4.56 mmol/L; diagnosed with severe liver disease, kidney disease or cancer; participating in another clinical trial within 30 days; other diseases or conditions for which the doctor does not agree to the participant's participation.		
	Setting: Jiaxing Women's and Children's Hospital, China.		
Interventions	Intervention group		
	Fish oil 2 g a day (220 mg EPA and 170 mg DHA per 1 g capsule), from the third trimester of pregnancy until the 4th week after birth.		
	Control group		
	Flaxseed oil 2 g a day (550 mg ALA per 1 g capsule), from the third trimester of pregnancy until the 4th week after birth.		
Outcomes	Primary outcomes: fasting glucose; fasting insulin.		



Li 2013 (Continued)	Secondary outcomes: birthweight; birth length; blood lipids.			
Starting date	August 2013			
Contact information	Professor Duo Li (Principal Investigator) and Huijuan Liu (contact provided), Zhejiang University, China.			
	E-mails: not provided.			
Notes	Funding and collaborators: Zhejiang University; National Natural Science Foundation of China.			
Makrides 2013 (ORIP)				
Trial name or title	Omega-3 fats to reduce the incidence of prematurity: the ORIP trial			
Methods	RCT: ACTRN12613001142729			
Participants	Recruitment target: 5540 women.			
	Inclusion criteria: < 20 weeks' gestation (singleton or multiple pregnancy).			
	Exclusion criteria : known fetal abnormality; taking dietary supplements containing LCPUFA 150mg/day; taking dietary supplements containing LCPUFA 150 mg/day and not willing to stop; bleeding disorders where fish oil is contraindicated or on anticoagulant therapy; and history of drug or alcohol abuse.			
	Setting: South Australia, Victoria and Queensland, Australia.			
Interventions	Intervention group			
	3 capsules of fish oil containing a total dose of approximately 800 mg of DHA daily from enrolment (12 weeks - 20 weeks' gestation) until 34 weeks of gestation or birth (whichever occurs first)			
	Control group			
	3 placebo vegetable oil capsules with a trace of DHA to aid masking.			
Outcomes	Primary outcome: early preterm birth (< 34 weeks).			
	Other outcomes : post-term induction; post-term pre labour caesarean birth; preterm birth (< 37 weeks); safety and tolerability of DHA supplementation; low birthweight (< 2500 g); small-for-gestational age (< 10th centile for corresponding GA and sex); neonatal complications (up to 28 days post birth); admission to NICU.			
Starting date	October 2013			
Contact information	Prof Maria Makrides, SAHMRI. E-mail: maria.makrides@sahmri.com			
Notes	Funding and collaborators: NHMRC; SAHMRI.			
	Recruitment ended 27 April 2017.			



Martini 2014 (CORDHA)			
Trial name or title	A randomised controlled trial for the optimization of the viability of stem cells derived from umbilical CORd blood after maternal supplementation with DHA during the second or third trimester of pregnancy (CORDHA)		
Methods	RCT: ISRCTN58396079.		
Participants	Recruitment target: 150 women.		
	Inclusion criteria : Caucasian, non-smoker, > 18 years of age, single pregnancy, absence of diabetes or hypertension or any other type of pathology requiring pharmacological therapy, absence of chromosome abnormalities and/or congenital malformations in the fetus, and HBV, HCV, HIV and CMV negative.		
	Exclusion criteria : impossible to collect cord blood (for bureaucratic reasons or because of emergencies regarding the health of the mother or the baby), taken other supplements containing DHA or fish oil, temperature of 39C during birth, and UCB samples with a volume of less than 80 mL and/or less than 70% cell vitality.		
	Setting: Rome, Italy.		
Interventions	Intervention group		
	DHA (250 mg/day), from the 20th or from the 28th week up to the 40th week of estimated gestational age.		
	Control group		
	Placebo (250 mg olive oil/day), from the 20th or from the 28th week up to the 40th week of estimated gestational age.		
Outcomes	Primary outcome : measure of the viability (%) and the number of CD34+ cells collected from the umbilical cord blood at birth.		
	No other outcomes specified.		
Starting date	September 2010		
Contact information	Irene Martini, Via Vittorio Locchi 9 00197 Rome, Italy. E-mail: martini@smartbank.it		
Notes	Funding and collaborators: SmartBank SRL; Biovault and Avantgarde SAS.		
	Recruitment ended September 2014.		

Mbayiwa 2016

Trial name or title	Improving Maternal and Child Health Through Prenatal Fatty Acid Supplementation (NAPS): A Randomized Controlled Study in African-American Women Living in Low-income Envrionments	
Methods	RCT (NCT02647723)	
Participants	Recruitment target: 162 women.	
	Inclusion criteria : women 18 to 34 years of age, household recipient of public assistance (e.g. Medicaid insurance) due to low income, low levels of DHA consumption defined as less than 2 fish servings per week	
	Exclusion criteria : reports of known medical complications, regular use of steroid medications or alcohol or cigarettes or illegal substances (by medical report), use of blood thinners or anti-coagulants, use of psychotropic medications, BMI > 40, allergy to iodine and/or soy	



Mbayiwa 2016 (Continued)			
	Setting: United States, Illinois		
Interventions	Intervention group		
	DHA (450 mg twice daily), by oral intake, for 24 weeks		
	Control group		
	Placebo (450 mg soybean oil twice daily, by oral intake, for 24 weeks)		
Outcomes	Primary outcome : average change in perceived stress scale (PSS) score (time frame 16 months), assessed at baseline and at 24, 30 and 36 weeks of pregnancy, and at 1, 4 and 9 months after birth		
	No other outcomes specified		
Starting date	January 2016		
Contact information	Kimberley Mbayiwa. E-mail: kmbayiwa@yoda.bsd.uchicago.edu		
Notes	Funding and collaborations: University of Chicago, Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD), University of Pittsburgh.		

Murff 2017 (FORTUNE)

Trial name or title	Fish Oil to Reduce Tobacco Use iN Expectant Mothers (FORTUNE)			
Methods	RCT: NCT03077724			
Participants	40 pregnant women aged 18 to 45 years, between 6 and 36 weeks' gestation, who are active smokers			
	Setting: Nashville, Tennessee, United States			
Interventions	Omega-3 (4.2 g/day) versus placebo (olive oil)			
Outcomes	Reduction in total cigarettes smoked per day			
Starting date	March 2017			
Contact information	Associate Professor Harvey Murff, Vanderbilt University Medical Center, USA.			
	email: harvey.j.murff@vanderbilt.edu			
Notes	Pilot feasibility trial			

Nishi 2015 (SYNCHRO)

Participants	Recruitment target: 108 women.	
Methods	RCT (multi centre): NCT02166424	
Trial name or title	SYNCHRO	



Nishi 2015 (SYNCHRO) (Continued)

Inclusion criteria: ≥ 20 years; 12-24 weeks' gestation, Japanese conversational ability in Japan site, Mandarin conversational ability in Taiwan site, willing to take assessments after childbirth, with EPDS 9 or more and in good physical health (judged by obstetricians).

Exclusion criteria: history and current suspicion of psychosis or bipolar disorder or substance-related disorder or eating disorder or personality disorder, serious phychiatric symptoms such as self-harm behaviour or in need of rapid psychiatric treatment, difficult to expect a normal birth, having a history of bleeding disorder such as von Willebrand's Disease, regular treatment with Asprin or warfarin within the last 3 months, a smoking habit of ≥ 40 cigarettes per day, regular treatment with ethyl icosapentate or regular consumption of omega-3 PUFA supplements within the last 3 months, and a habit of eating fish as a main dish ≥ times per week.

Setting: Japan and Taiwan.

Interventions Intervention group

Omega-3 polyunsaturated fatty acids (1200 mg eicosapentaenoic acid EPA and 600 mg DHA daily).

Control group

Placebo (2880 mg olive oil daily).

Outcomes

Primary outcome: total score on HAMD (12 weeks).

Other outcomes: total score on HAMD (4-6 weeks after childbirth); MDD as determined by the depression module of MINI (4-6 weeks after childbirth); total scores on EPDS (4-6 weeks after childbirth); total score on BDI- π (4-6 weeks after childbirth); omega-3 fatty acids concentrations in erythrocytes (4-6 weeks after childbirth); oestrogen in plasma (4-6 weeks after childbirth); oxytocin in plasmal (4-6 weeks after childbirth); progesterone in plasma (4-6 weeks after childbirth), hCG in plasma (4-6 weeks after childbirth), phospholipase A2 in plasma (4-6 weeks after childbirth); gestational age (at childbirth); GDM (4-6 weeks after childbirth); gestational hypertension or pre-eclampsia (4-6 weeks after childbirth); gestational hypertension or pre-eclampsia (4-6 weeks after childbirth); induced labour; blood loss at childbirth; caesarean section; operative vaginal birth; birthweight; Apgar score (at 1 minute and 5 minutes); NICU admission (4-6 weeks after childbirth); and cholesterol (4-6 weeks after childbirth).

Starting date	June 2014		
Contact information	Daisuke Nishi, Assistant Professor, Tokyo Medical University. E-mail: d-nishi@umin.ac.jp		
Notes	<u>Funding and collaborators</u> : Tokyo Medical University, China Medical University (Taiwan), University of Toyama, Chiba University, and National Center for Child Health and Development.		
	A non-randomised pilot for this study was registered as NCT01948596 (completed: Nishi 2016)		

Wang 2018

Trial name or title	Impact of DHA/Oat on metabolic health in gestational diabetes mellitus		
Methods	RCT		
Participants	80 women with a singleton pregnancy without any evidence of malformation and with a de nov diagnosis of gestational diabetes at 22-28 weeks gestation		
Interventions	1) DHA		
	2) oat		
	3) DHA + oat		



Wang 2018 (Continued)				
	4) placebo			
Outcomes	neonatal leptin; maternal fasting plasma glucose concentration,			
Starting date	August 2017			
Contact information	Wen_Juan Wang, Master			
	email:wangwe.njuean@163.com			
Notes	Xinhua Hospital, Shanghai Jiao Tong University School of Medicine			
Zielinsky 2015				
Trial name or title	Effect of mother's supplementation omega-3 in the dynamics of fetal ductus arteriosus: a randomized clinical trial			
Methods	RCT: NCT02565290			
Participants	Recruitment target: 74 women.			
	Inclusion criteria: at 28-32 weeks' gestation; and willing to participate			
	Exclusion criteria : hypertensive; diabetic; not using anti-inflammatory drugs; not HIV positive; not using mate, black or green tea; no inflammation in past 5 days; and not allergic to fish or soy.			
	Setting: Brazil			
Interventions	Intervention group			
	Omega 3 capsules (1g), 2 times a day, for 21 days.			
	Control group			
	Placebo soy oil capsules (1g), 2 times a day, for 21 days.			
Outcomes	Primary outcome: pulsatility index of fetal ductus arteriosus			
	Other outcomes : inflammatory biomarkers (interleukins, prostaglandins, cyclooxygenase); systolic and diastolic velocity of fetal ductus arteriosus; and biomarkers of oxidative stress.			
Starting date	May 2015			
Contact information	Dr Paulo Zielinsky, Instituto de Cardiologia do Rio Grande do Sul. E-mail: paulozie.voy@ter-ra.com.br			
Notes	Funding and collaborators: Instituto de Cardiologia do Rio Grande do Sul.			
Zimmermann 2018	Diotany supplementation of Omoga 2 and the participation in the placentary versitance			
Trial name or title Dietary supplementation of Omega 3 and the participation in the placentary vasc mechanism in pregnant people				
Methods	RCT			



Zimmermann 2018 (Continue	d)			
Participants	150 pregnant women ≥ 19 years; non-smokers			
Interventions	women without risk factors for pre-eclampsia:			
	1) 400 mg omega from 12 weeks gestation to one week prior to birth			
	2) no omega			
	women at risk of pre-eclampsia			
	1) AAS + 400 mg omega			
	2) 400 mg omega			
	women with thrombophilia			
	1) heparin + 400 mg omega			
	2) 400 mg omega			
Outcomes	placental vascular resistance; pre-eclampsia; oligohydramnios; intrauterine growth restriction			
Starting date	August 2018			
Contact information	Juliana Barroso Zimmermann			
	email: julianabz@uol,com,br			
Notes	Faculdade de Medicina de Barbacena			

Abbreviations: ADA: American Diabetes Associaton; ADORE: Assessment of DHA on Reducing Early Preterm Birth; ADP: Air Displacement Plethsmography; BDI: Beck Depression Inventory; BIS: Bioelectical Independence Spectroscopy; BMI: Body Mass Index; CMV: cytomegalovirus; DHA4PREG: DHA for PREGnant women; DHA: docosahexaenoic acid; DXA: dual x-ray absorptiometry; EPA: eicosapentaenoic acid; EPDS: Edinburgh Postnatal Depression Scale; FOPCHIN: Fish Oil Supplementation to Pregnant Women in China; GA: gestational age; GDM: gestational diabetes mellitus; HAMD: Hamilton Rating Scale for Depression; HBV: Hepatitis B virus; HCV: Hepatitis C virus; HIV: human immunodeficiency virus; HOMA-IR: Homeostatic Model Assessment of Insulin Resistance; hCG: human chorionic gonadotropin; IVGTT: Intravenous Glucose Tolerance Test; LCPUFA: long-chain polyunsaturated acids; MDD: major depressive disorder; MINI: Min-International Neuropsychiatric Interview; NHMRC: National Health and Medical Research Institute; NICU: neonatal intensive care unit; NSAIDs: nonsteroidal anti-inflammatory drugs; OGTT: oral glucose tolerance test; PUFA: polyunsaturated fatty acid; RCT: randomised controlled trial; SAHMRI: South Australian Health and Medical Research Institute; SYNCHRO:The Synchronized Trial on Expectant Mothers with Depressive Symptoms by Omega-3 PUFAs; UBC: umbilical cord blood

DATA AND ANALYSES

Comparison 1. Overall: omega-3 versus no omega-3

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Preterm birth (< 37 weeks)	26	10304	Risk Ratio (M-H, Fixed, 95% CI)	0.89 [0.81, 0.97]
2 Early preterm birth (< 34 weeks)	9	5204	Risk Ratio (M-H, Fixed, 95% CI)	0.58 [0.44, 0.77]
3 Prolonged gestation (> 42 weeks)	6	5141	Risk Ratio (M-H, Fixed, 95% CI)	1.61 [1.11, 2.33]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4 Maternal death	4	4830	Risk Ratio (M-H, Fixed, 95% CI)	1.69 [0.07, 39.30]
5 Pre-eclampsia (hypertension with proteinuria)	20	8306	Risk Ratio (M-H, Fixed, 95% CI)	0.84 [0.69, 1.01]
6 High blood pressure (without proteinuria)	7	4531	Risk Ratio (M-H, Fixed, 95% CI)	1.03 [0.89, 1.20]
7 Eclampsia	1	100	Risk Ratio (M-H, Fixed, 95% CI)	0.14 [0.01, 2.70]
8 Maternal antepartum hospi- talisation	5	2876	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.81, 1.04]
8.1 Any	4	2813	Risk Ratio (M-H, Fixed, 95% CI)	0.91 [0.80, 1.03]
8.2 Due to PIH or IUGR	1	63	Risk Ratio (M-H, Fixed, 95% CI)	1.23 [0.67, 2.28]
9 Mother's length of stay in hospital (days)	2	2290	Mean Difference (IV, Fixed, 95% CI)	0.18 [-0.20, 0.57]
10 Maternal anaemia	1	846	Risk Ratio (M-H, Fixed, 95% CI)	1.16 [0.91, 1.48]
11 Miscarriage (< 24 weeks)	9	4190	Risk Ratio (M-H, Fixed, 95% CI)	1.07 [0.80, 1.43]
12 Antepartum vaginal bleed- ing	2	2151	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.69, 1.48]
13 Rupture of membranes (PPROM; PROM)	4		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
13.1 Preterm prelabour rup- ture of membranes (PPROM)	3	925	Risk Ratio (M-H, Fixed, 95% CI)	0.53 [0.25, 1.10]
13.2 Premature rupture of membranes (PROM)	3	915	Risk Ratio (M-H, Fixed, 95% CI)	0.41 [0.21, 0.82]
14 Maternal admission to intensive care	2	2458	Risk Ratio (M-H, Fixed, 95% CI)	0.56 [0.12, 2.63]
15 Maternal adverse events	17		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
15.1 Severe adverse event	2	2690	Risk Ratio (M-H, Fixed, 95% CI)	1.04 [0.40, 2.72]
15.2 Severe enough for cessa- tion	6	1487	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.53, 1.93]
15.3 Any	5	1480	Risk Ratio (M-H, Fixed, 95% CI)	1.38 [1.16, 1.65]
15.4 Nausea	9	2929	Risk Ratio (M-H, Fixed, 95% CI)	1.05 [0.90, 1.22]
15.5 Unpleasant taste	5	2356	Risk Ratio (M-H, Fixed, 95% CI)	4.82 [3.35, 6.92]
15.6 Vomiting	7	3640	Risk Ratio (M-H, Fixed, 95% CI)	1.14 [0.95, 1.37]
15.7 Stomach pain	4	928	Risk Ratio (M-H, Fixed, 95% CI)	1.49 [0.62, 3.59]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
15.8 Reflux	1	26	Risk Ratio (M-H, Fixed, 95% CI)	1.0 [0.16, 6.07]
15.9 Belching or burping	5	2262	Risk Ratio (M-H, Fixed, 95% CI)	3.52 [2.86, 4.34]
15.10 Diarrhoea	6	1764	Risk Ratio (M-H, Fixed, 95% CI)	0.80 [0.52, 1.24]
15.11 Constipation	1	1077	Risk Ratio (M-H, Fixed, 95% CI)	0.42 [0.08, 2.15]
15.12 Nasal bleeding	2	1506	Risk Ratio (M-H, Fixed, 95% CI)	0.94 [0.71, 1.24]
15.13 Swelling/other reaction at injection site	1	852	Risk Ratio (M-H, Fixed, 95% CI)	1.10 [0.99, 1.22]
15.14 Insomnia	1	36	Risk Ratio (M-H, Fixed, 95% CI)	1.5 [0.28, 7.93]
15.15 Headache	1	301	Risk Ratio (M-H, Fixed, 95% CI)	1.61 [0.91, 2.86]
15.16 Gynaecological infec- tions	1	291	Risk Ratio (M-H, Fixed, 95% CI)	0.84 [0.45, 1.55]
15.17 Labour related	1	291	Risk Ratio (M-H, Fixed, 95% CI)	0.49 [0.27, 0.88]
15.18 Urinary tract infection	1	291	Risk Ratio (M-H, Fixed, 95% CI)	0.30 [0.06, 1.42]
16 Caesarean section	28	8481	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.91, 1.03]
17 Induction (post-term)	3	2900	Risk Ratio (M-H, Random, 95% CI)	0.82 [0.22, 2.98]
18 Blood loss at birth (mL)	6	2776	Mean Difference (IV, Fixed, 95% CI)	11.50 [-6.75, 29.76
19 Postpartum haemorrhage	4	4085	Risk Ratio (M-H, Fixed, 95% CI)	1.03 [0.82, 1.30]
20 Gestational diabetes	12	5235	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.83, 1.26]
21 Maternal insulin resistance (HOMA-IR)	3	176	Mean Difference (IV, Random, 95% CI)	-0.85 [-2.50, 0.80]
22 Excessive gestational weight gain	1	350	Risk Ratio (M-H, Fixed, 95% CI)	1.21 [0.95, 1.55]
23 Gestational weight gain (kg)	11	2297	Mean Difference (IV, Random, 95% CI)	-0.05 [-0.68, 0.59]
24 Depression during pregnancy: thresholds	3		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
24.1 HAMD 50% reduction (after 8 weeks)	1	24	Risk Ratio (M-H, Fixed, 95% CI)	2.26 [0.78, 6.49]
24.2 HAMD ≤ 7	1	24	Risk Ratio (M-H, Fixed, 95% CI)	2.12 [0.51, 8.84]
24.3 Unspecified	1	301	Risk Ratio (M-H, Fixed, 95% CI)	2.39 [0.47, 12.11]
24.4 EPDS ≥ 11	1	34	Risk Ratio (M-H, Fixed, 95% CI)	1.4 [0.55, 3.55]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
25 Depression during pregnancy: scores	5		Mean Difference (IV, Random, 95% CI)	Subtotals only
25.1 BDI	2	104	Mean Difference (IV, Random, 95% CI)	-5.86 [-8.32, -3.39]
25.2 HAMD	3	71	Mean Difference (IV, Random, 95% CI)	-0.92 [-5.91, 4.06]
25.3 EPDS	4	122	Mean Difference (IV, Random, 95% CI)	-0.40 [-3.70, 2.89]
25.4 MADRS	1	26	Mean Difference (IV, Random, 95% CI)	-1.60 [-7.80, 4.60]
26 Anxiety during pregnancy	1	301	Risk Ratio (M-H, Fixed, 95% CI)	0.95 [0.06, 15.12]
27 Difficult life circumstances (maternal)	1	51	Mean Difference (IV, Fixed, 95% CI)	0.32 [-0.15, 0.79]
28 Stress (maternal)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
28.1 Perceived Stress Scale (scores)	1	51	Mean Difference (IV, Fixed, 95% CI)	-1.82 [-3.68, 0.04]
29 Depressive symptoms post- partum: threshold	4		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
29.1 PDSS ≥ 80	1	42	Risk Ratio (M-H, Random, 95% CI)	0.37 [0.04, 3.25]
29.2 EPDS	2	2431	Risk Ratio (M-H, Random, 95% CI)	0.99 [0.56, 1.77]
29.3 Major depressive disorder	1	118	Risk Ratio (M-H, Random, 95% CI)	1.33 [0.27, 6.56]
30 Depressive symptoms post- partum: scores	2		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
30.1 BDI: 6-8 weeks postpar- tum	1	118	Mean Difference (IV, Fixed, 95% CI)	0.25 [-1.93, 2.43]
30.2 PDSS total (LS over 6 months)	1	42	Mean Difference (IV, Fixed, 95% CI)	-6.08 [-12.42, 0.26]
30.3 Disturbances sleep/eating (LS over 6 months)	1	42	Mean Difference (IV, Fixed, 95% CI)	-1.0 [-2.66, 0.66]
30.4 Anxiety/insecurity (LS over 6 months)	1	42	Mean Difference (IV, Fixed, 95% CI)	-1.30 [-2.96, 0.36]
30.5 Emotional lability (LS over 6 months)	1	42	Mean Difference (IV, Fixed, 95% CI)	-1.29 [-3.10, 0.52]
30.6 Mental confusion (LS over 6 months)	1	42	Mean Difference (IV, Fixed, 95% CI)	-1.30 [-2.92, 0.32]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
30.7 Loss of self (LS over 6 months)	1	42	Mean Difference (IV, Fixed, 95% CI)	-0.90 [-1.80, 0.00]
30.8 Guilt (LS over 6 months)	1	42	Mean Difference (IV, Fixed, 95% CI)	-0.30 [-1.13, 0.53]
30.9 Suicide (LS over 6 months)	1	42	Mean Difference (IV, Fixed, 95% CI)	-0.07 [-0.35, 0.21]
30.10 PDSS total at 6 months	1	42	Mean Difference (IV, Fixed, 95% CI)	-2.87 [-12.17, 6.43]
30.11 Disturbances sleep/eating at 6 months	1	42	Mean Difference (IV, Fixed, 95% CI)	-0.20 [-2.08, 1.68]
30.12 Anxiety/insecurity at 6 months	1	42	Mean Difference (IV, Fixed, 95% CI)	-0.46 [-2.65, 1.73]
30.13 Emotional lability at 6 months	1	42	Mean Difference (IV, Fixed, 95% CI)	-0.96 [-3.32, 1.40]
30.14 Mental confusion at 6 months	1	42	Mean Difference (IV, Fixed, 95% CI)	-0.13 [-2.15, 1.89]
30.15 Loss of self at 6 months	1	42	Mean Difference (IV, Fixed, 95% CI)	-0.97 [-2.18, 0.24]
30.16 Guilt at 6 months	1	42	Mean Difference (IV, Fixed, 95% CI)	0.21 [-0.69, 1.11]
30.17 Suicide at 6 months	1	42	Mean Difference (IV, Fixed, 95% CI)	-0.52 [-1.13, 0.09]
31 Gestational length (days)	43	12517	Mean Difference (IV, Random, 95% CI)	1.67 [0.95, 2.39]
32 Perinatal death	10	7416	Risk Ratio (M-H, Fixed, 95% CI)	0.75 [0.54, 1.03]
33 Stillbirth	16	7880	Risk Ratio (M-H, Fixed, 95% CI)	0.94 [0.62, 1.42]
34 Neonatal death	9	7448	Risk Ratio (M-H, Fixed, 95% CI)	0.61 [0.34, 1.11]
35 Infant death	4	3239	Risk Ratio (M-H, Fixed, 95% CI)	0.74 [0.25, 2.19]
36 Large-for-gestational age	6	3722	Risk Ratio (M-H, Fixed, 95% CI)	1.15 [0.97, 1.36]
37 Macrosomia	6	2008	Risk Ratio (M-H, Fixed, 95% CI)	0.69 [0.43, 1.13]
38 Low birthweight (< 2500 g)	15	8449	Risk Ratio (M-H, Fixed, 95% CI)	0.90 [0.82, 0.99]
39 Small-for-gestational age/ IUGR	8	6907	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.90, 1.13]
40 Birthweight (g)	44	11584	Mean Difference (IV, Random, 95% CI)	75.74 [38.05, 113.43]
41 Birthweight Z score	4	2792	Mean Difference (IV, Fixed, 95% CI)	0.06 [-0.02, 0.13]
42 Birth length (cm)	29	8128	Mean Difference (IV, Random, 95% CI)	0.11 [-0.10, 0.31]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
43 Head circumference at birth (cm)	24	7161	Mean Difference (IV, Random, 95% CI)	0.07 [-0.05, 0.19]
44 Head circumference at birth Z score	2	2462	Mean Difference (IV, Fixed, 95% CI)	-0.03 [-0.14, 0.07]
45 Length at birth Z score	2	2462	Mean Difference (IV, Random, 95% CI)	0.18 [-0.18, 0.54]
46 Baby admitted to neonatal care	9	6920	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.83, 1.03]
47 Infant length of stay in hospital (days)	1	2041	Mean Difference (IV, Fixed, 95% CI)	0.11 [-1.40, 1.62]
48 Congenital anomalies	3	1807	Risk Ratio (M-H, Fixed, 95% CI)	1.08 [0.61, 1.92]
49 Retinopathy of prematurity	1	837	Risk Ratio (M-H, Fixed, 95% CI)	1.20 [0.32, 4.44]
50 Bronchopulmonary dysplasia	2	3191	Risk Ratio (M-H, Fixed, 95% CI)	1.06 [0.45, 2.48]
51 Respiratory distress syndrome	2	1129	Risk Ratio (M-H, Random, 95% CI)	1.17 [0.54, 2.52]
52 Necrotising enterocolitis (NEC)	2	3198	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.26, 3.55]
53 Neonatal sepsis (proven)	3	3788	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.44, 2.14]
54 Convulsion	1	2361	Risk Ratio (M-H, Fixed, 95% CI)	0.09 [0.01, 1.63]
55 Intraventricular haemor- rhage	3		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
55.1 Any	3	5423	Risk Ratio (M-H, Random, 95% CI)	1.00 [0.29, 3.49]
55.2 Grade 3 or 4	1	837	Risk Ratio (M-H, Random, 95% CI)	1.60 [0.38, 6.65]
56 Neonatal/infant adverse events	3		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
56.1 Any adverse event	2	592	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.82, 1.02]
56.2 Serious adverse events	2	2690	Risk Ratio (M-H, Fixed, 95% CI)	0.72 [0.53, 0.99]
57 Neonatal/infant morbidity: cardiovascular	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
58 Neonatal/infant morbidity: respiratory	1	291	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.66, 1.57]
59 Neonatal/infant morbidity: due to pregnancy/birth events	1	291	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.67, 1.55]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
60 Neonatal/infant morbidity: other	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
60.1 Colds in past 15 days: at 1 month of age	1	849	Risk Ratio (M-H, Fixed, 95% CI)	0.85 [0.72, 1.00]
60.2 Colds in past 15 days: at 3 months of age	1	834	Risk Ratio (M-H, Fixed, 95% CI)	0.86 [0.73, 1.01]
60.3 Colds in past 15 days: at 6 months of age	1	834	Risk Ratio (M-H, Fixed, 95% CI)	0.99 [0.86, 1.15]
60.4 Fever in past 15 days: at 1 month of age	1	849	Risk Ratio (M-H, Fixed, 95% CI)	1.08 [0.53, 2.22]
60.5 Fever in past 15 days: at 3 months of age	1	834	Risk Ratio (M-H, Fixed, 95% CI)	0.80 [0.53, 1.23]
60.6 Fever in past 15 days: at 6 months of age	1	834	Risk Ratio (M-H, Fixed, 95% CI)	0.99 [0.74, 1.31]
60.7 Rash in past 15 days: at 1 month of age	1	849	Risk Ratio (M-H, Fixed, 95% CI)	1.11 [0.89, 1.38]
60.8 Rash in past 15 days: at 3 months of age	1	834	Risk Ratio (M-H, Fixed, 95% CI)	0.82 [0.54, 1.26]
60.9 Rash in past 15 days: at 6 months of age	1	834	Risk Ratio (M-H, Fixed, 95% CI)	1.14 [0.76, 1.71]
60.10 Vomiting in past 15 days: at 1 month of age	1	849	Risk Ratio (M-H, Fixed, 95% CI)	1.55 [0.82, 2.93]
60.11 Vomiting in past 15 days: at 3 months of age	1	834	Risk Ratio (M-H, Fixed, 95% CI)	1.43 [0.69, 2.96]
60.12 Vomiting in past 15 days: at 6 months of age	1	834	Risk Ratio (M-H, Fixed, 95% CI)	1.33 [0.72, 2.46]
60.13 Diarrhoea in past 15 days: at 1 month of age	1	849	Risk Ratio (M-H, Fixed, 95% CI)	0.83 [0.42, 1.67]
60.14 Diarrhoea in past 15 days: at 3 months of age	1	834	Risk Ratio (M-H, Fixed, 95% CI)	0.83 [0.46, 1.51]
60.15 Diarrhoea in past 15 days: at 6 months of age	1	834	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.63, 1.64]
60.16 Other illness in the past 15 days: at 1 month	1	849	Risk Ratio (M-H, Fixed, 95% CI)	1.40 [0.81, 2.41]
60.17 Other illness in the past 15 days: at 3 months	1	834	Risk Ratio (M-H, Fixed, 95% CI)	0.96 [0.54, 1.73]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
60.18 Other illness in the past 15 days: at 6 months	1	834	Risk Ratio (M-H, Fixed, 95% CI)	1.15 [0.68, 1.95]
61 Infant/child morbidity	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
61.1 ICU admissions	1	1396	Risk Ratio (M-H, Fixed, 95% CI)	0.58 [0.31, 1.06]
61.2 Medical diagnosis of at- tention deficit hyperactivity disorder (ADHD)	1	1526	Risk Ratio (M-H, Fixed, 95% CI)	2.96 [0.31, 28.40]
61.3 Medical diagnosis of autism spectrum disorder	1	1526	Risk Ratio (M-H, Fixed, 95% CI)	1.15 [0.54, 2.47]
61.4 Medical diagnosis of oth- er learning/behavioural disor- ders	1	1526	Risk Ratio (M-H, Fixed, 95% CI)	1.12 [0.78, 1.60]
61.5 Medical diagnosis of other chronic health conditions	1	1526	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.65, 1.44]
62 Ponderal index	6	887	Mean Difference (IV, Random, 95% CI)	0.05 [-0.01, 0.11]
63 Infant/child weight (kg)	11		Mean Difference (IV, Random, 95% CI)	Subtotals only
63.1 At < 3 months	2	863	Mean Difference (IV, Random, 95% CI)	0.01 [-0.07, 0.09]
63.2 At 3 to < 12 months	4	1028	Mean Difference (IV, Random, 95% CI)	0.01 [-0.18, 0.20]
63.3 At 1 to < 2 years	4	1084	Mean Difference (IV, Random, 95% CI)	0.01 [-0.19, 0.21]
63.4 At 2 to < 3 years	2	182	Mean Difference (IV, Random, 95% CI)	0.24 [-0.20, 0.68]
63.5 At 3 to < 4 years	2	1651	Mean Difference (IV, Random, 95% CI)	0.18 [-0.20, 0.57]
63.6 At 4 to < 5 years	2	631	Mean Difference (IV, Random, 95% CI)	0.38 [-0.29, 1.05]
63.7 At 5 to < 6 years	4	2618	Mean Difference (IV, Random, 95% CI)	0.23 [-0.18, 0.63]
63.8 At ≥ 6 years	3	508	Mean Difference (IV, Random, 95% CI)	-0.08 [-0.79, 0.64]
64 Infant/child length/height (cm)	11		Mean Difference (IV, Random, 95% CI)	Subtotals only



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
64.1 < 3 months	2	861	Mean Difference (IV, Random, 95% CI)	-0.01 [-0.69, 0.66]
64.2 3 to < 12 months	4	1115	Mean Difference (IV, Random, 95% CI)	0.11 [-0.20, 0.42]
64.3 1 to < 2 years	4	998	Mean Difference (IV, Random, 95% CI)	0.01 [-0.45, 0.48]
64.4 2 to < 3 years	2	182	Mean Difference (IV, Random, 95% CI)	0.18 [-0.73, 1.08]
64.5 3 to < 4 years	2	1651	Mean Difference (IV, Random, 95% CI)	0.18 [-0.21, 0.58]
64.6 4 to < 5 years	2	631	Mean Difference (IV, Random, 95% CI)	0.30 [-0.36, 0.95]
64.7 5 to < 6 years	5	2733	Mean Difference (IV, Random, 95% CI)	0.20 [-0.17, 0.57]
64.8 At ≥ 6 years	2	393	Mean Difference (IV, Random, 95% CI)	-1.22 [-2.29, -0.16]
65 Infant/child head circumference (cm)	10		Mean Difference (IV, Random, 95% CI)	Subtotals only
65.1 At < 3 months	2	863	Mean Difference (IV, Random, 95% CI)	-0.04 [-0.22, 0.14]
65.2 At 3 to < 12 months	5	1309	Mean Difference (IV, Random, 95% CI)	-0.03 [-0.19, 0.12]
65.3 At 1 to < 2 years	4	1084	Mean Difference (IV, Random, 95% CI)	0.06 [-0.18, 0.30]
65.4 At 2 to < 3 years	2	182	Mean Difference (IV, Random, 95% CI)	-0.04 [-0.47, 0.40]
65.5 At 3 to < 4 years	2	1651	Mean Difference (IV, Random, 95% CI)	-0.01 [-0.16, 0.14]
65.6 At 4 to < 5 years	1	107	Mean Difference (IV, Random, 95% CI)	0.0 [-0.47, 0.47]
65.7 At ≥ 5 years	3	1760	Mean Difference (IV, Random, 95% CI)	0.02 [-0.13, 0.17]
66 Infant/child length/height for age Z score (LAZ/HAZ)	3		Mean Difference (IV, Random, 95% CI)	Subtotals only
66.1 At < 3 months	2	875	Mean Difference (IV, Random, 95% CI)	-0.13 [-0.27, 0.02]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
66.2 At 3 to < 12 months	3	1085	Mean Difference (IV, Random, 95% CI)	-0.05 [-0.19, 0.09]
66.3 At 12 to < 24 months	2	897	Mean Difference (IV, Random, 95% CI)	-0.06 [-0.31, 0.18]
66.4 At 4 to < 5 years	1	524	Mean Difference (IV, Random, 95% CI)	0.0 [-0.15, 0.15]
66.5 At ≥ 5 years	1	802	Mean Difference (IV, Random, 95% CI)	0.0 [-0.12, 0.12]
67 Infant/child waist circum- ference (cm)	2		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
67.1 At 2 to < 3 years	1	101	Mean Difference (IV, Fixed, 95% CI)	-0.20 [-1.29, 0.89]
67.2 At 3 to < 4 years	2	1651	Mean Difference (IV, Fixed, 95% CI)	0.28 [-0.05, 0.60]
67.3 At 4 to < 5 years	1	106	Mean Difference (IV, Fixed, 95% CI)	0.70 [-0.40, 1.80]
67.4 At ≥ 5 years	2	1645	Mean Difference (IV, Fixed, 95% CI)	0.15 [-0.24, 0.55]
68 Infant/child weight-for-age Z score (WAZ)	2		Mean Difference (IV, Random, 95% CI)	Subtotals only
68.1 At < 3 months	2	874	Mean Difference (IV, Random, 95% CI)	-0.09 [-0.30, 0.12]
68.2 At 3 to < 12 months	2	834	Mean Difference (IV, Random, 95% CI)	-0.05 [-0.18, 0.08]
68.3 At 12 to < 24 months	2	883	Mean Difference (IV, Random, 95% CI)	-0.01 [-0.13, 0.12]
68.4 At ≥ 60 months	1	802	Mean Difference (IV, Random, 95% CI)	-0.1 [-0.25, 0.05]
69 Infant/child BMI Z score	5		Mean Difference (IV, Random, 95% CI)	Subtotals only
69.1 At 1 to < 2 years	2	801	Mean Difference (IV, Random, 95% CI)	-0.05 [-0.09, 0.00]
69.2 At 2 to < 3 years	1	63	Mean Difference (IV, Random, 95% CI)	-0.07 [-0.25, 0.11]
69.3 At 3 to < 4 years	1	1531	Mean Difference (IV, Random, 95% CI)	0.02 [-0.08, 0.12]
69.4 At 4 to < 5 years	2	587	Mean Difference (IV, Random, 95% CI)	0.15 [-0.16, 0.47]
69.5 At 5 to < 6 years	3	2504	Mean Difference (IV, Random, 95% CI)	0.03 [-0.05, 0.11]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
69.6 At 6 to < 7 years	1	115	Mean Difference (IV, Random, 95% CI)	0.01 [-0.02, 0.05]
69.7 At ≥ 7 years	1	250	Mean Difference (IV, Random, 95% CI)	0.18 [-0.10, 0.46]
70 Infant/child weight for length/height Z score (WHZ)	3		Mean Difference (IV, Random, 95% CI)	Subtotals only
70.1 At < 3 months	2	860	Mean Difference (IV, Random, 95% CI)	-0.03 [-0.41, 0.34]
70.2 At 3 to < 12 months	3	1083	Mean Difference (IV, Random, 95% CI)	-0.00 [-0.14, 0.14]
70.3 At 12 to < 24 months	2	883	Mean Difference (IV, Random, 95% CI)	-0.02 [-0.14, 0.10]
71 Infant/child BMI percentile	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
71.1 At 24 months	1	118	Mean Difference (IV, Fixed, 95% CI)	4.5 [-5.50, 14.50]
71.2 At 36 months	1	120	Mean Difference (IV, Fixed, 95% CI)	8.0 [-1.09, 17.09]
71.3 At 48 months	1	107	Mean Difference (IV, Fixed, 95% CI)	13.0 [3.19, 22.81]
71.4 At 60 months	1	114	Mean Difference (IV, Fixed, 95% CI)	4.80 [-4.70, 14.30]
72 Child/adult BMI	3		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
72.1 At 3 to 4 years	1	1531	Mean Difference (IV, Fixed, 95% CI)	0.01 [-0.14, 0.16]
72.2 At 5 to 6 years	1	1531	Mean Difference (IV, Fixed, 95% CI)	-0.01 [-0.18, 0.16]
72.3 At 7 to 9 years	2	393	Mean Difference (IV, Fixed, 95% CI)	0.16 [-0.25, 0.57]
72.4 At 19 years	1	243	Mean Difference (IV, Fixed, 95% CI)	0.0 [-0.83, 0.83]
73 Infant/child body fat (%)	3		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
73.1 At 1 year	1	165	Mean Difference (IV, Fixed, 95% CI)	0.0 [-0.88, 0.88]
73.2 At 2 to < 3 years	1	110	Mean Difference (IV, Fixed, 95% CI)	0.20 [-0.68, 1.08]
73.3 At 3 to < 4 years	2	1644	Mean Difference (IV, Fixed, 95% CI)	-0.18 [-0.74, 0.38]
73.4 At 4 to < 5 years	1	102	Mean Difference (IV, Fixed, 95% CI)	0.30 [-0.79, 1.39]
73.5 At 5 to < 6 years	3	1797	Mean Difference (IV, Fixed, 95% CI)	0.01 [-0.56, 0.58]
73.6 At ≥ 7 years: BIS	1	250	Mean Difference (IV, Fixed, 95% CI)	1.44 [-0.31, 3.19]
73.7 At ≥ 7 years: BOD POD	1	250	Mean Difference (IV, Fixed, 95% CI)	-0.42 [-2.23, 1.39]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
74 Infant/child total fat mass (kg)	3		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
74.1 At 1 year	1	164	Mean Difference (IV, Fixed, 95% CI)	0.0 [-0.07, 0.07]
74.2 At 2 to < 3 years	1	110	Mean Difference (IV, Fixed, 95% CI)	0.10 [-0.09, 0.29]
74.3 At 3 to < 4 years	2	1644	Mean Difference (IV, Fixed, 95% CI)	-0.01 [-0.12, 0.10]
74.4 At 4 to < 5 years	1	102	Mean Difference (IV, Fixed, 95% CI)	0.20 [-0.05, 0.45]
74.5 At 5 to < 6 years	3	1797	Mean Difference (IV, Fixed, 95% CI)	0.05 [-0.10, 0.21]
74.6 Up to 8 years: BOD POD	1	250	Mean Difference (IV, Fixed, 95% CI)	0.08 [-0.71, 0.87]
74.7 Up to 8 years: BIS	1	250	Mean Difference (IV, Fixed, 95% CI)	0.29 [-0.47, 1.05]
75 Cognition: thresholds	3		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
75.1 BSID III < 85 at 18 months	1	726	Risk Ratio (M-H, Fixed, 95% CI)	0.49 [0.24, 0.98]
75.2 BSID III > 115 at 18 months	1	726	Risk Ratio (M-H, Fixed, 95% CI)	0.84 [0.49, 1.44]
75.3 BSID II < 85 at 18 months	1	730	Risk Ratio (M-H, Fixed, 95% CI)	1.43 [0.97, 2.12]
75.4 BSID III cognitive score (highest quartile): at 18 months	1	154	Risk Ratio (M-H, Fixed, 95% CI)	0.90 [0.49, 1.65]
76 Cognition: scores	10		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
76.1 BSID II score < 24 months	4	1154	Mean Difference (IV, Fixed, 95% CI)	-0.37 [-1.49, 0.76]
76.2 BSID III score < 24 months	2	809	Mean Difference (IV, Fixed, 95% CI)	0.04 [-1.59, 1.68]
76.3 Fagan novelty preference < 24 months	2	274	Mean Difference (IV, Fixed, 95% CI)	-0.79 [-1.68, 0.11]
76.4 K-ABC mental processing composite at 2 to 5 years	1	84	Mean Difference (IV, Fixed, 95% CI)	4.10 [-0.14, 8.34]
76.5 K-ABC sequential processing at 5 to 6 years	1	96	Mean Difference (IV, Fixed, 95% CI)	0.0 [-1.80, 1.80]
76.6 GMDS general quotient score at 2 to 5 years	1	72	Mean Difference (IV, Fixed, 95% CI)	3.70 [-1.02, 8.42]
76.7 DAS II: General Conceptu- al Ability Scale at 2 to 5 years	1	646	Mean Difference (IV, Fixed, 95% CI)	0.13 [-1.53, 1.79]
76.8 DAS II: Non-verbal Rea- soning Scale at 2 to 5 years	1	646	Mean Difference (IV, Fixed, 95% CI)	-0.35 [-2.04, 1.34]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
76.9 DAS II: Verbal Scale at 2 to 5 years	1	646	Mean Difference (IV, Fixed, 95% CI)	-0.35 [-1.74, 1.04]
76.10 DAS II: Spatial Scale at 2 to 5 years	1	646	Mean Difference (IV, Fixed, 95% CI)	0.96 [-0.77, 2.69]
76.11 MCDS: scale index general cognitive at 5 years	1	797	Mean Difference (IV, Fixed, 95% CI)	-0.5 [-2.35, 1.35]
76.12 WASI full-scale IQ at 6 to 9 years	1	543	Mean Difference (IV, Fixed, 95% CI)	1.0 [-0.79, 2.79]
76.13 WISC-IV full scale IQ at > 12 years	1	50	Mean Difference (IV, Fixed, 95% CI)	1.0 [-5.16, 7.16]
77 Attention: scores	3		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
77.1 K-CPT omissions at 5 years	1	797	Mean Difference (IV, Fixed, 95% CI)	-1.90 [-3.39, -0.41]
77.2 K-CPT commissions at 5 years	1	797	Mean Difference (IV, Fixed, 95% CI)	0.10 [-1.37, 1.57]
77.3 K-CPT hit response time at 5 years	1	797	Mean Difference (IV, Fixed, 95% CI)	-0.60 [-2.06, 0.86]
77.4 Attention: single-object task: total time looking at toy(s) at 2 to 5 years	1	150	Mean Difference (IV, Fixed, 95% CI)	-7.80 [-22.59, 6.99]
77.5 Attention: multiple-object task; # times shifted looks between toys at 2 to 5 years	1	150	Mean Difference (IV, Fixed, 95% CI)	-0.40 [-4.28, 3.48]
77.6 Attention: distractibility: av latency to look when attention focused (s) at 2 to 5 years	1	150	Mean Difference (IV, Fixed, 95% CI)	-0.30 [-0.86, 0.26]
77.7 Attention: global speed (ms) at 8.5 years	1	130	Mean Difference (IV, Fixed, 95% CI)	-5.5 [-47.16, 36.16]
77.8 Attention: interference (ms) at 8.5 years	1	130	Mean Difference (IV, Fixed, 95% CI)	6.97 [-16.42, 30.36]
77.9 Attention: orienting (ms) at 8.5 years	1	130	Mean Difference (IV, Fixed, 95% CI)	3.99 [-16.90, 24.88]
77.10 Attention: alertness (ms) at 8.5 years	1	130	Mean Difference (IV, Fixed, 95% CI)	-5.69 [-27.88, 16.50]
78 Motor: thresholds	2		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
78.1 BSID II score < 85 at 18 months	1	730	Risk Ratio (M-H, Fixed, 95% CI)	0.88 [0.65, 1.19]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
78.2 Fine motor (highest quartile): at 18 months	1	154	Risk Ratio (M-H, Fixed, 95% CI)	1.19 [0.71, 1.99]
78.3 Gross motor (highest quartile): at 18 months	1	154	Risk Ratio (M-H, Fixed, 95% CI)	1.13 [0.68, 1.88]
79 Motor: scores	6		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
79.1 BSID II at < 24 months	4	1153	Mean Difference (IV, Fixed, 95% CI)	0.23 [-0.90, 1.36]
79.2 BSID III at < 24 months	1	726	Mean Difference (IV, Fixed, 95% CI)	0.06 [-1.52, 1.64]
79.3 BSID III fine motor score at < 24 months	1	49	Mean Difference (IV, Fixed, 95% CI)	0.05 [-1.20, 1.30]
79.4 BSID III gross motor score at < 24 months	1	49	Mean Difference (IV, Fixed, 95% CI)	0.05 [-0.68, 0.78]
80 Language: thresholds	2		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
80.1 BSID III < 85	1	726	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.74, 1.40]
80.2 BSID III > 115	1	726	Risk Ratio (M-H, Fixed, 95% CI)	0.82 [0.52, 1.29]
80.3 Receptive language (highest quartile)	1	154	Risk Ratio (M-H, Fixed, 95% CI)	1.82 [1.07, 3.10]
80.4 Expressive language (highest quartile)	1	154	Risk Ratio (M-H, Fixed, 95% CI)	1.65 [1.02, 2.68]
80.5 Infant CDI: words under- stood (highest quartile)	1	159	Risk Ratio (M-H, Fixed, 95% CI)	2.42 [1.33, 4.42]
80.6 Infant CDI: words pro- duced (highest quartile)	1	159	Risk Ratio (M-H, Fixed, 95% CI)	2.08 [1.15, 3.74]
80.7 Infant CDI: words under- stood (highest quartile)	1	134	Risk Ratio (M-H, Fixed, 95% CI)	1.97 [1.11, 3.48]
80.8 Infant CDI: words pro- duced (highest quartile)	1	134	Risk Ratio (M-H, Fixed, 95% CI)	1.97 [1.11, 3.48]
80.9 Toddler CDI: words produced (highest quartile)	1	134	Risk Ratio (M-H, Fixed, 95% CI)	2.09 [1.12, 3.90]
80.10 Non-native constant contrast discrimination	1	144	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.68, 1.40]
81 Language: scores	5		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
81.1 Receptive communication at < 24 months	1	49	Mean Difference (IV, Fixed, 95% CI)	0.55 [-0.77, 1.87]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
81.2 Receptive language (Peabody Picture Vocabulary Test IIIA) at 2 to 5 years	1	70	Mean Difference (IV, Fixed, 95% CI)	3.90 [-0.73, 8.53]
81.3 Expressive communication at < 24 months	1	49	Mean Difference (IV, Fixed, 95% CI)	0.21 [-0.86, 1.28]
81.4 BSID III at < 24 months	2	809	Mean Difference (IV, Fixed, 95% CI)	-0.84 [-2.77, 1.09]
81.5 CELF-P2 Core Language Score at 2 to 5 years	1	646	Mean Difference (IV, Fixed, 95% CI)	-0.93 [-2.92, 1.06]
81.6 CELF-P2 Core Language Score at 6 to 9 years	1	543	Mean Difference (IV, Fixed, 95% CI)	-0.21 [-2.51, 2.09]
81.7 Peabody Picture Vocabulary Test	1	97	Mean Difference (IV, Fixed, 95% CI)	4.0 [-3.11, 11.11]
82 Behaviour: thresholds	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
82.1 Behaviour Rating Scale scores < 26: at < 24 months	1	730	Risk Ratio (M-H, Fixed, 95% CI)	5.0 [0.24, 103.79]
83 Behaviour: scores	6		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
83.1 NBAS habituation	1	27	Mean Difference (IV, Fixed, 95% CI)	-1.45 [-8.49, 5.59]
83.2 NBAS orienting	1	27	Mean Difference (IV, Fixed, 95% CI)	3.65 [-9.09, 16.39]
83.3 NBAS motor	1	27	Mean Difference (IV, Fixed, 95% CI)	2.99 [-8.23, 14.21]
83.4 NBAS state organisation	1	27	Mean Difference (IV, Fixed, 95% CI)	1.63 [-7.21, 10.47]
83.5 NBAS state regulation	1	27	Mean Difference (IV, Fixed, 95% CI)	0.51 [-14.70, 15.72]
83.6 NBAS autonomic	1	27	Mean Difference (IV, Fixed, 95% CI)	3.30 [-8.75, 15.35]
83.7 NBAS reflexes	1	27	Mean Difference (IV, Fixed, 95% CI)	0.68 [-10.28, 11.64]
83.8 BehavioUr Rating Scale score 12 to < 24 months	1	730	Mean Difference (IV, Fixed, 95% CI)	0.0 [-0.94, 0.94]
83.9 Wolke: approach at < 12 months	1	249	Mean Difference (IV, Fixed, 95% CI)	-0.10 [-0.42, 0.22]
83.10 Wolke: activity at < 12 months	1	249	Mean Difference (IV, Fixed, 95% CI)	-0.10 [-0.45, 0.25]
83.11 Wolke: co-operation at < 12 months	1	249	Mean Difference (IV, Fixed, 95% CI)	0.0 [-0.39, 0.39]
83.12 Wolke: emotional tone at < 12 months	1	249	Mean Difference (IV, Fixed, 95% CI)	-0.10 [-0.49, 0.29]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
83.13 Wolke: vocalisation at < 12 months	1	249	Mean Difference (IV, Fixed, 95% CI)	-0.10 [-0.52, 0.32]
83.14 BSID III social-emotional score at < 24 months	2	809	Mean Difference (IV, Fixed, 95% CI)	-0.70 [-3.04, 1.64]
83.15 BSID III adaptive behaviour score at < 24 months	2	809	Mean Difference (IV, Fixed, 95% CI)	-1.20 [-3.12, 0.72]
83.16 SDQ Total Difficulties at 2 to 5 years	1	646	Mean Difference (IV, Fixed, 95% CI)	0.62 [-0.00, 1.24]
83.17 SDQ Total Difficulties at 6 to 9 years	1	543	Mean Difference (IV, Fixed, 95% CI)	1.08 [0.18, 1.98]
83.18 BASC-2: Behavioral Symptoms Index (%) at 5 years	1	797	Mean Difference (IV, Fixed, 95% CI)	-0.5 [-4.54, 3.54]
83.19 CBCL total problem be- haviour at 2 - 5 years	1	72	Mean Difference (IV, Fixed, 95% CI)	-1.0 [-3.41, 1.41]
83.20 CBCL parent report: total behaviours score at 12+ years	1	48	Mean Difference (IV, Fixed, 95% CI)	-0.80 [-5.23, 3.63]
83.21 CBCL parent report: to- tal competence score at > 12 years	1	48	Mean Difference (IV, Fixed, 95% CI)	-0.20 [-6.36, 5.96]
84 Vision: visual acuity (cy- cles/degree)	2		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
84.1 At 2 months	1	135	Mean Difference (IV, Fixed, 95% CI)	0.18 [-0.01, 0.37]
84.2 At 4 months	1	30	Mean Difference (IV, Fixed, 95% CI)	0.5 [-0.43, 1.43]
84.3 At 6 months	1	26	Mean Difference (IV, Fixed, 95% CI)	0.5 [-0.48, 1.48]
85 Vision: VEP acuity	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
85.1 Adjusted VEP acuity at 4 months (cpd)	1	182	Mean Difference (IV, Fixed, 95% CI)	-0.18 [-0.75, 0.39]
85.2 Unadjusted VEP acuity at 4 months (cpd)	1	182	Mean Difference (IV, Fixed, 95% CI)	-0.18 [-0.76, 0.40]
86 Vision: VEP latency	3		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
86.1 Peak latency N1 at birth	1	9	Mean Difference (IV, Fixed, 95% CI)	-12.60 [-29.40, 4.20
86.2 Peak latency P1 at birth	1	14	Mean Difference (IV, Fixed, 95% CI)	-6.80 [-20.44, 6.84]
86.3 Peak latency N2 at birth	1	49	Mean Difference (IV, Fixed, 95% CI)	3.60 [-12.39, 19.59]
86.4 Peak latency P2 at birth	1	55	Mean Difference (IV, Fixed, 95% CI)	0.10 [-16.28, 16.48]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
86.5 Peak latency N3 at birth	1	53	Mean Difference (IV, Fixed, 95% CI)	-6.20 [-36.15, 23.75
86.6 Latency N1 (ms) at 3 months	1	679	Mean Difference (IV, Fixed, 95% CI)	0.30 [-2.21, 2.81]
86.7 Latency P1 (ms) at 3 months	1	679	Mean Difference (IV, Fixed, 95% CI)	-0.5 [-3.19, 2.19]
86.8 Latency N3 (ms) at 3 months	1	679	Mean Difference (IV, Fixed, 95% CI)	-2.30 [-5.91, 1.31]
86.9 Latency (69 min of arc) at 4 months (ms)	1	182	Mean Difference (IV, Fixed, 95% CI)	-1.0 [-3.47, 1.47]
86.10 Latency (48 min of arc) at 4 months (ms)	1	182	Mean Difference (IV, Fixed, 95% CI)	0.0 [-3.20, 3.20]
86.11 Latency (20 min of arc) at 4 months (ms)	1	182	Mean Difference (IV, Fixed, 95% CI)	0.0 [-4.22, 4.22]
86.12 Latency N1 (ms) at 6 months	1	817	Mean Difference (IV, Fixed, 95% CI)	-1.40 [-3.44, 0.64]
86.13 Latency P1 (ms) at 6 months	1	817	Mean Difference (IV, Fixed, 95% CI)	-0.80 [-2.78, 1.18]
86.14 Latency N3 (ms) at 6 months	1	817	Mean Difference (IV, Fixed, 95% CI)	-0.70 [-3.45, 2.05]
87 Hearing: brainstem audito- ry-evoked responses	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
87.1 Latency 1 (ms) at 1 month	1	749	Mean Difference (IV, Fixed, 95% CI)	-0.01 [-0.03, 0.01]
87.2 Latency 3 (ms) at 1 month	1	749	Mean Difference (IV, Fixed, 95% CI)	-0.01 [-0.06, 0.04]
87.3 Latency 5 (ms) at 1 month	1	749	Mean Difference (IV, Fixed, 95% CI)	-0.03 [-0.09, 0.03]
87.4 Interpeak latency 1-3 (ms) at 1 month	1	749	Mean Difference (IV, Fixed, 95% CI)	-0.01 [-0.06, 0.04]
87.5 Interpeak latency 3-5 (ms) at 1 month	1	749	Mean Difference (IV, Fixed, 95% CI)	0.0 [-0.05, 0.05]
87.6 Interpeak latency 1-5 (ms) at 1 month	1	749	Mean Difference (IV, Fixed, 95% CI)	-0.02 [-0.07, 0.03]
87.7 Latency 1 (ms) at 3 months	1	664	Mean Difference (IV, Fixed, 95% CI)	0.0 [-0.02, 0.02]
87.8 Latency 3 (ms) at 3 months	1	664	Mean Difference (IV, Fixed, 95% CI)	0.01 [-0.04, 0.06]
87.9 Latency 5 (ms) at 3 months	1	664	Mean Difference (IV, Fixed, 95% CI)	-0.04 [-0.10, 0.02]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size		
87.10 Interpeak latency 1-3 (ms) at 3 months	1	664	Mean Difference (IV, Fixed, 95% CI)	0.01 [-0.03, 0.05]		
87.11 Interpeak latency 3-5 (ms) at 3 months	1	664	Mean Difference (IV, Fixed, 95% CI)	-0.03 [-0.08, 0.02]		
87.12 Interpeak latency 1-5 (ms) at 3 months	1	664	Mean Difference (IV, Fixed, 95% CI)	-0.03 [-0.09, 0.03]		
88 Neurodevelopment: thresholds	3		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only		
88.1 Hempel: simple minor neurological dysfunction at 18 months	1	114	Risk Ratio (M-H, Fixed, 95% CI)	1.11 [0.80, 1.53]		
88.2 Hempel: simple and com- plex minor neurological dys- function at 4 years	1	167	Risk Ratio (M-H, Fixed, 95% CI)	1.09 [0.37, 3.23]		
88.3 Hempel: complex minor neurological dysfunction at 18 months	1	114	Risk Ratio (M-H, Fixed, 95% CI)	0.68 [0.24, 1.93]		
88.4 ASQ total at 6 months (subnormal - below 2 SD less than mean scores)	1	146	Risk Ratio (M-H, Fixed, 95% CI)	0.54 [0.17, 1.77]		
88.5 Touwen: simple and com- plex minor neurological dys- function at 5.5 years	1	148	Risk Ratio (M-H, Fixed, 95% CI)	1.00 [0.61, 1.63]		
88.6 Neonatal neurological classification: mildly/definitely abnormal at 2 weeks	1	119	Risk Ratio (M-H, Fixed, 95% CI)	0.87 [0.38, 1.97]		
88.7 General movements: mildly/definitely abnormal at 2 weeks	1	119	Risk Ratio (M-H, Fixed, 95% CI)	1.27 [0.75, 2.14]		
88.8 General movements: mildly/definitely abnormal at 12 weeks	1	119	Risk Ratio (M-H, Fixed, 95% CI)	1.54 [0.89, 2.65]		
89 Neurodevelopment: scores	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only		
89.1 ASQ gross motor at 4 months	1	148	Mean Difference (IV, Fixed, 95% CI)	0.30 [-2.38, 2.98]		
89.2 ASQ gross motor at 6 months	1	146	Mean Difference (IV, Fixed, 95% CI)	1.20 [-2.31, 4.71]		
89.3 ASQ fine motor at 4 months	1	148	Mean Difference (IV, Fixed, 95% CI)	1.10 [-2.03, 4.23]		

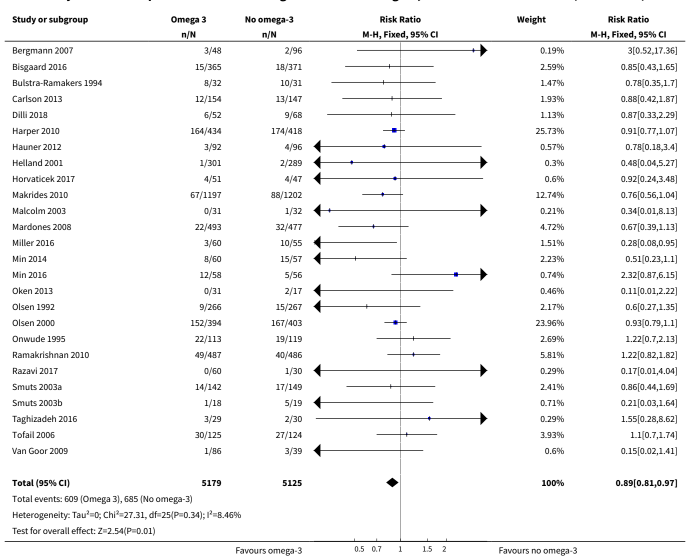


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
89.4 ASQ fine motor at 6 months	1	146	Mean Difference (IV, Fixed, 95% CI)	1.20 [-1.59, 3.99]
89.5 ASQ problem solving at 4 months	1	148	Mean Difference (IV, Fixed, 95% CI)	1.60 [-0.99, 4.19]
89.6 ASQ problem solving at 6 months	1	146	Mean Difference (IV, Fixed, 95% CI)	0.5 [-1.95, 2.95]
89.7 ASQ personal-social at 4 months	1	148	Mean Difference (IV, Fixed, 95% CI)	1.10 [-1.64, 3.84]
89.8 ASQ personal-social at 6 months	1	146	Mean Difference (IV, Fixed, 95% CI)	0.80 [-2.61, 4.21]
89.9 ASQ communication at 4 months	1	148	Mean Difference (IV, Fixed, 95% CI)	2.70 [0.41, 4.99]
89.10 ASQ communication at 6 months	1	146	Mean Difference (IV, Fixed, 95% CI)	0.40 [-1.55, 2.35]
90 Child Development Invento- ry	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
90.1 Social	1	130	Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]
90.2 Self help	1	130	Risk Ratio (M-H, Fixed, 95% CI)	0.29 [0.01, 6.90]
90.3 Gross motor	1	130	Risk Ratio (M-H, Fixed, 95% CI)	4.30 [0.21, 87.76]
90.4 Fine motor	1	130	Risk Ratio (M-H, Fixed, 95% CI)	4.30 [0.21, 87.76]
90.5 Expressive language	1	130	Risk Ratio (M-H, Fixed, 95% CI)	0.86 [0.05, 13.41]
90.6 Language comprehension	1	130	Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]
90.7 Letters	1	130	Risk Ratio (M-H, Fixed, 95% CI)	0.17 [0.01, 3.51]
90.8 Numbers	1	130	Risk Ratio (M-H, Fixed, 95% CI)	0.86 [0.05, 13.41]
90.9 General development	1	130	Risk Ratio (M-H, Fixed, 95% CI)	0.51 [0.13, 2.06]
91 Infant sleep behaviour (%)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
91.1 Arousals in quiet sleep: day 1	1	46	Mean Difference (IV, Fixed, 95% CI)	-3.19 [-6.07, -0.31]
91.2 Arousals in quiet sleep: day 2	1	39	Mean Difference (IV, Fixed, 95% CI)	-1.89 [-4.49, 0.71]
91.3 Quiet sleep: day 1	1	46	Mean Difference (IV, Fixed, 95% CI)	0.74 [-1.97, 3.45]
91.4 Quiet sleep: day 2	1	39	Mean Difference (IV, Fixed, 95% CI)	-1.0 [-4.36, 2.36]
91.5 Active sleep: day 1	1	46	Mean Difference (IV, Fixed, 95% CI)	-2.42 [-8.51, 3.67]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size		
91.6 Active sleep: day 2	1	39	Mean Difference (IV, Fixed, 95% CI)	-0.13 [-8.23, 7.97]		
91.7 Arousals in active sleep: day 1	1	46	Mean Difference (IV, Fixed, 95% CI)	-3.0 [-5.66, -0.34]		
91.8 Arousals in active sleep: day 2	1	46	Mean Difference (IV, Fixed, 95% CI)	-0.63 [-4.12, 2.86]		
92 Cerebral palsy	1	114	Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]		

Analysis 1.1. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 1 Preterm birth (< 37 weeks).





Analysis 1.2. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 2 Early preterm birth (< 34 weeks).

Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
3/32	6/31		4.92%	0.48[0.13,1.77]
1/154	7/147	4 +	5.78%	0.14[0.02,1.09]
4/224	7/121		7.34%	0.31[0.09,1.03]
1/51	0/47		0.42%	2.77[0.12,66.36]
13/1197	27/1202		21.74%	0.48[0.25,0.93]
2/493	10/477	+	8.2%	0.19[0.04,0.88]
4/60	4/57		3.31%	0.95[0.25,3.62]
2/58	0/56		0.41%	4.83[0.24,98.44]
42/394	60/403	-	47.88%	0.72[0.5,1.04]
2663	2541	•	100%	0.58[0.44,0.77]
omega-3)				
=8(P=0.27); I ² =19.12%	6			
	n/N 3/32 1/154 4/224 1/51 13/1197 2/493 4/60 2/58 42/394 2663 omega-3)	n/N n/N 3/32 6/31 1/154 7/147 4/224 7/121 1/51 0/47 13/1197 27/1202 2/493 10/477 4/60 4/57 2/58 0/56 42/394 60/403 2663 2541	n/N	n/N

Analysis 1.3. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 3 Prolonged gestation (> 42 weeks).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
Harris 2015	2/224	0/121	+	1.54%	2.71[0.13,56.02]
Hauner 2012	1/92	0/96		1.16%	3.13[0.13,75.84]
Makrides 2010	6/1184	3/1183		7.14%	2[0.5,7.97]
Mulder 2014	0/68	0/67			Not estimable
Olsen 1992	32/266	27/267	- -	64.13%	1.19[0.73,1.93]
Olsen 2000	26/782	11/791		26.02%	2.39[1.19,4.8]
Total (95% CI)	2616	2525	•	100%	1.61[1.11,2.33]
Total events: 67 (Omega-3), 41	(No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =3.	11, df=4(P=0.54); I ² =0%				
Test for overall effect: Z=2.49(P	=0.01)				
		Favours omega-3	0.1 0.2 0.5 1 2 5 10	Favours no omega-3	

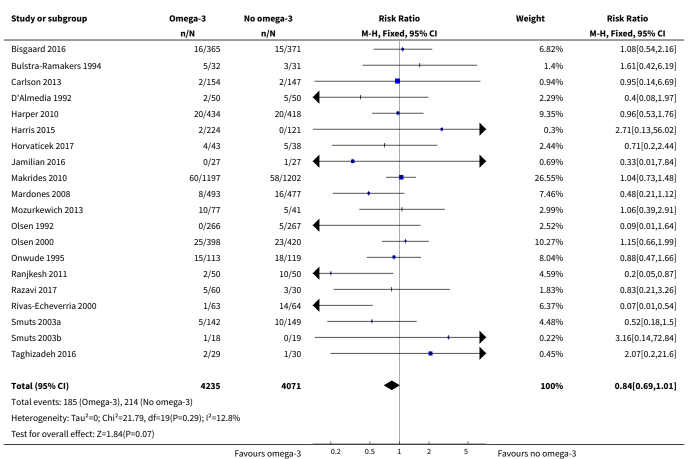
Analysis 1.4. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 4 Maternal death.

Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI						M-H, Fixed, 95% CI	
Bisgaard 2016	0/365	0/371							Not estimable	
Makrides 2010	0/1197	0/1202							Not estimable	
Oken 2013	1/31	0/17		-	-		_	100%	1.69[0.07,39.3]	
Olsen 2000	0/818	0/829							Not estimable	
Total (95% CI)	2411	2419					-	100%	1.69[0.07,39.3]	
Total events: 1 (Omega-3), 0 (No om	iega-3)									
Heterogeneity: Not applicable										
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3		



Study or subgroup	Omega-3 n/N	No omega-3 n/N		Risk Ratio M-H, Fixed, 95% CI				Weight	Risk Ratio M-H, Fixed, 95% CI
Test for overall effect: Z=0.33(P=0.74)									
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3	

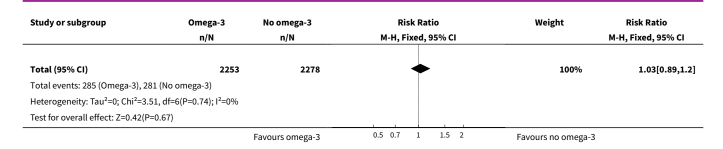
Analysis 1.5. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 5 Pre-eclampsia (hypertension with proteinuria).



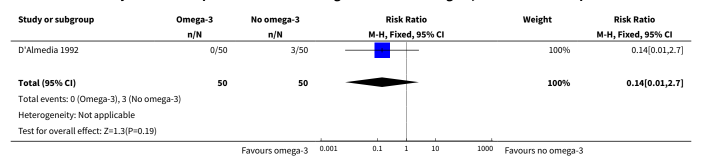
Analysis 1.6. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 6 High blood pressure (without proteinuria).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
n/N n/N		M-H, Fixed, 95% CI		M-H, Fixed, 95% CI	
Bulstra-Ramakers 1994	7/32	4/31		1.46%	1.7[0.55,5.22]
Carlson 2013	32/154	25/147		9.19%	1.22[0.76,1.96]
D'Almedia 1992	9/50	13/50		4.67%	0.69[0.33,1.47]
Makrides 2010	98/1197	107/1202		38.35%	0.92[0.71,1.2]
Olsen 1992	8/266	7/267		2.51%	1.15[0.42,3.12]
Olsen 2000	93/441	90/462	- • -	31.57%	1.08[0.84,1.4]
Onwude 1995	38/113	35/119		12.25%	1.14[0.78,1.67]
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	





Analysis 1.7. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 7 Eclampsia.



Analysis 1.8. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 8 Maternal antepartum hospitalisation.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
1.8.1 Any					
Carlson 2013	14/154	15/147		3.9%	0.89[0.45,1.78]
Jamilian 2016	0/27	3/27	+	- 0.89%	0.14[0.01,2.64]
Makrides 2010	332/1197	362/1202		91.75%	0.92[0.81,1.04]
Taghizadeh 2016	0/29	2/30	+	0.62%	0.21[0.01,4.13]
Subtotal (95% CI)	1407	1406	•	97.16%	0.91[0.8,1.03]
Total events: 346 (Omega-3), 382 (No	omega-3)				
Heterogeneity: Tau ² =0; Chi ² =2.53, df=3	B(P=0.47); I ² =0%				
Test for overall effect: Z=1.53(P=0.13)					
1.8.2 Due to PIH or IUGR					
Bulstra-Ramakers 1994	14/32	11/31		2.84%	1.23[0.67,2.28]
Subtotal (95% CI)	32	31		2.84%	1.23[0.67,2.28]
Total events: 14 (Omega-3), 11 (No om	nega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.67(P=0.51)					
Total (95% CI)	1439	1437	•	100%	0.92[0.81,1.04]
Total events: 360 (Omega-3), 393 (No o	omega-3)				
Heterogeneity: Tau ² =0; Chi ² =3.41, df= ²	4(P=0.49); I ² =0%				
Test for overall effect: Z=1.4(P=0.16)					
Test for subgroup differences: Chi ² =0.9	91, df=1 (P=0.34), I ²	=0%			
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	



Analysis 1.9. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 9 Mother's length of stay in hospital (days).

Study or subgroup	0	mega-3	No	omega-3	a-3 Mean Difference		Mean Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fixed	, 95% CI			Fixed, 95% CI
Bisgaard 2016	365	2.9 (2.6)	371	2.7 (2.8)		_	-		96.79%	0.2[-0.19,0.59]
Olsen 2000	772	6.1 (18.8)	782	6.5 (24)	←	+		—	3.21%	-0.33[-2.47,1.81]
Total ***	1137		1153			-			100%	0.18[-0.2,0.57]
Heterogeneity: Tau ² =0; Chi ² =	=0.23, df=1(P=0.6	3); I ² =0%								
Test for overall effect: Z=0.93	B(P=0.35)									
					-1	-0.5	0 0.5	1	Favours no	omega-3

Analysis 1.10. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 10 Maternal anaemia.

Study or subgroup	Omega-3	No omega-3		Risk Ratio			Weight	Risk Ratio				
	n/N	n/N	M-H, Fixed, 95% CI								M-H, Fixed, 95% CI	
Olsen 2000	101/407	94/439				+	-			100%	1.16[0.91,1.48]	
Total (95% CI)	407	439				•	•			100%	1.16[0.91,1.48]	
Total events: 101 (Omega-3), 94	4 (No omega-3)											
Heterogeneity: Not applicable												
Test for overall effect: Z=1.17(P	=0.24)			1								
		Favours omega-3	0.1	0.2	0.5	1	2	5	10	Favours no omega-3		

Analysis 1.11. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 11 Miscarriage (< 24 weeks).

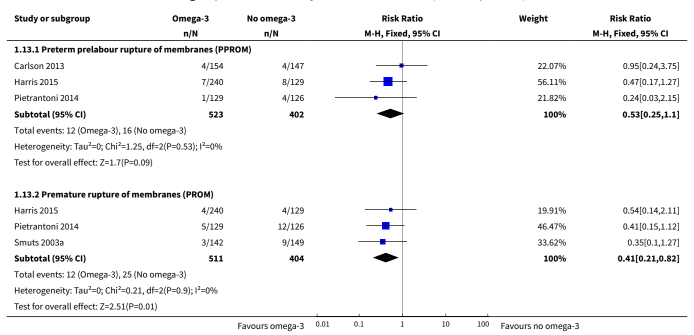
Study or subgroup	Omega-3	No omega-3		Ri	sk Ratio		Weight	Risk Ratio
	n/N	n/N		М-Н, F	ixed, 95% CI			M-H, Fixed, 95% CI
Bulstra-Ramakers 1994	0/34	1/34	→	+			1.86%	0.33[0.01,7.91]
Carlson 2013	4/154	3/147		-	+		3.8%	1.27[0.29,5.59]
Haghiac 2015	1/36	1/36	+				1.24%	1[0.07,15.38]
Helland 2001	1/301	1/289	+		-		1.26%	0.96[0.06,15.28]
Horvaticek 2017	4/58	3/53			+		3.88%	1.22[0.29,5.19]
Mardones 2008	63/589	52/552			_		66.44%	1.14[0.8,1.61]
Min 2014	11/86	9/87			+		11.07%	1.24[0.54,2.83]
Min 2016	0/57	1/58	+				1.84%	0.34[0.01,8.15]
Olsen 2000	4/804	7/815		-			8.6%	0.58[0.17,1.97]
Total (95% CI)	2119	2071			•		100%	1.07[0.8,1.43]
Total events: 88 (Omega-3), 78 (No om	ega-3)							
Heterogeneity: Tau ² =0; Chi ² =2.3, df=8(P=0.97); I ² =0%							
Test for overall effect: Z=0.48(P=0.63)				1				
		Favours omega-3	0.1	0.2 0.5	1 2	5 10	Favours no omega-3	



Analysis 1.12. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 12 Antepartum vaginal bleeding.

Study or subgroup	Omega-3	No omega-3			Ri	sk Rat	io			Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI								M-H, Fixed, 95% CI	
Olsen 1992	15/266	12/267			_	-				23.65%	1.25[0.6,2.63]	
Olsen 2000	36/802	39/816			-	-	-			76.35%	0.94[0.6,1.46]	
Total (95% CI)	1068	1083				•	-			100%	1.01[0.69,1.48]	
Total events: 51 (Omega-3), 51	(No omega-3)											
Heterogeneity: Tau ² =0; Chi ² =0.	43, df=1(P=0.51); I ² =0%					İ						
Test for overall effect: Z=0.07(F	P=0.94)											
		Favours omega-3	0.1	0.2	0.5	1	2	5	10	Favours no omega-3		

Analysis 1.13. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 13 Rupture of membranes (PPROM; PROM).



Analysis 1.14. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 14 Maternal admission to intensive care.

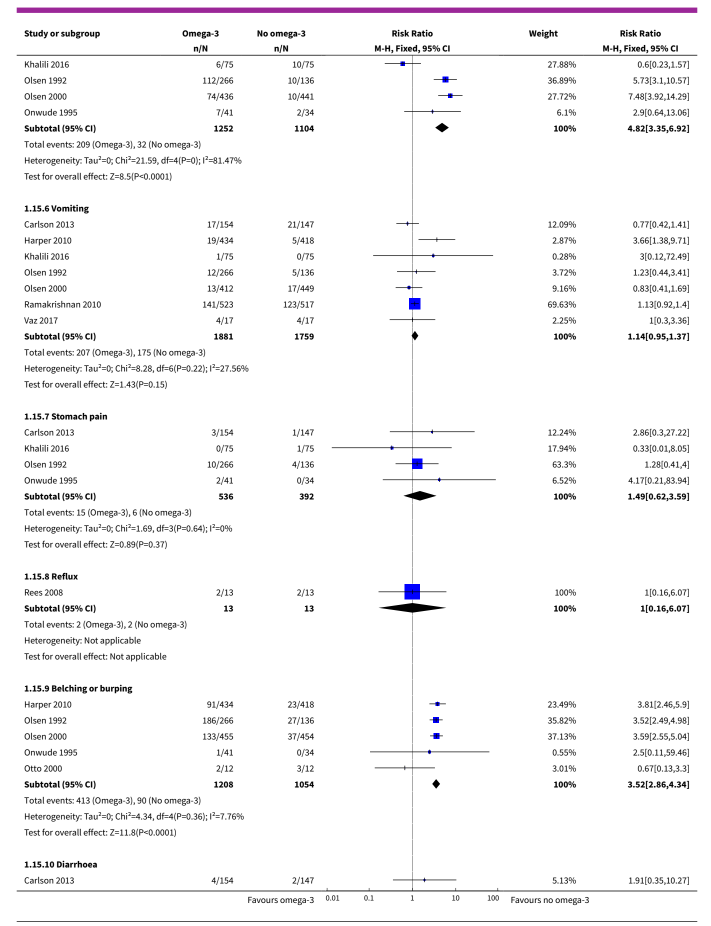
Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		М-Н	Fixed, 95	% CI			M-H, Fixed, 95% CI
Makrides 2010	2/1197	2/1202			-			44.8%	1[0.14,7.12]
Taghizadeh 2016	0/29	2/30		-		_		55.2%	0.21[0.01,4.13]
Total (95% CI)	1226	1232		-				100%	0.56[0.12,2.63]
Total events: 2 (Omega-3), 4 (No on	nega-3)								
Heterogeneity: Tau ² =0; Chi ² =0.77, d	f=1(P=0.38); I ² =0%								
Test for overall effect: Z=0.73(P=0.4	7)								
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3	



Analysis 1.15. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 15 Maternal adverse events.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
1.15.1 Severe adverse event	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
	2/1107	2/1202		25 420/	1[0 14 7 12]
Makrides 2010 Smuts 2003a	2/1197	2/1202		25.42%	1[0.14,7.12]
Subtotal (95% CI)	6/142 1339	6/149 1351		74.58% 100%	1.05[0.35,3.18] 1.04[0.4,2.72]
Total events: 8 (Omega-3), 8 (No on		1331		100%	1.04[0.4,2.72]
Heterogeneity: Tau ² =0; Chi ² =0, df=1					
Test for overall effect: Z=0.08(P=0.9					
1656101 0Verall effect. 2 0.00(1 0.5	•,				
1.15.2 Severe enough for cessation	on				
Bulstra-Ramakers 1994	2/34	2/34		12.39%	1[0.15,6.7]
Freeman 2008	0/12	0/9			Not estimable
Khalili 2016	1/75	0/75	-	3.1%	3[0.12,72.49]
Mozurkewich 2013	9/77	7/41		56.62%	0.68[0.27,1.7]
Ramakrishnan 2010	6/547	3/547	-	18.59%	2[0.5,7.96]
Su 2008	0/18	1/18 —	•	9.3%	0.33[0.01,7.68]
Subtotal (95% CI)	763	724	*	100%	1.01[0.53,1.93]
Total events: 18 (Omega-3), 13 (No	omega-3)				
Heterogeneity: Tau ² =0; Chi ² =2.57, d	If=4(P=0.63); I ² =0%				
Test for overall effect: Z=0.02(P=0.9	8)				
1.15.3 Any					
Noakes 2012	0/54	0/54			Not estimable
Olsen 2000	152/475	84/495	.	52.86%	1.89[1.49,2.38]
Onwude 1995	17/41	8/34	-	5.62%	1.76[0.87,3.57]
Smuts 2003a	38/142	58/149	-	36.37%	0.69[0.49,0.96]
Su 2008	6/18	8/18		5.14%	0.75[0.33,1.72]
Subtotal (95% CI)	730	750	•	100%	1.38[1.16,1.65]
Total events: 213 (Omega-3), 158 (N	lo omega-3)				
Heterogeneity: Tau ² =0; Chi ² =25.62,	df=3(P<0.0001); I ² =88	.29%			
Test for overall effect: Z=3.59(P=0)					
1.15.4 Nausea					
Carlson 2013	16/154	21/147		8.7%	0.73[0.4,1.34]
Khalili 2016	7/75	10/75		4.05%	0.7[0.28,1.74]
Olsen 1992	28/266	7/136	 	3.75%	2.05[0.92,4.56]
Olsen 2000	29/417	40/458	-+	15.44%	0.8[0.5,1.26]
Onwude 1995	4/41	1/34	-	0.44%	3.32[0.39,28.3]
Otto 2000	1/12	4/12		1.62%	0.25[0.03,1.92]
Ramakrishnan 2010	176/523	157/517	<u> </u>	63.96%	1.11[0.93,1.32]
Rees 2008	1/13	1/13		0.41%	1[0.07,14.34]
Su 2008	6/18	4/18		1.62%	1.5[0.51,4.43]
Subtotal (95% CI)	1519	1410	•	100%	1.05[0.9,1.22]
Total events: 268 (Omega-3), 245 (N	lo omega-3)				
Heterogeneity: Tau ² =0; Chi ² =9.98, d	If=8(P=0.27); I ² =19.87 ⁰	%			
Test for overall effect: Z=0.59(P=0.5	5)				
1.15.5 Unpleasant taste					
Harper 2010	10/434	0/418		1.42%	20.23[1.19,344.09]
		Favours omega-3 0.01	0.1 1 10	100 Favours no omega-3	

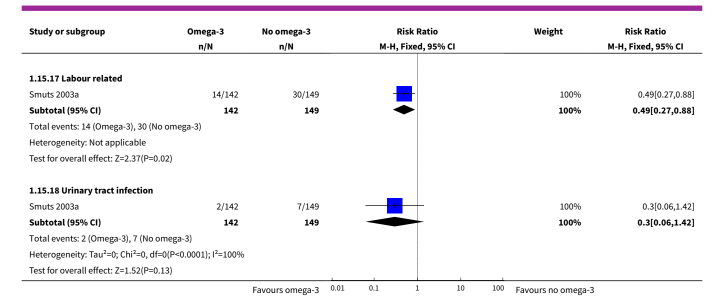






Study or subgroup	Omega-3 n/N	No omega-3 n/N	Risk Ratio M-H, Fixed, 95% CI	Weight	Risk Ratio M-H, Fixed, 95% CI
Khalili 2016	1/75	0/75		1.25%	3[0.12,72.49
Olsen 1992	24/266	18/136	-	59.68%	0.68[0.38,1.21
Olsen 2000	7/407	11/442		26.42%	0.69[0.27,1.77
Rees 2008	2/13	1/13		2.51%	2[0.21,19.44
Su 2008	1/18	2/18		5.01%	0.5[0.05,5.04
Subtotal (95% CI)	933	831		100%	0.8[0.52,1.24
Total events: 39 (Omega-3), 34 (No omeg				20070	010[010_j_1_
Heterogeneity: Tau ² =0; Chi ² =2.86, df=5(P					
Test for overall effect: Z=0.99(P=0.32)	-0.12),1 -070				
1.15.11 Constipation					
Olsen 2000	2/526	5/551		100%	0.42[0.08,2.15
Subtotal (95% CI)	526	551		100%	0.42[0.08,2.1
Total events: 2 (Omega-3), 5 (No omega-3	3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=1.04(P=0.3)					
1.15.12 Nasal bleeding					
Olsen 1992	24/266	26/267	-	28.77%	0.93[0.55,1.5
Olsen 2000	60/481	65/492		71.23%	0.94[0.68,1.3
Subtotal (95% CI)	747	759	*	100%	0.94[0.71,1.2
Fotal events: 84 (Omega-3), 91 (No omega	a-3)				
Heterogeneity: Tau ² =0; Chi ² =0, df=1(P=0.	95); I ² =0%				
Test for overall effect: Z=0.44(P=0.66)					
1.15.13 Swelling/other reaction at inje	ction site				
Harper 2010	279/434	245/418	+	100%	1.1[0.99,1.2
Subtotal (95% CI)	434	418	>	100%	1.1[0.99,1.2
Total events: 279 (Omega-3), 245 (No om	ega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=1.7(P=0.09)					
1.15.14 Insomnia					
Su 2008	3/18	2/18	- 1	100%	1.5[0.28,7.9
Subtotal (95% CI)	18	18		100%	1.5[0.28,7.9
Total events: 3 (Omega-3), 2 (No omega-3	3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.48(P=0.63)					
1.15.15 Headache					
Carlson 2013	27/154	16/147	 	100%	1.61[0.91,2.8
Subtotal (95% CI)	154	147	•	100%	1.61[0.91,2.8
Total events: 27 (Omega-3), 16 (No omeg	a-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=1.62(P=0.1)					
1.15.16 Gynaecological infections					
Smuts 2003a	16/142	20/149	-	100%	0.84[0.45,1.5
Subtotal (95% CI)	142	149	•	100%	0.84[0.45,1.5
Total events: 16 (Omega-3), 20 (No omega	a-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.56(P=0.58)					

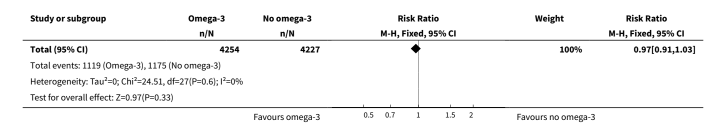




Analysis 1.16. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 16 Caesarean section.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
Ali 2017	17/34	19/34		1.64%	0.89[0.57,1.4]
Bergmann 2007	33/43	55/77		3.4%	1.07[0.86,1.33]
Bisgaard 2016	77/365	71/371		6.08%	1.1[0.83,1.47]
Bulstra-Ramakers 1994	8/32	10/31	-	0.88%	0.78[0.35,1.7]
Carlson 2013	46/154	44/147		3.89%	1[0.71,1.41]
de Groot 2004	3/29	4/29	<u> </u>	0.35%	0.75[0.18,3.06]
Dilli 2018	34/52	54/68	-+-	4.04%	0.82[0.65,1.04]
Dunstan 2008	11/40	8/43	-	0.67%	1.48[0.66,3.3]
Hauner 2012	31/96	31/92		2.73%	0.96[0.64,1.44]
Helland 2001	28/175	14/166		1.24%	1.9[1.04,3.48]
Jamilian 2016	12/26	18/27		1.53%	0.69[0.42,1.13]
Judge 2007	8/27	5/21		0.49%	1.24[0.48,3.25]
Khalili 2016	30/75	33/75		2.85%	0.91[0.62,1.33]
Makrides 2010	326/1197	350/1202		30.17%	0.94[0.82,1.06]
Mardones 2008	17/493	22/477		1.93%	0.75[0.4,1.39]
Miller 2016	20/60	12/55	+	1.08%	1.53[0.83,2.83]
Min 2014	25/60	24/57		2.13%	0.99[0.65,1.52]
Min 2016	28/58	29/56		2.55%	0.93[0.65,1.35]
Mozurkewich 2013	22/77	11/41		1.24%	1.06[0.57,1.97]
Noakes 2012	8/53	9/54		0.77%	0.91[0.38,2.17]
Olsen 1992	16/266	20/267		1.72%	0.8[0.43,1.52]
Onwude 1995	36/113	25/119	 	2.1%	1.52[0.98,2.36]
Ramakrishnan 2010	216/429	234/440		19.96%	0.95[0.83,1.08]
Ranjkesh 2011	23/50	22/50		1.9%	1.05[0.68,1.61]
Razavi 2017	17/60	11/30	+	1.27%	0.77[0.42,1.43]
Smuts 2003a	18/142	21/149		1.77%	0.9[0.5,1.62]
Smuts 2003b	2/18	6/19		0.5%	0.35[0.08,1.52]
	7/30	13/30	ı İ	1.12%	0.54[0.25,1.16]





Analysis 1.17. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 17 Induction (post-term).

Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio	
	n/N	n/N	n/N M-H, Random			% CI			M-H, Random, 95% CI	
Harris 2015	4/224	6/121		_	-			39.31%	0.36[0.1,1.25]	
Hauner 2012	0/96	0/92							Not estimable	
Makrides 2010	153/1184	110/1183			-			60.69%	1.39[1.1,1.75]	
Total (95% CI)	1504	1396		-				100%	0.82[0.22,2.98]	
Total events: 157 (Omega-3), 1	16 (No omega-3)									
Heterogeneity: Tau ² =0.7; Chi ² =	4.37, df=1(P=0.04); I ² =77.1	%								
Test for overall effect: Z=0.31(F	P=0.76)									
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3		

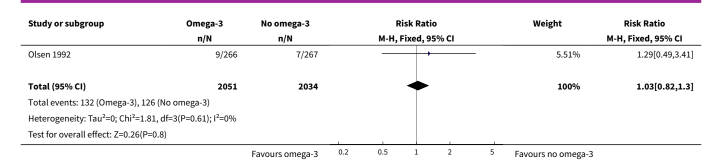
Analysis 1.18. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 18 Blood loss at birth (mL).

Study or subgroup	Oı	nega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
Furuhjelm 2009	54	473 (232)	66	492 (232)	+	4.79%	-19[-102.44,64.44]
Hauner 2012	92	377 (153)	96	366 (124)		20.92%	11[-28.91,50.91]
Helland 2001	175	362 (219)	166	354 (324)	+ • • • • • • • • • • • • • • • • • • •	9.57%	8[-51.01,67.01]
Mozurkewich 2013	77	507.5 (411.5)	41	454 (296)	-	2%	53.49[-75.57,182.55]
Olsen 1992	266	316 (260)	267	290 (213)	-	20.46%	26[-14.36,66.36]
Olsen 2000	725	351.7 (282.7)	751	344.7 (267.1)		42.27%	7[-21.08,35.08]
Total ***	1389		1387			100%	11.5[-6.75,29.76]
Heterogeneity: Tau ² =0; Chi ² =	=1.53, df=5(P=0.9	1); I ² =0%					
Test for overall effect: Z=1.24	1(P=0.22)						
			Fav	ours omega-3	-40 -20 0 20 40	Favours no	omega-3

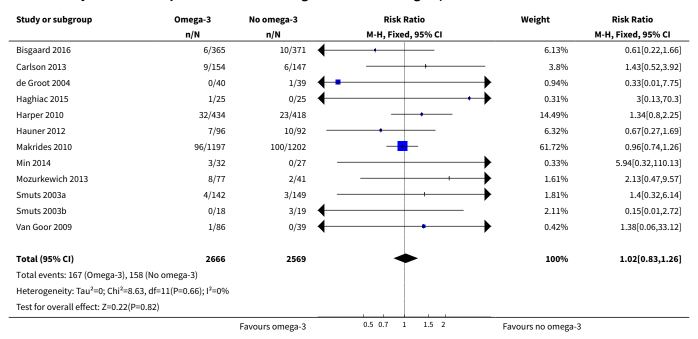
Analysis 1.19. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 19 Postpartum haemorrhage.

Study or subgroup	Omega-3	No omega-3		R	isk Rati	0		Weight	Risk Ratio
	n/N	n/N		М-Н,	Fixed, 9	5% CI			M-H, Fixed, 95% CI
Carlson 2013	6/154	3/147				1	\longrightarrow	2.42%	1.91[0.49,7.49]
Harper 2010	60/434	52/418			-	_		41.75%	1.11[0.79,1.57]
Makrides 2010	57/1197	64/1202			-			50.33%	0.89[0.63,1.27]
		Favours omega-3	0.2	0.5	1	2	5	Favours no omega-3	





Analysis 1.20. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 20 Gestational diabetes.



Analysis 1.21. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 21 Maternal insulin resistance (HOMA-IR).

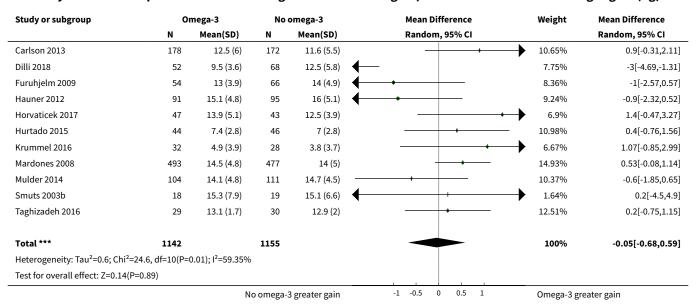
Study or subgroup	Oı	mega-3	No	omega-3		Mea	n Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rand	lom, 95% CI			Random, 95% CI
Krummel 2016	32	3.7 (2.1)	28	3.2 (1)			+		35.64%	0.53[-0.29,1.35]
Samimi 2015	28	3.1 (1.5)	28	4.3 (3.3)					31.05%	-1.2[-2.54,0.14]
Taghizadeh 2016	30	2.5 (1)	30	4.5 (2.9)		-			33.31%	-2[-3.1,-0.9]
Total ***	90		86						100%	-0.85[-2.5,0.8]
Heterogeneity: Tau ² =1.82; Chi ² =	:14.31, df=2(P:	=0); I ² =86.02%								
Test for overall effect: Z=1.01(P=	0.31)									
			Fav	ours omega-3	-5	-2.5	0 2.5	5	Favours no c	mega-3



Analysis 1.22. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 22 Excessive gestational weight gain.

Study or subgroup	Omega-3	No omega-3			Weight	Risk Ratio			
	n/N	n/N		M-H, I	Fixed, 9	5% CI			M-H, Fixed, 95% CI
Carlson 2013	84/178	67/172			+			100%	1.21[0.95,1.55]
Total (95% CI)	178	172			•	•		100%	1.21[0.95,1.55]
Total events: 84 (Omega-3), 67 (No or	mega-3)								
Heterogeneity: Not applicable									
Test for overall effect: Z=1.55(P=0.12)			1	1					
		Favours omega-3	0.2	0.5	1	2	5	Favours no omega-3	

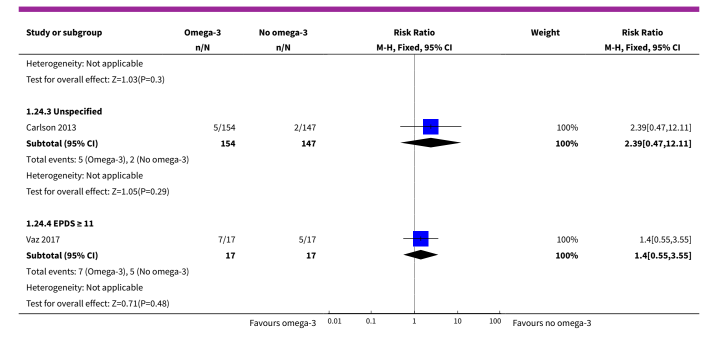
Analysis 1.23. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 23 Gestational weight gain (kg).



Analysis 1.24. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 24 Depression during pregnancy: thresholds.

Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio	
	n/N			M-F	I, Fixed, 95%	6 CI			M-H, Fixed, 95% CI	
1.24.1 HAMD 50% reduction (after 8 v	weeks)									
Su 2008	8/13	3/11			+	_		100%	2.26[0.78,6.49]	
Subtotal (95% CI)	13	11				-		100%	2.26[0.78,6.49]	
Total events: 8 (Omega-3), 3 (No omega	a-3)									
Heterogeneity: Not applicable										
Test for overall effect: Z=1.51(P=0.13)										
1.24.2 HAMD ≤ 7										
Su 2008	5/13	2/11			-			100%	2.12[0.51,8.84]	
Subtotal (95% CI)	13	11				-		100%	2.12[0.51,8.84]	
Total events: 5 (Omega-3), 2 (No omega	a-3)									
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3		

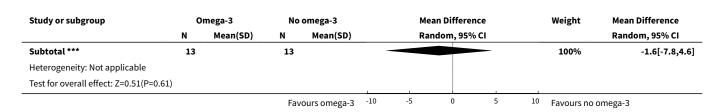




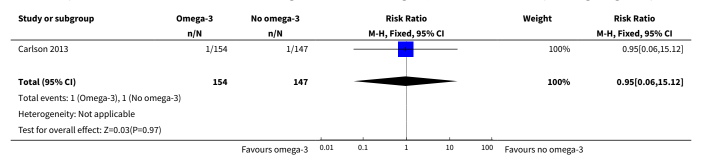
Analysis 1.25. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 25 Depression during pregnancy: scores.

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
1.25.1 BDI							
Kaviani 2014	40	9.2 (5.3)	40	14.7 (6.5)		90.54%	-5.53[-8.12,-2.94]
Su 2008	13	10.8 (8.3)	11	19.8 (11.2)		9.46%	-9[-17.01,-0.99]
Subtotal ***	53		51	-	•	100%	-5.86[-8.32,-3.39]
Heterogeneity: Tau ² =0; Chi ² =0.65	s, df=1(P=0.4	2); I ² =0%					
Test for overall effect: Z=4.66(P<0	0.0001)						
1.25.2 HAMD							
Freeman 2008	12	14.2 (5.8)	9	10.2 (3.7)		32.84%	3.95[-0.13,8.03]
Rees 2008	13	7.9 (5.1)	13	9.7 (5.1)		33.39%	-1.8[-5.72,2.12]
Su 2008	13	9 (4)	11	13.8 (5.3)		33.77%	-4.8[-8.61,-0.99]
Subtotal ***	38		33			100%	-0.92[-5.91,4.06]
Heterogeneity: Tau ² =15.38; Chi ² =	9.63, df=2(P	=0.01); I ² =79.23%	6				
Test for overall effect: Z=0.36(P=0	0.72)						
1.25.3 EPDS							
Freeman 2008	12	11.2 (6.8)	9	7.8 (4.2)	-	22.28%	3.39[-1.31,8.09]
Keenan 2014	34	12.1 (4.8)	17	11.4 (5.1)		30.87%	0.77[-2.14,3.68]
Rees 2008	13	8.5 (5.5)	13	9 (5.2)		24.9%	-0.5[-4.61,3.61]
Su 2008	13	8.5 (5.5)	11	14.3 (6.3)		21.95%	-5.8[-10.57,-1.03]
Subtotal ***	72		50			100%	-0.4[-3.7,2.89]
Heterogeneity: Tau ² =6.96; Chi ² =7	7.97, df=3(P=	0.05); I ² =62.37%					
Test for overall effect: Z=0.24(P=0	0.81)						
1.25.4 MADRS							
Rees 2008	13	13.5 (8.6)	13	15.1 (7.5)		100%	-1.6[-7.8,4.6]





Analysis 1.26. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 26 Anxiety during pregnancy.



Analysis 1.27. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 27 Difficult life circumstances (maternal).

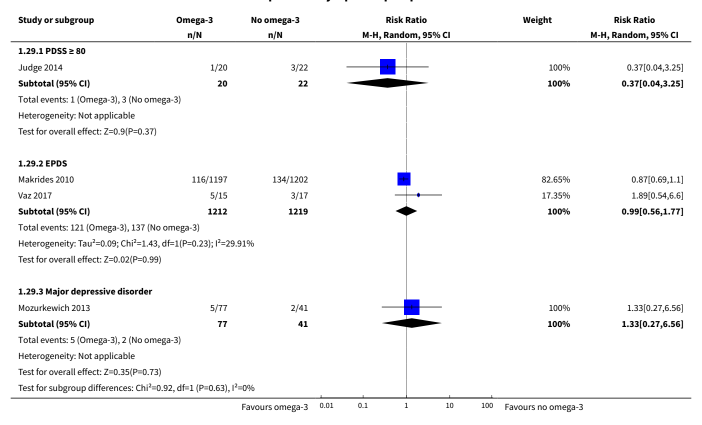
Study or subgroup	Omega-3		No omega-3			Mea	n Differen	ce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fix	ked, 95% C	I			Fixed, 95% CI
Keenan 2014	34	3.9 (0.6)	17	3.6 (0.9)			1			100%	0.32[-0.15,0.79]
Total ***	34		17				•			100%	0.32[-0.15,0.79]
Heterogeneity: Not applicable											
Test for overall effect: Z=1.33(P=0.18)										
			Favo	ours omega-3	-4	-2	0	2	4	Favours no	omega-3

Analysis 1.28. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 28 Stress (maternal).

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
1.28.1 Perceived Stress Scal	e (scores)						
Keenan 2014	34	27.5 (3.4)	17	29.3 (3.1)		100%	-1.82[-3.68,0.04]
Subtotal ***	34		17			100%	-1.82[-3.68,0.04]
Heterogeneity: Tau ² =0; Chi ² =0	, df=0(P<0.0001	L); I ² =100%					
Test for overall effect: Z=1.91(P=0.06)						
			Fav	ours omega-3	-5 -2.5 0 2.5 5	Favours no	omega-3



Analysis 1.29. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 29 Depressive symptoms postpartum: threshold.



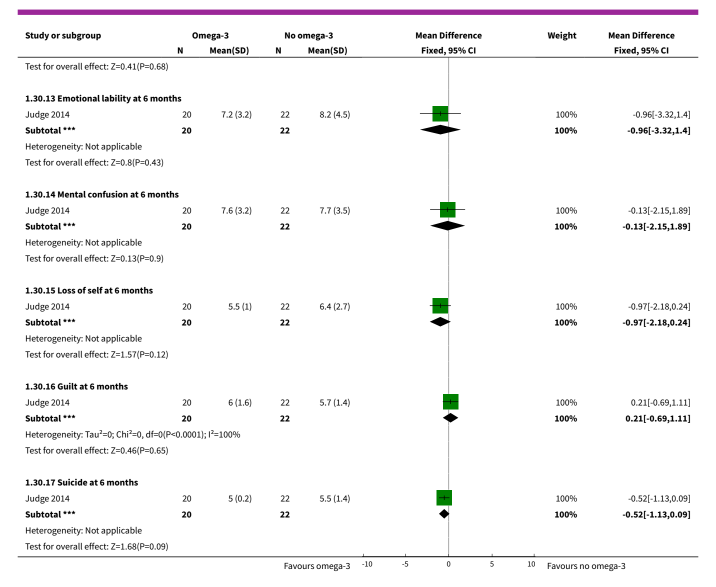
Analysis 1.30. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 30 Depressive symptoms postpartum: scores.

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
1.30.1 BDI: 6-8 weeks postpartum							
Mozurkewich 2013	77	6.2 (5)	41	5.9 (6.1)	— —	100%	0.25[-1.93,2.43]
Subtotal ***	77		41			100%	0.25[-1.93,2.43]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.23(P=0.82)						
1.30.2 PDSS total (LS over 6 mont)	ıs)						
Judge 2014	20	46 (9.7)	22	52.1 (11.3)	 	100%	-6.08[-12.42,0.26]
Subtotal ***	20		22			100%	-6.08[-12.42,0.26]
Heterogeneity: Tau ² =0; Chi ² =0, df=0	P<0.000	L); I ² =100%					
Test for overall effect: Z=1.88(P=0.06)						
1.30.3 Disturbances sleep/eating (LS over (5 months)					
Judge 2014	20	8.1 (2.7)	22	9.1 (2.8)	-	100%	-1[-2.66,0.66]
Subtotal ***	20		22			100%	-1[-2.66,0.66]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.18(P=0.24	.)						
			Fav	ours omega-3	-10 -5 0 5	10 Favours no	omega-3



Study or subgroup	Oı N	mega-3 Mean(SD)	No N	omega-3 Mean(SD)	Mean Difference Fixed, 95% CI	Weight	Mean Difference Fixed, 95% CI
1.30.4 Anxiety/insecurity (LS over				(02)	1 1.102,00 % 0.1		
Judge 2014	20	7.7 (2.7)	22	9 (2.8)		100%	-1.3[-2.96,0.3
Subtotal ***	20	(=,	22	- (=,		100%	-1.3[-2.96,0.3
Heterogeneity: Not applicable							,
Test for overall effect: Z=1.53(P=0.13)						
1.30.5 Emotional lability (LS over	6 months	s)					
Judge 2014	20	7 (2.7)	22	8.3 (3.3)	-	100%	-1.29[-3.1,0.52
Subtotal ***	20		22			100%	-1.29[-3.1,0.52
Heterogeneity: Not applicable							
Test for overall effect: Z=1.4(P=0.16)							
1.30.6 Mental confusion (LS over 6	months)					
Judge 2014	20	7.1 (2.5)	22	8.4 (2.8)		100%	-1.3[-2.92,0.3
Subtotal ***	20		22		•	100%	-1.3[-2.92,0.32
Heterogeneity: Not applicable							
Test for overall effect: Z=1.57(P=0.12)						
1.30.7 Loss of self (LS over 6 mont	hs)						
Judge 2014	20	5.5 (1.3)	22	6.4 (1.6)	-	100%	-0.9[-1.8,0
Subtotal ***	20		22		•	100%	-0.9[-1.8,0
Heterogeneity: Tau²=0; Chi²=0, df=0	P<0.0001	.); I ² =100%					
Test for overall effect: Z=1.95(P=0.05)						
1.30.8 Guilt (LS over 6 months)							
Judge 2014	20	5.4 (1.3)	22	5.7 (1.4)		100%	-0.3[-1.13,0.5
Subtotal ***	20		22		◆	100%	-0.3[-1.13,0.5
Heterogeneity: Not applicable							
Test for overall effect: Z=0.71(P=0.48)						
1.30.9 Suicide (LS over 6 months)							
Judge 2014	20	5.1 (0.4)	22	5.2 (0.5)	+	100%	-0.07[-0.35,0.2]
Subtotal ***	20		22		▼	100%	-0.07[-0.35,0.21
Heterogeneity: Not applicable							
Test for overall effect: Z=0.5(P=0.62)							
1.30.10 PDSS total at 6 months							
Judge 2014	20	45.6 (13.5)	22	48.4 (17.2)		100%	-2.87[-12.17,6.43
Subtotal ***	20		22	•		100%	-2.87[-12.17,6.43
Heterogeneity: Not applicable							
Test for overall effect: Z=0.6(P=0.55)							
1.30.11 Disturbances sleep/eating	at 6 mor	nths			<u> </u>		
Judge 2014	20	6.8 (3.4)	22	7 (2.7)		100%	-0.2[-2.08,1.68
Subtotal ***	20		22		•	100%	-0.2[-2.08,1.68
Heterogeneity: Not applicable							
Test for overall effect: Z=0.21(P=0.83)						
1.30.12 Anxiety/insecurity at 6 mo	nths						
Judge 2014	20	7.7 (3.7)	22	8.1 (3.5)		100%	-0.46[-2.65,1.7
Subtotal ***	20		22			100%	-0.46[-2.65,1.73
Heterogeneity: Not applicable							

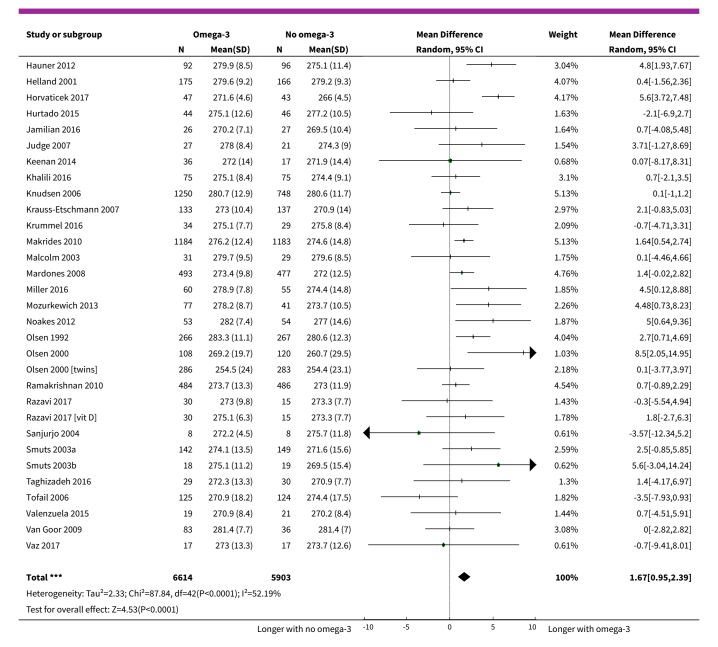




Analysis 1.31. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 31 Gestational length (days).

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Ali 2017	34	252 (4.9)	34	252 (6.3)		3.23%	0[-2.68,2.68]
Bergmann 2007	43	273.7 (11.5)	74	276.5 (9.7)		2.04%	-2.8[-6.88,1.28]
Carlson 2013	154	275.7 (11.2)	147	272.8 (17)	 	2.66%	2.9[-0.37,6.17]
de Groot 2004	29	281 (7.4)	29	276.5 (12.2)	+	1.45%	4.5[-0.69,9.69]
Dilli 2018	52	266 (12.6)	68	261.8 (14)	+	1.64%	4.2[-0.58,8.98]
Dunstan 2008	40	275 (6.3)	43	274 (6.6)	- +	3.13%	1[-1.77,3.77]
Furuhjelm 2009	54	280 (9.8)	66	280 (11.2)		2.26%	0[-3.76,3.76]
Giorlandino 2013	21	268.8 (9.1)	21	255.5 (12.6)		0.98%	13.3[6.65,19.95]
Gustafson 2013	22	275.8 (7.7)	24	279.3 (7.7)		1.81%	-3.5[-7.95,0.95]
Haghiac 2015	25	274.4 (11.2)	24	270.9 (8.4)	+	1.32%	3.5[-2.03,9.03]
Harper 2010	434	263.9 (28.4)	418	261.8 (28.4)		2.22%	2.1[-1.71,5.91]
Harris 2015	224	275.3 (19.5)	121	271.6 (13.2)		2.49%	3.7[0.23,7.17]
		L	onger wit	th no omega-3	-10 -5 0 5	10 Longer with	n omega-3

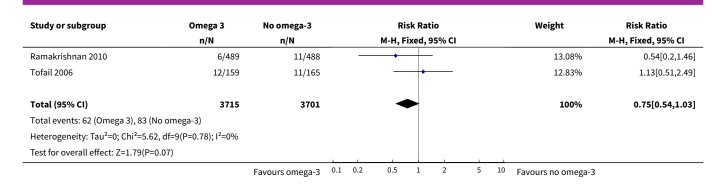




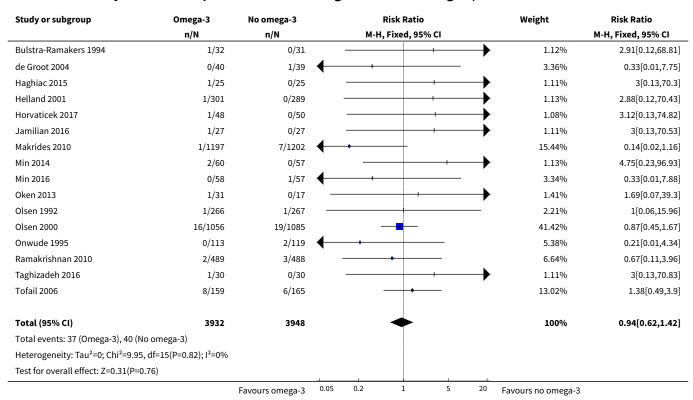
Analysis 1.32. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 32 Perinatal death.

Study or subgroup	Omega 3	No omega-3		Risk Ratio)		Weight	Risk Ratio
	n/N	n/N		M-H, Fixed, 95	5% CI			M-H, Fixed, 95% CI
Ali 2017	2/34	3/34	_	+			3.56%	0.67[0.12,3.74]
Bulstra-Ramakers 1994	2/32	3/31	_	+			3.62%	0.65[0.12,3.61]
Harper 2010	16/434	17/418			-		20.58%	0.91[0.46,1.77]
Horvaticek 2017	1/56	0/43	+		-	\rightarrow	0.67%	2.32[0.1,55.48]
Khalili 2016	0/75	1/75	+	+		_	1.78%	0.33[0.01,8.05]
Makrides 2010	3/1197	12/1202	+				14.23%	0.25[0.07,0.89]
Olsen 2000	19/1126	23/1126					27.33%	0.83[0.45,1.51]
Onwude 1995	1/113	2/119	+	+ +			2.31%	0.53[0.05,5.73]
		Favours omega-3	0.1	0.2 0.5 1	2 5	10	Favours no omega-3	





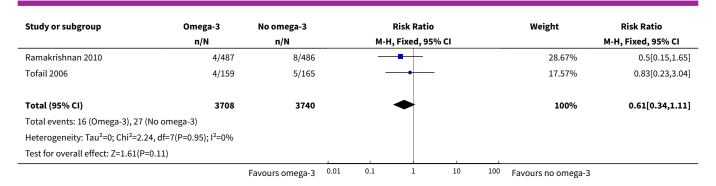
Analysis 1.33. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 33 Stillbirth.



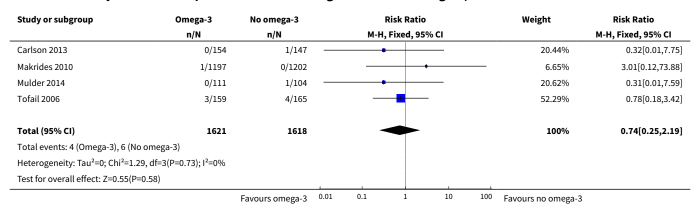
Analysis 1.34. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 34 Neonatal death.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI	
Bisgaard 2016	0/365	0/371			Not estimable	
Bulstra-Ramakers 1994	1/32	3/31		10.91%	0.32[0.04,2.94]	
Carlson 2013	1/154	1/147		3.66%	0.95[0.06,15.12]	
Khalili 2016	0/75	1/75		5.37%	0.33[0.01,8.05]	
Makrides 2010	2/1197	5/1202		17.86%	0.4[0.08,2.07]	
Olsen 2000	3/1126	4/1144		14.21%	0.76[0.17,3.4]	
Onwude 1995	1/113	0/119	· · · · · · · · · · · · · · · · · · ·	1.74%	3.16[0.13,76.73]	
		Favours omega-3	0.01 0.1 1 10 100	Favours no omega-3		





Analysis 1.35. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 35 Infant death.



Analysis 1.36. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 36 Large-for-gestational age.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
Dilli 2018	7/52	18/68		7.25%	0.51[0.23,1.13]
Harper 2010	21/427	15/410		7.11%	1.34[0.7,2.57]
Hauner 2012	9/96	7/92		3.32%	1.23[0.48,3.17]
Makrides 2010	204/1197	173/1202		80.25%	1.18[0.98,1.43]
Min 2014	1/60	0/59	+	0.23%	2.95[0.12,71.01]
Taghizadeh 2016	4/29	4/30		1.83%	1.03[0.29,3.75]
Total (95% CI)	1861	1861	•	100%	1.15[0.97,1.36]
Total events: 246 (Omega-3), 2	17 (No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =4.	75, df=5(P=0.45); I ² =0%				
Test for overall effect: Z=1.61(P	P=0.11)				
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	



Analysis 1.37. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 37 Macrosomia.

Study or subgroup	Omega-3	No omega-3		Risk Ratio	Weight	Risk Ratio	
	n/N	n/N		M-H, Fixed, 95% CI		M-H, Fixed, 95% CI	
Horvaticek 2017	8/47	7/43			20.69%	1.05[0.41,2.64]	
Jamilian 2016	0/27	3/27	•		9.9%	0.14[0.01,2.64]	
Min 2016	3/58	3/56		+	8.64%	0.97[0.2,4.58]	
Olsen 2000	9/792	10/809			28%	0.92[0.38,2.25]	
Razavi 2017	5/60	5/30			18.86%	0.5[0.16,1.59]	
Taghizadeh 2016	1/29	5/30	-		13.91%	0.21[0.03,1.67]	
Total (95% CI)	1013	995		•	100%	0.69[0.43,1.13]	
Total events: 26 (Omega-3), 33	(No omega-3)						
Heterogeneity: Tau ² =0; Chi ² =4.	.03, df=5(P=0.54); I ² =0%						
Test for overall effect: Z=1.46(F	P=0.14)						
		Favours omega-3	0.2	0.5 1 2	5 Favours no omega-3		

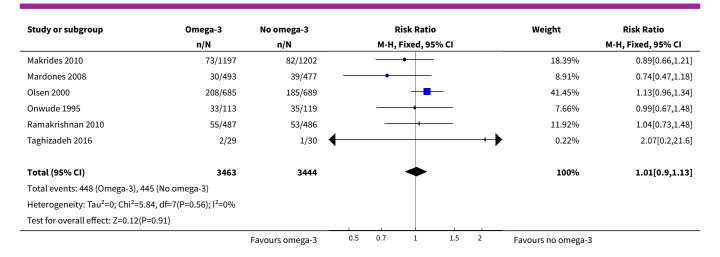
Analysis 1.38. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 38 Low birthweight (< 2500 g).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
Bulstra-Ramakers 1994	11/32	9/31		- 1.39%	1.18[0.57,2.46]
Carlson 2013	6/154	13/147	4	2.03%	0.44[0.17,1.13]
D'Almedia 1992	2/50	5/50	+	0.76%	0.4[0.08,1.97]
Harper 2010	94/427	112/410	+-	17.42%	0.81[0.63,1.02]
Khalili 2016	0/75	5/75	—	0.84%	0.09[0.01,1.62]
Makrides 2010	41/1197	63/1202		9.58%	0.65[0.44,0.96]
Mardones 2008	27/493	37/477		5.73%	0.71[0.44,1.14]
Min 2014	8/60	8/57	+	1.25%	0.95[0.38,2.36]
Min 2016	8/58	4/56		0.62%	1.93[0.62,6.05]
Olsen 2000	283/799	287/817	-	43.26%	1.01[0.88,1.15]
Onwude 1995	33/113	35/119		5.2%	0.99[0.67,1.48]
Ramakrishnan 2010	27/487	27/486		4.12%	1[0.59,1.68]
Smuts 2003a	13/142	16/149		2.38%	0.85[0.43,1.71]
Smuts 2003b	0/18	5/19	—	0.82%	0.1[0.01,1.62]
Tofail 2006	36/125	30/124		4.59%	1.19[0.79,1.8]
Total (95% CI)	4230	4219	•	100%	0.9[0.82,0.99]
Total events: 589 (Omega-3), 656	6 (No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =19.	75, df=14(P=0.14); l ² =29.1	11%			
Test for overall effect: Z=2.12(P=	0.03)				
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	

Analysis 1.39. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 39 Small-for-gestational age/IUGR.

Study or subgroup	Omega-3	No omega-3	Risk Ratio					Weight	Risk Ratio
	n/N	n/N		M-H,	Fixed,	95% CI			M-H, Fixed, 95% CI
Bulstra-Ramakers 1994	12/32	9/31						2.05%	1.29[0.64,2.63]
Harper 2010	35/427	41/410						9.4%	0.82[0.53,1.26]
		Favours omega-3	0.5	0.7	1	1.5	2	Favours no omega-3	

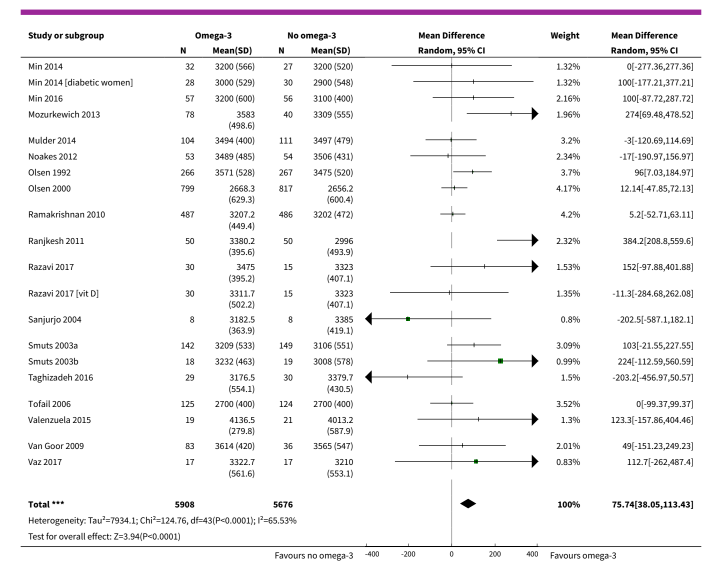




Analysis 1.40. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 40 Birthweight (g).

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Ali 2017	34	2324 (110.8)	32	2022 (141.4)		4.15%	302[240.46,363.54]
Carlson 2013	154	3359 (524)	147	3187 (602)		3.04%	172[44.25,299.75]
de Groot 2004	29	3662.8 (568)	29	3298 (456.4)		1.41%	364.8[99.6,630]
Dilli 2018	52	3288 (641)	68	3538 (671)		1.65%	-250[-486.2,-13.8]
Dunstan 2008	40	3503.4 (337.7)	43	3430.1 (371.8)		2.64%	73.3[-79.36,225.96]
England 1989	17	2200 (700)	18	2000 (500)	-	0.73%	200[-205.07,605.07]
Furuhjelm 2009	52	3500 (500)	65	3600 (600)		2.02%	-100[-299.36,99.36]
Gustafson 2013	22	3416.8 (552.9)	24	3435.5 (404.8)		1.29%	-18.7[-300.85,263.45]
Haghiac 2015	25	3278 (448)	24	2935 (356)		1.74%	343[116.89,569.11]
Harper 2010	427	2990 (1005.8)	410	2923 (1252.8)		2.62%	67[-87.29,221.29]
Harris 2015	224	3215.5 (506.2)	121	3165 (494.6)	+-	3.33%	50.5[-59.77,160.77]
Hauner 2012	96	3534 (465)	92	3357 (557)		2.73%	177[30.01,323.99]
Helland 2001	175	3609 (493)	166	3618 (527)		3.36%	-9[-117.45,99.45]
Horvaticek 2017	47	3580.9 (568)	43	3456.9 (575.8)		1.64%	124[-112.62,360.62]
Hurtado 2015	44	3300 (500)	46	3200 (500)		1.94%	100[-106.65,306.65]
Jamilian 2016	26	3418.8 (344.7)	27	3405.2 (465.1)		1.8%	13.6[-206.23,233.43]
Judge 2007	27	3394 (430)	21	3224.6 (431.2)	-	1.56%	169.38[-76.22,414.98]
Keenan 2014	36	3074.4 (582.3)	17	2919.6 (537.7)		1.08%	154.83[-163.81,473.47]
Khalili 2016	75	3260 (360)	75	3230 (430)		3.05%	30[-96.92,156.92]
Krauss-Etschmann 2007	96	3317.3 (574.3)	99	3298.4 (502.4)		2.66%	18.9[-132.73,170.53]
Krummel 2016	34	3502 (433)	29	3484 (411)		1.92%	18[-190.71,226.71]
Makrides 2010	1197	3475 (564)	1202	3407 (576)		4.36%	68[22.38,113.62]
Malcolm 2003	31	3507.7 (500.8)	29	3645.1 — (495)		1.51%	-137.4[-389.46,114.66]
Mardones 2008	493	3265.9 (452.1)	477	3200.5 (506)		4.16%	65.4[4.95,125.85]



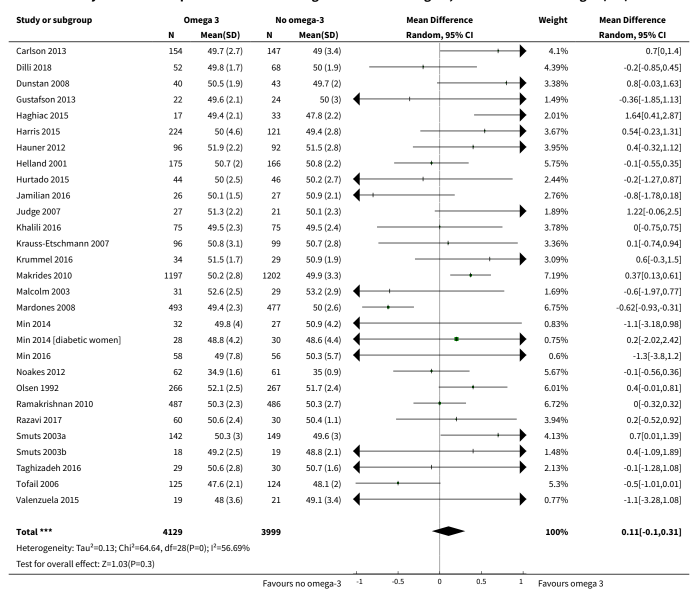


Analysis 1.41. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 41 Birthweight Z score.

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
Bergmann 2007	41	1 (0.1)	74	1 (0.9)		12.02%	0[-0.21,0.21]
Krummel 2016	34	0.8 (1)	29	0.7 (0.8)		2.75%	0.19[-0.25,0.63]
Makrides 2010	1197	0.3 (1.1)	1202	0.2 (1)		76.19%	0.06[-0.02,0.14]
Mulder 2014	104	0.4 (0.8)	111	0.4 (1)	•	9.03%	0.06[-0.18,0.3]
Total ***	1376		1416		•	100%	0.06[-0.02,0.13]
Heterogeneity: Tau ² =0; Chi ² =	=0.61, df=3(P=0.9); I ² =0%					
Test for overall effect: Z=1.53	B(P=0.13)						
			Lower	with omega-3	-0.2 -0.1 0 0.1 0.2	Higher with	omega-3



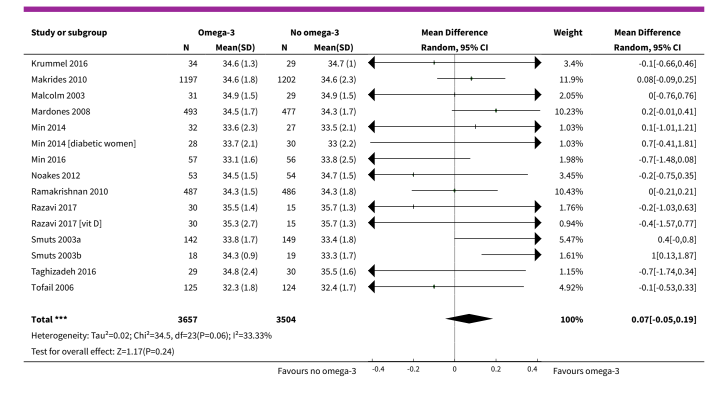
Analysis 1.42. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 42 Birth length (cm).



Analysis 1.43. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 43 Head circumference at birth (cm).

Study or subgroup	0	Omega-3		omega-3		Mean Difference			Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rar	ndom, 95% CI			Random, 95% CI
Carlson 2013	154	34.2 (1.7)	147	33.7 (2)				——	5.16%	0.5[0.08,0.92]
Dilli 2018	52	35.1 (1.2)	68	35.5 (1.4)	\leftarrow				4.47%	-0.4[-0.87,0.07]
Harris 2015	224	34.5 (3.1)	121	33.9 (2.2)					3.38%	0.6[0.04,1.16]
Hauner 2012	96	35.1 (1.4)	92	34.8 (1.7)					4.75%	0.3[-0.15,0.75]
Helland 2001	175	35.3 (1.5)	166	35.2 (1.6)			+		6.95%	0.1[-0.23,0.43]
Hurtado 2015	44	34 (1.5)	46	34.1 (1.4)	\leftarrow				3.04%	-0.1[-0.7,0.5]
Jamilian 2016	26	34.9 (1)	27	35.3 (1)	\leftarrow				3.45%	-0.4[-0.95,0.15]
Judge 2007	25	34.4 (1)	20	34.3 (1.1)	\leftarrow			·	2.96%	0.18[-0.43,0.79]
Khalili 2016	75	34.7 (1.4)	75	34.7 (1.5)	•				4.49%	0[-0.46,0.46]
			Favour	s no omega-3	-0.4	-0.2	0	0.2 0.4	Favours omega	-3





Analysis 1.44. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 44 Head circumference at birth Z score.

Study or subgroup	0	mega-3	No	omega-3		Mea	an Difference		Weight	Mean Difference	
	N	Mean(SD)	N	Mean(SD)		Fi	xed, 95% CI			Fixed, 95% CI	
Krummel 2016	34	0.7 (1.1)	29	0.6 (0.9)			+		4.4%	0.08[-0.41,0.57]	
Makrides 2010	1197	0 (1.1)	1202	0.1 (1.5)					95.6%	-0.04[-0.15,0.07]	
Total ***	1231		1231				•		100%	-0.03[-0.14,0.07]	
Heterogeneity: Tau ² =0; Chi ² =	=0.22, df=1(P=0.6	4); I ² =0%									
Test for overall effect: Z=0.66	S(P=0.51)										
			Fav	ours omega-3	-1	-0.5	0 0.5	1	Favours no	omega-3	

Analysis 1.45. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 45 Length at birth Z score.

Study or subgroup	0	mega-3	No	omega-3		Mea	n Difference	Weight		Mean Difference
	N	Mean(SD)	N	Mean(SD)		Ran	dom, 95% CI			Random, 95% CI
Krummel 2016	34	1.5 (1.1)	29	1 (0.9)			-		30.24%	0.46[-0.04,0.96]
Makrides 2010	1197	-0.1 (0.9)	1202	-0.2 (1)			+		69.76%	0.06[-0.02,0.14]
Total ***	1231		1231				•		100%	0.18[-0.18,0.54]
Heterogeneity: Tau ² =0.05; Ch	ni ² =2.41, df=1(P=	0.12); I ² =58.55%								
Test for overall effect: Z=0.98	s(P=0.32)									
			Lower	with omega-3	-2	-1	0 1	2	Lower with	no omega-3



Analysis 1.46. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 46 Baby admitted to neonatal care.

Study or subgroup	Omega-3	No omega-3		R	isk Ratio		Weight	Risk Ratio
	n/N	n/N		M-H,	Fixed, 95% CI			M-H, Fixed, 95% CI
Ali 2017	10/32	15/31	$\overline{}$	-			2.93%	0.65[0.34,1.21]
Bisgaard 2016	40/365	41/371					7.82%	0.99[0.66,1.5]
Carlson 2013	13/154	13/147			-		2.56%	0.95[0.46,1.99]
Harper 2010	110/427	99/410					19.43%	1.07[0.84,1.35]
Makrides 2010	21/1197	37/1202	\leftarrow	+			7.1%	0.57[0.34,0.97]
Mozurkewich 2013	8/78	4/40	\leftarrow				1.02%	1.03[0.33,3.2]
Olsen 2000	258/1062	283/1076		_	-		54.08%	0.92[0.8,1.07]
Smuts 2003a	21/142	21/149			+		3.94%	1.05[0.6,1.84]
Smuts 2003b	2/18	6/19	•				1.12%	0.35[0.08,1.52]
Total (95% CI)	3475	3445			•		100%	0.92[0.83,1.03]
Total events: 483 (Omega-3), 519 (No	omega-3)							
Heterogeneity: Tau ² =0; Chi ² =7.9, df=8	3(P=0.44); I ² =0%							
Test for overall effect: Z=1.41(P=0.16)								
		Favours omega-3	0.	5 0.7	1 1.	5 2	Favours no omega-3	

Analysis 1.47. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 47 Infant length of stay in hospital (days).

Study or subgroup	0	mega-3	No omega-3			Mean Difference			Weight		Mean Difference	
	N	Mean(SD)	N	Mean(SD)		F	ixed, 95% CI				Fixed, 95% CI	
Olsen 2000	1017	7.7 (16.3)	1024	7.6 (18.5)						100%	0.11[-1.4,1.62]	
Total ***	1017		1024				•			100%	0.11[-1.4,1.62]	
Heterogeneity: Not applicable												
Test for overall effect: Z=0.14(P=0.89	9)											
			Fave	nurs amaga_3	-10	-5	0	5	10	Favours no	nmeαa-3	

Analysis 1.48. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 48 Congenital anomalies.

Study or subgroup	Omega-3	No omega-3			Ri	sk Rat	io			Weight	Risk Ratio
	n/N	n/N			M-H, F	ixed, 9	95% CI				M-H, Fixed, 95% CI
Carlson 2013	5/154	2/147					+		→	9.28%	2.39[0.47,12.11]
Olsen 1992	3/266	5/267			-					22.63%	0.6[0.15,2.49]
Ramakrishnan 2010	16/487	15/486			-	-				68.09%	1.06[0.53,2.13]
Total (95% CI)	907	900			4		-			100%	1.08[0.61,1.92]
Total events: 24 (Omega-3), 22	(No omega-3)										
Heterogeneity: Tau ² =0; Chi ² =1.	57, df=2(P=0.46); I ² =0%										
Test for overall effect: Z=0.27(P	=0.79)										
		Favours omega-3	0.1	0.2	0.5	1	2	5	10	Favours no omega-3	



Analysis 1.49. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 49 Retinopathy of prematurity.

Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		M-H	H, Fixed, 95%	6 CI			M-H, Fixed, 95% CI
Harper 2010	5/427	4/410			-	-		100%	1.2[0.32,4.44]
Total (95% CI)	427	410				-		100%	1.2[0.32,4.44]
Total events: 5 (Omega-3), 4 (No omeg	a-3)								
Heterogeneity: Not applicable									
Test for overall effect: Z=0.27(P=0.78)									
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3	

Analysis 1.50. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 50 Bronchopulmonary dysplasia.

Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		M-	H, Fixed, 95%	CI			M-H, Fixed, 95% CI
Harper 2010	9/425	6/403			-			60.58%	1.42[0.51,3.96]
Makrides 2010	2/1184	4/1179			-			39.42%	0.5[0.09,2.71]
Total (95% CI)	1609	1582			•			100%	1.06[0.45,2.48]
Total events: 11 (Omega-3), 10	(No omega-3)								
Heterogeneity: Tau ² =0; Chi ² =1.	.08, df=1(P=0.3); I ² =7.42%								
Test for overall effect: Z=0.13(F	P=0.9)								
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3	

Analysis 1.51. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 51 Respiratory distress syndrome.

Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		М-Н,	Random, 95%	6 CI			M-H, Random, 95% CI
Carlson 2013	9/154	12/147						39.12%	0.72[0.31,1.65]
Harper 2010	59/425	35/403			-			60.88%	1.6[1.08,2.37]
Total (95% CI)	579	550			•			100%	1.17[0.54,2.52]
Total events: 68 (Omega-3), 47	(No omega-3)								
Heterogeneity: Tau ² =0.21; Chi ²	!=2.91, df=1(P=0.09); I ² =65.6	62%							
Test for overall effect: Z=0.39(P	P=0.69)					1			
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3	

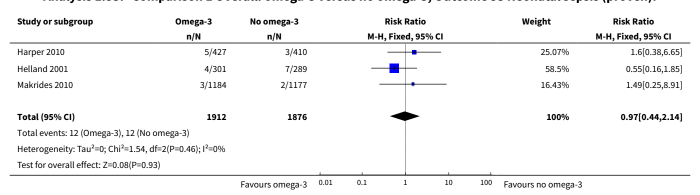
Analysis 1.52. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 52 Necrotising enterocolitis (NEC).

Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		М-Н	Fixed, 95	% CI			M-H, Fixed, 95% CI
Harper 2010	3/427	4/410				-		89.06%	0.72[0.16,3.2]
Makrides 2010	1/1184	0/1177			 '	-		10.94%	2.98[0.12,73.13]
Total (95% CI)	1611	1587		-		-		100%	0.97[0.26,3.55]
Total events: 4 (Omega-3), 4 (No	o omega-3)								
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3	

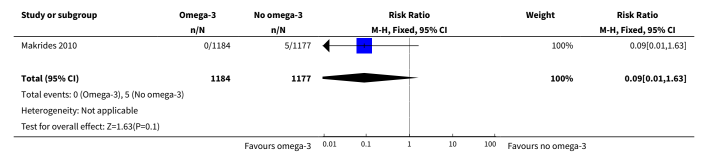


Study or subgroup	Omega-3	No omega-3		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		М-Н	, Fixed, 95%	CI			M-H, Fixed, 95% CI
Heterogeneity: Tau ² =0; Chi ² =0.63, df	=1(P=0.43); I ² =0%								
Test for overall effect: Z=0.05(P=0.96))								
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3	

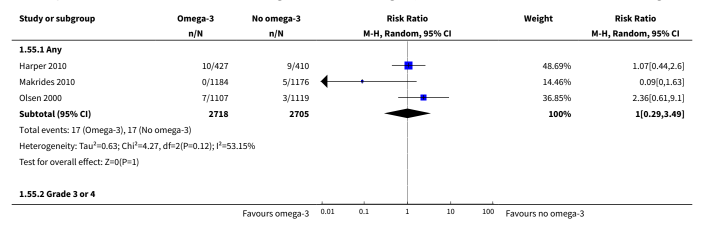
Analysis 1.53. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 53 Neonatal sepsis (proven).



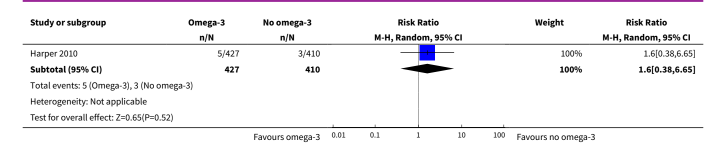
Analysis 1.54. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 54 Convulsion.



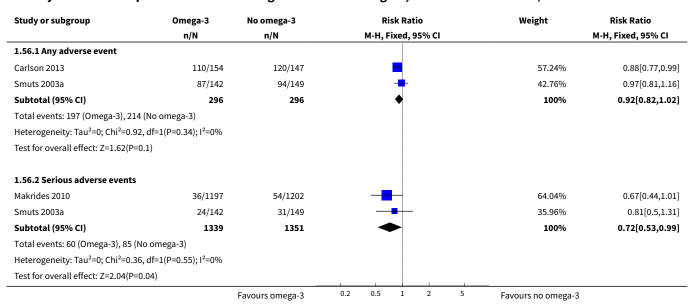
Analysis 1.55. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 55 Intraventricular haemorrhage.







Analysis 1.56. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 56 Neonatal/infant adverse events.



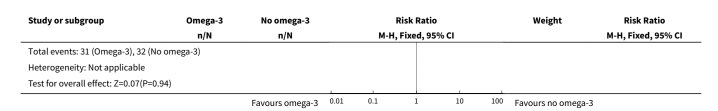
Analysis 1.57. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 57 Neonatal/infant morbidity: cardiovascular.

Study or subgroup	Omega-3 No omega-3		Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
Smuts 2003a	48/142	42/149	++-	0%	1.2[0.85,1.69]
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	

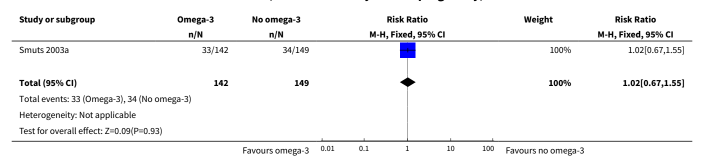
Analysis 1.58. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 58 Neonatal/infant morbidity: respiratory.

Study or subgroup	Omega-3	No omega-3	No omega-3					Weight	Risk Ratio	
	n/N	n/N		М-Н	l, Fixed, 95	% CI			M-H, Fixed, 95% CI	
Smuts 2003a	31/142	32/149			-			100%	1.02[0.66,1.57]	
Total (95% CI)	142	149			•	,		100%	1.02[0.66,1.57]	
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3		





Analysis 1.59. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 59 Neonatal/infant morbidity: due to pregnancy/birth events.



Analysis 1.60. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 60 Neonatal/infant morbidity: other.

Study or subgroup	Omega-3	No omega-3	ı	Risk Ratio	V	Veight	Risk Ratio
	n/N	n/N	м-н,	Fixed, 95% CI			M-H, Fixed, 95% CI
1.60.1 Colds in past 15 days: at 1 mor	nth of age						
Ramakrishnan 2010	159/422	190/427		+		100%	0.85[0.72,1]
Subtotal (95% CI)	422	427		•		100%	0.85[0.72,1]
Total events: 159 (Omega-3), 190 (No o	mega-3)						
Heterogeneity: Not applicable							
Test for overall effect: Z=2.01(P=0.04)							
1.60.2 Colds in past 15 days: at 3 mor	nths of age						
Ramakrishnan 2010	157/415	185/419		+		100%	0.86[0.73,1.01]
Subtotal (95% CI)	415	419		•		100%	0.86[0.73,1.01]
Total events: 157 (Omega-3), 185 (No o	mega-3)						
Heterogeneity: Tau ² =0; Chi ² =0, df=0(P<	(0.0001); I ² =100%						
Test for overall effect: Z=1.85(P=0.06)							
1.60.3 Colds in past 15 days: at 6 mor	nths of age						
Ramakrishnan 2010	194/420	193/414		+		100%	0.99[0.86,1.15]
Subtotal (95% CI)	420	414		 		100%	0.99[0.86,1.15]
Total events: 194 (Omega-3), 193 (No o	mega-3)						
Heterogeneity: Not applicable							
Test for overall effect: Z=0.12(P=0.9)							
1.60.4 Fever in past 15 days: at 1 moi	nth of age						
Ramakrishnan 2010	15/422	14/427				100%	1.08[0.53,2.22]
Subtotal (95% CI)	422	427		•		100%	1.08[0.53,2.22]
Total events: 15 (Omega-3), 14 (No ome	ega-3)						
		Favours omega-3	0.01 0.1	1 10	¹⁰⁰ Favou	rs no omega-3	



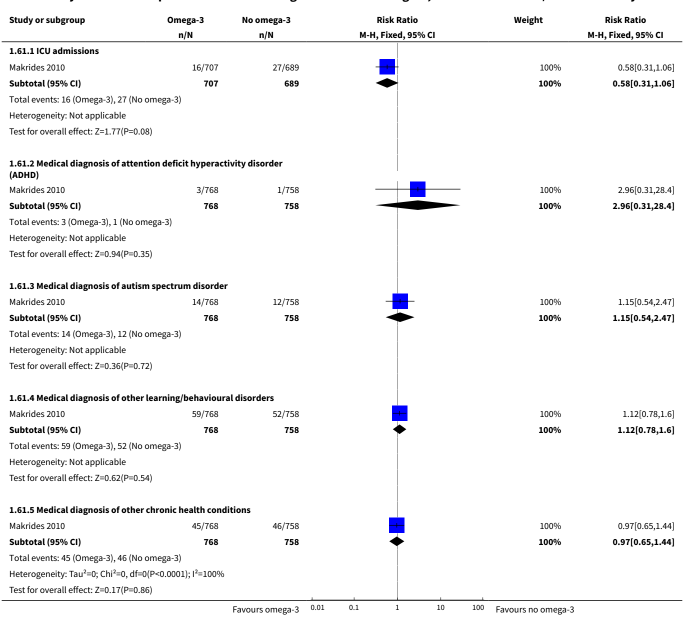
Study or subgroup	Omega-3 n/N	No omega-3 n/N	Risk Ratio M-H, Fixed, 95% CI	Weight	Risk Ratio M-H, Fixed, 95% CI
Heterogeneity: Not applicable					
Test for overall effect: Z=0.22(P=0.82)					
1.60.5 Fever in past 15 days: at 3 mor	nths of age				
Ramakrishnan 2010	35/415	44/419		100%	0.8[0.53,1.23]
Subtotal (95% CI)	415	419	•	100%	0.8[0.53,1.23]
Total events: 35 (Omega-3), 44 (No ome	ega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=1.02(P=0.31)					
1.60.6 Fever in past 15 days: at 6 mor	nths of age				
Ramakrishnan 2010	77/420	77/414	-	100%	0.99[0.74,1.31]
Subtotal (95% CI)	420	414	+	100%	0.99[0.74,1.31]
Total events: 77 (Omega-3), 77 (No ome	ega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.1(P=0.92)					
1.60.7 Rash in past 15 days: at 1 mon	th of age				
Ramakrishnan 2010	122/422	111/427	-	100%	1.11[0.89,1.38]
Subtotal (95% CI)	422	427	*	100%	1.11[0.89,1.38]
Total events: 122 (Omega-3), 111 (No o	mega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.95(P=0.34)					
1.60.8 Rash in past 15 days: at 3 mon	ths of age				
Ramakrishnan 2010	35/415	43/419		100%	0.82[0.54,1.26]
Subtotal (95% CI)	415	419	•	100%	0.82[0.54,1.26]
Total events: 35 (Omega-3), 43 (No ome	ega-3)				
Heterogeneity: Tau ² =0; Chi ² =0, df=0(P<	(0.0001); I ² =100%				
Test for overall effect: Z=0.9(P=0.37)					
1.60.9 Rash in past 15 days: at 6 mon	ths of age				
Ramakrishnan 2010	45/420	39/414		100%	1.14[0.76,1.71]
Subtotal (95% CI)	420	414	*	100%	1.14[0.76,1.71]
Total events: 45 (Omega-3), 39 (No ome	ega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.62(P=0.54)					
1.60.10 Vomiting in past 15 days: at 1	L month of age				
Ramakrishnan 2010	23/422	15/427		100%	1.55[0.82,2.93]
Subtotal (95% CI)	422	427	•	100%	1.55[0.82,2.93]
Total events: 23 (Omega-3), 15 (No ome	ega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=1.35(P=0.18)					
1.60.11 Vomiting in past 15 days: at 3	3 months of age				
Ramakrishnan 2010	17/415	12/419	-	100%	1.43[0.69,2.96]
Subtotal (95% CI)	415	419	*	100%	1.43[0.69,2.96]
Total events: 17 (Omega-3), 12 (No ome	ega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.97(P=0.33)					
		Favours omega-3	0.01 0.1 1 10	100 Favours no omega-3	



Study or subgroup	Omega-3 n/N	No omega-3 n/N	Risk Ratio M-H, Fixed, 95% CI	Weight	Risk Ratio M-H, Fixed, 95% CI
1.60.12 Vomiting in past 15 days: at					, ,
Ramakrishnan 2010	23/420	17/414	-	100%	1.33[0.72,2.46]
Subtotal (95% CI)	420	414	•	100%	1.33[0.72,2.46]
Total events: 23 (Omega-3), 17 (No om	iega-3)				
Heterogeneity: Not applicable	,				
Test for overall effect: Z=0.92(P=0.36)					
1.60.13 Diarrhoea in past 15 days: at	t 1 month of age				
Ramakrishnan 2010	14/422	17/427	-	100%	0.83[0.42,1.67]
Subtotal (95% CI)	422	427	•	100%	0.83[0.42,1.67]
Total events: 14 (Omega-3), 17 (No om	iega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.51(P=0.61)					
1.60.14 Diarrhoea in past 15 days: at	t 3 months of age				
Ramakrishnan 2010	19/415	23/419	<u> </u>	100%	0.83[0.46,1.51]
Subtotal (95% CI)	415	419	•	100%	0.83[0.46,1.51]
Total events: 19 (Omega-3), 23 (No om	iega-3)				- , -
Heterogeneity: Not applicable	,				
Test for overall effect: Z=0.6(P=0.55)					
1.60.15 Diarrhoea in past 15 days: at	t 6 months of age				
Ramakrishnan 2010	32/420	31/414	-	100%	1.02[0.63,1.64]
Subtotal (95% CI)	420	414	<u> </u>	100%	1.02[0.63,1.64]
Total events: 32 (Omega-3), 31 (No om			Ţ		
Heterogeneity: Not applicable	67				
Test for overall effect: Z=0.07(P=0.94)					
1.60.16 Other illness in the past 15 d	lavs: at 1 month				
Ramakrishnan 2010	29/422	21/427		100%	1.4[0.81,2.41]
Subtotal (95% CI)	422	427	_	100%	1.4[0.81,2.41]
Total events: 29 (Omega-3), 21 (No om				20070	
Heterogeneity: Not applicable	6.2				
Test for overall effect: Z=1.2(P=0.23)					
1.60.17 Other illness in the past 15 d	lavs: at 3 months				
Ramakrishnan 2010	21/415	22/419		100%	0.96[0.54,1.73]
Subtotal (95% CI)	415	419	<u> </u>	100%	0.96[0.54,1.73]
Total events: 21 (Omega-3), 22 (No om			Ī		,,
Heterogeneity: Not applicable	67				
Test for overall effect: Z=0.12(P=0.9)					
1.60.18 Other illness in the past 15 d	lays: at 6 months				
Ramakrishnan 2010	28/420	24/414	<u> </u>	100%	1.15[0.68,1.95]
Subtotal (95% CI)	420	414	<u></u>	100%	1.15[0.68,1.95]
Total events: 28 (Omega-3), 24 (No om				200,0	
Heterogeneity: Not applicable	00/				
obeniety. Hot applicable					



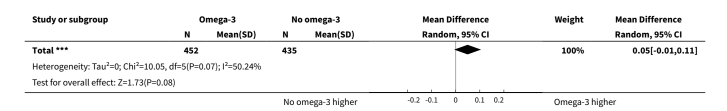
Analysis 1.61. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 61 Infant/child morbidity.



Analysis 1.62. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 62 Ponderal index.

Study or subgroup	Oi	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Carlson 2013	154	2.7 (0.3)	147	2.7 (0.4)	-	21.53%	0[-0.08,0.08]
Haghiac 2015	17	2.7 (0.3)	16	2.6 (0.2)	+	7.73%	0.12[-0.06,0.3]
Hauner 2012	96	2.5 (0.2)	92	2.4 (0.2)	_ 	24.79%	0.08[0.01,0.15]
Jamilian 2016	26	1.7 (1)	27	1.7 (1.4)	•	0.77%	-0.02[-0.65,0.61]
Krummel 2016	34	2.6 (0.2)	29	2.6 (0.2)		15.56%	-0.06[-0.17,0.05]
Tofail 2006	125	2.5 (0.2)	124	2.4 (0.2)		29.62%	0.1[0.05,0.15]
			No or	nega-3 higher	-0.2 -0.1 0 0.1 0.2	Omega-3 hi	gher

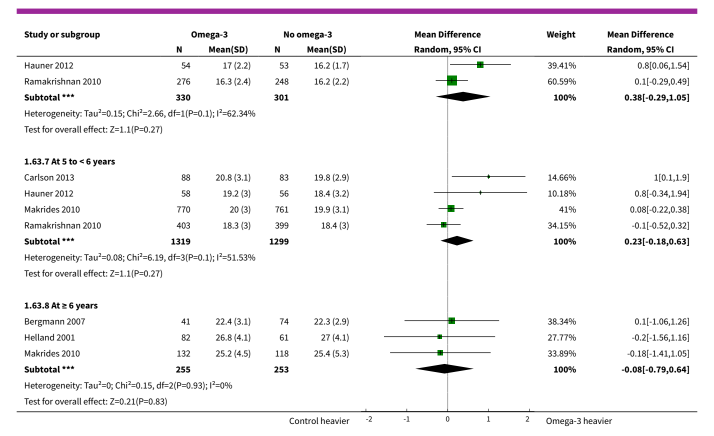




Analysis 1.63. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 63 Infant/child weight (kg).

Study or subgroup	On	nega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
1.63.1 At < 3 months							
Hauner 2012	89	4.8 (0.6)	91	4.7 (0.6)		20.03%	0.06[-0.12,0.24]
Ramakrishnan 2010	343	4.3 (0.6)	340	4.3 (0.6)	+	79.97%	0[-0.09,0.09]
Subtotal ***	432		431		\rightarrow	100%	0.01[-0.07,0.09]
Heterogeneity: Tau²=0; Chi²=0.3	31, df=1(P=0.58	s); I ² =0%					
Test for overall effect: Z=0.28(P=	=0.78)						
1.63.2 At 3 to < 12 months							
Hauner 2012	87	6.5 (0.7)	87	6.3 (0.7)	-	29.96%	0.17[-0.04,0.38]
Khalili 2016	75	8 (0.8)	71	7.8 (0.9)	-	23.88%	0.14[-0.13,0.41]
Malcolm 2003	28	8.3 (1)	27	8.6 (1.1)		9.52%	-0.36[-0.91,0.19]
Ramakrishnan 2010	336	8.3 (0.9)	317	8.4 (1)	-	36.64%	-0.1[-0.25,0.05]
Subtotal ***	526		502		*	100%	0.01[-0.18,0.2]
Heterogeneity: Tau²=0.02; Chi²=	7.12, df=3(P=0	.07); I ² =57.85%					
Test for overall effect: Z=0.14(P=	=0.89)						
1.63.3 At 1 to < 2 years							
Hauner 2012	87	9.7 (1)	83	9.4 (1)		25.45%	0.27[-0.04,0.58]
Hurtado 2015	32	6.3 (0.9)	29	6.6 (0.9)	-+-	14.94%	-0.3[-0.75,0.15]
Ramakrishnan 2010	369	10.4 (1.1)	370	10.4 (1.2)	-	45.7%	0[-0.17,0.17]
Van Goor 2009	80	11.4 (1.4)	34	11.5 (1.1)		13.9%	-0.1[-0.57,0.37]
Subtotal ***	568		516		*	100%	0.01[-0.19,0.21]
Heterogeneity: Tau²=0.02; Chi²=	-4.74, df=3(P=0	.19); I ² =36.77%					
Test for overall effect: Z=0.1(P=0	0.92)						
1.63.4 At 2 to < 3 years							
Dunstan 2008	28	14.5 (2)	36	14.1 (2)		19.57%	0.4[-0.59,1.39]
Hauner 2012	61	12.5 (1.4)	57	12.3 (1.3)		80.43%	0.2[-0.29,0.69]
Subtotal ***	89		93			100%	0.24[-0.2,0.68]
Heterogeneity: Tau ² =0; Chi ² =0.1 Test for overall effect: Z=1.07(P=); I ² =0%					
1.63.5 At 3 to < 4 years							
Hauner 2012	61	14.8 (1.9)	59	14.3 (1.5)	-	27.61%	0.5[-0.11,1.11]
Makrides 2010	770	15.4 (2)	761	15.3 (2)	-	72.39%	0.06[-0.14,0.26]
Subtotal ***	831	. ,	820			100%	0.18[-0.2,0.57]
Heterogeneity: Tau²=0.04; Chi²=		.18); I ² =44.25%					- · ·
Test for overall effect: Z=0.92(P=		••					
1.63.6 At 4 to < 5 years							

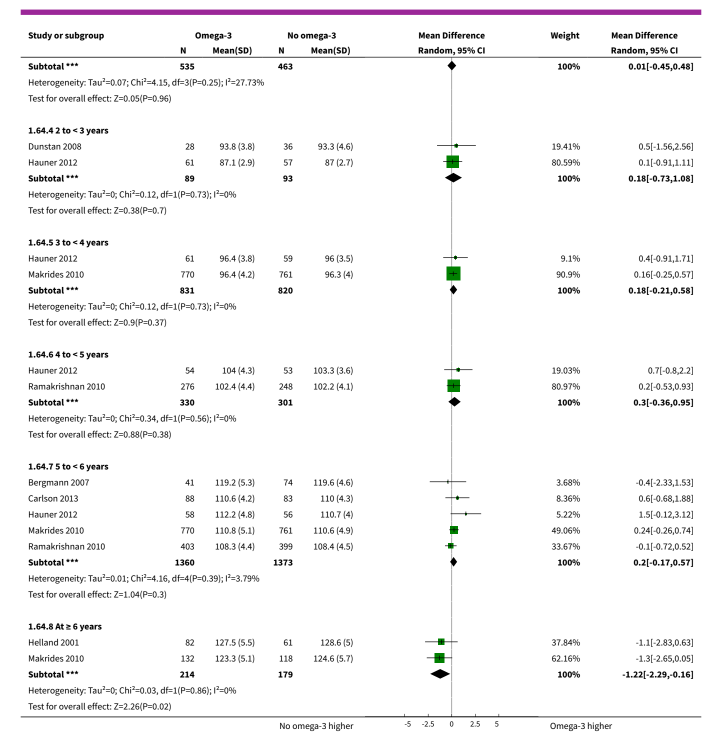




Analysis 1.64. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 64 Infant/child length/height (cm).

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
1.64.1 < 3 months							
Hauner 2012	89	56 (2)	91	55.6 (2.6)	-	40.87%	0.4[-0.28,1.08]
Ramakrishnan 2010	343	53.5 (2.3)	338	53.8 (2.2)	=	59.13%	-0.3[-0.64,0.04]
Subtotal ***	432		429		*	100%	-0.01[-0.69,0.66]
Heterogeneity: Tau ² =0.17; Chi ²	=3.29, df=1(P=	0.07); I ² =69.6%					
Test for overall effect: Z=0.04(P	P=0.97)						
1.64.2 3 to < 12 months							
Hauner 2012	88	62.6 (2)	87	62.4 (2.2)	-	24.46%	0.2[-0.42,0.82]
Khalili 2016	75	67.1 (2.9)	71	66.9 (2.4)	+	12.8%	0.2[-0.66,1.06]
Malcolm 2003	28	60 (2.6)	27	60.5 (2.9)	-+-	4.47%	-0.5[-1.96,0.96]
Ramakrishnan 2010	369	79.6 (2.8)	370	79.5 (2.8)	•	58.27%	0.1[-0.3,0.5]
Subtotal ***	560		555		•	100%	0.11[-0.2,0.42]
Heterogeneity: Tau ² =0; Chi ² =0.	.8, df=3(P=0.85); I ² =0%					
Test for overall effect: Z=0.7(P=	=0.48)						
1.64.3 1 to < 2 years							
Hauner 2012	87	75.5 (2.4)	83	74.9 (2.8)	-	24.51%	0.6[-0.19,1.39]
Hurtado 2015	32	24.7 (2.4)	29	25.3 (2.6)	+	11.66%	-0.6[-1.86,0.66]
Ramakrishnan 2010	336	69.6 (2.3)	317	69.6 (2.3)	•	55.65%	0[-0.35,0.35]
Van Goor 2009	80	83.2 (3.9)	34	84 (3.8)		8.18%	-0.81[-2.35,0.73]
			No or	nega-3 higher	-5 -2.5 0 2.5 5	Omega-3 hi	gher





Analysis 1.65. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 65 Infant/child head circumference (cm).

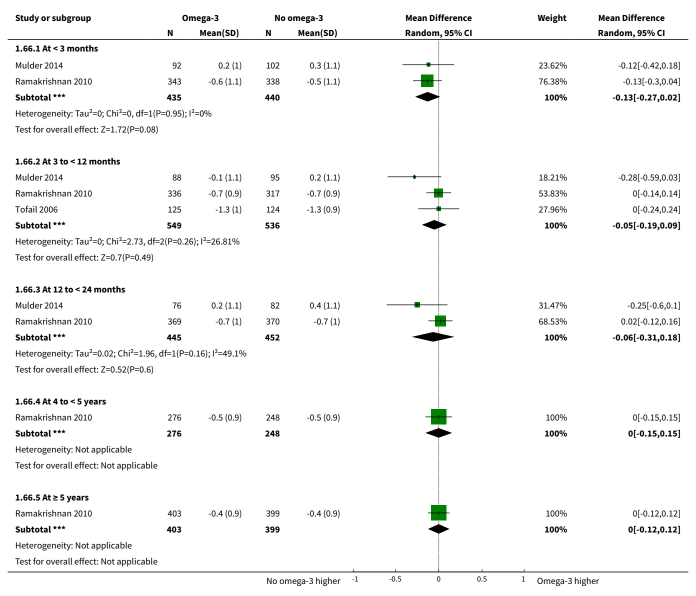
Study or subgroup	c	Omega-3 No omega-3		Mean Difference					Weight	Mean Difference	
	N	Mean(SD)	N	Mean(SD)		Ran	dom, 95	% CI			Random, 95% CI
1.65.1 At < 3 months											
			No omega-3 greater		-2	-1	0	1	2	Omega-3 gre	ater



Study or subgroup		mega-3		omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Hauner 2012	89	38.4 (1.1)	90	38.3 (1.2)	-	29.56%	0.1[-0.24,0.4
Ramakrishnan 2010	343	37.2 (1.6)	341	37.3 (1.3)		70.44%	-0.1[-0.32,0.1
Subtotal ***	432		431		*	100%	-0.04[-0.22,0.1
Heterogeneity: Tau²=0; Chi²=0.95,	df=1(P=0.3	3); I ² =0%					
Test for overall effect: Z=0.44(P=0.	66)						
1.65.2 At 3 to < 12 months							
Hauner 2012	87	41.2 (1.3)	87	41 (1.3)	+-	15.31%	0.2[-0.19,0.5
Khalili 2016	75	43 (1.2)	71	43.2 (1.5)	-+-	11.69%	-0.2[-0.64,0.2
Malcolm 2003	28	39.9 (1.5)	27	40.1 (2.3)		2.15%	-0.2[-1.23,0.8
Ramakrishnan 2010	343	42.9 (1.4)	342	42.9 (1.4)	+	51.96%	0[-0.21,0.2
Tofail 2006	125	43 (1.4)	124	43.2 (1.4)		18.89%	-0.2[-0.55,0.1
Subtotal ***	658		651		•	100%	-0.03[-0.19,0.1
Heterogeneity: Tau²=0; Chi²=3.03,	df=4(P=0.5	5); I ² =0%					
Test for overall effect: Z=0.45(P=0.	•						
1.65.3 At 1 to < 2 years							
Hauner 2012	87	46.5 (1.6)	83	46.1 (1.5)		21.79%	0.4[-0.07,0.8
Hurtado 2015	32	12.6 (6.3)	29	11.9 (2)		1.09%	0.68[-1.62,2.9
Ramakrishnan 2010	369	47 (1.5)	370	47 (1.4)	<u> </u>	61.8%	0[-0.21,0.2
Van Goor 2009	80	47.6 (1.3)	34	47.8 (1.5)	<u>-</u>	15.32%	-0.24[-0.81,0.3
Subtotal ***	568	(=12)	516	(=12)	•	100%	0.06[-0.18,0
Heterogeneity: Tau²=0.01; Chi²=3.		0 3)· l ² =17 61%				20070	0.00[0.20,0
Test for overall effect: Z=0.47(P=0.		0.5/,1 17.01/0					
1.65.4 At 2 to < 3 years							
Dunstan 2008	28	49.4 (1.6)	36	49.8 (1.7)		27.13%	-0.4[-1.21,0.4
Hauner 2012	61	48.7 (1.3)	57	48.6 (1.3)	_	72.87%	0.1[-0.37,0.5
Subtotal ***	89	10.1 (1.5)	93	10.0 (1.3)		100%	-0.04[-0.47,0
Heterogeneity: Tau ² =0.01; Chi ² =1.		0 3)·12=8 38%	55		T	200 /0	0.04[0.11,0
Test for overall effect: Z=0.16(P=0.		0.57,1 0.5070					
1.65.5 At 3 to < 4 years							
Hauner 2012	61	49.9 (1.4)	59	49.8 (1.3)		9.47%	0.1[-0.38,0.5
Makrides 2010	770	50 (1.6)	761	50.1 (1.6)		90.53%	-0.02[-0.18,0.1
Subtotal ***	831	30 (1.0)	820	30.1 (1.0)		100%	-0.01[-0.16,0.1
Heterogeneity: Tau ² =0; Chi ² =0.21,		4)· 1²-0%	020		Y	100 /0	-0.01[-0.10,0.1
Test for overall effect: Z=0.11(P=0.	•	4),1 -0 /0					
1.65.6 At 4 to < 5 years							
Hauner 2012	54	50.6 (1.3)	53	50.6 (1.2)	_	100%	0[-0.47,0.4
Subtotal ***	54	55.5 (1.5)	53	55.5 (I.Z)		100%	0[-0.47,0.4
Heterogeneity: Not applicable	34		33			250 70	٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠٠
Test for overall effect: Not applica	ble						
1.65.7 At ≥ 5 years							
Bergmann 2007	41	52.5 (16)	74	52.7 (1.3)		0.09%	-0.2[-5.11,4.
Hauner 2012	58	51.2 (1.3)	56	51.2 (1.3)		9.51%	0[-0.48,0.4
Makrides 2010	770	51.2 (1.5)	761	51.3 (1.6)		90.4%	0.02[-0.13,0.
Subtotal ***		J1.+ (1.J)		31.3 (1.0)			
Subtotal """ Heterogeneity: Tau²=0; Chi²=0.01,	869	0). 12-00/	891		Y	100%	0.02[-0.13,0.
neterogeneity, rau -0; CNI-=0.01,	u1-2(r=0.9	J), I -U70					



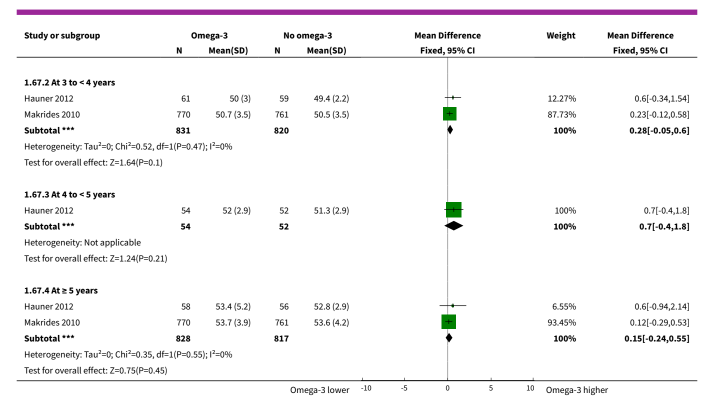
Analysis 1.66. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 66 Infant/child length/height for age Z score (LAZ/HAZ).



Analysis 1.67. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 67 Infant/child waist circumference (cm).

Study or subgroup	Omega-3		No omega-3			Mean Difference			Weight	Mean Difference	
	N	Mean(SD)	N	Mean(SD)		F	Fixed, 95% CI				Fixed, 95% CI
1.67.1 At 2 to < 3 years											
Hauner 2012	50	47.8 (2.9)	51	48 (2.7)						100%	-0.2[-1.29,0.89]
Subtotal ***	50		51				•			100%	-0.2[-1.29,0.89]
Heterogeneity: Not applicable											
Test for overall effect: Z=0.36(P=0.72)										
			O	mega-3 lower	-10	-5	0	5	10	Omega-3 highe	r

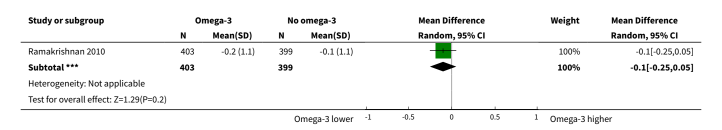




Analysis 1.68. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 68 Infant/child weight-for-age Z score (WAZ).

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
1.68.1 At < 3 months							
Mulder 2014	90	-0.2 (1.1)	101	0.1 (1.1)		31.83%	-0.25[-0.56,0.06]
Ramakrishnan 2010	343	-0.3 (0.9)	340	-0.3 (1)	- 	68.17%	-0.02[-0.16,0.12]
Subtotal ***	433		441		-	100%	-0.09[-0.3,0.12]
Heterogeneity: Tau ² =0.01; Chi ² =1	.78, df=1(P=	0.18); I ² =43.68%					
Test for overall effect: Z=0.87(P=0).38)						
1.68.2 At 3 to < 12 months							
Mulder 2014	87	0 (1.1)	94	0 (1)		17.47%	0.01[-0.3,0.32]
Ramakrishnan 2010	336	-0.3 (0.9)	317	-0.2 (1)	- 	82.53%	-0.06[-0.2,0.08]
Subtotal ***	423		411		•	100%	-0.05[-0.18,0.08]
Heterogeneity: Tau ² =0; Chi ² =0.16	, df=1(P=0.6	9); I ² =0%					
Test for overall effect: Z=0.73(P=0).47)						
1.68.3 At 12 to < 24 months							
Mulder 2014	74	0.2 (1)	70	0.3 (1)		13.93%	-0.06[-0.39,0.27]
Ramakrishnan 2010	369	-0.2 (0.9)	370	-0.2 (0.9)	-	86.07%	0[-0.13,0.13]
Subtotal ***	443		440		*	100%	-0.01[-0.13,0.12]
Heterogeneity: Tau ² =0; Chi ² =0.11	, df=1(P=0.7	4); I ² =0%					
Test for overall effect: Z=0.13(P=0).89)						
1.68.4 At ≥ 60 months							
			0	mega-3 lower ⁻¹	-0.5 0 0.5	¹ Omega-3 hi	igher

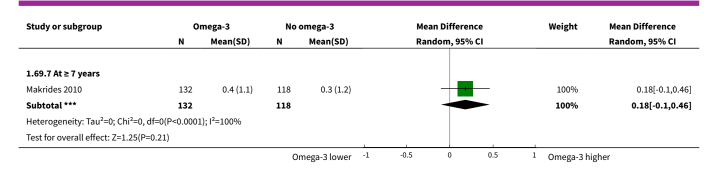




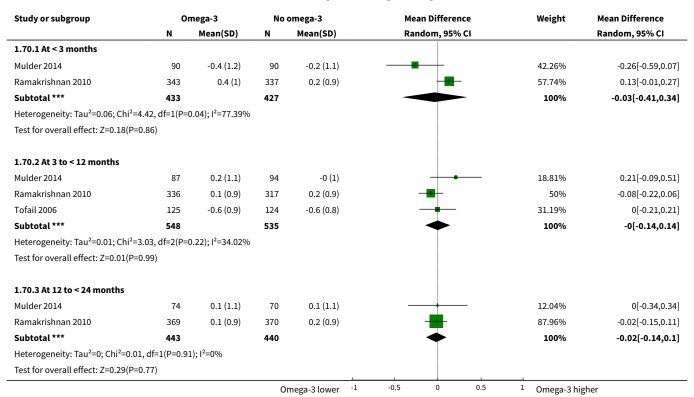
Analysis 1.69. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 69 Infant/child BMI Z score.

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
1.69.1 At 1 to < 2 years							
Bergmann 2007	20	1 (0.1)	42	1 (0.1)	_	84.14%	-0.05[-0.11,0]
Ramakrishnan 2010	369	0.3 (0.8)	370	0.3 (0.9)	-	15.86%	-0.01[-0.13,0.11]
Subtotal ***	389		412		•	100%	-0.05[-0.09,0]
Heterogeneity: Tau ² =0; Chi ² =0.38,	df=1(P=0.5	4); I ² =0%					
Test for overall effect: Z=1.82(P=0.	07)						
1.69.2 At 2 to < 3 years							
Krummel 2016	34	0.5 (0.4)	29	0.6 (0.3)	_	100%	-0.07[-0.25,0.11]
Subtotal ***	34		29			100%	-0.07[-0.25,0.11]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.78(P=0.	43)						
1.69.3 At 3 to < 4 years							
Makrides 2010	770	0.7 (1)	761	0.7 (1.1)		100%	0.02[-0.08,0.12]
Subtotal ***	770		761		•	100%	0.02[-0.08,0.12]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.39(P=0.	7)						
1.69.4 At 4 to < 5 years							
Krummel 2016	34	1.2 (0.4)	29	0.9 (0.5)	_ 	47.49%	0.32[0.1,0.54]
Ramakrishnan 2010	276	0.1 (1)	248	0.1 (1)	_ 	52.51%	0[-0.17,0.17]
Subtotal ***	310		277			100%	0.15[-0.16,0.47]
Heterogeneity: Tau ² =0.04; Chi ² =5,	df=1(P=0.0	3); I ² =80.02%					
Test for overall effect: Z=0.95(P=0.	34)						
1.69.5 At 5 to < 6 years							
Carlson 2013	88	0.7 (0.9)	83	0.5 (1)	-	7.91%	0.19[-0.1,0.48]
Makrides 2010	770	0.6 (1)	761	0.5 (1)	#	64.24%	0.02[-0.08,0.12]
Ramakrishnan 2010	403	0.1 (1.1)	399	0.1 (1.1)	-	27.85%	0[-0.15,0.15]
Subtotal ***	1261		1243		*	100%	0.03[-0.05,0.11]
Heterogeneity: Tau ² =0; Chi ² =1.39,	df=2(P=0.5); I ² =0%					
Test for overall effect: Z=0.68(P=0.	5)						
1.69.6 At 6 to < 7 years							
Bergmann 2007	41	1 (0.1)	74	1 (0.1)	<u> </u>	100%	0.01[-0.02,0.05]
Subtotal ***	41		74			100%	0.01[-0.02,0.05]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.61(P=0.	54)						





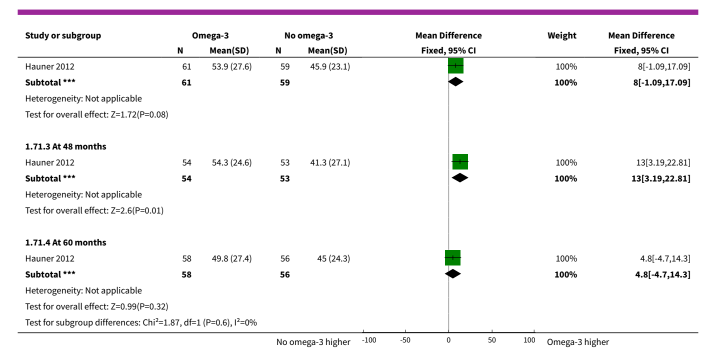
Analysis 1.70. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 70 Infant/child weight for length/height Z score (WHZ).



Analysis 1.71. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 71 Infant/child BMI percentile.

Omega-3		No omega-3			Mean Difference			Weight	Mean Difference	
N	Mean(SD)	N	Mean(SD)		Fi	ixed, 95% CI				Fixed, 95% CI
61	56.7 (27.5)	57	52.2 (27.9)			-			100%	4.5[-5.5,14.5]
61		57				◆			100%	4.5[-5.5,14.5]
8)										
		No or	nega-3 higher	-100	-50	0	50	100	Omega-3 highe	r
	N 61	N Mean(SD) 61 56.7 (27.5) 61	N Mean(SD) N 61 56.7 (27.5) 57 61 57	N Mean(SD) N Mean(SD) 61 56.7 (27.5) 57 52.2 (27.9) 61 57 57 57	N Mean(SD) N Mean(SD) 61 56.7 (27.5) 57 52.2 (27.9) 61 57	N Mean(SD) N Mean(SD) Fi 61 56.7 (27.5) 57 52.2 (27.9) 61 57	N Mean(SD) N Mean(SD) Fixed, 95% CI 61 56.7 (27.5) 57 52.2 (27.9) 61 57	N Mean(SD) N Mean(SD) Fixed, 95% CI 61 56.7 (27.5) 57 52.2 (27.9) 61 57	N Mean(SD) N Mean(SD) Fixed, 95% CI 61 56.7 (27.5) 57 52.2 (27.9) 61 57	N Mean(SD) N Mean(SD) Fixed, 95% CI 61 56.7 (27.5) 57 52.2 (27.9) 100% 61 57 100%





Analysis 1.72. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 72 Child/adult BMI.

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
1.72.1 At 3 to 4 years							
Makrides 2010	770	16.5 (1.4)	761	16.5 (1.5)	-	100%	0.01[-0.14,0.16]
Subtotal ***	770		761		→	100%	0.01[-0.14,0.16]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.13(P=0.8	39)						
1.72.2 At 5 to 6 years							
Makrides 2010	770	16.2 (1.6)	761	16.2 (1.7)	+	100%	-0.01[-0.18,0.16]
Subtotal ***	770		761		*	100%	-0.01[-0.18,0.16]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.12(P=0.9	91)						
1.72.3 At 7 to 9 years							
Helland 2001	82	16.4 (1.7)	61	16.3 (1.7)	- 	53.65%	0.1[-0.46,0.66]
Makrides 2010	132	16.5 (2.3)	118	16.3 (2.6)		46.35%	0.23[-0.38,0.84]
Subtotal ***	214		179			100%	0.16[-0.25,0.57]
Heterogeneity: Tau ² =0; Chi ² =0.09, o	df=1(P=0.7	6); I ² =0%					
Test for overall effect: Z=0.76(P=0.4	15)						
1.72.4 At 19 years							
Olsen 1992	108	22.5 (3.5)	135	22.5 (3)		100%	0[-0.83,0.83]
Subtotal ***	108		135			100%	0[-0.83,0.83]
Heterogeneity: Not applicable							
Test for overall effect: Not applicab	le						



Analysis 1.73. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 73 Infant/child body fat (%).

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
1.73.1 At 1 year							
Hauner 2012	85	19.7 (3)	80	19.7 (2.8)		100%	0[-0.88,0.88]
Subtotal ***	85		80			100%	0[-0.88,0.88]
Heterogeneity: Not applicable							
Test for overall effect: Not applicable	е						
1.73.2 At 2 to < 3 years							
Hauner 2012	57	19.2 (2.3)	53	19 (2.4)		100%	0.2[-0.68,1.08]
Subtotal ***	57		53			100%	0.2[-0.68,1.08]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.45(P=0.66	5)						
1.73.3 At 3 to < 4 years							
Hauner 2012	58	18.4 (2.6)	55	18.3 (2.6)		34.07%	0.1[-0.86,1.06]
Makrides 2010	770	24.5 (7.1)	761	24.9 (6.7)		65.93%	-0.33[-1.02,0.36]
Subtotal ***	828		816			100%	-0.18[-0.74,0.38]
Heterogeneity: Tau ² =0; Chi ² =0.51, df	f=1(P=0.4	8); I ² =0%					
Test for overall effect: Z=0.64(P=0.52	2)						
1.73.4 At 4 to < 5 years							
Hauner 2012	50	18.2 (2.6)	52	17.9 (3)		100%	0.3[-0.79,1.39]
Subtotal ***	50		52			100%	0.3[-0.79,1.39]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.54(P=0.59))						
1.73.5 At 5 to < 6 years							
Carlson 2013	78	25.1 (4.9)	76	24.9 (6.5)	•	9.69%	0.2[-1.62,2.02]
Hauner 2012	57	17.9 (3.4)	55	18.1 (3.6)		19.08%	-0.2[-1.5,1.1]
Makrides 2010	770	23.5 (6.8)	761	23.4 (6.6)		71.23%	0.04[-0.63,0.71]
Subtotal ***	905		892			100%	0.01[-0.56,0.58]
Heterogeneity: Tau²=0; Chi²=0.15, di	f=2(P=0.9	3); I ² =0%					
Test for overall effect: Z=0.03(P=0.97	")						
1.73.6 At ≥ 7 years: BIS							
Makrides 2010	132	24.2 (7.4)	118	22.8 (6.7)	-	100%	1.44[-0.31,3.19]
Subtotal ***	132		118			100%	1.44[-0.31,3.19]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.61(P=0.11	.)						
1.73.7 At ≥ 7 years: BOD POD					_		
Makrides 2010	132	19.2 (6.9)	118	19.6 (7.6)	1	100%	-0.42[-2.23,1.39]
Subtotal ***	132		118			100%	-0.42[-2.23,1.39]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.45(P=0.65	:)						

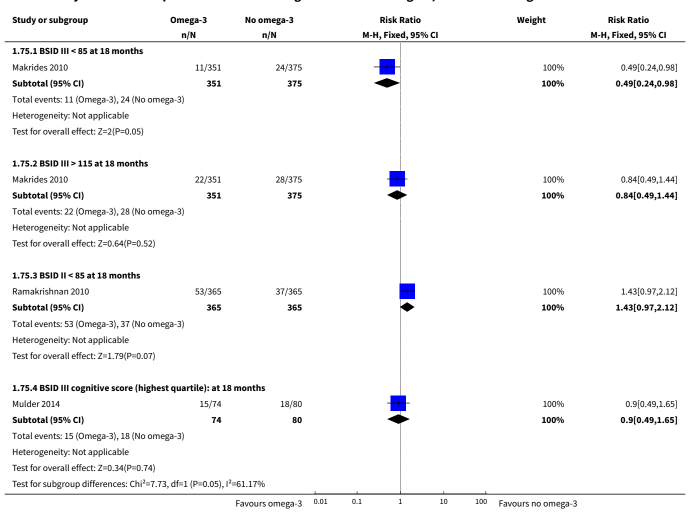


Analysis 1.74. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 74 Infant/child total fat mass (kg).

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
1.74.1 At 1 year							
Hauner 2012	84	2.1 (0.2)	80	2.1 (0.2)	+	100%	0[-0.07,0.07
Subtotal ***	84		80		→	100%	0[-0.07,0.07
Heterogeneity: Not applicable							
Test for overall effect: Not applical	ble						
1.74.2 At 2 to < 3 years							
Hauner 2012	57	2.4 (0.5)	53	2.3 (0.5)	-	100%	0.1[-0.09,0.29
Subtotal ***	57		53		*	100%	0.1[-0.09,0.29
Heterogeneity: Not applicable							
Test for overall effect: Z=1.05(P=0.	29)						
1.74.3 At 3 to < 4 years							
Hauner 2012	58	2.7 (0.7)	55	2.6 (0.5)		24.05%	0.1[-0.12,0.32
Makrides 2010	770	3.8 (1.3)	761	3.8 (1.3)	#	75.95%	-0.05[-0.18,0.08
Subtotal ***	828		816		♦	100%	-0.01[-0.12,0.1
Heterogeneity: Tau ² =0; Chi ² =1.32,	df=1(P=0.2	5); I ² =23.96%					
Test for overall effect: Z=0.25(P=0.	8)						
1.74.4 At 4 to < 5 years							
Hauner 2012	50	3.1 (0.7)	52	2.9 (0.6)	-	100%	0.2[-0.05,0.45
Subtotal ***	50		52		•	100%	0.2[-0.05,0.45
Heterogeneity: Not applicable							
Test for overall effect: Z=1.55(P=0.	12)						
1.74.5 At 5 to < 6 years							
Carlson 2013	78	5.2 (1.7)	76	4.9 (1.6)	+	8.8%	0.3[-0.22,0.82
Hauner 2012	57	3.5 (1.1)	55	3.4 (0.8)	-	18.94%	0.1[-0.26,0.46
Makrides 2010	770	4.8 (1.8)	761	4.7 (1.9)	#	72.26%	0.01[-0.17,0.19
Subtotal ***	905		892		*	100%	0.05[-0.1,0.21
Heterogeneity: Tau²=0; Chi²=1.14,	df=2(P=0.5	6); I ² =0%					
Test for overall effect: Z=0.67(P=0.	51)						
1.74.6 Up to 8 years: BOD POD					<u>L</u>		
Makrides 2010	132	5.2 (3)	118	5.1 (3.3)		100%	0.08[-0.71,0.87
Subtotal ***	132		118			100%	0.08[-0.71,0.87
Heterogeneity: Not applicable							
Test for overall effect: Z=0.2(P=0.8	4)						
1.74.7 Up to 8 years: BIS							
Makrides 2010	132	6.3 (2.9)	118	6 (3.2)		100%	0.29[-0.47,1.05
Subtotal ***	132		118			100%	0.29[-0.47,1.0
Heterogeneity: Not applicable							
Test for overall effect: Z=0.75(P=0.	45)						



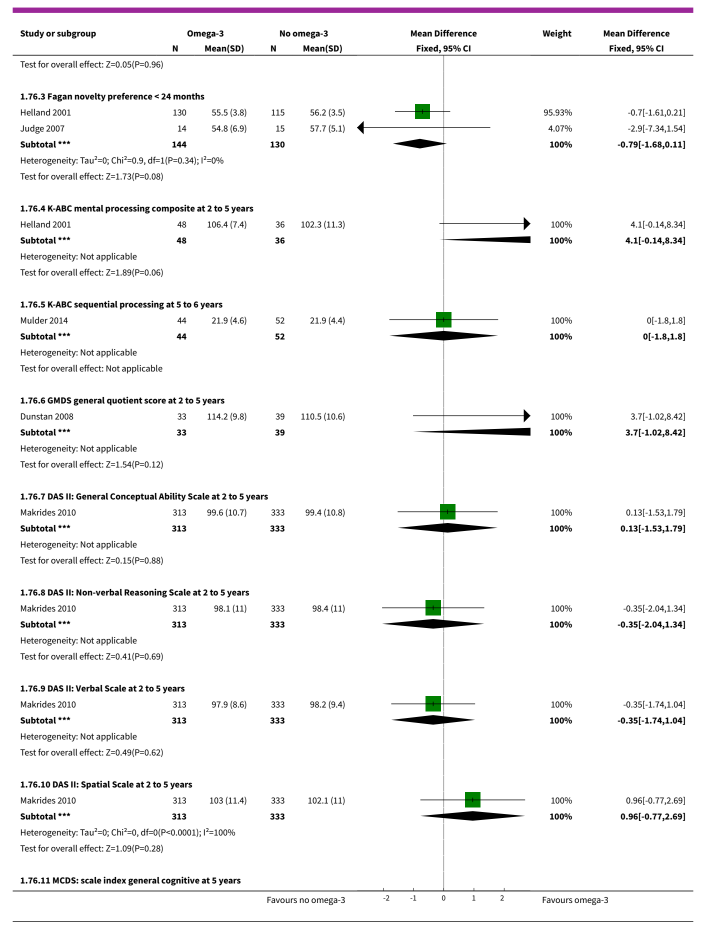
Analysis 1.75. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 75 Cognition: thresholds.



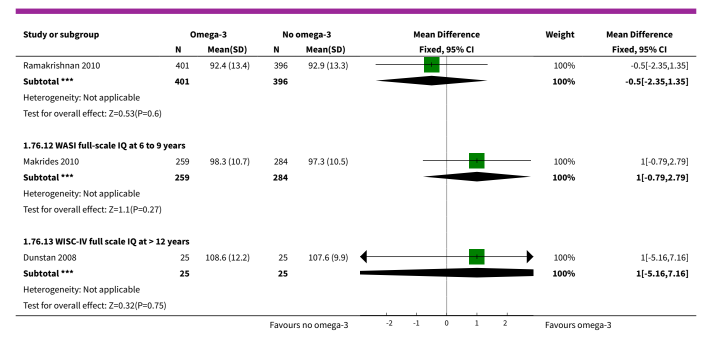
Analysis 1.76. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 76 Cognition: scores.

Study or subgroup	Omega-3		No	omega-3		Mean I	Difference	Weight		Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fixed	, 95% CI			Fixed, 95% CI
1.76.1 BSID II score < 24 months										
Hurtado 2015	32	99.9 (17.5)	29	102.1 (12.2)	+ +				2.23%	-2.2[-9.72,5.32]
Ramakrishnan 2010	365	94.3 (10.7)	365	95.2 (9.3)	_	-	 		59.59%	-0.9[-2.35,0.55]
Tofail 2006	125	102.5 (8)	124	101.5 (7.8)			-	\rightarrow	32.72%	1[-0.96,2.96]
Van Goor 2009	80	113.2 (12.8)	34	115.2 (11.6)	\leftarrow				5.46%	-2[-6.8,2.8]
Subtotal ***	602		552						100%	-0.37[-1.49,0.76]
Heterogeneity: Tau ² =0; Chi ² =3.05,	df=3(P=0.3	8); I ² =1.7%								
Test for overall effect: Z=0.64(P=0.	52)									
1.76.2 BSID III score < 24 months										
Makrides 2010	351	101.8 (11.1)	375	101.8 (12.6)			-		90.35%	0.06[-1.66,1.78]
Miller 2016	48	109.7 (11.3)	35	109.8 (12.6)	←		+	\rightarrow	9.65%	-0.1[-5.36,5.16]
Subtotal ***	399		410						100%	0.04[-1.59,1.68]
Heterogeneity: Tau ² =0; Chi ² =0, df=	1(P=0.95);	I ² =0%								
			Favou	rs no omega-3	-2	2 -1	0 1 2		Favours om	ega-3





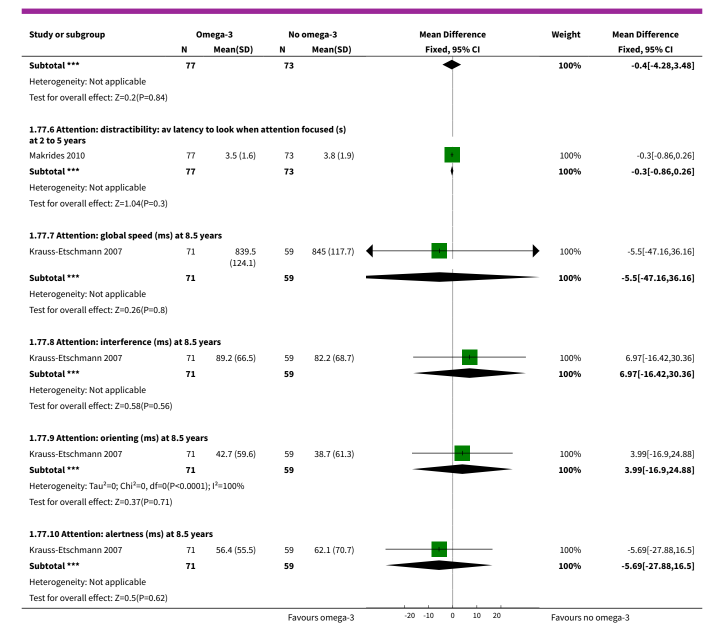




Analysis 1.77. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 77 Attention: scores.

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
1.77.1 K-CPT omissions at 5 years							
Ramakrishnan 2010	401	47.6 (10.3)	396	49.5 (11.2)	+	100%	-1.9[-3.39,-0.41]
Subtotal ***	401		396		◆	100%	-1.9[-3.39,-0.41]
Heterogeneity: Tau ² =0; Chi ² =0, df=0	(P<0.000	1); I ² =100%					
Test for overall effect: Z=2.49(P=0.0	1)						
1.77.2 K-CPT commissions at 5 years	ars						
Ramakrishnan 2010	401	51.2 (11)	396	51.1 (10.2)	+	100%	0.1[-1.37,1.57]
Subtotal ***	401		396		•	100%	0.1[-1.37,1.57]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.13(P=0.8	9)						
1.77.3 K-CPT hit response time at	5 years						
Ramakrishnan 2010	401	55.5 (10.5)	396	56.1 (10.6)	+	100%	-0.6[-2.06,0.86]
Subtotal ***	401		396		♦	100%	-0.6[-2.06,0.86]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.8(P=0.42))						
1.77.4 Attention: single-object tas	sk: total t	ime looking at t	toy(s) at	2 to 5 years			
Makrides 2010	77	238.8 (51.1)	73	246.6 (41)	- 	100%	-7.8[-22.59,6.99]
Subtotal ***	77		73			100%	-7.8[-22.59,6.99]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.03(P=0.3))						
1.77.5 Attention: multiple-object to 5 years	task; # ti	mes shifted lool	ks betwe	en toys at 2			
Makrides 2010	77	41.6 (11.8)	73	42 (12.4)	.	100%	-0.4[-4.28,3.48]
			Fav	ours omega-3	-20 -10 0 10 20	Favours no	omega-3

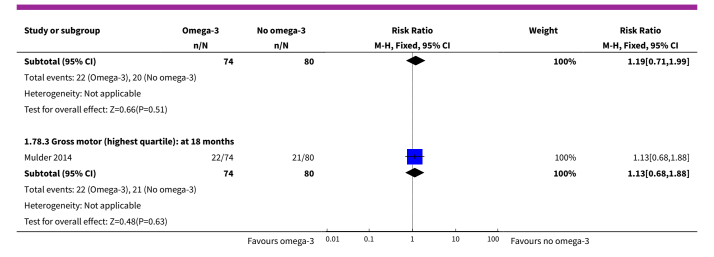




Analysis 1.78. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 78 Motor: thresholds.

Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio	
n/N		n/N	M-H, Fixed, 95% CI						M-H, Fixed, 95% CI	
1.78.1 BSID II score < 85 at 18 mon	iths									
Ramakrishnan 2010	65/365	74/365			-			100%	0.88[0.65,1.19]	
Subtotal (95% CI)	365	365			*			100%	0.88[0.65,1.19]	
Total events: 65 (Omega-3), 74 (No o	omega-3)									
Heterogeneity: Not applicable										
Test for overall effect: Z=0.85(P=0.4)										
1.78.2 Fine motor (highest quartil	e): at 18 months									
Mulder 2014	22/74	20/80						100%	1.19[0.71,1.99]	
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3		





Analysis 1.79. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 79 Motor: scores.

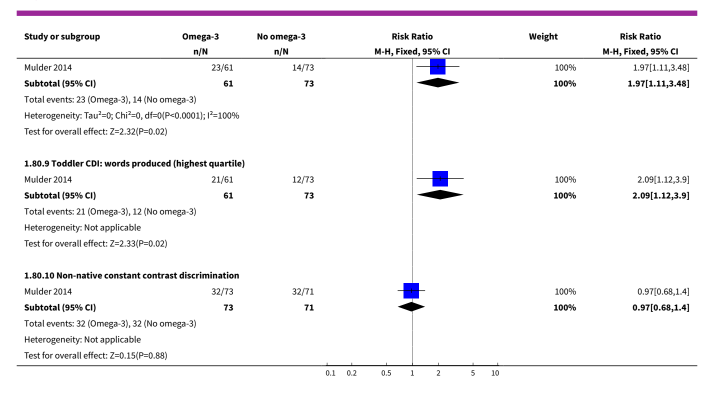
94.6 (17.5) 93 (8.9) 101.7 (10.9) 94.2 (10.2)	N 29 365 124	96.2 (15.3) 93.3 (9.8)	Fixed, 95% CI	1.89%	Fixed, 95% CI -1.6[-9.83,6.63]
93 (8.9) 101.7 (10.9)	365				-1.6[-9.83,6.63]
93 (8.9) 101.7 (10.9)	365				-1.6[-9.83,6.63]
101.7 (10.9)		93.3 (9.8)	—		
	124			69.38%	-0.3[-1.66,1.06]
94.2 (10.2)		100.5 (10.1)		18.79%	1.2[-1.41,3.81]
	34	91.7 (8.3)		9.94%	2.46[-1.13,6.05]
	552		*	100%	0.23[-0.9,1.36]
43); I ² =0%					
102.6 (10.2)	375	102.6 (11.5)	-	100%	0.06[-1.52,1.64]
	375		→	100%	0.06[-1.52,1.64]
onths					
2.4 (1.9)	15	2.3 (2.1)	-	100%	0.05[-1.2,1.3]
	15		→	100%	0.05[-1.2,1.3]
nonths					
1.7 (1.2)	15	1.6 (1.2)	-	100%	0.05[-0.68,0.78]
	15		→	100%	0.05[-0.68,0.78]
	2.4 (1.9) nonths 1.7 (1.2)	375 conths 2.4 (1.9) 15 15 15 1.7 (1.2) 15 15	375 conths 2.4 (1.9) 15 2.3 (2.1) 15 nonths 1.7 (1.2) 15 1.6 (1.2)	375 conths 2.4 (1.9) 15 2.3 (2.1) 15 conths 1.7 (1.2) 15 1.6 (1.2)	375 100% 100% 15 100% 100% 100% 100% 100% 100% 100%



Analysis 1.80. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 80 Language: thresholds.

Study or subgroup	Omega-3 n/N	No omega-3 n/N	Risk Ratio M-H, Fixed, 95% CI	Weight	Risk Ratio M-H, Fixed, 95% CI
1.80.1 BSID III < 85					
Makrides 2010	62/351	65/375	-	100%	1.02[0.74,1.4]
Subtotal (95% CI)	351	375	→	100%	1.02[0.74,1.4]
Total events: 62 (Omega-3), 65 (No on	nega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.12(P=0.91)					
1.80.2 BSID III > 115					
Makrides 2010	30/351	39/375		100%	0.82[0.52,1.29]
Subtotal (95% CI)	351	375	—	100%	0.82[0.52,1.29]
Total events: 30 (Omega-3), 39 (No on	nega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.85(P=0.4)					
1.80.3 Receptive language (highest	quartile)				
Mulder 2014	27/74	16/80		100%	1.82[1.07,3.1]
Subtotal (95% CI)	74	80		100%	1.82[1.07,3.1]
Total events: 27 (Omega-3), 16 (No or	nega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=2.22(P=0.03)					
1.80.4 Expressive language (highes	t quartile)		_		
Mulder 2014	29/74	19/80		100%	1.65[1.02,2.68]
Subtotal (95% CI)	74	80		100%	1.65[1.02,2.68]
Total events: 29 (Omega-3), 19 (No on	nega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=2.03(P=0.04)					
1.80.5 Infant CDI: words understoo	d (highest quartile)		_		
Mulder 2014	28/78	12/81	- - 	100%	2.42[1.33,4.42]
Subtotal (95% CI)	78	81		100%	2.42[1.33,4.42]
Total events: 28 (Omega-3), 12 (No or	nega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=2.89(P=0)					
1.80.6 Infant CDI: words produced (_		
Mulder 2014	26/78	13/81		100%	2.08[1.15,3.74]
Subtotal (95% CI)	78	81		100%	2.08[1.15,3.74]
Total events: 26 (Omega-3), 13 (No or	nega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=2.43(P=0.01)					
1.80.7 Infant CDI: words understoo			_		
Mulder 2014	23/61	14/73		100%	1.97[1.11,3.48]
Subtotal (95% CI)	61	73		100%	1.97[1.11,3.48]
Total events: 23 (Omega-3), 14 (No on					
Heterogeneity: Tau ² =0; Chi ² =0, df=0(F					
Test for overall effect: Z=2.32(P=0.02)					
1.80.8 Infant CDI: words produced ((highest quartile)			<u> </u>	
		0.1	0.2 0.5 1 2 5 1	.0	

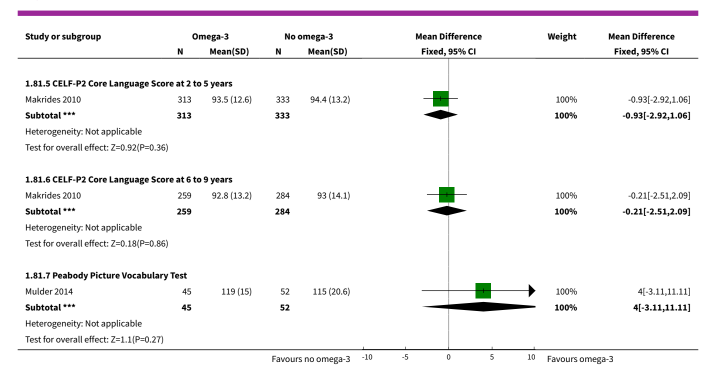




Analysis 1.81. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 81 Language: scores.

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
1.81.1 Receptive communication	at < 24 m	onths					
Keenan 2014	34	4.9 (2.7)	15	4.3 (1.9)	-	100%	0.55[-0.77,1.87]
Subtotal ***	34		15		•	100%	0.55[-0.77,1.87]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.82(P=0.4	1)						
1.81.2 Receptive language (Peabo	ody Pictui	re Vocabulary T	est IIIA) a	t 2 to 5 years			
Dunstan 2008	31	101.3 (9.9)	39	97.4 (9.7)	-	100%	3.9[-0.73,8.53]
Subtotal ***	31		39			100%	3.9[-0.73,8.53]
Heterogeneity: Tau ² =0; Chi ² =0, df=0	(P<0.0001	.); I²=100%					
Test for overall effect: Z=1.65(P=0.1)						
1.81.3 Expressive communication	n at < 24 m	nonths					
Keenan 2014	34	6.2 (1.4)	15	6 (1.9)		100%	0.21[-0.86,1.28]
Subtotal ***	34		15		*	100%	0.21[-0.86,1.28]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.39(P=0.7)						
1.81.4 BSID III at < 24 months							
Makrides 2010	351	96.5 (13.6)	375	97.9 (15.3)		83.71%	-1.47[-3.58,0.64]
Miller 2016	48	98.5 (12.7)	35	96.1 (9.5)	-	16.29%	2.4[-2.38,7.18]
Subtotal ***	399		410		-	100%	-0.84[-2.77,1.09]
Heterogeneity: Tau ² =0; Chi ² =2.11, c	lf=1(P=0.1	5); I ² =52.63%					
Test for overall effect: Z=0.85(P=0.3	9)						





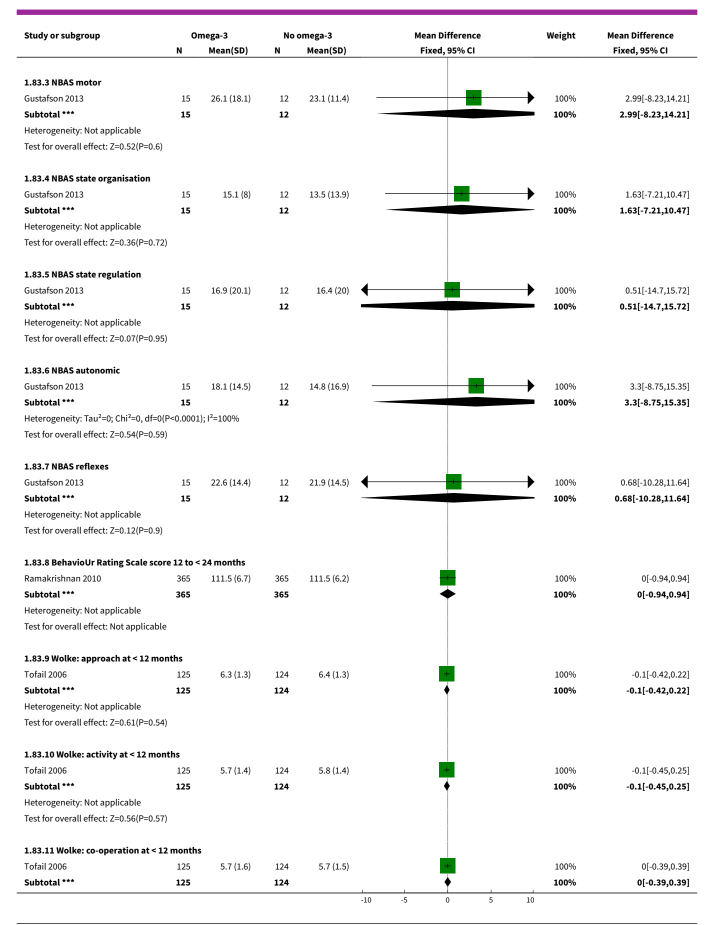
Analysis 1.82. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 82 Behaviour: thresholds.

Study or subgroup	Omega-3	No omega-3			Risk Ratio		Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI					M-H, Fixed, 95% CI	
1.82.1 Behaviour Rating Scale scor	es < 26: at < 24 mon	iths							
Ramakrishnan 2010	2/365	0/365		_			100%	5[0.24,103.79]	
Subtotal (95% CI)	365	365		-			100%	5[0.24,103.79]	
Total events: 2 (Omega-3), 0 (No ome	ega-3)								
Heterogeneity: Not applicable									
Test for overall effect: Z=1.04(P=0.3)									
		Favours omega-3	0.01	0.1	1 1	0 100	Favours no omega-3		

Analysis 1.83. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 83 Behaviour: scores.

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
		Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
1.83.1 NBAS habituation							
Gustafson 2013	15	8.5 (9.3)	12	9.9 (9.3)		100%	-1.45[-8.49,5.59]
Subtotal ***	15		12			100%	-1.45[-8.49,5.59]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.4(P=0.69)							
1.83.2 NBAS orienting							
Gustafson 2013	15	23.4 (18.3)	12	19.8 (15.5)	-	100%	3.65[-9.09,16.39]
Subtotal ***	15		12			100%	3.65[-9.09,16.39]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.56(P=0.57)							

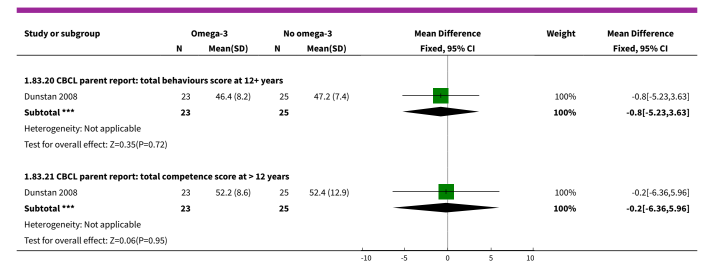






Study or subgroup	O N	mega-3 Mean(SD)	No N	omega-3 Mean(SD)	Mean Difference Fixed, 95% CI	Weight	Mean Difference Fixed, 95% CI
Heterogeneity: Not applicable					i incluyed to the		1 11.02,00 /0 01
Test for overall effect: Not applica	ble						
1.83.12 Wolke: emotional tone a	nt < 12 mon	ths					
Tofail 2006	125	5.6 (1.6)	124	5.7 (1.5)	+	100%	-0.1[-0.49,0.29
Subtotal ***	125		124		•	100%	-0.1[-0.49,0.29
Heterogeneity: Not applicable							
Test for overall effect: Z=0.51(P=0	.61)						
1.83.13 Wolke: vocalisation at <	12 months						
Tofail 2006	125	5 (1.7)	124	5.1 (1.7)	+	100%	-0.1[-0.52,0.32
Subtotal ***	125		124		♦	100%	-0.1[-0.52,0.32
Heterogeneity: Not applicable Test for overall effect: Z=0.46(P=0	.64)						
1.83.14 BSID III social-emotiona	l score at <	24 months			_		
Makrides 2010	351	106.3 (17.1)	375	107.3 (17.7)		85.89%	-0.95[-3.48,1.58
Miller 2016	48	108.4 (15.8)	35	107.6 (13.1)	+	14.11%	0.8[-5.43,7.03
Subtotal ***	399		410			100%	-0.7[-3.04,1.64
Heterogeneity: Tau ² =0; Chi ² =0.26, Test for overall effect: Z=0.59(P=0		1); I ² =0%					
1.83.15 BSID III adaptive behavi	our score a	t < 24 months					
Makrides 2010	351	99.2 (14)	375	100.8 (14.5)		86.43%	-1.58[-3.65,0.49
Miller 2016	48	108.5 (13.3)	35	107.3 (10.9)		13.57%	1.2[-4.02,6.42
Subtotal ***	399		410			100%	-1.2[-3.12,0.72
Heterogeneity: Tau²=0; Chi²=0.94,	df=1(P=0.3	3); I ² =0%					,
Test for overall effect: Z=1.23(P=0							
1.83.16 SDQ Total Difficulties at	2 to 5 year	s					
Makrides 2010	313	8.8 (4)	333	8.1 (4)	+	100%	0.62[-0,1.24
Subtotal ***	313		333		◆	100%	0.62[-0,1.24
Heterogeneity: Not applicable							
Test for overall effect: Z=1.96(P=0	.05)						
1.83.17 SDQ Total Difficulties at	6 to 9 year	s					
Makrides 2010	259	9.7 (5.2)	284	8.6 (5.5)		100%	1.08[0.18,1.98
Subtotal ***	259		284		•	100%	1.08[0.18,1.98
Heterogeneity: Not applicable							
Test for overall effect: Z=2.35(P=0	.02)						
1.83.18 BASC-2: Behavioral Sym	-						
Ramakrishnan 2010	401	51 (29.8)	396	51.5 (28.4)		100%	-0.5[-4.54,3.54 -
Subtotal ***	401		396			100%	-0.5[-4.54,3.54
Heterogeneity: Not applicable							
Test for overall effect: Z=0.24(P=0	.81)						
1.83.19 CBCL total problem beh		-			_		_
Dunstan 2008	33	35 (5.2)	39	36 (5.2)		100%	-1[-3.41,1.41
Subtotal ***	33		39			100%	-1[-3.41,1.41
Heterogeneity: Not applicable	10)						
Test for overall effect: Z=0.81(P=0	.42)						





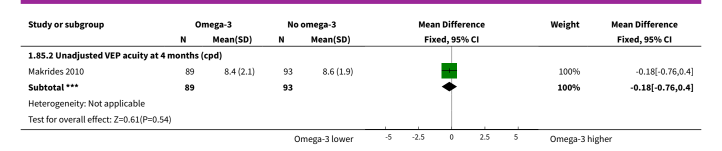
Analysis 1.84. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 84 Vision: visual acuity (cycles/degree).

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
1.84.1 At 2 months							
Mulder 2014	68	2.6 (0.6)	67	2.4 (0.5)	+	100%	0.18[-0.01,0.37]
Subtotal ***	68		67		→	100%	0.18[-0.01,0.37]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.84(P=0.07))						
1.84.2 At 4 months							
Judge 2007	16	3.7 (1.3)	14	3.2 (1.3)	-	100%	0.5[-0.43,1.43]
Subtotal ***	16		14		•	100%	0.5[-0.43,1.43]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.05(P=0.29))						
1.84.3 At 6 months							
Judge 2007	15	5.9 (1.2)	11	5.4 (1.3)	-	100%	0.5[-0.48,1.48]
Subtotal ***	15		11		•	100%	0.5[-0.48,1.48]
Heterogeneity: Not applicable							
Test for overall effect: Z=1(P=0.32)							
			Fav	ours omega-3	5 -2.5 0 2.5	5 Favours no	omega-3

Analysis 1.85. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 85 Vision: VEP acuity.

Study or subgroup	0	mega-3	No	omega-3		Mea	n Difference	•	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fix	ked, 95% CI			Fixed, 95% CI
1.85.1 Adjusted VEP acuity at 4 mo	nths (cp	d)								
Makrides 2010	89	8.4 (2)	93	8.6 (2)			-		100%	-0.18[-0.75,0.39]
Subtotal ***	89		93				*		100%	-0.18[-0.75,0.39]
Heterogeneity: Not applicable										
Test for overall effect: Z=0.62(P=0.54)									
			0	mega-3 lower	-5	-2.5	0 2	.5 5	Omega-3 high	er





Analysis 1.86. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 86 Vision: VEP latency.

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
1.86.1 Peak latency N1 at birth							
Malcolm 2003	5	62.2 (3.8)	4	74.8 (16.8)		100%	-12.6[-29.4,4.2]
Subtotal ***	5		4		•	100%	-12.6[-29.4,4.2]
Heterogeneity: Tau ² =0; Chi ² =0, df=0(P<0.000	1); I ² =100%					
Test for overall effect: Z=1.47(P=0.14							
1.86.2 Peak latency P1 at birth							
Malcolm 2003	9	101 (13.6)	5	107.8 (11.8)		100%	-6.8[-20.44,6.84]
Subtotal ***	9		5		•	100%	-6.8[-20.44,6.84]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.98(P=0.33)						
1.86.3 Peak latency N2 at birth							
Malcolm 2003	27	153.5 (28.9)	22	149.9 (28)	-	100%	3.6[-12.39,19.59]
Subtotal ***	27		22		*	100%	3.6[-12.39,19.59]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.44(P=0.66)						
1.86.4 Peak latency P2 at birth							
Malcolm 2003	28	201.9 (28.4)	27	201.8 (33.3)		100%	0.1[-16.28,16.48]
Subtotal ***	28		27		*	100%	0.1[-16.28,16.48]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.01(P=0.99)						
1.86.5 Peak latency N3 at birth							
Malcolm 2003	26	292.2 (58.2)	27	298.4 (52.8)		100%	-6.2[-36.15,23.75]
Subtotal ***	26		27			100%	-6.2[-36.15,23.75]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.41(P=0.68)						
1.86.6 Latency N1 (ms) at 3 month	s						
Ramakrishnan 2010	337	94.2 (16.3)	342	93.9 (17.1)	+	100%	0.3[-2.21,2.81]
Subtotal ***	337		342		→	100%	0.3[-2.21,2.81]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.23(P=0.81)						
1.86.7 Latency P1 (ms) at 3 months	5						
Ramakrishnan 2010	337	125.8 (17.5)	342	126.3 (18.3)	+	100%	-0.5[-3.19,2.19]



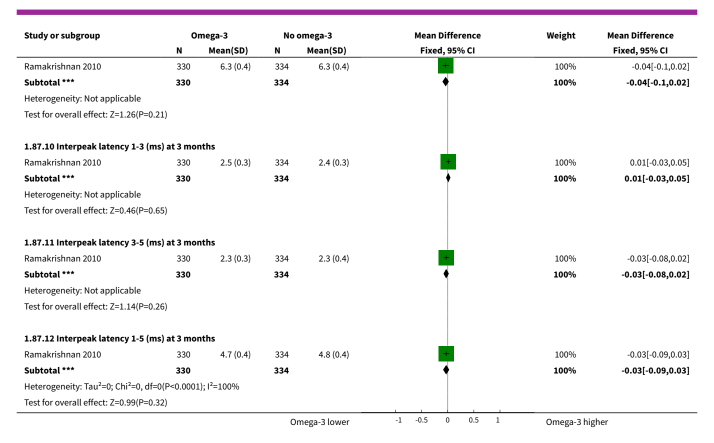
Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
Subtotal ***	337		342		†	100%	-0.5[-3.19,2.19
Heterogeneity: Not applicable							
Test for overall effect: Z=0.36(P=0.72)						
1.86.8 Latency N3 (ms) at 3 month	s						
Ramakrishnan 2010	337	154.8 (23.9)	342	157.1 (24.1)	+	100%	-2.3[-5.91,1.31
Subtotal ***	337		342		•	100%	-2.3[-5.91,1.31
Heterogeneity: Not applicable							
Test for overall effect: Z=1.25(P=0.21)						
1.86.9 Latency (69 min of arc) at 4	months	(ms)					
Makrides 2010	89	115 (8)	93	116 (9)	+	100%	-1[-3.47,1.47
Subtotal ***	89		93			100%	-1[-3.47,1.47
Heterogeneity: Not applicable							
Test for overall effect: Z=0.79(P=0.43)						
1.86.10 Latency (48 min of arc) at 4	l month	s (ms)					
Makrides 2010	89	121 (10)	93	121 (12)	+	100%	0[-3.2,3.2
Subtotal ***	89		93		▼	100%	0[-3.2,3.2
Heterogeneity: Not applicable							
Test for overall effect: Not applicable	<u> </u>						
1.86.11 Latency (20 min of arc) at 4	l month:	s (ms)					
Makrides 2010	89	133 (15)	93	133 (14)	+	100%	0[-4.22,4.22
Subtotal ***	89	, ,	93	. ,	▼	100%	0[-4.22,4.22
Heterogeneity: Not applicable							- ,
Test for overall effect: Not applicable	<u> </u>						
1.86.12 Latency N1 (ms) at 6 mont	hs						
Ramakrishnan 2010	407	90.5 (14.6)	410	91.9 (15.1)	+	100%	-1.4[-3.44,0.64
Subtotal ***	407	(=,	410	()	▼	100%	-1.4[-3.44,0.64
Heterogeneity: Not applicable							
Test for overall effect: Z=1.35(P=0.18)						
1.86.13 Latency P1 (ms) at 6 montl	ns						
Ramakrishnan 2010	407	122.7 (14.6)	410	123.5 (14.3)	+	100%	-0.8[-2.78,1.18
Subtotal ***	407		410		—	100%	-0.8[-2.78,1.18
Heterogeneity: Not applicable							
Test for overall effect: Z=0.79(P=0.43)						
1.86.14 Latency N3 (ms) at 6 mont	hs						
Ramakrishnan 2010	407	154.2 (19.9)	410	154.9 (20.2)	+	100%	-0.7[-3.45,2.05
Subtotal ***	407	10 (10.0)	410		—	100%	-0.7[-3.45,2.05
Heterogeneity: Not applicable			0		Ĭ	_30 /0	5.10,2.00
Test for overall effect: Z=0.5(P=0.62)							
Test for subgroup differences: Chi ² =5	٠ عد عد-	I (D=0.07) 12=00/					



Analysis 1.87. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 87 Hearing: brainstem auditory-evoked responses.

Study or subgroup	On	nega-3	No o	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI	_	Fixed, 95% CI
1.87.1 Latency 1 (ms) at 1 month							
Ramakrishnan 2010	372	1.6 (0.2)	377	1.6 (0.1)	+	100%	-0.01[-0.03,0.01]
Subtotal ***	372		377		$\overline{}$	100%	-0.01[-0.03,0.01]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.94(P=0.35)							
1.87.2 Latency 3 (ms) at 1 month							
Ramakrishnan 2010	372	4.2 (0.3)	377	4.2 (0.3)	+	100%	-0.01[-0.06,0.04]
Subtotal ***	372		377			100%	-0.01[-0.06,0.04]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.42(P=0.67)							
1.87.3 Latency 5 (ms) at 1 month							
Ramakrishnan 2010	372	6.5 (0.5)	377	6.6 (0.4)	+	100%	-0.03[-0.09,0.03]
Subtotal ***	372		377		→	100%	-0.03[-0.09,0.03]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.91(P=0.36)							
1.87.4 Interpeak latency 1-3 (ms) at	t 1 montl	h					
Ramakrishnan 2010	372	2.6 (0.3)	377	2.6 (0.4)	+	100%	-0.01[-0.06,0.04]
Subtotal ***	372		377			100%	-0.01[-0.06,0.04]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.43(P=0.67)							
1.87.5 Interpeak latency 3-5 (ms) at	t 1 montl	'n					
Ramakrishnan 2010	372	2.4 (0.3)	377	2.4 (0.3)	+	100%	0[-0.05,0.05]
Subtotal ***	372		377		→	100%	0[-0.05,0.05]
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
1.87.6 Interpeak latency 1-5 (ms) at	t 1 montl	h					
Ramakrishnan 2010	372	4.9 (0.4)	377	4.9 (0.4)	+	100%	-0.02[-0.07,0.03]
Subtotal ***	372		377		▼	100%	-0.02[-0.07,0.03]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.73(P=0.47)							
1.87.7 Latency 1 (ms) at 3 months							
Ramakrishnan 2010	330	1.6 (0.2)	334	1.6 (0.2)	÷	100%	0[-0.02,0.02]
Subtotal ***	330		334		$\overline{}$	100%	0[-0.02,0.02]
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
1.87.8 Latency 3 (ms) at 3 months							
Ramakrishnan 2010	330	4 (0.3)	334	4 (0.3)	+	100%	0.01[-0.04,0.06]
Subtotal ***	330		334		→	100%	0.01[-0.04,0.06]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.4(P=0.69)							
1.87.9 Latency 5 (ms) at 3 months							
			On	nega-3 lower	-1 -0.5 0 0.5 1	Omega-3 hi	gher
				<u> </u>			<u>-</u>

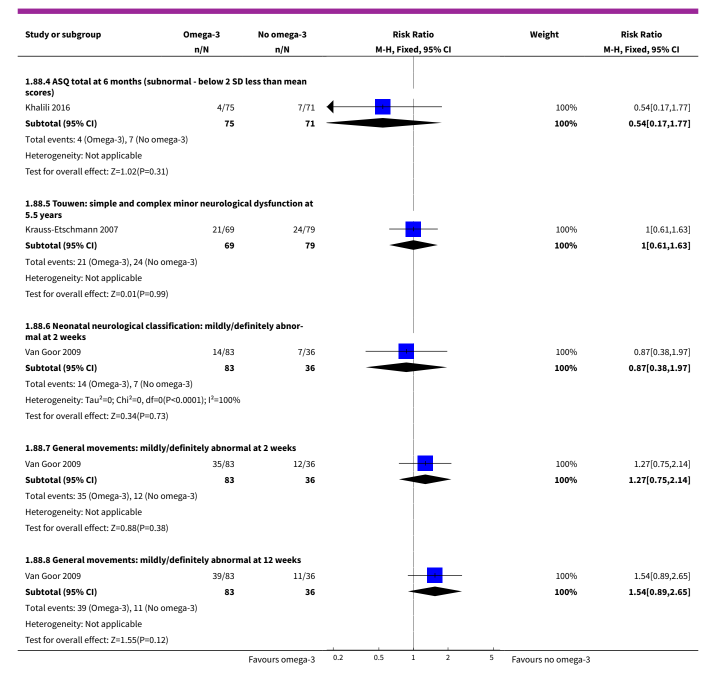




Analysis 1.88. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 88 Neurodevelopment: thresholds.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N		M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
1.88.1 Hempel: simple minor neurolog	gical dysfunction	n at 18 months			
Van Goor 2009	52/80	20/34	-	100%	1.11[0.8,1.53]
Subtotal (95% CI)	80	34	*	100%	1.11[0.8,1.53]
Total events: 52 (Omega-3), 20 (No omeg	ga-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.6(P=0.55)					
1.88.2 Hempel: simple and complex m 4 years	inor neurologic	al dysfunction at			
Krauss-Etschmann 2007	6/80	6/87		100%	1.09[0.37,3.23]
Subtotal (95% CI)	80	87		100%	1.09[0.37,3.23]
Total events: 6 (Omega-3), 6 (No omega-	-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.15(P=0.88)					
1.88.3 Hempel: complex minor neurol	ogical dysfuncti	on at 18 months			
Van Goor 2009	8/80	5/34		100%	0.68[0.24,1.93]
Subtotal (95% CI)	80	34		100%	0.68[0.24,1.93]
Total events: 8 (Omega-3), 5 (No omega-	-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.72(P=0.47)					
		Favours omega-3 0.2	0.5 1 2	5 Favours no omega-3	





Analysis 1.89. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 89 Neurodevelopment: scores.

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
1.89.1 ASQ gross motor at 4 months	s						
Khalili 2016	75	55.2 (7.9)	73	54.9 (8.7)		100%	0.3[-2.38,2.98]
Subtotal ***	75		73		—	100%	0.3[-2.38,2.98]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.22(P=0.83)							
			Favour	s no omega-3	-10 -5 0 5 10	Favours ome	ega-3



Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI	-	Fixed, 95% CI
1.89.2 ASQ gross motor at 6 months							
Khalili 2016	75	50.1 (10.4)	71	48.9 (11.2)		100%	1.2[-2.31,4.71]
Subtotal ***	75		71		•	100%	1.2[-2.31,4.71]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.67(P=0.5)							
1.89.3 ASQ fine motor at 4 months							
Khalili 2016	75	53.2 (8.8)	73	52.1 (10.5)	_	100%	1.1[-2.03,4.23]
Subtotal ***	75		73		•	100%	1.1[-2.03,4.23]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.69(P=0.49)							
1.89.4 ASQ fine motor at 6 months							
Khalili 2016	75	55.8 (8.8)	71	54.6 (8.4)	-	100%	1.2[-1.59,3.99]
Subtotal ***	75		71		•	100%	1.2[-1.59,3.99]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.84(P=0.4)							
1.89.5 ASQ problem solving at 4 mon	ths						
Khalili 2016	75	56.5 (6)	73	54.9 (9.6)	-	100%	1.6[-0.99,4.19]
Subtotal ***	75		73			100%	1.6[-0.99,4.19]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.21(P=0.23)							
1.89.6 ASQ problem solving at 6 mon	ths						
Khalili 2016	75	56.2 (8.2)	71	55.7 (6.9)	-	100%	0.5[-1.95,2.95]
Subtotal ***	75		71		→	100%	0.5[-1.95,2.95]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.4(P=0.69)							
1.89.7 ASQ personal-social at 4 mont	hs						
Khalili 2016	75	54.2 (8.3)	73	53.1 (8.7)	-	100%	1.1[-1.64,3.84]
Subtotal ***	75		73		•	100%	1.1[-1.64,3.84]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.79(P=0.43)							
1.89.8 ASQ personal-social at 6 mont	hs						
Khalili 2016	75	52.1 (10.2)	71	51.3 (10.8)	-	100%	0.8[-2.61,4.21]
Subtotal ***	75		71		*	100%	0.8[-2.61,4.21]
Heterogeneity: Tau ² =0; Chi ² =0, df=0(P<	0.0001	L); I ² =100%					
Test for overall effect: Z=0.46(P=0.65)							
1.89.9 ASQ communication at 4 mon	ths						
Khalili 2016	75	54.8 (5.3)	73	52.1 (8.5)	-	100%	2.7[0.41,4.99]
Subtotal ***	75		73		•	100%	2.7[0.41,4.99]
Heterogeneity: Not applicable							
Test for overall effect: Z=2.31(P=0.02)							
1.89.10 ASQ communication at 6 mo	nths						
Khalili 2016	75	55.8 (5.9)	71	55.4 (6.1)	-	100%	0.4[-1.55,2.35]
Subtotal ***	75		71		*	100%	0.4[-1.55,2.35]
Heterogeneity: Not applicable							

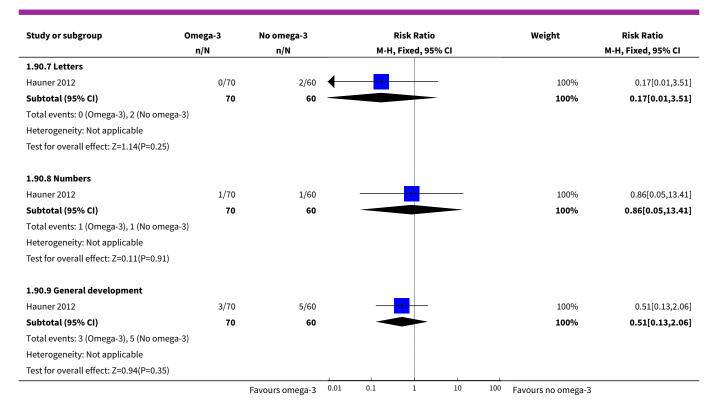


Study or subgroup	(Omega-3 No omega-3		Mean Difference					Weight Mean Difference	
	N Mean(SD)		N	Mean(SD)	Fixed, 95% CI			Fixed, 95% CI		
Test for overall effect: Z=0.4(P=0.69)				_						
			Favou	rs no omega-3	-10	-5	0	5	10	Favours omega-3

Analysis 1.90. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 90 Child Development Inventory.

Subtotal (95% CI) Total events: 0 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Not applicable 1.90.2 Self help Hauner 2012 0/ Subtotal (95% CI) Total events: 0 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.77(P=0.44) 1.90.3 Gross motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/	n/N 70 0/60 70 60	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
Subtotal (95% CI) Total events: 0 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Not applicable 1.90.2 Self help Hauner 2012 0/ Subtotal (95% CI) Total events: 0 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.77(P=0.44) 1.90.3 Gross motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)	*			
Total events: 0 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Not applicable 1.90.2 Self help Hauner 2012	70 60			Not estimable
Heterogeneity: Not applicable Test for overall effect: Not applicable 1.90.2 Self help Hauner 2012 0/ Subtotal (95% CI) Total events: 0 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.77(P=0.44) 1.90.3 Gross motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				Not estimable
Test for overall effect: Not applicable 1.90.2 Self help Hauner 2012 O/ Subtotal (95% CI) Total events: 0 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.77(P=0.44) 1.90.3 Gross motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
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Hauner 2012 Subtotal (95% CI) Total events: 0 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.77(P=0.44) 1.90.3 Gross motor Hauner 2012 Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
Subtotal (95% CI) Total events: 0 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.77(P=0.44) 1.90.3 Gross motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
Total events: 0 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.77(P=0.44) 1.90.3 Gross motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)	70 1/60 -		100%	0.29[0.01,6.9]
Heterogeneity: Not applicable Test for overall effect: Z=0.77(P=0.44) 1.90.3 Gross motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)	70 60 -		100%	0.29[0.01,6.9]
Test for overall effect: Z=0.77(P=0.44) 1.90.3 Gross motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
1.90.3 Gross motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
Hauner 2012 Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)	70 0/60		100%	4.3[0.21,87.76]
Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)	70 60		100%	4.3[0.21,87.76]
Test for overall effect: Z=0.95(P=0.34) 1.90.4 Fine motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
1.90.4 Fine motor Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
Hauner 2012 2/ Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
Subtotal (95% CI) Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
Total events: 2 (Omega-3), 0 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)	70 0/60	- 		4.3[0.21,87.76]
Heterogeneity: Not applicable Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)	70 60		100%	4.3[0.21,87.76]
Test for overall effect: Z=0.95(P=0.34) 1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
1.90.5 Expressive language Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
Hauner 2012 1/ Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
Subtotal (95% CI) Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)				
Total events: 1 (Omega-3), 1 (No omega-3) Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)	70 1/60		100%	0.86[0.05,13.41]
Heterogeneity: Not applicable Test for overall effect: Z=0.11(P=0.91)	70 60		100%	0.86[0.05,13.41]
Test for overall effect: Z=0.11(P=0.91)				
1.90.6 Language comprehension				
Hauner 2012 0/	70 0/60			Not estimable
Subtotal (95% CI)	70 60			Not estimable
Total events: 0 (Omega-3), 0 (No omega-3)				
Heterogeneity: Not applicable				
Test for overall effect: Not applicable				

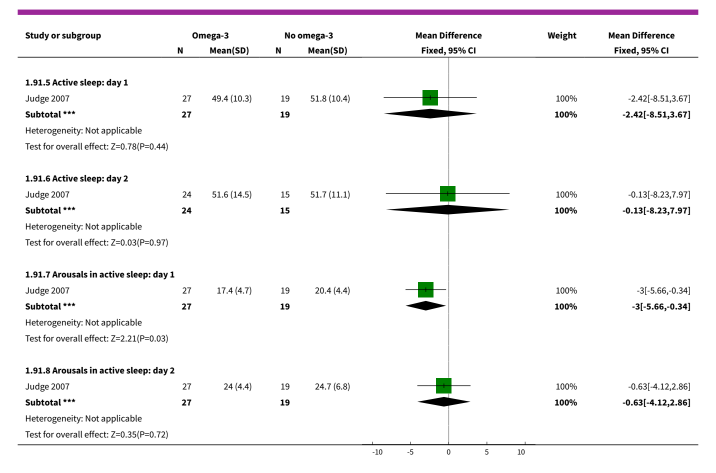




Analysis 1.91. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 91 Infant sleep behaviour (%).

Study or subgroup	Omega-3		No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
1.91.1 Arousals in quiet sleep: day 1	L						
Judge 2007	27	2.7 (2.7)	19	5.9 (6)		100%	-3.19[-6.07,-0.31]
Subtotal ***	27		19			100%	-3.19[-6.07,-0.31]
Heterogeneity: Not applicable							
Test for overall effect: Z=2.17(P=0.03)							
1.91.2 Arousals in quiet sleep: day 2	<u>!</u>						
Judge 2007	24	3.6 (4)	15	5.4 (4.1)		100%	-1.89[-4.49,0.71]
Subtotal ***	24		15			100%	-1.89[-4.49,0.71]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.42(P=0.15)							
1.91.3 Quiet sleep: day 1							
Judge 2007	27	15.9 (5.1)	19	15.1 (4.3)		100%	0.74[-1.97,3.45]
Subtotal ***	27		19			100%	0.74[-1.97,3.45]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.53(P=0.59)							
1.91.4 Quiet sleep: day 2							
Judge 2007	24	12.7 (5.9)	15	13.7 (4.8)		100%	-1[-4.36,2.36]
Subtotal ***	24		15			100%	-1[-4.36,2.36]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.58(P=0.56)							
					10 -5 0 5	10	





Analysis 1.92. Comparison 1 Overall: omega-3 versus no omega-3, Outcome 92 Cerebral palsy.

Study or subgroup	Omega-3	Control	Risk R		lisk Ratio			Weight	Risk Ratio
	n/N	n/N		М-Н,	Fixed, 95	% CI			M-H, Fixed, 95% CI
Van Goor 2009	0/80	0/34							Not estimable
Total (95% CI)	80	34							Not estimable
Total events: 0 (Omega-3), 0 (Control)									
Heterogeneity: Not applicable									
Test for overall effect: Not applicable						1			
		Favours omega-3	0.01	0.1	1	10	100	Favours control	

Comparison 2. Type of omega-3 intervention

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Preterm birth (< 37 weeks)	27	10304	Risk Ratio (M-H, Fixed, 95% CI)	0.88 [0.81, 0.97]
1.1 Omega-3 supplements only	18	7608	Risk Ratio (M-H, Fixed, 95% CI)	0.90 [0.80, 1.01]



Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1.2 Omega-3 supplements/enrich- ment + food/diet advice	3	516	Risk Ratio (M-H, Fixed, 95% CI)	0.73 [0.41, 1.29]
1.3 Omega-3 food/diet advice	1	48	Risk Ratio (M-H, Fixed, 95% CI)	0.11 [0.01, 2.22]
1.4 Omega-3 supplements + other agents	6	2132	Risk Ratio (M-H, Fixed, 95% CI)	0.89 [0.76, 1.04]
2 Early preterm birth (< 34 weeks)	9	5204	Risk Ratio (M-H, Fixed, 95% CI)	0.58 [0.44, 0.77]
2.1 Omega-3 supplements only	8	4234	Risk Ratio (M-H, Fixed, 95% CI)	0.62 [0.46, 0.82]
2.2 Omega-3 supplements + other agents	1	970	Risk Ratio (M-H, Fixed, 95% CI)	0.19 [0.04, 0.88]
3 Prolonged gestation (> 42 weeks)	6	5141	Risk Ratio (M-H, Fixed, 95% CI)	1.61 [1.11, 2.33]
3.1 Omega-3 supplements only	5	4953	Risk Ratio (M-H, Fixed, 95% CI)	1.59 [1.09, 2.31]
3.2 Omega-3 supplements + food/diet advice	1	188	Risk Ratio (M-H, Fixed, 95% CI)	3.13 [0.13, 75.84]
4 Maternal death	4	4830	Risk Ratio (M-H, Fixed, 95% CI)	1.69 [0.07, 39.30]
4.1 Omega-3 supplements only	3	4782	Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]
4.2 Omega-3 food/diet advice	1	48	Risk Ratio (M-H, Fixed, 95% CI)	1.69 [0.07, 39.30]
5 Pre-eclampsia (hypertension with proteinuria)	21	8306	Risk Ratio (M-H, Fixed, 95% CI)	0.84 [0.69, 1.01]
5.1 Omega-3 supplements only	13	5825	Risk Ratio (M-H, Fixed, 95% CI)	0.95 [0.76, 1.19]
5.2 Omega-3 supplements/enrich- ment + food/dietary advice	2	328	Risk Ratio (M-H, Fixed, 95% CI)	0.65 [0.25, 1.69]
5.3 Omega-3 supplements + other agents	6	2153	Risk Ratio (M-H, Fixed, 95% CI)	0.58 [0.39, 0.88]
6 High blood pressure (without proteinuria)	7	4531	Risk Ratio (M-H, Fixed, 95% CI)	1.03 [0.89, 1.20]
6.1 Omega-3 supplements only	6	4431	Risk Ratio (M-H, Fixed, 95% CI)	1.05 [0.90, 1.22]
6.2 Omega-3 supplements + other agents	1	100	Risk Ratio (M-H, Fixed, 95% CI)	0.69 [0.33, 1.47]
7 Eclampsia	1	100	Risk Ratio (M-H, Fixed, 95% CI)	0.14 [0.01, 2.70]
7.1 Omega-3 supplements + other agents	1	100	Risk Ratio (M-H, Fixed, 95% CI)	0.14 [0.01, 2.70]
8 Maternal antepartum hospitalisa- tion	5	2876	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.81, 1.04]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
8.1 Omega-3 supplements only	4	2817	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.82, 1.04]
8.2 Omega-3 supplementation + other agents	1	59	Risk Ratio (M-H, Fixed, 95% CI)	0.21 [0.01, 4.13]
9 Mother's length of stay in hospital (days)	2	2290	Mean Difference (IV, Fixed, 95% CI)	0.18 [-0.20, 0.57]
9.1 Omega-3 supplements only	2	2290	Mean Difference (IV, Fixed, 95% CI)	0.18 [-0.20, 0.57]
10 Maternal anaemia	1	846	Risk Ratio (M-H, Fixed, 95% CI)	1.16 [0.91, 1.48]
10.1 Omega-3 supplements only	1	846	Risk Ratio (M-H, Fixed, 95% CI)	1.16 [0.91, 1.48]
11 Miscarriage (< 24 weeks)	9	4190	Risk Ratio (M-H, Fixed, 95% CI)	1.07 [0.80, 1.43]
11.1 Omega-3 supplements only	8	3049	Risk Ratio (M-H, Fixed, 95% CI)	0.95 [0.56, 1.60]
11.2 Omega-3 supplements + other agents	1	1141	Risk Ratio (M-H, Fixed, 95% CI)	1.14 [0.80, 1.61]
12 Antepartum vaginal bleeding	2	2151	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.69, 1.48]
12.1 Omega-3 supplements only	2	2151	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.69, 1.48]
13 Preterm prelabour rupture of membranes	3	925	Risk Ratio (M-H, Fixed, 95% CI)	0.53 [0.25, 1.10]
13.1 Omega-3 supplements only	2	670	Risk Ratio (M-H, Fixed, 95% CI)	0.61 [0.28, 1.34]
13.2 Omega-3 supplementation/enrichment + food/diet advice	1	255	Risk Ratio (M-H, Fixed, 95% CI)	0.24 [0.03, 2.15]
14 Prelabour rupture of membranes	3	915	Risk Ratio (M-H, Fixed, 95% CI)	0.41 [0.21, 0.82]
14.1 Omega-3 supplements only	1	369	Risk Ratio (M-H, Fixed, 95% CI)	0.54 [0.14, 2.11]
14.2 Omega-3 supplementation/enrichment + food/diet advice	2	546	Risk Ratio (M-H, Fixed, 95% CI)	0.38 [0.17, 0.85]
15 Maternal admission to intensive care	2	2458	Risk Ratio (M-H, Fixed, 95% CI)	0.56 [0.12, 2.63]
15.1 Omega-3 supplements only	1	2399	Risk Ratio (M-H, Fixed, 95% CI)	1.00 [0.14, 7.12]
15.2 Omega-3 supplements + other agent	1	59	Risk Ratio (M-H, Fixed, 95% CI)	0.21 [0.01, 4.13]
16 Maternal severe adverse effects (including cessation)	8	4177	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.59, 1.75]
16.1 Omega-3 supplements only	7	3886	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.54, 1.87]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
16.2 Omega-3 supplementation/enrichment + food/diet advice	1	291	Risk Ratio (M-H, Fixed, 95% CI)	1.05 [0.35, 3.18]
17 Caesarean section	29	8481	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.91, 1.03]
17.1 Omega-3 supplements only	19	6537	Risk Ratio (M-H, Fixed, 95% CI)	0.98 [0.92, 1.06]
17.2 Omega-3 supplements/enrichment +food/diet advice	4	574	Risk Ratio (M-H, Fixed, 95% CI)	0.87 [0.63, 1.19]
17.3 Omega-3 food/diet advice	1	107	Risk Ratio (M-H, Fixed, 95% CI)	0.91 [0.38, 2.17]
17.4 Omega-3 supplements + other agents	5	1263	Risk Ratio (M-H, Fixed, 95% CI)	0.88 [0.72, 1.08]
18 Induction (post-term)	3	2900	Risk Ratio (M-H, Random, 95% CI)	0.82 [0.22, 2.98]
18.1 Omega-3 supplements only	2	2712	Risk Ratio (M-H, Random, 95% CI)	0.82 [0.22, 2.98]
18.2 Omega-3 supplements + food/ diet advice	1	188	Risk Ratio (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
19 Blood loss at birth (mL)	6	2776	Mean Difference (IV, Fixed, 95% CI)	11.50 [-6.75, 29.76
19.1 Omega-3 supplements only	5	2588	Mean Difference (IV, Fixed, 95% CI)	11.64 [-8.89, 32.17
19.2 Omega-3 supplements + food/ diet advice	1	188	Mean Difference (IV, Fixed, 95% CI)	11.0 [-28.91, 50.91
20 Postpartum haemorrhage	4	4085	Risk Ratio (M-H, Fixed, 95% CI)	1.03 [0.82, 1.30]
20.1 Omega-3 supplements only	3	3233	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.71, 1.34]
20.2 Omega-3 supplements + other agent	1	852	Risk Ratio (M-H, Fixed, 95% CI)	1.11 [0.79, 1.57]
21 Gestational diabetes	12	5235	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.83, 1.26]
21.1 Omega-3 supplements only	7	3726	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.80, 1.30]
21.2 Omega-3 supplements/enrichment + food/diet advice	4	595	Risk Ratio (M-H, Fixed, 95% CI)	0.66 [0.33, 1.34]
21.3 Omega-3 supplements + other agents	2	914	Risk Ratio (M-H, Fixed, 95% CI)	1.34 [0.80, 2.24]
22 Maternal insulin resistance (HOMA-IR)	3	176	Mean Difference (IV, Random, 95% CI)	-0.85 [-2.50, 0.80]
22.1 Omega-3 supplements only	2	116	Mean Difference (IV, Random, 95% CI)	-0.25 [-1.94, 1.44]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
22.2 Omega-3 supplements + other agents	1	60	Mean Difference (IV, Random, 95% CI)	-2.0 [-3.10, -0.90]
23 Excessive gestational weight gain	1	350	Risk Ratio (M-H, Fixed, 95% CI)	1.21 [0.95, 1.55]
23.1 Omega-3 supplements only	1	350	Risk Ratio (M-H, Fixed, 95% CI)	1.21 [0.95, 1.55]
24 Gestational weight gain (kg)	11	2297	Mean Difference (IV, Random, 95% CI)	-0.05 [-0.68, 0.59]
24.1 Omega-3 supplements only	6	955	Mean Difference (IV, Random, 95% CI)	-0.22 [-1.47, 1.03]
24.2 Omega-3 supplements/enrichment + food/diet advice	3	313	Mean Difference (IV, Random, 95% CI)	-0.11 [-0.99, 0.78]
24.3 Omega-3 supplements + other agents	2	1029	Mean Difference (IV, Random, 95% CI)	0.43 [-0.08, 0.95]
25 Depression during pregnancy: scores	5		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
25.1 Omega-3 supplements only: BDI	2	104	Mean Difference (IV, Fixed, 95% CI)	-5.86 [-8.32, -3.39]
25.2 Omega-3 supplements only: HAMD	3	71	Mean Difference (IV, Fixed, 95% CI)	-1.08 [-3.35, 1.19]
25.3 Omega-3 supplements only: EPDS	4	122	Mean Difference (IV, Fixed, 95% CI)	-0.15 [-2.09, 1.79]
25.4 Omega-3 supplements only: MADRS	1	26	Mean Difference (IV, Fixed, 95% CI)	-1.60 [-7.80, 4.60]
26 Depression during pregnancy: thresholds	3		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
26.1 Omega-3 supplements only: HAMD 50% reduction (after 8 weeks)	1	24	Risk Ratio (M-H, Fixed, 95% CI)	2.26 [0.78, 6.49]
26.2 Omega-3 supplements only: HAMD ≤ 7	1	24	Risk Ratio (M-H, Fixed, 95% CI)	2.12 [0.51, 8.84]
26.3 Omega-3 supplements only: unspecified	1	301	Risk Ratio (M-H, Fixed, 95% CI)	2.39 [0.47, 12.11]
26.4 Omega-3 supplements only: EPDS ≥ 11	1	34	Risk Ratio (M-H, Fixed, 95% CI)	1.4 [0.55, 3.55]
27 Depressive symptoms postpar- tum: thresholds	4		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
27.1 Omega-3 supplements only: PDSS ≥80	1	42	Risk Ratio (M-H, Fixed, 95% CI)	0.37 [0.04, 3.25]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
27.2 Omega-3 supplements only: EPDS	2	2431	Risk Ratio (M-H, Fixed, 95% CI)	0.89 [0.71, 1.12]
27.3 Omega-3 supplements only: major depressive disorder	1	118	Risk Ratio (M-H, Fixed, 95% CI)	1.33 [0.27, 6.56]
28 Depressive symptoms postpartum: scores	2		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
28.1 Omega-3 supplements only: BD: 6-8 weeks postpartum	1	118	Mean Difference (IV, Fixed, 95% CI)	0.25 [-1.93, 2.43]
28.2 Omega-3 supplements only: PDSS total (LS over 6 months)	1	42	Mean Difference (IV, Fixed, 95% CI)	-6.08 [-12.42, 0.26]
29 Length of gestation (days)	43	12517	Mean Difference (IV, Random, 95% CI)	1.65 [0.94, 2.37]
29.1 Omega-3 supplements only	29	9290	Mean Difference (IV, Random, 95% CI)	1.67 [0.76, 2.59]
29.2 Omega-3 supplements/enrichment + food/diet advice	6	680	Mean Difference (IV, Random, 95% CI)	2.45 [-0.14, 5.04]
29.3 Omega-3 food/diet advice	1	107	Mean Difference (IV, Random, 95% CI)	5.00 [0.64, 9.36]
29.4 Omega-3 supplements + other agents	8	2440	Mean Difference (IV, Random, 95% CI)	1.04 [0.05, 2.03]
30 Perinatal death	10	7416	Risk Ratio (M-H, Fixed, 95% CI)	0.75 [0.54, 1.03]
30.1 Omega-3 supplements only	8	6496	Risk Ratio (M-H, Fixed, 95% CI)	0.71 [0.48, 1.03]
30.2 Omega-3 supplements + other agents	2	920	Risk Ratio (M-H, Fixed, 95% CI)	0.87 [0.47, 1.62]
31 Stillbirth	16	7880	Risk Ratio (M-H, Fixed, 95% CI)	0.94 [0.62, 1.42]
31.1 Omega-3 supplements only	13	7693	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.60, 1.42]
31.2 Omega-3 supplements + food/ diet advice	1	79	Risk Ratio (M-H, Fixed, 95% CI)	0.33 [0.01, 7.75]
31.3 Omega-3 food/diet advice	1	48	Risk Ratio (M-H, Fixed, 95% CI)	1.69 [0.07, 39.30]
31.4 Omega-3 supplements + other agents	1	60	Risk Ratio (M-H, Fixed, 95% CI)	3.0 [0.13, 70.83]
32 Neonatal death	9	7448	Risk Ratio (M-H, Fixed, 95% CI)	0.61 [0.34, 1.11]
32.1 Omega-3 supplements only	9	7448	Risk Ratio (M-H, Fixed, 95% CI)	0.61 [0.34, 1.11]
33 Infant death	4	3239	Risk Ratio (M-H, Fixed, 95% CI)	0.74 [0.25, 2.19]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size	
33.1 Omega-3 supplements only	4	3239	Risk Ratio (M-H, Fixed, 95% CI)	0.74 [0.25, 2.19]	
34 Large-for-gestational age	5	3602	Risk Ratio (M-H, Fixed, 95% CI)	1.20 [1.01, 1.43]	
34.1 Omega-3 supplements only	2	2518	Risk Ratio (M-H, Fixed, 95% CI)	1.19 [0.99, 1.43]	
34.2 Omega-3 supplements + food/ diet advice	1	188	Risk Ratio (M-H, Fixed, 95% CI)	1.23 [0.48, 3.17]	
34.3 Omega-3 supplements + other agent	2	896	Risk Ratio (M-H, Fixed, 95% CI)	1.28 [0.72, 2.29]	
35 Macrosomia	7	2008	Risk Ratio (M-H, Fixed, 95% CI)	0.69 [0.43, 1.13]	
35.1 Omega-3 supplements only	5	1904	Risk Ratio (M-H, Fixed, 95% CI)	0.80 [0.47, 1.36]	
35.2 Omega-3 supplements + other agent	2	104	Risk Ratio (M-H, Fixed, 95% CI)	0.31 [0.08, 1.23]	
36 Low birthweight (< 2500 g)	15	8449	Risk Ratio (M-H, Fixed, 95% CI)	0.90 [0.82, 0.99]	
36.1 Omega-3 supplements only	10	6214	Risk Ratio (M-H, Fixed, 95% CI)	0.96 [0.86, 1.07]	
36.2 Omega-3 supplements/enrich- ment + food/diet advice	2	328	Risk Ratio (M-H, Fixed, 95% CI)	0.66 [0.34, 1.26]	
36.3 Omega-3 supplements + other agents	3	1907	Risk Ratio (M-H, Fixed, 95% CI)	0.77 [0.62, 0.95]	
37 Small-for-gestational age/IUGR	8	6907	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.90, 1.13]	
37.1 Omega-3 supplements only	5	5041	Risk Ratio (M-H, Fixed, 95% CI)	1.05 [0.93, 1.20]	
37.2 Omega-3 supplements + other agents	3	1866	Risk Ratio (M-H, Fixed, 95% CI)	0.80 [0.59, 1.09]	
38 Birthweight (g)	44	11584	Mean Difference (IV, Random, 95% CI)	75.74 [38.05, 113.43]	
38.1 Omega-3 supplements only	31	8522	Mean Difference (IV, Random, 95% CI)	59.41 [23.23, 95.59]	
38.2 Omega-3 supplements/enrichment + food/diet advice	6	859	Mean Difference (IV, Random, 95% CI)	129.42 [49.52, 209.31]	
38.3 Omega-3 food/diet advice	1	107	Mean Difference (IV, Random, 95% CI)	-17.0 [-190.97, 156.97]	
38.4 Omega-3 supplements + other agents	6	2096	Mean Difference (IV, Random, 95% CI)	69.14 [-72.81, 211.10]	
39 Birthweight Z score	4	2792	Mean Difference (IV, Fixed, 95% CI)	0.06 [-0.02, 0.13	



Outcome or subgroup title	tcome or subgroup title No. of studies		Statistical method	Effect size	
39.1 Omega-3 supplements only	3	2677	Mean Difference (IV, Fixed, 95% CI)	0.06 [-0.01, 0.14]	
39.2 Omega-3 supplements + other agent	1	115	Mean Difference (IV, Fixed, 95% CI)	0.00 [-0.21, 0.21]	
40 Birth length (cm)	29	8008	Mean Difference (IV, Random, 95% CI)	0.13 [-0.08, 0.34]	
40.1 Omega-3 supplements only	20	6010	Mean Difference (IV, Random, 95% CI)	0.21 [-0.03, 0.45]	
40.2 Omega-3 supplements/enrichment + food/diet advice	4	606	Mean Difference (IV, Random, 95% CI)	0.42 [-0.01, 0.85]	
40.3 Omega-3 food/diet advice	1	123	Mean Difference (IV, Random, 95% CI)	-0.10 [-0.56, 0.36]	
40.4 Omega-3 supplements + other agent	4	1269	Mean Difference (IV, Random, 95% CI)	-0.51 [-0.78, -0.23]	
41 Length at birth Z score	2	2462	Mean Difference (IV, Random, 95% CI)	0.18 [-0.18, 0.54]	
41.1 Omega-3 supplements only	2	2462	Mean Difference (IV, Random, 95% CI)	0.18 [-0.18, 0.54]	
42 Head circumference at birth (cm)	23	7041	Mean Difference (IV, Fixed, 95% CI)	0.10 [0.01, 0.18]	
42.1 Omega-3 supplements only	16	5442	Mean Difference (IV, Fixed, 95% CI)	0.07 [-0.03, 0.17]	
42.2 Omega-3 supplements/enrichment + food/diet advice	3	418	Mean Difference (IV, Fixed, 95% CI)	0.34 [0.03, 0.65]	
42.3 Omega-3 food/diet advice only	1	107	Mean Difference (IV, Fixed, 95% CI)	-0.20 [-0.75, 0.35]	
42.4 Omega-3 supplements + other agent	3	1074	Mean Difference (IV, Fixed, 95% CI)	0.15 [-0.06, 0.35]	
43 Head circumference at birth Z score	2	2462	Mean Difference (IV, Fixed, 95% CI)	-0.03 [-0.14, 0.07]	
43.1 Omega-3 supplementation only	2	2462	Mean Difference (IV, Fixed, 95% CI)	-0.03 [-0.14, 0.07]	
44 Baby admitted to neonatal care	9	6920	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.83, 1.03]	
44.1 Omega-3 supplements only	5	5692	Risk Ratio (M-H, Fixed, 95% CI)	0.90 [0.79, 1.02]	
44.2 Omega-3 supplements/enrichment + food/diet advice	2	328	Risk Ratio (M-H, Fixed, 95% CI)	0.89 [0.54, 1.50]	



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size	
44.3 Omega-3 supplements + other agents	2	900	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.81, 1.26]	
45 Infant length of stay in hospital (days)	1	2041	Mean Difference (IV, Fixed, 95% CI)	0.11 [-1.40, 1.62]	
45.1 Omega-3 supplementation only	1	2041	Mean Difference (IV, Fixed, 95% CI)	0.11 [-1.40, 1.62]	
46 Congenital anomalies	3	1807	Risk Ratio (M-H, Fixed, 95% CI)	1.08 [0.61, 1.92]	
46.1 Omega-3 supplements only	3	1807	Risk Ratio (M-H, Fixed, 95% CI)	1.08 [0.61, 1.92]	
47 Retinopathy of prematurity	1	837	Risk Ratio (M-H, Fixed, 95% CI)	1.20 [0.32, 4.44]	
47.1 Omega-3 supplementation + other agent only	1	837	Risk Ratio (M-H, Fixed, 95% CI)	1.20 [0.32, 4.44]	
48 Bronchopulmonary dysplasia	2	3191	Risk Ratio (M-H, Fixed, 95% CI)	1.06 [0.45, 2.48]	
48.1 Omega-3 supplementation only	1	2363	Risk Ratio (M-H, Fixed, 95% CI)	0.50 [0.09, 2.71]	
48.2 Omega-3 supplementation + other agent	1	828	Risk Ratio (M-H, Fixed, 95% CI)	1.42 [0.51, 3.96]	
49 Respiratory distress syndrome	2	1129	Risk Ratio (M-H, Random, 95% CI)	1.17 [0.54, 2.52]	
49.1 Omega-3 supplementation only	1	301	Risk Ratio (M-H, Random, 95% CI)	0.72 [0.31, 1.65]	
49.2 Omega-3 supplementation + other agent	1	828	Risk Ratio (M-H, Random, 95% CI)	1.60 [1.08, 2.37]	
50 Necrotising enterocolitis (NEC)	2	3198	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.26, 3.55]	
50.1 Omega-3 supplementation only	1	2361	Risk Ratio (M-H, Fixed, 95% CI)	2.98 [0.12, 73.13]	
50.2 Omega-3 supplementation + other agent	1	837	Risk Ratio (M-H, Fixed, 95% CI)	0.72 [0.16, 3.20]	
51 Neonatal sepsis (proven)	3	3788	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.44, 2.14]	
51.1 Omega-3 supplements only	3	3788	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.44, 2.14]	
52 Convulsion	1	2361	Risk Ratio (M-H, Fixed, 95% CI)	0.09 [0.01, 1.63]	
52.1 Omega-3 supplementation only	1	2361	Risk Ratio (M-H, Fixed, 95% CI)	0.09 [0.01, 1.63]	
53 Intraventricular haemorrhage	3	5423	Risk Ratio (M-H, Random, 95% CI)	1.00 [0.29, 3.49]	
53.1 Omega-3 supplements only	2	4586	Risk Ratio (M-H, Random, 95% CI)	0.59 [0.02, 16.16]	

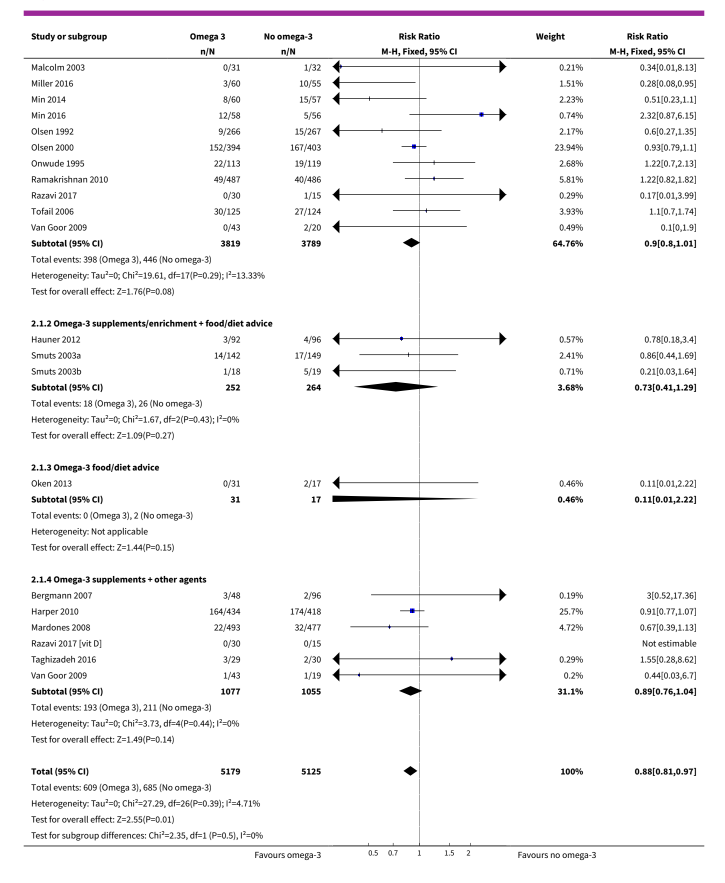


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size	
53.2 Omega-3 supplementation + other agent	1	837	Risk Ratio (M-H, Random, 95% CI)	1.07 [0.44, 2.60]	
54 Neonatal/infant serious adverse events	2	2690	Risk Ratio (M-H, Fixed, 95% CI)	0.72 [0.53, 0.99]	
54.1 Omega-3 supplementation	1	2399	Risk Ratio (M-H, Fixed, 95% CI)	0.67 [0.44, 1.01]	
54.2 Omega-3 supplements/enrichment + food/diet advice	1	291	Risk Ratio (M-H, Fixed, 95% CI)	0.81 [0.50, 1.31]	
55 Neonatal/infant morbidity: cardiovascular	1	291	Risk Ratio (M-H, Fixed, 95% CI)	1.20 [0.85, 1.69]	
55.1 Omega-3 supplements/enrich- ment + food/diet advice	1	291	Risk Ratio (M-H, Fixed, 95% CI)	1.20 [0.85, 1.69]	
56 Neonatal/infant morbidity: respiratory	1	291	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.66, 1.57]	
56.1 Omega-3 supplements/enrich- ment + food/diet advice	1	291	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.66, 1.57]	
57 Neonatal/infant morbidity: caused by pregnancy/birth	1	291	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.67, 1.55]	
57.1 Omega-3 supplements/enrichment + food/diet advice	1	291	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.67, 1.55]	
58 Ponderal index	6	887	Mean Difference (IV, Random, 95% CI)	0.05 [-0.01, 0.11]	
58.1 Omega-3 supplements only	5	699	Mean Difference (IV, Random, 95% CI)	0.04 [-0.04, 0.11]	
58.2 Omega-3 supplements + food/ diet advice	1	188	Mean Difference (IV, Random, 95% CI)	0.08 [0.01, 0.15]	

Analysis 2.1. Comparison 2 Type of omega-3 intervention, Outcome 1 Preterm birth (< 37 weeks).

Study or subgroup	Omega 3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.1.1 Omega-3 supplements only					
Bisgaard 2016	15/365	18/371		2.59%	0.85[0.43,1.65]
Bulstra-Ramakers 1994	8/32	10/31	+	1.47%	0.78[0.35,1.7]
Carlson 2013	12/154	13/147	+	1.93%	0.88[0.42,1.87]
Dilli 2018	6/52	9/68		1.13%	0.87[0.33,2.29]
Helland 2001	1/301	2/289	←	0.3%	0.48[0.04,5.27]
Horvaticek 2017	4/51	4/47	← →	0.6%	0.92[0.24,3.48]
Makrides 2010	67/1197	88/1202		12.73%	0.76[0.56,1.04]
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	







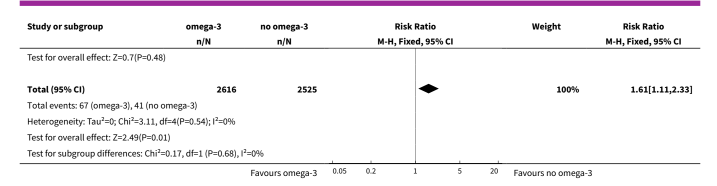
Analysis 2.2. Comparison 2 Type of omega-3 intervention, Outcome 2 Early preterm birth (< 34 weeks).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI	
2.2.1 Omega-3 supplements only						
Bulstra-Ramakers 1994	3/32	6/31	+	4.92%	0.48[0.13,1.77]	
Carlson 2013	1/154	7/147	+	5.78%	0.14[0.02,1.09]	
Harris 2015	4/224	7/121		7.34%	0.31[0.09,1.03]	
Horvaticek 2017	1/51	0/47		0.42%	2.77[0.12,66.36]	
Makrides 2010	13/1197	27/1202		21.74%	0.48[0.25,0.93]	
Min 2014	4/60	4/57		3.31%	0.95[0.25,3.62]	
Min 2016	2/58	0/56		0.41%	4.83[0.24,98.44]	
Olsen 2000	42/394	60/403		47.88%	0.72[0.5,1.04]	
Subtotal (95% CI)	2170	2064	◆	91.8%	0.62[0.46,0.82]	
Total events: 70 (Omega-3), 111 (No	omega-3)		İ			
Heterogeneity: Tau ² =0; Chi ² =7.62, d	f=7(P=0.37); I ² =8.12%	1	İ			
Test for overall effect: Z=3.35(P=0)						
2.2.2 Omega-3 supplements + other	er agents					
Mardones 2008	2/493	10/477		8.2%	0.19[0.04,0.88]	
Subtotal (95% CI)	493	477		8.2%	0.19[0.04,0.88]	
Total events: 2 (Omega-3), 10 (No or	nega-3)					
Heterogeneity: Not applicable						
Test for overall effect: Z=2.13(P=0.03	3)					
Total (95% CI)	2663	2541	•	100%	0.58[0.44,0.77]	
Total events: 72 (Omega-3), 121 (No	omega-3)					
Heterogeneity: Tau ² =0; Chi ² =9.89, d	f=8(P=0.27); I ² =19.129	%				
Test for overall effect: Z=3.84(P=0)						
Test for subgroup differences: Chi ² =	2.17. df=1 (P=0.14). I ²	=53.96%				

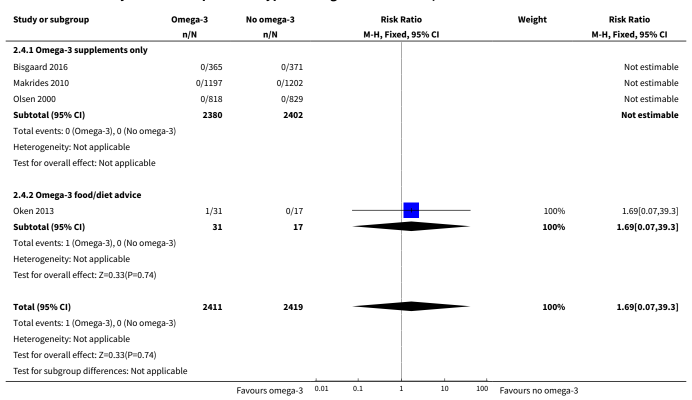
Analysis 2.3. Comparison 2 Type of omega-3 intervention, Outcome 3 Prolonged gestation (> 42 weeks).

Study or subgroup	omega-3	no omega-3	Risk Ratio	Weight	Risk Ratio
	n/N n/N		M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.3.1 Omega-3 supplements only					
Harris 2015	2/224	0/121		1.54%	2.71[0.13,56.02]
Makrides 2010	6/1184	3/1183		7.14%	2[0.5,7.97]
Mulder 2014	0/68	0/67			Not estimable
Olsen 1992	32/266	27/267	— <mark>—</mark>	64.13%	1.19[0.73,1.93]
Olsen 2000	26/782	11/791		26.02%	2.39[1.19,4.8]
Subtotal (95% CI)	2524	2429	•	98.84%	1.59[1.09,2.31]
Total events: 66 (omega-3), 41 (no om	ega-3)				
Heterogeneity: Tau ² =0; Chi ² =2.92, df=3	3(P=0.4); I ² =0%				
Test for overall effect: Z=2.42(P=0.02)					
2.3.2 Omega-3 supplements + food/	diet advice				
Hauner 2012	1/92	0/96		1.16%	3.13[0.13,75.84]
Subtotal (95% CI)	92	96		1.16%	3.13[0.13,75.84]
Total events: 1 (omega-3), 0 (no omeg	a-3)				
Heterogeneity: Not applicable					
		Favours omega-3 0.0	5 0.2 1 5 2	⁰ Favours no omega-3	





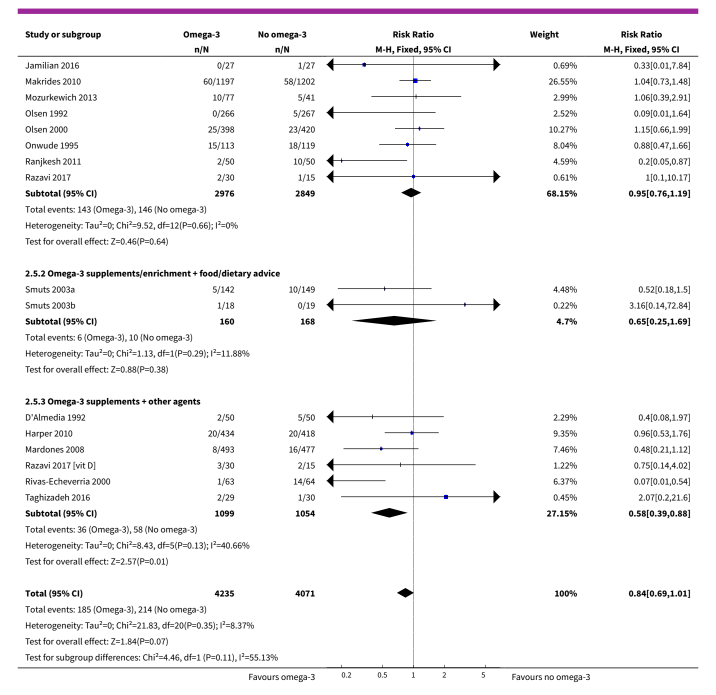
Analysis 2.4. Comparison 2 Type of omega-3 intervention, Outcome 4 Maternal death.



Analysis 2.5. Comparison 2 Type of omega-3 intervention, Outcome 5 Pre-eclampsia (hypertension with proteinuria).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.5.1 Omega-3 supplements only					
Bisgaard 2016	16/365	15/371		6.82%	1.08[0.54,2.16]
Bulstra-Ramakers 1994	5/32	3/31	+	1.4%	1.61[0.42,6.19]
Carlson 2013	2/154	2/147	+ • • • • • • • • • • • • • • • • • • •	0.94%	0.95[0.14,6.69]
Harris 2015	2/224	0/121	←	0.3%	2.71[0.13,56.02]
Horvaticek 2017	4/43	5/38		2.44%	0.71[0.2,2.44]
		Favours omega-3	0.2 0.5 1 2 5 F	avours no omega-3	

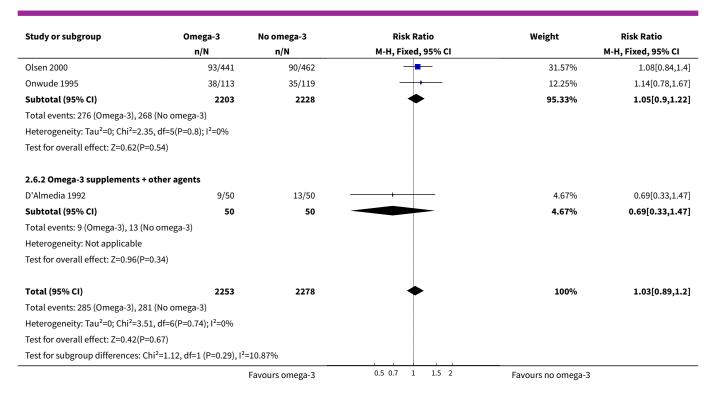




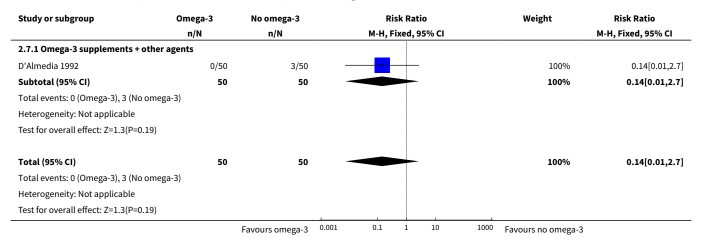
Analysis 2.6. Comparison 2 Type of omega-3 intervention, Outcome 6 High blood pressure (without proteinuria).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI	
2.6.1 Omega-3 supplements only						
Bulstra-Ramakers 1994	7/32	4/31	+	1.46%	1.7[0.55,5.22]	
Carlson 2013	32/154	25/147		9.19%	1.22[0.76,1.96]	
Makrides 2010	98/1197	107/1202		38.35%	0.92[0.71,1.2]	
Olsen 1992	8/266	7/267		2.51%	1.15[0.42,3.12]	
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3		





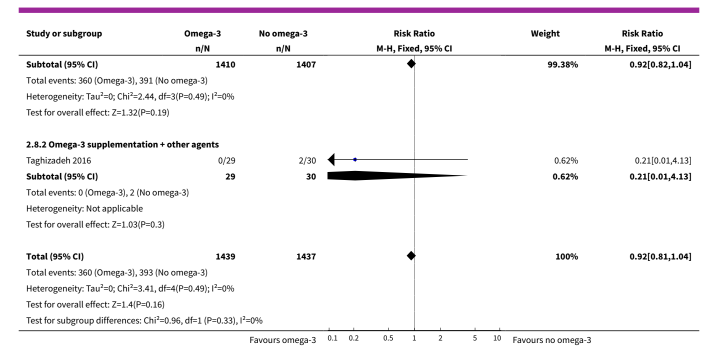
Analysis 2.7. Comparison 2 Type of omega-3 intervention, Outcome 7 Eclampsia.



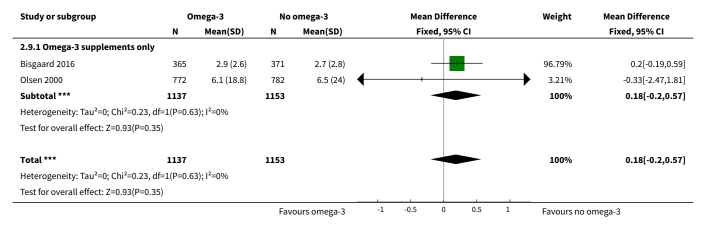
Analysis 2.8. Comparison 2 Type of omega-3 intervention, Outcome 8 Maternal antepartum hospitalisation.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.8.1 Omega-3 supplements only					
Bulstra-Ramakers 1994	14/32	11/31		2.84%	1.23[0.67,2.28]
Carlson 2013	14/154	15/147		3.9%	0.89[0.45,1.78]
Jamilian 2016	0/27	3/27	4	0.89%	0.14[0.01,2.64]
Makrides 2010	332/1197	362/1202	<u> </u>	91.75%	0.92[0.81,1.04]
		Favours omega-3	0.1 0.2 0.5 1 2	5 10 Favours no omega-3	

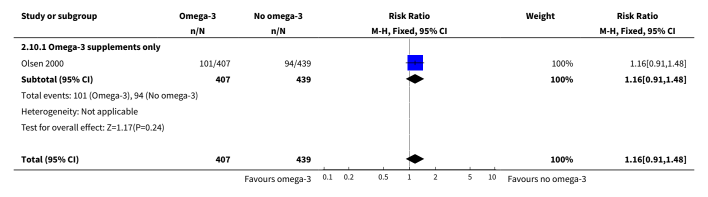




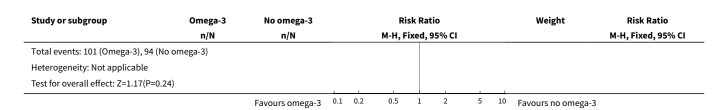
Analysis 2.9. Comparison 2 Type of omega-3 intervention, Outcome 9 Mother's length of stay in hospital (days).



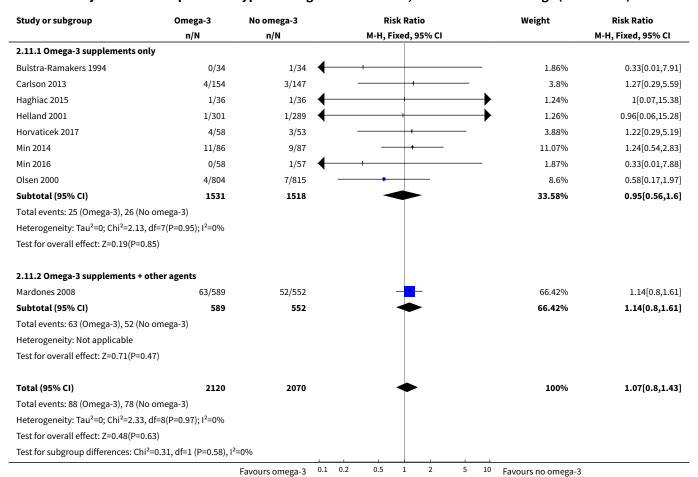
Analysis 2.10. Comparison 2 Type of omega-3 intervention, Outcome 10 Maternal anaemia.







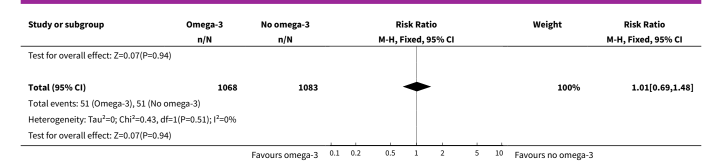
Analysis 2.11. Comparison 2 Type of omega-3 intervention, Outcome 11 Miscarriage (< 24 weeks).



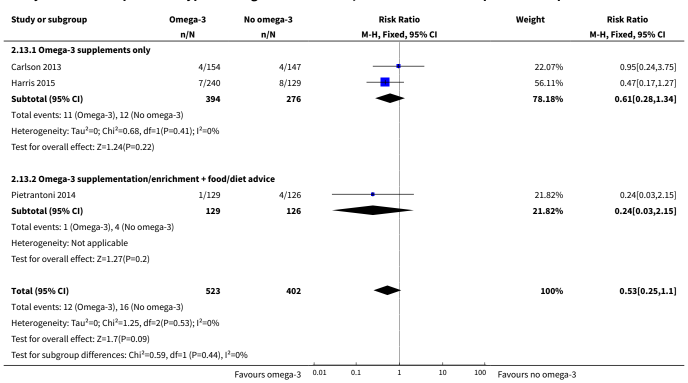
Analysis 2.12. Comparison 2 Type of omega-3 intervention, Outcome 12 Antepartum vaginal bleeding.

Study or subgroup	Omega-3	No omega-3			Ri	sk Rat	io			Weight	Risk Ratio
	n/N	n/N			M-H, F	ixed, 9	95% CI				M-H, Fixed, 95% CI
2.12.1 Omega-3 supplements	only										
Olsen 1992	15/266	12/267			_	+				23.65%	1.25[0.6,2.63]
Olsen 2000	36/802	39/816			_		-			76.35%	0.94[0.6,1.46]
Subtotal (95% CI)	1068	1083				*	-			100%	1.01[0.69,1.48]
Total events: 51 (Omega-3), 51	(No omega-3)										
Heterogeneity: Tau ² =0; Chi ² =0.	43, df=1(P=0.51); I ² =0%										
		Favours omega-3	0.1	0.2	0.5	1	2	5	10	Favours no omega-3	





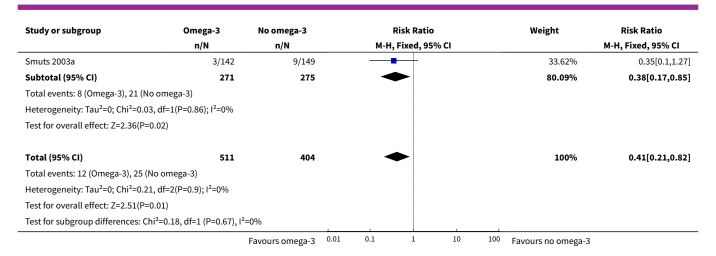
Analysis 2.13. Comparison 2 Type of omega-3 intervention, Outcome 13 Preterm prelabour rupture of membranes.



Analysis 2.14. Comparison 2 Type of omega-3 intervention, Outcome 14 Prelabour rupture of membranes.

Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio	
n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI	
,					
4/240	4/129		19.91%	0.54[0.14,2.11]	
240	129		19.91%	0.54[0.14,2.11]	
iega-3)					
7)					
/enrichment + food/	diet advice				
5/129	12/126		46.47%	0.41[0.15,1.12]	
	Favours omega-3 0.01	0.1 1 10	100 Favours no omega-3		
	n/N / 4/240 240 nega-3) //enrichment + food/	n/N n/N 4/240 4/129 240 129 nega-3) 7) /enrichment + food/diet advice 5/129 12/126	n/N n/N M-H, Fixed, 95% CI 4/240 4/129 240 129 nega-3) // /enrichment + food/diet advice 5/129 12/126	n/N n/N M-H, Fixed, 95% CI 4/240 4/129 19.91% 240 129 19.91% nega-3) // //enrichment + food/diet advice 5/129 12/126 46.47%	





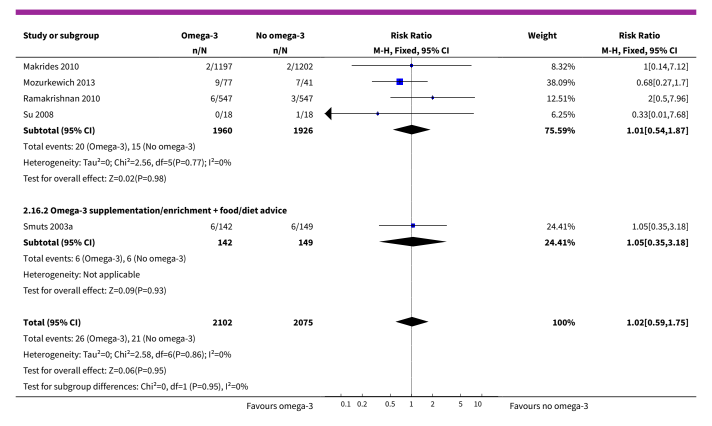
Analysis 2.15. Comparison 2 Type of omega-3 intervention, Outcome 15 Maternal admission to intensive care.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI	
2.15.1 Omega-3 supplements only						
Makrides 2010	2/1197	2/1202		44.8%	1[0.14,7.12]	
Subtotal (95% CI)	1197	1202		44.8%	1[0.14,7.12]	
Total events: 2 (Omega-3), 2 (No ome	ga-3)					
Heterogeneity: Not applicable						
Test for overall effect: Z=0(P=1)						
2.15.2 Omega-3 supplements + other	er agent					
Taghizadeh 2016	0/29	2/30 —		55.2%	0.21[0.01,4.13]	
Subtotal (95% CI)	29	30 —		55.2%	0.21[0.01,4.13	
Total events: 0 (Omega-3), 2 (No ome	ga-3)					
Heterogeneity: Not applicable						
Test for overall effect: Z=1.03(P=0.3)						
Total (95% CI)	1226	1232		100%	0.56[0.12,2.63]	
Total events: 2 (Omega-3), 4 (No ome	ga-3)					
Heterogeneity: Tau ² =0; Chi ² =0.77, df=	=1(P=0.38); I ² =0%					
Test for overall effect: Z=0.73(P=0.47)						
Test for subgroup differences: Chi ² =0.	.75, df=1 (P=0.39), I ²	=0%				
		Favours omega-3 0.01	0.1 1 10	100 Favours no omega-3		

Analysis 2.16. Comparison 2 Type of omega-3 intervention, Outcome 16 Maternal severe adverse effects (including cessation).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.16.1 Omega-3 supplements only					
Bulstra-Ramakers 1994	2/34	2/34		8.34%	1[0.15,6.7]
Freeman 2008	0/12	0/9			Not estimable
Khalili 2016	1/75	0/75		2.08%	3[0.12,72.49]
		Favours omega-3	0.1 0.2 0.5 1 2 5 10	Favours no omega-3	

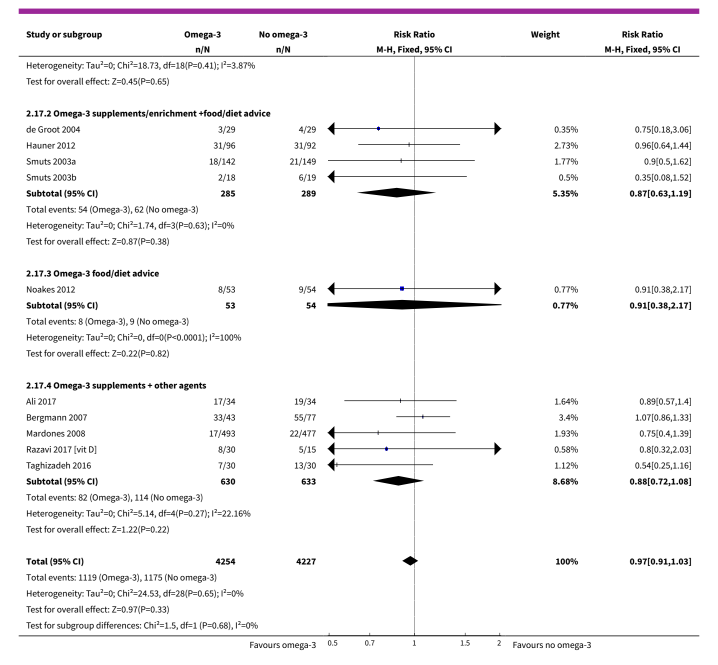




Analysis 2.17. Comparison 2 Type of omega-3 intervention, Outcome 17 Caesarean section.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.17.1 Omega-3 supplements only					
Bisgaard 2016	77/365	71/371		6.08%	1.1[0.83,1.47]
Bulstra-Ramakers 1994	8/32	10/31	—	0.88%	0.78[0.35,1.7]
Carlson 2013	46/154	44/147		3.89%	1[0.71,1.41]
Dilli 2018	34/52	54/68		4.04%	0.82[0.65,1.04]
Dunstan 2008	11/40	8/43	-	0.67%	1.48[0.66,3.3]
Helland 2001	28/175	14/166		1.24%	1.9[1.04,3.48]
Jamilian 2016	12/26	18/27		1.53%	0.69[0.42,1.13]
Judge 2007	8/27	5/21	+ + + +	0.49%	1.24[0.48,3.25]
Khalili 2016	30/75	33/75		2.85%	0.91[0.62,1.33]
Makrides 2010	326/1197	350/1202		30.17%	0.94[0.82,1.06]
Miller 2016	20/60	12/55		1.08%	1.53[0.83,2.83]
Min 2014	25/60	24/57		2.13%	0.99[0.65,1.52]
Min 2016	28/58	29/56		2.55%	0.93[0.65,1.35]
Mozurkewich 2013	22/77	11/41		1.24%	1.06[0.57,1.97]
Olsen 1992	16/266	20/267	+	1.72%	0.8[0.43,1.52]
Onwude 1995	36/113	25/119	+	2.1%	1.52[0.98,2.36]
Ramakrishnan 2010	216/429	234/440		19.96%	0.95[0.83,1.08]
Ranjkesh 2011	23/50	22/50		1.9%	1.05[0.68,1.61]
Razavi 2017	9/30	6/15	+	0.69%	0.75[0.33,1.71]
Subtotal (95% CI)	3286	3251	*	85.2%	0.98[0.92,1.06]
Total events: 975 (Omega-3), 990 (No	omega-3)				
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	}

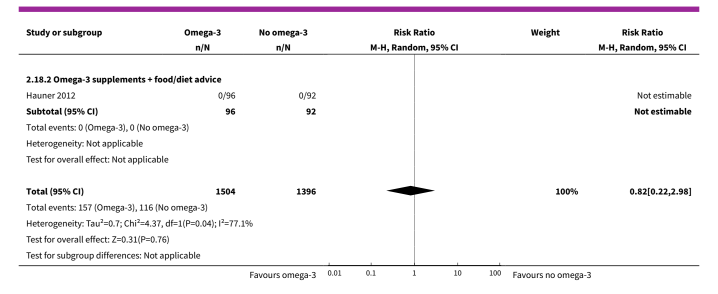




Analysis 2.18. Comparison 2 Type of omega-3 intervention, Outcome 18 Induction (post-term).

Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		М-Н, І	Random, 95%	CI			M-H, Random, 95% CI
2.18.1 Omega-3 supplements or	nly								
Harris 2015	4/224	6/121			-			39.31%	0.36[0.1,1.25]
Makrides 2010	153/1184	110/1183			-			60.69%	1.39[1.1,1.75]
Subtotal (95% CI)	1408	1304		-				100%	0.82[0.22,2.98]
Total events: 157 (Omega-3), 116	(No omega-3)								
Heterogeneity: Tau ² =0.7; Chi ² =4.3	37, df=1(P=0.04); I ² =77.1	%							
Test for overall effect: Z=0.31(P=0	0.76)								
		Favours omega-3	0.01	0.1	i	10	100	Favours no omega-3	





Analysis 2.19. Comparison 2 Type of omega-3 intervention, Outcome 19 Blood loss at birth (mL).

Study or subgroup	0	mega-3	No	omega-3		Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fixed, 95% CI		Fixed, 95% CI
2.19.1 Omega-3 supplements on	ly							
Furuhjelm 2009	54	473 (232)	66	492 (232)	•	+	4.79%	-19[-102.44,64.44]
Helland 2001	175	362 (219)	166	354 (324)			9.57%	8[-51.01,67.01]
Mozurkewich 2013	77	507.5 (411.5)	41	454 (296)		+	2%	53.49[-75.57,182.55]
Olsen 1992	266	316 (260)	267	290 (213)			20.46%	26[-14.36,66.36]
Olsen 2000	725	351.7 (282.7)	751	344.7 (267.1)			42.27%	7[-21.08,35.08]
Subtotal ***	1297		1291				79.08%	11.64[-8.89,32.17]
Heterogeneity: Tau ² =0; Chi ² =1.53,	df=4(P=0.8	2); I ² =0%						
Test for overall effect: Z=1.11(P=0.2	27)							
2.19.2 Omega-3 supplements + fo	ood/diet a	dvice						
Hauner 2012	92	377 (153)	96	366 (124)			20.92%	11[-28.91,50.91]
Subtotal ***	92		96				20.92%	11[-28.91,50.91]
Heterogeneity: Not applicable								
Test for overall effect: Z=0.54(P=0.5	59)							
Total ***	1389		1387			•	100%	11.5[-6.75,29.76]
Heterogeneity: Tau ² =0; Chi ² =1.53,	df=5(P=0.9	1); I ² =0%						
Test for overall effect: Z=1.24(P=0.2	22)							
Test for subgroup differences: Chi ²	=0, df=1 (P	=0.98), I ² =0%						
			Fav	ours omega-3	-100	-50 0 50	100 Favours no	omega-3



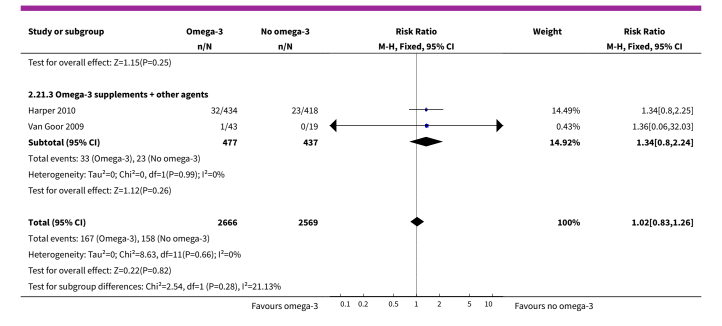
Analysis 2.20. Comparison 2 Type of omega-3 intervention, Outcome 20 Postpartum haemorrhage.

n/N 6/154	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
				,,
6/154				
	3/147		2.42%	1.91[0.49,7.49]
57/1197	64/1202		50.33%	0.89[0.63,1.27]
9/266	7/267		5.51%	1.29[0.49,3.41]
1617	1616	•	58.25%	0.97[0.71,1.34]
omega-3)				
=2(P=0.48); I ² =0%				
")				
ner agent				
60/434	52/418	—	41.75%	1.11[0.79,1.57]
434	418	*	41.75%	1.11[0.79,1.57]
omega-3)				
2051	2034	•	100%	1.03[0.82,1.3]
o omega-3)				
f=3(P=0.61); I ² =0%				
0.3, df=1 (P=0.58), I ² =	0%			
f	1617 omega-3) f=2(P=0.48); l ² =0% 7) her agent 60/434 434 omega-3) 2051 o omega-3) f=3(P=0.61); l ² =0%	1617 1616 pmega-3) f=2(P=0.48); l²=0% 7) her agent 60/434 52/418 434 418 pmega-3) 2051 2034 o omega-3) f=3(P=0.61); l²=0% 0.3, df=1 (P=0.58), l²=0%	1617 1616 pmega-3) f=2(P=0.48); I²=0% 7) her agent 60/434 52/418 434 418 pmega-3) 2051 2034 o omega-3) f=3(P=0.61); I²=0% 0.3, df=1 (P=0.58), I²=0%	1617 1616 58.25% comega-3) f=2(P=0.48); l²=0% 7) her agent 60/434 52/418 41.75% 434 418 comega-3) 2051 2034 co omega-3) f=3(P=0.61); l²=0% 0.3, df=1 (P=0.58), l²=0%

Analysis 2.21. Comparison 2 Type of omega-3 intervention, Outcome 21 Gestational diabetes.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.21.1 Omega-3 supplements	only				
Bisgaard 2016	6/365	10/371		6.13%	0.61[0.22,1.66]
Carlson 2013	9/154	6/147	+	3.8%	1.43[0.52,3.92]
Haghiac 2015	1/25	0/25		0.31%	3[0.13,70.3]
Makrides 2010	96/1197	100/1202	-	61.72%	0.96[0.74,1.26]
Min 2014	3/32	0/27	-	0.33%	5.94[0.32,110.13]
Mozurkewich 2013	8/77	2/41	+	1.61%	2.13[0.47,9.57]
Van Goor 2009	0/43	0/20			Not estimable
Subtotal (95% CI)	1893	1833	*	73.91%	1.02[0.8,1.3]
Total events: 123 (Omega-3), 11	L8 (No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =4.	38, df=5(P=0.5); I ² =0%				
Test for overall effect: Z=0.12(P	=0.9)				
2.21.2 Omega-3 supplements	/enrichment + food/diet	advice			
de Groot 2004	0/40	1/39		0.94%	0.33[0.01,7.75]
Hauner 2012	7/96	10/92		6.32%	0.67[0.27,1.69]
Smuts 2003a	4/142	3/149		1.81%	1.4[0.32,6.14]
Smuts 2003b	0/18	3/19	+	2.11%	0.15[0.01,2.72]
Subtotal (95% CI)	296	299		11.18%	0.66[0.33,1.34]
Total events: 11 (Omega-3), 17	(No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =2.	18, df=3(P=0.54); I ² =0%				





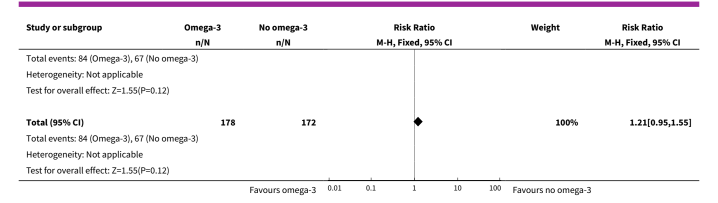
Analysis 2.22. Comparison 2 Type of omega-3 intervention, Outcome 22 Maternal insulin resistance (HOMA-IR).

Study or subgroup	0	mega-3	No	omega-3	Mea	an Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Rar	ndom, 95% CI		Random, 95% CI
2.22.1 Omega-3 supplements only	у							
Krummel 2016	32	3.7 (2.1)	28	3.2 (1)		 ■	35.64%	0.53[-0.29,1.35]
Samimi 2015	28	3.1 (1.5)	28	4.3 (3.3)		- 	31.05%	-1.2[-2.54,0.14]
Subtotal ***	60		56		-		66.69%	-0.25[-1.94,1.44]
Heterogeneity: Tau ² =1.17; Chi ² =4.6	5, df=1(P=	0.03); I ² =78.51%						
Test for overall effect: Z=0.29(P=0.7	7)							
2.22.2 Omega-3 supplements + ot	her agen	ts						
Taghizadeh 2016	30	2.5 (1)	30	4.5 (2.9)		_	33.31%	-2[-3.1,-0.9]
Subtotal ***	30		30		•	>	33.31%	-2[-3.1,-0.9]
Heterogeneity: Not applicable								
Test for overall effect: Z=3.57(P=0)								
Total ***	90		86		•		100%	-0.85[-2.5,0.8]
Heterogeneity: Tau ² =1.82; Chi ² =14.	31, df=2(P	=0); I ² =86.02%						
Test for overall effect: Z=1.01(P=0.3	1)							
Test for subgroup differences: Chi ²		L (P=0.09), I ² =65.	58%					
			Fav	ours omega-3	-5 -2.5	0 2.5	5 Favours no	omega-3

Analysis 2.23. Comparison 2 Type of omega-3 intervention, Outcome 23 Excessive gestational weight gain.

Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		М-Н	, Fixed, 95%	% CI			M-H, Fixed, 95% CI
2.23.1 Omega-3 supplements only									
Carlson 2013	84/178	67/172			+			100%	1.21[0.95,1.55]
Subtotal (95% CI)	178	172			◆			100%	1.21[0.95,1.55]
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3	



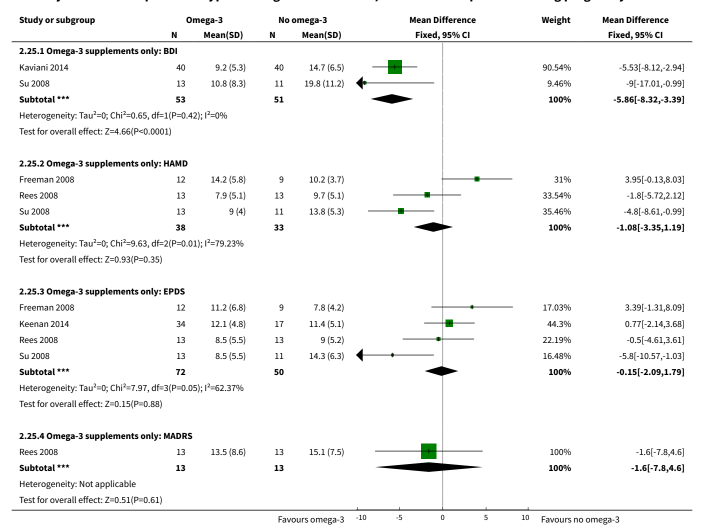


Analysis 2.24. Comparison 2 Type of omega-3 intervention, Outcome 24 Gestational weight gain (kg).

Study or subgroup	0	mega-3	No	omega-3		Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Random, 95% CI		Random, 95% CI
2.24.1 Omega-3 supplemer	its only							
Carlson 2013	178	12.5 (6)	172	11.6 (5.5)		+	10.65%	0.9[-0.31,2.11]
Dilli 2018	52	9.5 (3.6)	68	12.5 (5.8)	◀		7.75%	-3[-4.69,-1.31]
Furuhjelm 2009	54	13 (3.9)	66	14 (4.9)	+		8.36%	-1[-2.57,0.57]
Horvaticek 2017	47	13.9 (5.1)	43	12.5 (3.9)			6.9%	1.4[-0.47,3.27]
Krummel 2016	32	4.9 (3.9)	28	3.8 (3.7)		+	6.67%	1.07[-0.85,2.99]
Mulder 2014	104	14.1 (4.8)	111	14.7 (4.5)	\leftarrow	+	10.37%	-0.6[-1.85,0.65]
Subtotal ***	467		488				50.7%	-0.22[-1.47,1.03]
Heterogeneity: Tau ² =1.78; Cl	ni²=19.62, df=5(P	=0); I ² =74.51%						
Test for overall effect: Z=0.35	5(P=0.73)							
2.24.2 Omega-3 supplemer	nts/enrichment -	+ food/diet advi	ice					
Hauner 2012	91	15.1 (4.8)	95	16 (5.1)	←	<u> </u>	9.24%	-0.9[-2.32,0.52]
Hurtado 2015	44	7.4 (2.8)	46	7 (2.8)		+	10.98%	0.4[-0.76,1.56]
Smuts 2003b	18	15.3 (7.9)	19	15.1 (6.6)	←	+	1.64%	0.2[-4.5,4.9]
Subtotal ***	153		160		-		21.86%	-0.11[-0.99,0.78]
Heterogeneity: Tau ² =0; Chi ² =	1.95, df=2(P=0.3	8); I ² =0%						
Test for overall effect: Z=0.24	(P=0.81)							
2.24.3 Omega-3 supplemen	its + other agent	ts						
Mardones 2008	493	14.5 (4.8)	477	14 (5)		+	14.93%	0.53[-0.08,1.14]
Taghizadeh 2016	29	13.1 (1.7)	30	12.9 (2)		+	12.51%	0.2[-0.75,1.15]
Subtotal ***	522		507				27.44%	0.43[-0.08,0.95]
Heterogeneity: Tau ² =0; Chi ² =	0.33, df=1(P=0.5	7); I ² =0%						
Test for overall effect: Z=1.64	P(P=0.1)							
Total ***	1142		1155				100%	-0.05[-0.68,0.59]
Heterogeneity: Tau ² =0.6; Chi	² =24.6, df=10(P=	0.01); I ² =59.35%						
Test for overall effect: Z=0.14	(P=0.89)							
Test for subgroup difference	s: Chi²=1.65, df=1	(P=0.44), I ² =0%)					



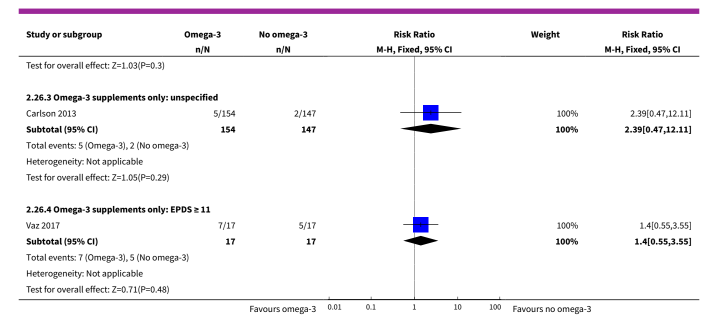
Analysis 2.25. Comparison 2 Type of omega-3 intervention, Outcome 25 Depression during pregnancy: scores.



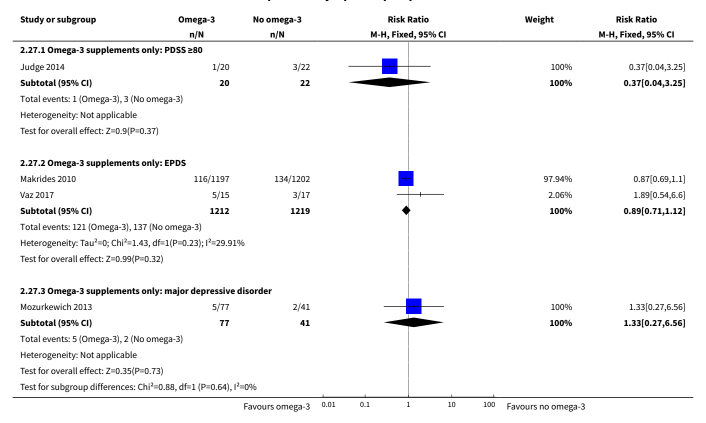
Analysis 2.26. Comparison 2 Type of omega-3 intervention, Outcome 26 Depression during pregnancy: thresholds.

Study or subgroup	idy or subgroup Omega-3 No omej n/N n/N				Risk Ratio			Weight	Risk Ratio
				M-H, Fixed, 95% CI					M-H, Fixed, 95% CI
2.26.1 Omega-3 supplements only: HA weeks)	MD 50% reduct	ion (after 8							
Su 2008	8/13	3/11			+	_		100%	2.26[0.78,6.49]
Subtotal (95% CI)	13	11				-		100%	2.26[0.78,6.49]
Total events: 8 (Omega-3), 3 (No omega-	3)								
Heterogeneity: Not applicable									
Test for overall effect: Z=1.51(P=0.13)									
2.26.2 Omega-3 supplements only: HA	.MD ≤ 7								
Su 2008	5/13	2/11			-			100%	2.12[0.51,8.84]
Subtotal (95% CI)	13	11				-		100%	2.12[0.51,8.84]
Total events: 5 (Omega-3), 2 (No omega-	3)								
Heterogeneity: Not applicable									
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3	





Analysis 2.27. Comparison 2 Type of omega-3 intervention, Outcome 27 Depressive symptoms postpartum: thresholds.





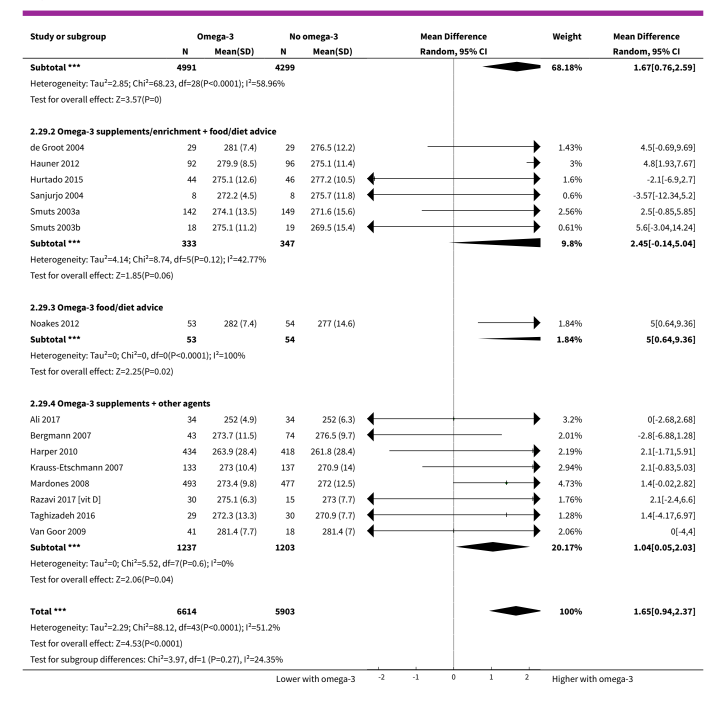
Analysis 2.28. Comparison 2 Type of omega-3 intervention, Outcome 28 Depressive symptoms postpartum: scores.

Study or subgroup	Oı	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
2.28.1 Omega-3 supplements onl	y: BD: 6-8	weeks postpar	tum				
Mozurkewich 2013	77	6.2 (5)	41	5.9 (6.1)		100%	0.25[-1.93,2.43
Subtotal ***	77		41			100%	0.25[-1.93,2.43
Heterogeneity: Not applicable							
Test for overall effect: Z=0.23(P=0.8	32)						
2.28.2 Omega-3 supplements onl	y: PDSS to	otal (LS over 6 m	onths)				
Judge 2014	20	46 (9.7)	22	52.1 (11.3)	 	100%	-6.08[-12.42,0.26
Subtotal ***	20		22			100%	-6.08[-12.42,0.26
Heterogeneity: Tau ² =0; Chi ² =0, df=0	0(P<0.0001); I ² =100%					
Test for overall effect: Z=1.88(P=0.0	06)						
Test for subgroup differences: Chi ² :	=3.43, df=1	(P=0.06), I ² =70.	81%				
			Fav	ours omega-3 -1	0 -5 0 5	10 Favours no	omega-3

Analysis 2.29. Comparison 2 Type of omega-3 intervention, Outcome 29 Length of gestation (days).

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
2.29.1 Omega-3 supplement	s only						
Carlson 2013	154	275.7 (11.2)	147	272.8 (17)		2.62%	2.9[-0.37,6.17]
Dilli 2018	52	266 (12.6)	68	261.8 (14)		1.62%	4.2[-0.58,8.98]
Dunstan 2008	40	275 (6.3)	43	274 (6.6)		3.1%	1[-1.77,3.77]
Furuhjelm 2009	54	280 (9.8)	66	280 (11.2)		2.23%	0[-3.76,3.76]
Giorlandino 2013	21	268.8 (9.1)	21	255.5 (12.6)		0.96%	13.3[6.65,19.95]
Gustafson 2013	22	275.8 (7.7)	24	279.3 (7.7)		1.79%	-3.5[-7.95,0.95]
Haghiac 2015	25	274.4 (11.2)	24	270.9 (8.4)		1.3%	3.5[-2.03,9.03]
Harris 2015	224	275.3 (19.5)	121	271.6 (13.2)		2.46%	3.7[0.23,7.17]
Helland 2001	175	279.6 (9.2)	166	279.2 (9.3)		4.04%	0.4[-1.56,2.36]
Horvaticek 2017	47	271.6 (4.6)	43	266 (4.5)		4.14%	5.6[3.72,7.48]
Jamilian 2016	26	270.2 (7.1)	27	269.5 (10.4)	+	1.62%	0.7[-4.08,5.48]
Judge 2007	27	278 (8.4)	21	274.3 (9)		1.52%	3.71[-1.27,8.69]
Keenan 2014	36	272 (14)	17	271.9 (14.4)	•	0.67%	0.07[-8.17,8.31]
Khalili 2016	75	275.1 (8.4)	75	274.4 (9.1)	+	3.07%	0.7[-2.1,3.5]
Knudsen 2006	1250	280.7 (12.9)	748	280.6 (11.7)		5.11%	0.1[-1,1.2]
Krummel 2016	34	275.1 (7.7)	29	275.8 (8.4)	+	2.06%	-0.7[-4.71,3.31]
Makrides 2010	1184	276.2 (12.4)	1183	274.6 (14.8)		5.11%	1.64[0.54,2.74]
Malcolm 2003	31	279.7 (9.5)	29	279.6 (8.5)	+	1.73%	0.1[-4.46,4.66]
Miller 2016	60	278.9 (7.8)	55	274.4 (14.8)		1.83%	4.5[0.12,8.88]
Mozurkewich 2013	77	278.2 (8.7)	41	273.7 (10.5)		2.23%	4.48[0.73,8.23]
Olsen 1992	266	283.3 (11.1)	267	280.6 (12.3)		4.01%	2.7[0.71,4.69]
Olsen 2000	108	269.2 (19.7)	120	260.7 (29.5)		1.01%	8.5[2.05,14.95]
Olsen 2000 [twins]	286	254.5 (24)	283	254.4 (23.1)	+	2.15%	0.1[-3.77,3.97]
Ramakrishnan 2010	484	273.7 (13.3)	486	273 (11.9)		4.52%	0.7[-0.89,2.29]
Razavi 2017	30	273 (9.8)	15	273.7 (7.7)	+ +	1.41%	-0.7[-5.94,4.54]
Tofail 2006	125	270.9 (18.2)	124	274.4 (17.5)	<u> </u>	1.8%	-3.5[-7.93,0.93]
Valenzuela 2015	19	270.9 (8.4)	21	270.2 (8.4)		1.42%	0.7[-4.51,5.91]
Van Goor 2009	42	281.4 (7.7)	18	281.4 (7)		2.07%	0[-3.98,3.98]
Vaz 2017	17	273 (13.3)	17	273.7 (12.6)	+	0.6%	-0.7[-9.41,8.01]

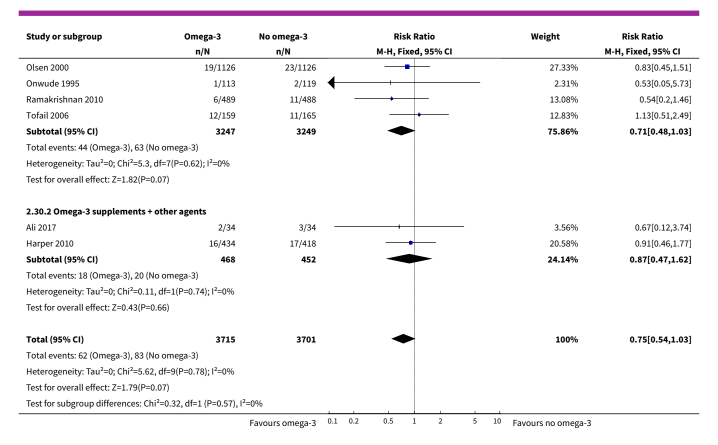




Analysis 2.30. Comparison 2 Type of omega-3 intervention, Outcome 30 Perinatal death.

Study or subgroup	Omega-3 No omega-3		Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI	
2.30.1 Omega-3 supplements only						
Bulstra-Ramakers 1994	2/32	3/31		3.62%	0.65[0.12,3.61]	
Horvaticek 2017	1/56	0/43	←	0.67%	2.32[0.1,55.48]	
Khalili 2016	0/75	1/75	+	1.78%	0.33[0.01,8.05]	
Makrides 2010	3/1197	12/1202	★	14.23%	0.25[0.07,0.89]	
		Favours omega-3	0.1 0.2 0.5 1 2 5 10	Favours no omega-3	 	

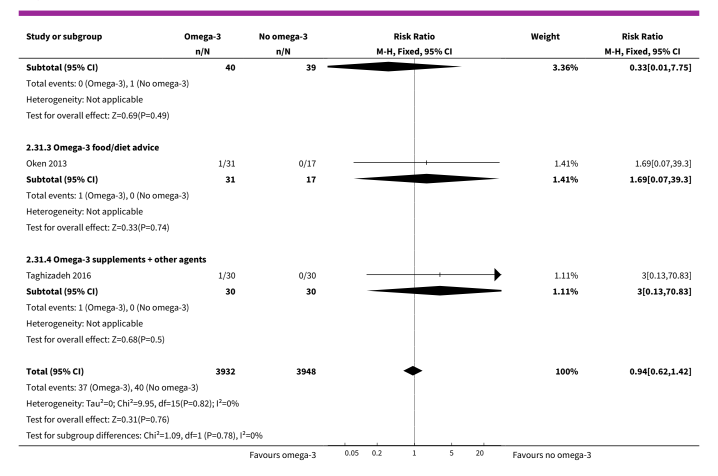




Analysis 2.31. Comparison 2 Type of omega-3 intervention, Outcome 31 Stillbirth.

Study or subgroup	Omega-3	No omega-3		Risk Ratio	Weight	Risk Ratio
	n/N	n/N		M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.31.1 Omega-3 supplements or	nly					
Bulstra-Ramakers 1994	1/32	0/31			1.12%	2.91[0.12,68.81]
Haghiac 2015	1/25	0/25			1.11%	3[0.13,70.3]
Helland 2001	1/301	0/289			1.13%	2.88[0.12,70.43]
Horvaticek 2017	1/48	0/50		+	1.08%	3.12[0.13,74.82]
Jamilian 2016	1/27	0/27			1.11%	3[0.13,70.53]
Makrides 2010	1/1197	7/1202	-	+	15.44%	0.14[0.02,1.16]
Min 2014	2/60	0/57			1.13%	4.75[0.23,96.93]
Min 2016	0/58	1/57	•		3.34%	0.33[0.01,7.88]
Olsen 1992	1/266	1/267			2.21%	1[0.06,15.96]
Olsen 2000	16/1056	19/1085			41.42%	0.87[0.45,1.67]
Onwude 1995	0/113	2/119	-		5.38%	0.21[0.01,4.34]
Ramakrishnan 2010	2/489	3/488			6.64%	0.67[0.11,3.96]
Tofail 2006	8/159	6/165			13.02%	1.38[0.49,3.9]
Subtotal (95% CI)	3831	3862		•	94.12%	0.92[0.6,1.42]
Total events: 35 (Omega-3), 39 (N	o omega-3)					
Heterogeneity: Tau ² =0; Chi ² =8.88	, df=12(P=0.71); I ² =0%					
Test for overall effect: Z=0.37(P=0	.71)					
2.31.2 Omega-3 supplements +	food/diet advice					
de Groot 2004	0/40	1/39	←	+	3.36%	0.33[0.01,7.75]
		Favours omega-3	0.05	0.2 1 5 2	Favours no omega-3	





Analysis 2.32. Comparison 2 Type of omega-3 intervention, Outcome 32 Neonatal death.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI	
2.32.1 Omega-3 supplements onl	у					
Bisgaard 2016	0/365	0/371			Not estimable	
Bulstra-Ramakers 1994	1/32	3/31		10.91%	0.32[0.04,2.94]	
Carlson 2013	1/154	1/147		3.66%	0.95[0.06,15.12]	
Khalili 2016	0/75	1/75 -		5.37%	0.33[0.01,8.05]	
Makrides 2010	2/1197	5/1202		17.86%	0.4[0.08,2.07]	
Olsen 2000	3/1126	4/1144	+ 	14.21%	0.76[0.17,3.4]	
Onwude 1995	1/113	0/119		1.74%	3.16[0.13,76.73]	
Ramakrishnan 2010	4/487	8/486		28.67%	0.5[0.15,1.65]	
Tofail 2006	4/159	5/165		17.57%	0.83[0.23,3.04]	
Subtotal (95% CI)	3708	3740	•	100%	0.61[0.34,1.11]	
Total events: 16 (Omega-3), 27 (No	omega-3)					
Heterogeneity: Tau ² =0; Chi ² =2.24, c	df=7(P=0.95); I ² =0%					
Test for overall effect: Z=1.61(P=0.1	11)					
Total (95% CI)	3708	3740	•	100%	0.61[0.34,1.11]	
Total events: 16 (Omega-3), 27 (No	omega-3)					
Heterogeneity: Tau ² =0; Chi ² =2.24, c	df=7(P=0.95); I ² =0%					
Test for overall effect: Z=1.61(P=0.1	11)					
		Favours omega-3 0.0	1 0.1 1 10	100 Favours no omega-3		



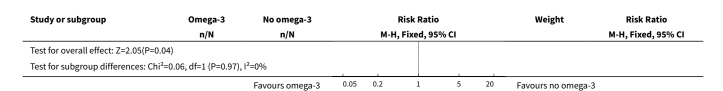
Analysis 2.33. Comparison 2 Type of omega-3 intervention, Outcome 33 Infant death.

Study or subgroup	Omega-3	No omega-3		Ri	sk Ratio		Weight	Risk Ratio
	n/N	n/N		М-Н, F	ixed, 95% CI			M-H, Fixed, 95% CI
2.33.1 Omega-3 supplements only								
Carlson 2013	0/154	1/147		•			20.44%	0.32[0.01,7.75]
Makrides 2010	1/1197	0/1202		-	+-		6.65%	3.01[0.12,73.88]
Mulder 2014	0/111	1/104		•			20.62%	0.31[0.01,7.59]
Tofail 2006	3/159	4/165			-		52.29%	0.78[0.18,3.42]
Subtotal (95% CI)	1621	1618		•			100%	0.74[0.25,2.19]
Total events: 4 (Omega-3), 6 (No ome	ga-3)							
Heterogeneity: Tau ² =0; Chi ² =1.29, df=	=3(P=0.73); I ² =0%							
Test for overall effect: Z=0.55(P=0.58)								
Total (95% CI)	1621	1618		•			100%	0.74[0.25,2.19]
Total events: 4 (Omega-3), 6 (No ome	ga-3)							
Heterogeneity: Tau ² =0; Chi ² =1.29, df=	=3(P=0.73); I ² =0%							
Test for overall effect: Z=0.55(P=0.58)								
		Favours omega-3	0.01	0.1	1 10	100	Favours no omega-3	

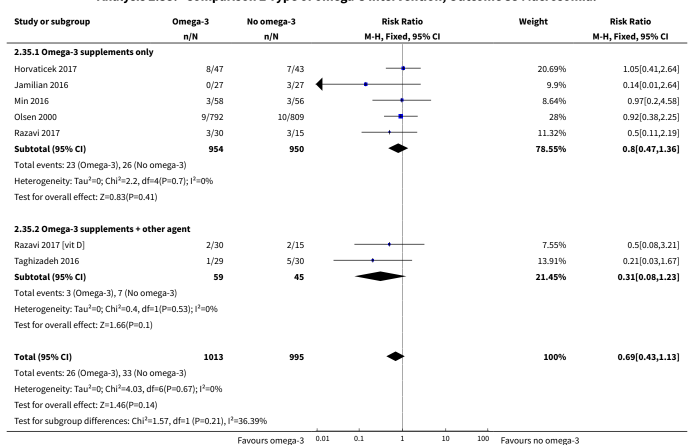
Analysis 2.34. Comparison 2 Type of omega-3 intervention, Outcome 34 Large-for-gestational age.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.34.1 Omega-3 supplements o	nly				
Makrides 2010	204/1197	173/1202		86.52%	1.18[0.98,1.43]
Min 2014	1/60	0/59		0.25%	2.95[0.12,71.01]
Subtotal (95% CI)	1257	1261	•	86.78%	1.19[0.99,1.43]
Total events: 205 (Omega-3), 173	(No omega-3)				
Heterogeneity: Tau²=0; Chi²=0.32	2, df=1(P=0.57); I ² =0%				
Test for overall effect: Z=1.83(P=0	0.07)				
2.34.2 Omega-3 supplements +	food/diet advice				
Hauner 2012	9/96	7/92		3.58%	1.23[0.48,3.17]
Subtotal (95% CI)	96	92		3.58%	1.23[0.48,3.17]
Total events: 9 (Omega-3), 7 (No	omega-3)				
Heterogeneity: Tau²=0; Chi²=0, d	f=0(P<0.0001); I ² =100%				
Test for overall effect: Z=0.43(P=0	0.67)				
2.34.3 Omega-3 supplements +	other agent				
Harper 2010	21/427	15/410	+	7.67%	1.34[0.7,2.57]
Taghizadeh 2016	4/29	4/30		1.97%	1.03[0.29,3.75]
Subtotal (95% CI)	456	440	*	9.64%	1.28[0.72,2.29]
Total events: 25 (Omega-3), 19 (N	lo omega-3)				
Heterogeneity: Tau²=0; Chi²=0.13	3, df=1(P=0.72); I ² =0%				
Test for overall effect: Z=0.84(P=0	0.4)				
Total (95% CI)	1809	1793	•	100%	1.2[1.01,1.43]
Total events: 239 (Omega-3), 199	(No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =0.5,	df=4(P=0.97)·12=0%				





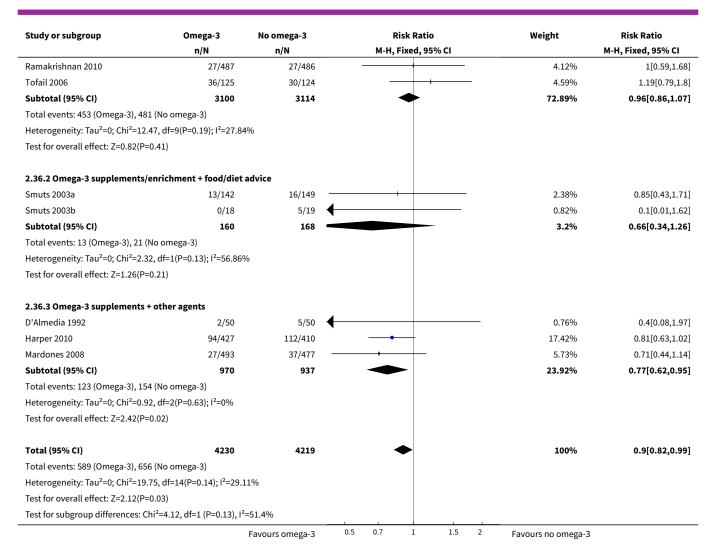
Analysis 2.35. Comparison 2 Type of omega-3 intervention, Outcome 35 Macrosomia.



Analysis 2.36. Comparison 2 Type of omega-3 intervention, Outcome 36 Low birthweight (< 2500 g).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.36.1 Omega-3 supplements only					
Bulstra-Ramakers 1994	11/32	9/31		1.39%	1.18[0.57,2.46]
Carlson 2013	6/154	13/147	+	2.03%	0.44[0.17,1.13]
Khalili 2016	0/75	5/75	←	0.84%	0.09[0.01,1.62]
Makrides 2010	41/1197	63/1202		9.58%	0.65[0.44,0.96]
Min 2014	8/60	8/57	+	1.25%	0.95[0.38,2.36]
Min 2016	8/58	4/56		0.62%	1.93[0.62,6.05]
Olsen 2000	283/799	287/817		43.26%	1.01[0.88,1.15]
Onwude 1995	33/113	35/119		5.2%	0.99[0.67,1.48]
		Favours omega-3	0.5 0.7 1 1.5 2 p	avours no omega-3	

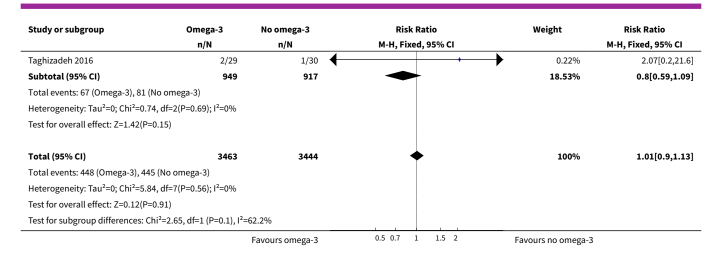




Analysis 2.37. Comparison 2 Type of omega-3 intervention, Outcome 37 Small-for-gestational age/IUGR.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.37.1 Omega-3 supplements on	ly				
Bulstra-Ramakers 1994	12/32	9/31		2.05%	1.29[0.64,2.63]
Makrides 2010	73/1197	82/1202	+-	18.39%	0.89[0.66,1.21]
Olsen 2000	208/685	185/689	 -	41.45%	1.13[0.96,1.34]
Onwude 1995	33/113	35/119		7.66%	0.99[0.67,1.48]
Ramakrishnan 2010	55/487	53/486		11.92%	1.04[0.73,1.48]
Subtotal (95% CI)	2514	2527	*	81.47%	1.05[0.93,1.2]
Total events: 381 (Omega-3), 364 (No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =2.21,	df=4(P=0.7); I ² =0%				
Test for overall effect: Z=0.82(P=0.	41)				
2.37.2 Omega-3 supplements + o	other agents				
Harper 2010	35/427	41/410		9.4%	0.82[0.53,1.26]
Mardones 2008	30/493	39/477		8.91%	0.74[0.47,1.18]
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	

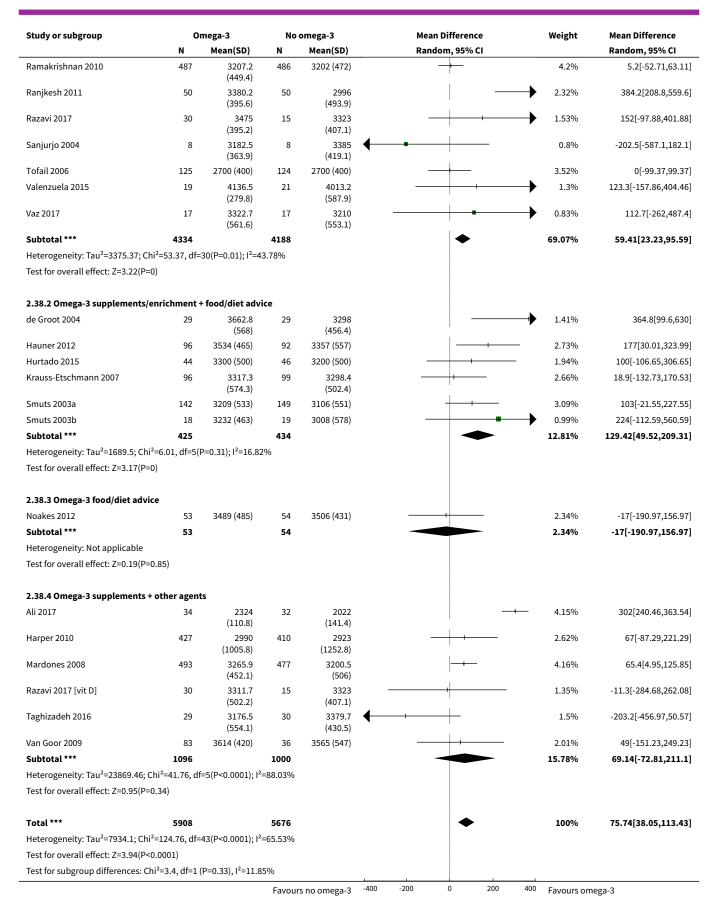




Analysis 2.38. Comparison 2 Type of omega-3 intervention, Outcome 38 Birthweight (g).

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference Random, 95% CI
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		
2.38.1 Omega-3 supplements or	ıly						
Carlson 2013	154	3359 (524)	147	3187 (602)		3.04%	172[44.25,299.75]
Dilli 2018	52	3288 (641)	68	3538 (671)		1.65%	-250[-486.2,-13.8]
Dunstan 2008	40	3503.4 (337.7)	43	3430.1 (371.8)		2.64%	73.3[-79.36,225.96]
England 1989	17	2200 (700)	18	2000 (500)	-	0.73%	200[-205.07,605.07]
Furuhjelm 2009	52	3500 (500)	65	3600 (600)		2.02%	-100[-299.36,99.36]
Gustafson 2013	22	3416.8 (552.9)	24	3435.5 (404.8)	+	1.29%	-18.7[-300.85,263.45]
Haghiac 2015	25	3278 (448)	24	2935 (356)		1.74%	343[116.89,569.11]
Harris 2015	224	3215.5 (506.2)	121	3165 (494.6)		3.33%	50.5[-59.77,160.77]
Helland 2001	175	3609 (493)	166	3618 (527)		3.36%	-9[-117.45,99.45]
Horvaticek 2017	47	3580.9 (568)	43	3456.9 (575.8)		1.64%	124[-112.62,360.62]
Jamilian 2016	26	3418.8 (344.7)	27	3405.2 (465.1)		1.8%	13.6[-206.23,233.43]
Judge 2007	27	3394 (430)	21	3224.6 (431.2)		1.56%	169.38[-76.22,414.98]
Keenan 2014	36	3074.4 (582.3)	17	2919.6 (537.7)		1.08%	154.83[-163.81,473.47]
Khalili 2016	75	3260 (360)	75	3230 (430)		3.05%	30[-96.92,156.92]
Krummel 2016	34	3502 (433)	29	3484 (411)		1.92%	18[-190.71,226.71]
Makrides 2010	1197	3475 (564)	1202	3407 (576)		4.36%	68[22.38,113.62]
Malcolm 2003	31	3507.7 (500.8)	29	3645.1 — (495)	+ -	1.51%	-137.4[-389.46,114.66]
Min 2014	32	3200 (566)	27	3200 (520)		1.32%	0[-277.36,277.36]
Min 2014 [diabetic women]	28	3000 (529)	30	2900 (548)		1.32%	100[-177.21,377.21]
Min 2016	57	3200 (600)	56	3100 (400)		2.16%	100[-87.72,287.72]
Mozurkewich 2013	78	3583 (498.6)	40	3309 (555)		1.96%	274[69.48,478.52]
Mulder 2014	104	3494 (400)	111	3497 (479)		3.2%	-3[-120.69,114.69]
Olsen 1992	266	3571 (528)	267	3475 (520)		3.7%	96[7.03,184.97]
Olsen 2000	799	2668.3 (629.3)	817	2656.2 (600.4)	-	4.17%	12.14[-47.85,72.13]







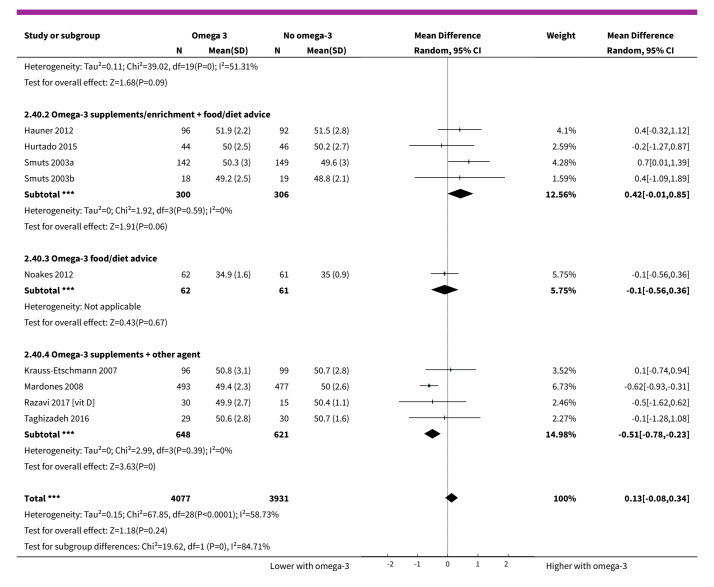
Analysis 2.39. Comparison 2 Type of omega-3 intervention, Outcome 39 Birthweight Z score.

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
2.39.1 Omega-3 supplements on	ıly						
Krummel 2016	34	0.8 (1)	29	0.7 (0.8)	+	2.75%	0.19[-0.25,0.63]
Makrides 2010	1197	0.3 (1.1)	1202	0.2 (1)		76.19%	0.06[-0.02,0.14]
Mulder 2014	104	0.4 (0.8)	111	0.4 (1)		9.03%	0.06[-0.18,0.3]
Subtotal ***	1335		1342		•	87.98%	0.06[-0.01,0.14]
Heterogeneity: Tau ² =0; Chi ² =0.33,	df=2(P=0.8	5); I ² =0%					
Test for overall effect: Z=1.62(P=0.	11)						
2.39.2 Omega-3 supplements + o	other agen	t					
Bergmann 2007	41	1 (0.1)	74	1 (0.9)		12.02%	0[-0.21,0.21]
Subtotal ***	41		74			12.02%	0[-0.21,0.21]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.04(P=0.	97)						
Total ***	1376		1416		•	100%	0.06[-0.02,0.13]
Heterogeneity: Tau ² =0; Chi ² =0.61,	df=3(P=0.9); I ² =0%					
Test for overall effect: Z=1.53(P=0.	13)						
Test for subgroup differences: Chi	² =0.28, df=1	. (P=0.6), I ² =0%					
			Lower	with omega-3	-0.5 -0.25 0 0.25 0.5	Higher with	omega-3

Analysis 2.40. Comparison 2 Type of omega-3 intervention, Outcome 40 Birth length (cm).

Study or subgroup	0	mega 3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
2.40.1 Omega-3 supplements of	nly						
Carlson 2013	154	49.7 (2.7)	147	49 (3.4)	 	4.24%	0.7[0,1.4]
Dunstan 2008	40	50.5 (1.9)	43	49.7 (2)	 	3.54%	0.8[-0.03,1.63]
Gustafson 2013	22	49.6 (2.1)	24	50 (3)		1.6%	-0.36[-1.85,1.13]
Haghiac 2015	17	49.4 (2.1)	33	47.8 (2.2)		2.15%	1.64[0.41,2.87]
Harris 2015	224	50 (4.6)	121	49.4 (2.8)	 	3.83%	0.54[-0.23,1.31]
Helland 2001	175	50.7 (2)	166	50.8 (2.2)		5.82%	-0.1[-0.55,0.35]
Jamilian 2016	26	50.1 (1.5)	27	50.9 (2.1)		2.92%	-0.8[-1.78,0.18]
Judge 2007	27	51.3 (2.2)	21	50.1 (2.3)	 	2.02%	1.22[-0.06,2.5]
Khalili 2016	75	49.5 (2.3)	75	49.5 (2.4)		3.94%	0[-0.75,0.75]
Krummel 2016	34	51.5 (1.7)	29	50.9 (1.9)	-	3.25%	0.6[-0.3,1.5]
Makrides 2010	1197	50.2 (2.8)	1202	49.9 (3.3)		7.13%	0.37[0.13,0.61]
Malcolm 2003	31	52.6 (2.5)	29	53.2 (2.9)		1.81%	-0.6[-1.97,0.77]
Min 2014	32	49.8 (4)	27	50.9 (4.2)		0.91%	-1.1[-3.18,0.98]
Min 2014 [diabetic women]	28	48.8 (4.2)	30	48.6 (4.4)		- 0.81%	0.2[-2.02,2.42]
Min 2016	58	49 (7.8)	56	50.3 (5.7)		0.65%	-1.3[-3.8,1.2]
Olsen 1992	266	52.1 (2.5)	267	51.7 (2.4)		6.06%	0.4[-0.01,0.81]
Ramakrishnan 2010	487	50.3 (2.3)	486	50.3 (2.7)		6.7%	0[-0.32,0.32]
Razavi 2017	30	51.3 (2.1)	15	50.4 (1.1)		3.09%	0.9[-0.04,1.84]
Tofail 2006	125	47.6 (2.1)	124	48.1 (2)		5.4%	-0.5[-1.01,0.01]
Valenzuela 2015	19	48 (3.6)	21	49.1 (3.4)		0.84%	-1.1[-3.28,1.08]
Subtotal ***	3067		2943		•	66.71%	0.21[-0.03,0.45]





Analysis 2.41. Comparison 2 Type of omega-3 intervention, Outcome 41 Length at birth Z score.

Study or subgroup	0	mega-3	No o	mega-3		Mean Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rar	ndom, 95% CI		Random, 95% CI
2.41.1 Omega-3 suppleme	ents only								
Krummel 2016	34	1.5 (1.1)	29	1 (0.9)			-	30.24%	0.46[-0.04,0.96]
Makrides 2010	1197	-0.1 (0.9)	1202	-0.2 (1)			+	69.76%	0.06[-0.02,0.14]
Subtotal ***	1231		1231				•	100%	0.18[-0.18,0.54]
Heterogeneity: Tau ² =0.05;	Chi ² =2.41, df=1(P=	0.12); I ² =58.55%)						
Test for overall effect: Z=0.	98(P=0.32)								
Total ***	1231		1231				•	100%	0.18[-0.18,0.54]
Heterogeneity: Tau ² =0.05;	Chi ² =2.41, df=1(P=	0.12); I ² =58.55%)						
Test for overall effect: Z=0.	98(P=0.32)								
			Lower w	rith omega-3	-2	-1	0 1	² Higher with	omega-3



Analysis 2.42. Comparison 2 Type of omega-3 intervention, Outcome 42 Head circumference at birth (cm).

Study or subgroup	Omega-3		No omega-3		Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
2.42.1 Omega-3 supplements only							
Carlson 2013	154	34.2 (1.7)	147	33.7 (2)		3.89%	0.5[0.08,0.92
Harris 2015	224	34.5 (3.1)	121	33.9 (2.2)		2.18%	0.6[0.04,1.10
Hauner 2012	96	35.1 (1.4)	92	34.8 (1.7)	++-	3.45%	0.3[-0.15,0.7
Helland 2001	175	35.3 (1.5)	166	35.2 (1.6)		6.33%	0.1[-0.23,0.4
Jamilian 2016	26	34.9 (1)	27	35.3 (1)		2.24%	-0.4[-0.95,0.1
Judge 2007	25	34.4 (1)	20	34.3 (1.1)		1.85%	0.18[-0.43,0.7
Khalili 2016	75	34.7 (1.4)	75	34.7 (1.5)		3.19%	0[-0.46,0.4
Krummel 2016	34	34.6 (1.3)	29	34.7 (1)		2.2%	-0.1[-0.66,0.4
Makrides 2010	1197	34.6 (1.8)	1202	34.6 (2.3)	- -	25.24%	0.08[-0.09,0.2
Malcolm 2003	31	34.9 (1.5)	29	34.9 (1.5)		1.19%	0[-0.76,0.7
Min 2014	32	33.6 (2.3)	27	33.5 (2.1)		0.56%	0.1[-1.01,1.2
Min 2014 [diabetic women]	28	33.7 (2.1)	30	33 (2.2)		0.56%	0.7[-0.41,1.8
Min 2016	57	33.1 (1.6)	56	33.8 (2.5)		1.14%	-0.7[-1.48,0.0
Ramakrishnan 2010	487	34.3 (1.5)	486	34.3 (1.8)		15.86%	0[-0.21,0.2
Razavi 2017	30	35.5 (1.4)	15	35.7 (1.3)		1.01%	-0.2[-1.03,0.6
Tofail 2006	125	32.3 (1.8)	124	32.4 (1.7)		3.64%	-0.1[-0.53,0.3
Subtotal ***	2796	()	2646			74.54%	0.07[-0.03,0.1
Heterogeneity: Tau²=0; Chi²=18.33, c) 25)· l²=18 19%				14.5476	0.01[0.003,0.2
Test for overall effect: Z=1.44(P=0.15		7.25),1 -10.1570					
	,						
2.42.2 Omega-3 supplements/enri							
Hurtado 2015	44	34 (1.5)	46	34.1 (1.4)		1.91%	-0.1[-0.7,0
Smuts 2003a	142	33.8 (1.7)	149	33.4 (1.8)		4.25%	0.4[-0,0.
Smuts 2003b	18	34.3 (0.9)	19	33.3 (1.7)		0.91%	1[0.13,1.8
Subtotal ***	204		214			7.07%	0.34[0.03,0.6
Heterogeneity: Tau ² =0; Chi ² =4.36, df		1); I ² =54.13%					
Test for overall effect: Z=2.15(P=0.03)						
2.42.3 Omega-3 food/diet advice o	nly						
Noakes 2012	53	34.5 (1.5)	54	34.7 (1.5)		2.24%	-0.2[-0.75,0.3
Subtotal ***	53		54			2.24%	-0.2[-0.75,0.3
Heterogeneity: Not applicable							
Test for overall effect: Z=0.71(P=0.48)						
2.42.4 Omega-3 supplements + oth	er agent	t					
Mardones 2008	493	34.5 (1.7)	477	34.3 (1.7)		15.01%	0.2[-0.01,0.4
Razavi 2017 [vit D]	30	35.3 (2.7)	15	35.7 (1.3)	+	0.5%	-0.4[-1.57,0.7
Taghizadeh 2016	29	34.8 (2.4)	30	35.5 (1.6)		0.63%	-0.7[-1.74,0.3
Subtotal ***	552		522			16.15%	0.15[-0.06,0.3
Heterogeneity: Tau ² =0; Chi ² =3.6, df=); ² =44.49%	- ==				
Test for overall effect: Z=1.39(P=0.17		,,					
Total ***	3605		3436		•	100%	0.1[0.01,0.1
Heterogeneity: Tau²=0; Chi²=30.28, c		111112-27 2504	3730			100%	0.1[0.01,0.1
		,.±1,, 1 –21.35%					
Fest for overall effect: Z=2.26(P=0.02 Fest for subgroup differences: Chi ² =3		/D 0.26\ 12 = :	740/				
Test for sungroup differences ('hi'=	s.99, at=1	L (P=U.26), I^=24.	14%				



Analysis 2.43. Comparison 2 Type of omega-3 intervention, Outcome 43 Head circumference at birth Z score.

Study or subgroup	o	Omega-3		omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
2.43.1 Omega-3 suppleme	entation only						
Krummel 2016	34	0.7 (1.1)	29	0.6 (0.9)		4.4%	0.08[-0.41,0.57]
Makrides 2010	1197	0 (1.1)	1202	0.1 (1.5)		95.6%	-0.04[-0.15,0.07]
Subtotal ***	1231		1231		•	100%	-0.03[-0.14,0.07]
Heterogeneity: Tau ² =0; Chi	i ² =0.22, df=1(P=0.6	4); I ² =0%					
Test for overall effect: Z=0.	66(P=0.51)						
Total ***	1231		1231		•	100%	-0.03[-0.14,0.07]
Heterogeneity: Tau ² =0; Chi	i²=0.22, df=1(P=0.6	4); I ² =0%					
Test for overall effect: Z=0.	66(P=0.51)						
			Higher	with omega-3	-1 -0.5 0 0.5 1	Lower with	omega-3

Analysis 2.44. Comparison 2 Type of omega-3 intervention, Outcome 44 Baby admitted to neonatal care.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.44.1 Omega-3 supplements on	ıly				
Bisgaard 2016	40/365	41/371		7.82%	0.99[0.66,1.5]
Carlson 2013	13/154	13/147		2.56%	0.95[0.46,1.99]
Makrides 2010	21/1197	37/1202		7.1%	0.57[0.34,0.97]
Mozurkewich 2013	8/78	4/40		1.02%	1.03[0.33,3.2]
Olsen 2000	258/1062	283/1076		54.08%	0.92[0.8,1.07]
Subtotal (95% CI)	2856	2836	•	72.58%	0.9[0.79,1.02]
Total events: 340 (Omega-3), 378 ((No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =3.27,	df=4(P=0.51); I ² =0%				
Test for overall effect: Z=1.6(P=0.1	1)				
2.44.2 Omega-3 supplements/en	nrichment + food/diet	advice			
Smuts 2003a	21/142	21/149		3.94%	1.05[0.6,1.84]
Smuts 2003b	2/18	6/19	+	1.12%	0.35[0.08,1.52]
Subtotal (95% CI)	160	168		5.06%	0.89[0.54,1.5]
Total events: 23 (Omega-3), 27 (No	o omega-3)				
Heterogeneity: Tau ² =0; Chi ² =1.87,	df=1(P=0.17); I ² =46.56 ^c	%			
Test for overall effect: Z=0.42(P=0.	.67)				
2.44.3 Omega-3 supplements + c	other agents				
Ali 2017	10/32	15/31		2.93%	0.65[0.34,1.21]
Harper 2010	110/427	99/410	-	19.43%	1.07[0.84,1.35]
Subtotal (95% CI)	459	441	*	22.36%	1.01[0.81,1.26]
Total events: 120 (Omega-3), 114 ((No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =2.15,	df=1(P=0.14); I ² =53.46 ^o	%			
Test for overall effect: Z=0.1(P=0.9	2)				
Total (95% CI)	3475	3445	•	100%	0.92[0.83,1.03]
Total events: 483 (Omega-3), 519 ((No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =7.9, d	df=8(P=0.44); I ² =0%				
Test for overall effect: Z=1.41(P=0.	.16)				
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	1

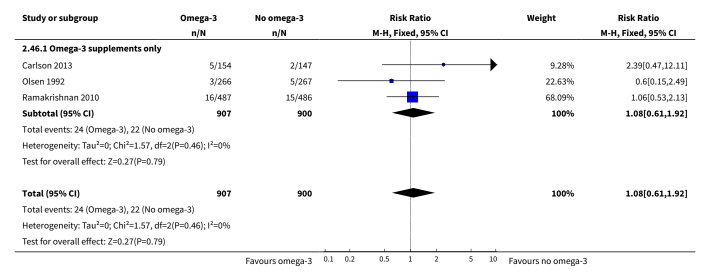


Study or subgroup	Omega-3 n/N	No omega-3 n/N	Risk Ratio M-H, Fixed, 95% CI	Weight	Risk Ratio M-H, Fixed, 95% CI
Test for subgroup differences:	Chi ² =0.84, df=1 (P=0.66),	I ² =0%			
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	

Analysis 2.45. Comparison 2 Type of omega-3 intervention, Outcome 45 Infant length of stay in hospital (days).

Study or subgroup	0	mega-3	No omega-3			Me	an Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fi	ixed, 95% CI		Fixed, 95% CI
2.45.1 Omega-3 supplementat	ion only								
Olsen 2000	1017	7.7 (16.3)	1024	7.6 (18.5)				100%	0.11[-1.4,1.62]
Subtotal ***	1017		1024				→	100%	0.11[-1.4,1.62]
Heterogeneity: Not applicable									
Test for overall effect: Z=0.14(P=	0.89)								
Total ***	1017		1024				•	100%	0.11[-1.4,1.62]
Heterogeneity: Not applicable									
Test for overall effect: Z=0.14(P=	0.89)								
			Favo	ours omega-3	-10	-5	0 5	10 Favours no	omega-3

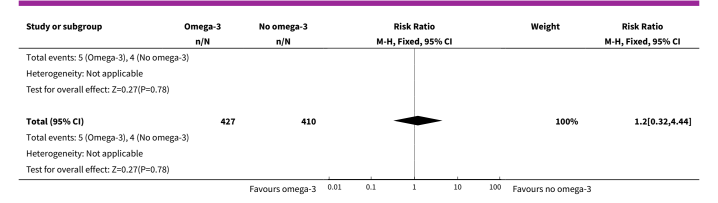
Analysis 2.46. Comparison 2 Type of omega-3 intervention, Outcome 46 Congenital anomalies.



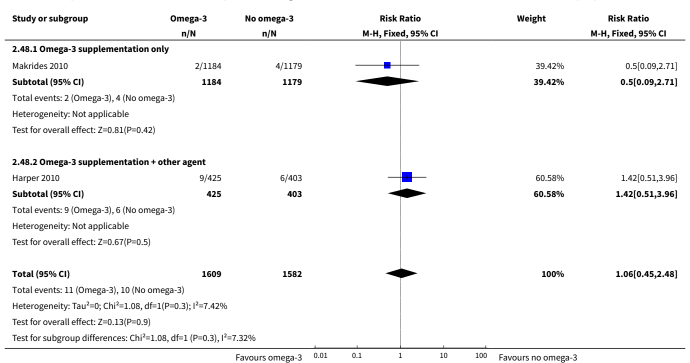
Analysis 2.47. Comparison 2 Type of omega-3 intervention, Outcome 47 Retinopathy of prematurity.

Study or subgroup	Omega-3	mega-3 No omega-3			Risk Ratio			Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI					M-H, Fixed, 95% CI		
2.47.1 Omega-3 supplement	ation + other agent only									
Harper 2010	5/427	4/410			-	_		100%	1.2[0.32,4.44]	
Subtotal (95% CI)	427	410			—	-		100%	1.2[0.32,4.44]	
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3		





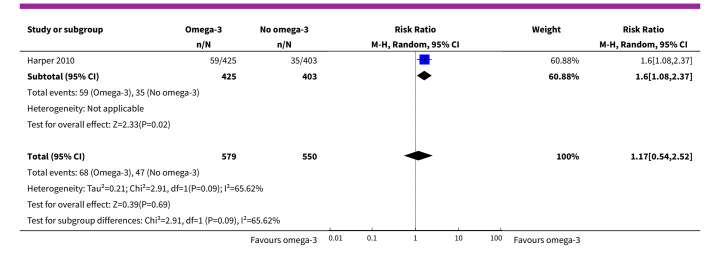
Analysis 2.48. Comparison 2 Type of omega-3 intervention, Outcome 48 Bronchopulmonary dysplasia.



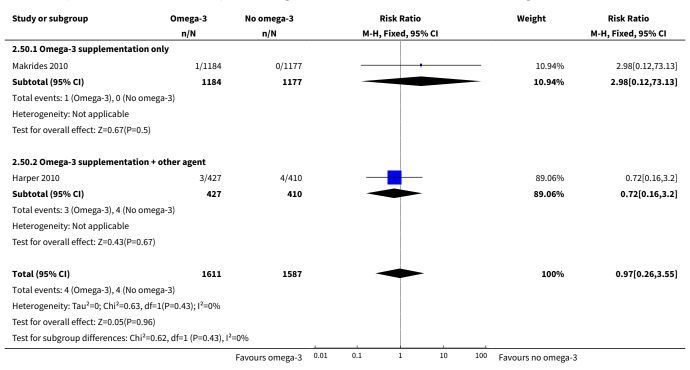
Analysis 2.49. Comparison 2 Type of omega-3 intervention, Outcome 49 Respiratory distress syndrome.

Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio	
	n/N	n/N	M-H, Random, 95% CI						M-H, Random, 95% CI	
2.49.1 Omega-3 supplementation of	only									
Carlson 2013	9/154	12/147						39.12%	0.72[0.31,1.65]	
Subtotal (95% CI)	154	147						39.12%	0.72[0.31,1.65]	
Total events: 9 (Omega-3), 12 (No om	nega-3)									
Heterogeneity: Not applicable										
Test for overall effect: Z=0.79(P=0.43))									
2.49.2 Omega-3 supplementation +	other agent									
		Favours omega-3	0.01	0.1	1	10	100	Favours omega-3		





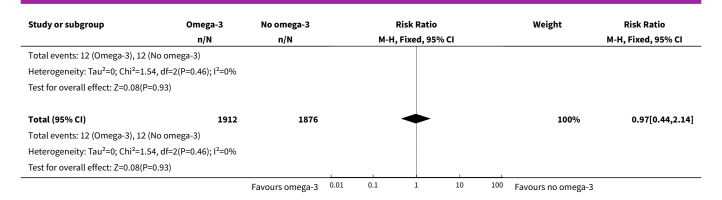
Analysis 2.50. Comparison 2 Type of omega-3 intervention, Outcome 50 Necrotising enterocolitis (NEC).



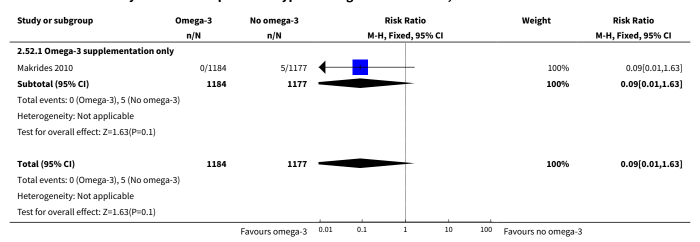
Analysis 2.51. Comparison 2 Type of omega-3 intervention, Outcome 51 Neonatal sepsis (proven).

Study or subgroup	Omega-3 No omega-3 n/N n/N			Risk Ratio				Weight	Risk Ratio
			M-H, Fixed, 95% CI						M-H, Fixed, 95% CI
2.51.1 Omega-3 supplements only									
Harper 2010	5/427	3/410						25.07%	1.6[0.38,6.65]
Helland 2001	4/301	7/289		_	-			58.5%	0.55[0.16,1.85]
Makrides 2010	3/1184	2/1177		-	+			16.43%	1.49[0.25,8.91]
Subtotal (95% CI)	1912	1876						100%	0.97[0.44,2.14]
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3	

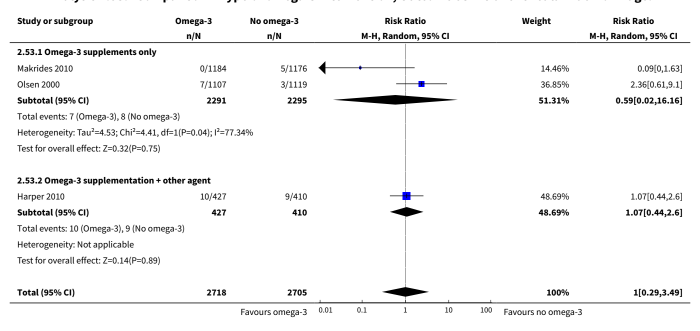




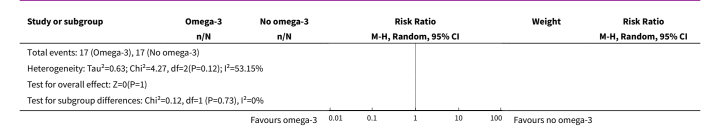
Analysis 2.52. Comparison 2 Type of omega-3 intervention, Outcome 52 Convulsion.



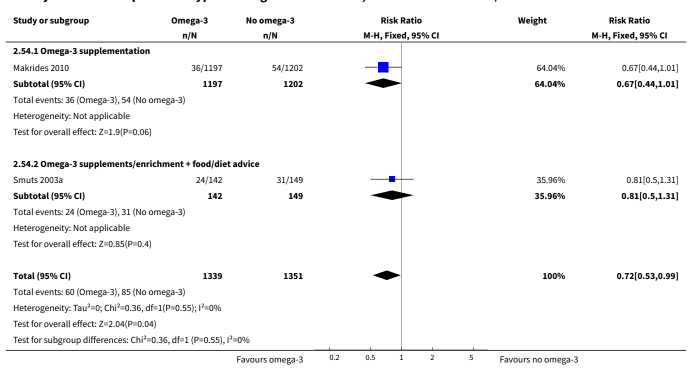
Analysis 2.53. Comparison 2 Type of omega-3 intervention, Outcome 53 Intraventricular haemorrhage.







Analysis 2.54. Comparison 2 Type of omega-3 intervention, Outcome 54 Neonatal/infant serious adverse events.



Analysis 2.55. Comparison 2 Type of omega-3 intervention, Outcome 55 Neonatal/infant morbidity: cardiovascular.

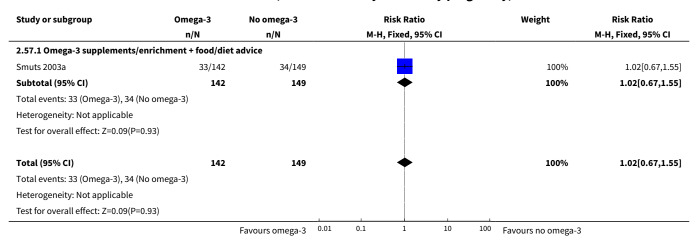
Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.55.1 Omega-3 supplements/enric	hment + food/diet	advice			
Smuts 2003a	48/142	42/149	- - - - - - - - - - 	100%	1.2[0.85,1.69]
Subtotal (95% CI)	142	149		100%	1.2[0.85,1.69]
Total events: 48 (Omega-3), 42 (No on	nega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=1.03(P=0.3)					
Total (95% CI)	142	149		100%	1.2[0.85,1.69]
Total events: 48 (Omega-3), 42 (No on	nega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=1.03(P=0.3)					
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	



Analysis 2.56. Comparison 2 Type of omega-3 intervention, Outcome 56 Neonatal/infant morbidity: respiratory.

Study or subgroup	Omega-3	No omega-3		Risk Rati	o		Weight	Risk Ratio
	n/N	n/N		M-H, Fixed, 9	5% CI			M-H, Fixed, 95% CI
2.56.1 Omega-3 supplements/enricl	hment + food/diet	advice						
Smuts 2003a	31/142	32/149		-	_		100%	1.02[0.66,1.57]
Subtotal (95% CI)	142	149			-		100%	1.02[0.66,1.57]
Total events: 31 (Omega-3), 32 (No on	nega-3)							
Heterogeneity: Not applicable								
Test for overall effect: Z=0.07(P=0.94)								
Total (95% CI)	142	149			-		100%	1.02[0.66,1.57]
Total events: 31 (Omega-3), 32 (No on	nega-3)							
Heterogeneity: Not applicable								
Test for overall effect: Z=0.07(P=0.94)								
		Favours omega-3	0.2	0.5 1	2	5	Favours no omega-3	

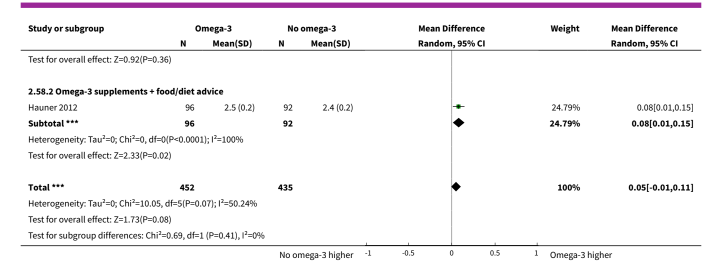
Analysis 2.57. Comparison 2 Type of omega-3 intervention, Outcome 57 Neonatal/infant morbidity: caused by pregnancy/birth.



Analysis 2.58. Comparison 2 Type of omega-3 intervention, Outcome 58 Ponderal index.

Study or subgroup	Oi	mega-3	No	No omega-3		Mean Difference		Weight		Mean Difference	
	N	Mean(SD)	N	Mean(SD)		Rai	ndom, 95% C	1			Random, 95% CI
2.58.1 Omega-3 supplemen	nts only										
Carlson 2013	154	2.7 (0.3)	147	2.7 (0.4)			+			21.53%	0[-0.08,0.08]
Haghiac 2015	17	2.7 (0.3)	16	2.6 (0.2)			+			7.73%	0.12[-0.06,0.3]
Jamilian 2016	26	1.7 (1)	27	1.7 (1.4)			-			0.77%	-0.02[-0.65,0.61]
Krummel 2016	34	2.6 (0.2)	29	2.6 (0.2)			+			15.56%	-0.06[-0.17,0.05]
Tofail 2006	125	2.5 (0.2)	124	2.4 (0.2)			-			29.62%	0.1[0.05,0.15]
Subtotal ***	356		343				•			75.21%	0.04[-0.04,0.11]
Heterogeneity: Tau ² =0; Chi ² =	=9.73, df=4(P=0.0	5); I ² =58.9%									
			No on	nega-3 higher	1	-0.5	0	0.5	1	Omega-3 highe	r





Comparison 3. Dose (DHA/EPA) subgroups

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Preterm birth (< 37 weeks)	26	10294	Risk Ratio (M-H, Fixed, 95% CI)	0.88 [0.80, 0.97]
1.1 Low: < 500 mg/day	6	1604	Risk Ratio (M-H, Fixed, 95% CI)	0.88 [0.65, 1.20]
1.2 Mid: 500 mg-1 g/day	9	4343	Risk Ratio (M-H, Fixed, 95% CI)	0.79 [0.64, 0.98]
1.3 High: > 1 g/day	9	4240	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.83, 1.03]
1.4 Other	2	107	Risk Ratio (M-H, Fixed, 95% CI)	0.66 [0.19, 2.32]
2 Early preterm birth (< 34 weeks)	9	5204	Risk Ratio (M-H, Fixed, 95% CI)	0.58 [0.44, 0.77]
2.1 Low: < 500 mg/day	1	168	Risk Ratio (M-H, Fixed, 95% CI)	0.29 [0.05, 1.51]
2.2 Mid: 500 mg-1 g/day	7	4176	Risk Ratio (M-H, Fixed, 95% CI)	0.47 [0.30, 0.75]
2.3 High: > 1 g/day	2	860	Risk Ratio (M-H, Fixed, 95% CI)	0.69 [0.49, 0.99]
3 Prolonged gestation (> 42 weeks)	6	5141	Risk Ratio (M-H, Fixed, 95% CI)	1.59 [1.10, 2.30]
3.1 Low: < 500 mg/day	2	303	Risk Ratio (M-H, Fixed, 95% CI)	1.72 [0.07, 41.64]
3.2 Mid: 500 mg-1 g/day	2	2544	Risk Ratio (M-H, Fixed, 95% CI)	1.92 [0.54, 6.81]
3.3 High: > 1 g/day	3	2294	Risk Ratio (M-H, Fixed, 95% CI)	1.56 [1.05, 2.30]
4 Pre-eclampsia (hyper- tension with proteinuria)	20	8306	Risk Ratio (M-H, Fixed, 95% CI)	0.83 [0.69, 1.01]
4.1 Low: < 500 mg/day	5	650	Risk Ratio (M-H, Fixed, 95% CI)	0.59 [0.28, 1.26]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4.2 Mid: 500 mg-1 g/day	7	4118	Risk Ratio (M-H, Fixed, 95% CI)	0.83 [0.62, 1.11]
4.3 High: > 1 g/day	8	3479	Risk Ratio (M-H, Fixed, 95% CI)	0.87 [0.66, 1.14]
4.4 Other	1	59	Risk Ratio (M-H, Fixed, 95% CI)	2.07 [0.20, 21.60]
5 Caesarean section	28	8481	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.91, 1.03]
5.1 Low: < 500 g/day	8	1670	Risk Ratio (M-H, Fixed, 95% CI)	0.94 [0.84, 1.06]
5.2 Mid: 500 mg-1 g/day	10	4399	Risk Ratio (M-H, Fixed, 95% CI)	0.93 [0.85, 1.02]
5.3 High: > 1 g/day	8	2294	Risk Ratio (M-H, Fixed, 95% CI)	1.15 [0.97, 1.37]
5.4 Other	2	118	Risk Ratio (M-H, Fixed, 95% CI)	0.59 [0.30, 1.15]
6 Length of gestation (days)	42	12517	Mean Difference (IV, Random, 95% CI)	1.67 [0.95, 2.39]
6.1 Low: < 500 mg/day	12	2117	Mean Difference (IV, Random, 95% CI)	1.05 [0.07, 2.03]
6.2 Mid: 500 mg-1 g/day	15	4881	Mean Difference (IV, Random, 95% CI)	1.97 [0.56, 3.38]
6.3 High: > 1 g/day	12	3364	Mean Difference (IV, Random, 95% CI)	1.86 [0.45, 3.27]
6.4 Mixed	1	1998	Mean Difference (IV, Random, 95% CI)	0.10 [-1.00, 1.20]
6.5 Other	3	157	Mean Difference (IV, Random, 95% CI)	2.24 [-0.83, 5.31]
7 Perinatal death	10	7416	Risk Ratio (M-H, Fixed, 95% CI)	0.75 [0.54, 1.03]
7.1 Low: < 500 mg/day	2	1127	Risk Ratio (M-H, Fixed, 95% CI)	0.52 [0.20, 1.33]
7.2 Mid: 500 mg-1 g/day	3	2566	Risk Ratio (M-H, Fixed, 95% CI)	0.41 [0.16, 1.02]
7.3 High: > 1 g/day	5	3723	Risk Ratio (M-H, Fixed, 95% CI)	0.89 [0.61, 1.29]
8 Stillbirth	16	7880	Risk Ratio (M-H, Fixed, 95% CI)	0.94 [0.62, 1.42]
8.1 Low: < 500 mg/day	1	977	Risk Ratio (M-H, Fixed, 95% CI)	0.67 [0.11, 3.96]
8.2 Mid: 500 mg/day-1 g/ day	5	2783	Risk Ratio (M-H, Fixed, 95% CI)	0.70 [0.27, 1.83]
8.3 High: > 1 g/day	7	3933	Risk Ratio (M-H, Fixed, 95% CI)	1.03 [0.62, 1.69]
8.4 Other	3	187	Risk Ratio (M-H, Fixed, 95% CI)	1.16 [0.23, 5.94]
9 Neonatal death	9	7448	Risk Ratio (M-H, Fixed, 95% CI)	0.61 [0.34, 1.11]
9.1 Low: < 500 mg/day	2	1123	Risk Ratio (M-H, Fixed, 95% CI)	0.47 [0.15, 1.44]
9.2 Mid: 500 mg/day-1 g/ day	2	2700	Risk Ratio (M-H, Fixed, 95% CI)	0.50 [0.12, 1.98]

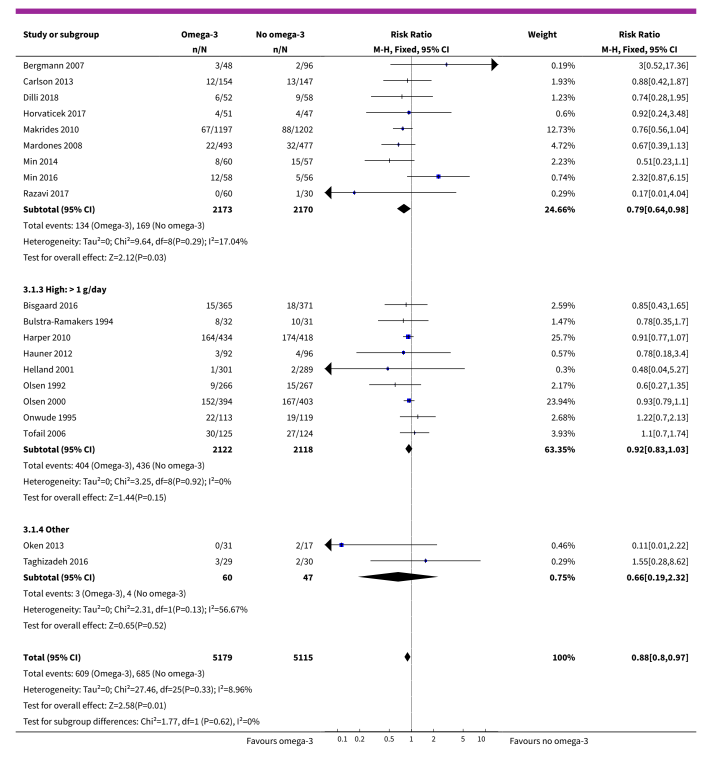


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
9.3 High: > 1 g/day	5	3625	Risk Ratio (M-H, Fixed, 95% CI)	0.78 [0.34, 1.78]
10 Low birthweight (< 2500 g)	15	8449	Risk Ratio (M-H, Fixed, 95% CI)	0.90 [0.82, 0.99]
10.1 Low: < 500 mg/day	5	1551	Risk Ratio (M-H, Fixed, 95% CI)	0.74 [0.51, 1.08]
10.2 Mid: 500 mg-1 g/day	5	3901	Risk Ratio (M-H, Fixed, 95% CI)	0.71 [0.54, 0.92]
10.3 High: > 1 g/day	5	2997	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.88, 1.08]
11 Small-for-gestational age/IUGR	8	6907	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.90, 1.13]
11.1 Low: < 500 mg/day	1	973	Risk Ratio (M-H, Fixed, 95% CI)	1.04 [0.73, 1.48]
11.2 Mid: 500 mg-1 g/day	2	3369	Risk Ratio (M-H, Fixed, 95% CI)	0.85 [0.66, 1.09]
11.3 High: > 1 g/day	4	2506	Risk Ratio (M-H, Fixed, 95% CI)	1.07 [0.93, 1.23]
11.4 Other	1	59	Risk Ratio (M-H, Fixed, 95% CI)	2.07 [0.20, 21.60]
12 Birthweight (g)	44	11584	Mean Difference (IV, Random, 95% CI)	75.30 [38.09, 112.50]
12.1 Low: < 500 mg/day	12	2220	Mean Difference (IV, Random, 95% CI)	26.32 [-12.74, 65.39]
12.2 Mid: 500 mg-1 g/day	18	5007	Mean Difference (IV, Random, 95% CI)	91.49 [24.34, 158.64]
12.3 High: > 1 g/day	14	4298	Mean Difference (IV, Random, 95% CI)	88.31 [29.61, 147.01]
12.4 Other	1	59	Mean Difference (IV, Random, 95% CI)	-203.20 [-456.97, 50.57]

Analysis 3.1. Comparison 3 Dose (DHA/EPA) subgroups, Outcome 1 Preterm birth (< 37 weeks).

Study or subgroup	Omega-3	No omega-3			Ris	k Ratio			Weight	Risk Ratio
	n/N	n/N		M-H, Fixed, 95% CI					M-H, Fixed, 95% CI	
3.1.1 Low: < 500 mg/day										
Malcolm 2003	0/31	1/32	\leftarrow		+	_			0.21%	0.34[0.01,8.13]
Miller 2016	3/60	10/55	_			_			1.51%	0.28[0.08,0.95]
Ramakrishnan 2010	49/487	40/486				+-			5.81%	1.22[0.82,1.82]
Smuts 2003a	14/142	17/149				+			2.41%	0.86[0.44,1.69]
Smuts 2003b	1/18	5/19	\leftarrow						0.71%	0.21[0.03,1.64]
Van Goor 2009	1/86	3/39	\leftarrow	•		+			0.6%	0.15[0.02,1.41]
Subtotal (95% CI)	824	780			4	•			11.24%	0.88[0.65,1.2]
Total events: 68 (Omega-3), 76 (N	o omega-3)									
Heterogeneity: Tau ² =0; Chi ² =10.6	1, df=5(P=0.06); I ² =52.87	7%								
Test for overall effect: Z=0.8(P=0.4	42)									
3.1.2 Mid: 500 mg-1 g/day										
		Favours omega-3	0.1	0.2	0.5	1 2	5 1	.0	Favours no omega-3	

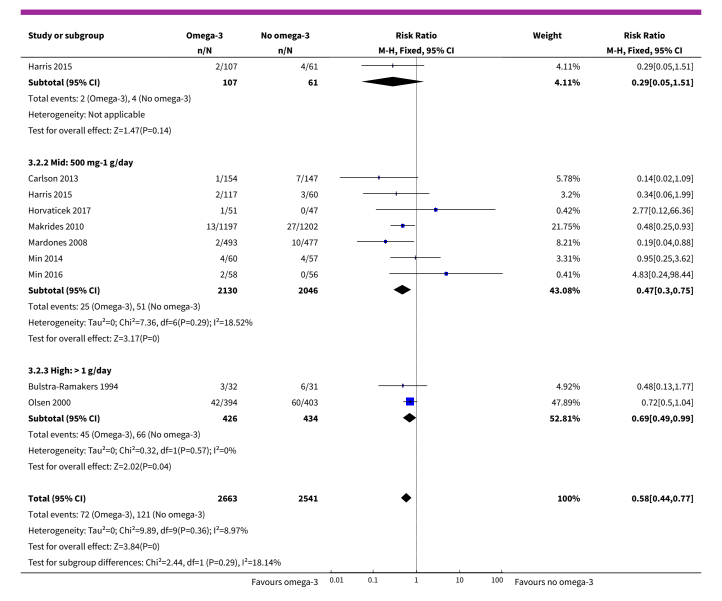




Analysis 3.2. Comparison 3 Dose (DHA/EPA) subgroups, Outcome 2 Early preterm birth (< 34 weeks).

Study or subgroup	Omega-3	No omega-3		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		М-Н,	Fixed, 95	% CI			M-H, Fixed, 95% CI
3.2.1 Low: < 500 mg/day									
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3	

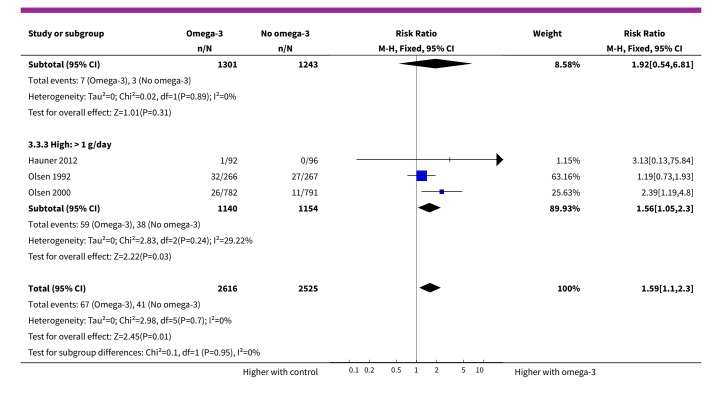




Analysis 3.3. Comparison 3 Dose (DHA/EPA) subgroups, Outcome 3 Prolonged gestation (> 42 weeks).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
3.3.1 Low: < 500 mg/day					
Harris 2015	1/107	0/61	-	1.49%	1.72[0.07,41.64]
Mulder 2014	0/68	0/67			Not estimable
Subtotal (95% CI)	175	128		1.49%	1.72[0.07,41.64]
Total events: 1 (Omega-3), 0 (No	omega-3)				
Heterogeneity: Tau ² =0; Chi ² =0, di	f=0(P<0.0001); I ² =100%				
Test for overall effect: Z=0.33(P=0).74)				
3.3.2 Mid: 500 mg-1 g/day					
Harris 2015	1/117	0/60	—	1.54%	1.55[0.06,37.5]
Makrides 2010	6/1184	3/1183		7.03%	2[0.5,7.97]
	Н	igher with control	0.1 0.2 0.5 1 2 5 10	Higher with omega-3	

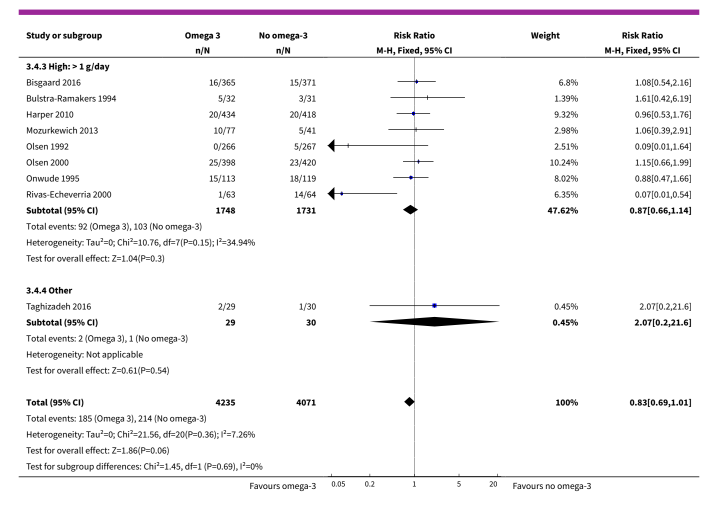




Analysis 3.4. Comparison 3 Dose (DHA/EPA) subgroups, Outcome 4 Pre-eclampsia (hypertension with proteinuria).

Study or subgroup	Omega 3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
3.4.1 Low: < 500 mg/day					
D'Almedia 1992	2/50	5/50		2.29%	0.4[0.08,1.97]
Harris 2015	1/107	0/61	•	0.29%	1.72[0.07,41.64]
Jamilian 2016	0/27	1/27	•	0.69%	0.33[0.01,7.84]
Smuts 2003a	5/142	10/149		4.46%	0.52[0.18,1.5]
Smuts 2003b	1/18	0/19	-	0.22%	3.16[0.14,72.84]
Subtotal (95% CI)	344	306		7.95%	0.59[0.28,1.26]
Total events: 9 (Omega 3), 16 (No	o omega-3)				
Heterogeneity: Tau ² =0; Chi ² =1.93	3, df=4(P=0.75); I ² =0%				
Test for overall effect: Z=1.37(P=	0.17)				
3.4.2 Mid: 500 mg-1 g/day					
Carlson 2013	2/154	2/147		0.94%	0.95[0.14,6.69]
Harris 2015	1/117	0/60 -	•	0.3%	1.55[0.06,37.5]
Horvaticek 2017	4/43	5/38		2.43%	0.71[0.2,2.44]
Makrides 2010	60/1197	58/1202	-	26.47%	1.04[0.73,1.48]
Mardones 2008	8/493	16/477		7.44%	0.48[0.21,1.12]
Ranjkesh 2011	2/50	10/50 —		4.57%	0.2[0.05,0.87]
Razavi 2017	5/60	3/30		1.83%	0.83[0.21,3.26]
Subtotal (95% CI)	2114	2004	•	43.98%	0.83[0.62,1.11]
Total events: 82 (Omega 3), 94 (N	lo omega-3)				
Heterogeneity: Tau ² =0; Chi ² =6.99	9, df=6(P=0.32); I ² =14.17 ⁹	%	į		
Test for overall effect: Z=1.24(P=	0.21)				
		Favours omega-3 0.0	5 0.2 1 5	20 Favours no omega-3	<u> </u>

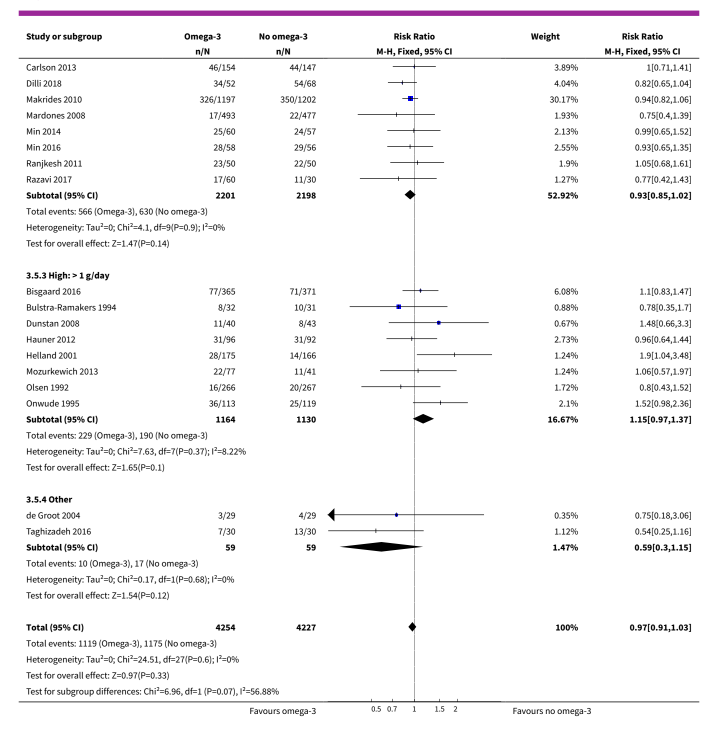




Analysis 3.5. Comparison 3 Dose (DHA/EPA) subgroups, Outcome 5 Caesarean section.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
3.5.1 Low: < 500 g/day					
Jamilian 2016	12/26	18/27		1.53%	0.69[0.42,1.13]
Judge 2007	8/27	5/21		0.49%	1.24[0.48,3.25]
Khalili 2016	30/75	33/75		2.85%	0.91[0.62,1.33]
Miller 2016	20/60	12/55		1.08%	1.53[0.83,2.83]
Noakes 2012	8/53	9/54		0.77%	0.91[0.38,2.17]
Ramakrishnan 2010	216/429	234/440		19.96%	0.95[0.83,1.08]
Smuts 2003a	18/142	21/149		1.77%	0.9[0.5,1.62]
Smuts 2003b	2/18	6/19		0.5%	0.35[0.08,1.52]
Subtotal (95% CI)	830	840	•	28.94%	0.94[0.84,1.06]
Total events: 314 (Omega-3), 338	(No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =6, d	f=7(P=0.54); I ² =0%				
Test for overall effect: Z=1.03(P=0	0.3)				
3.5.2 Mid: 500 mg-1 g/day					
Ali 2017	17/34	19/34		1.64%	0.89[0.57,1.4]
Bergmann 2007	33/43	55/77	+	3.4%	1.07[0.86,1.33]
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	

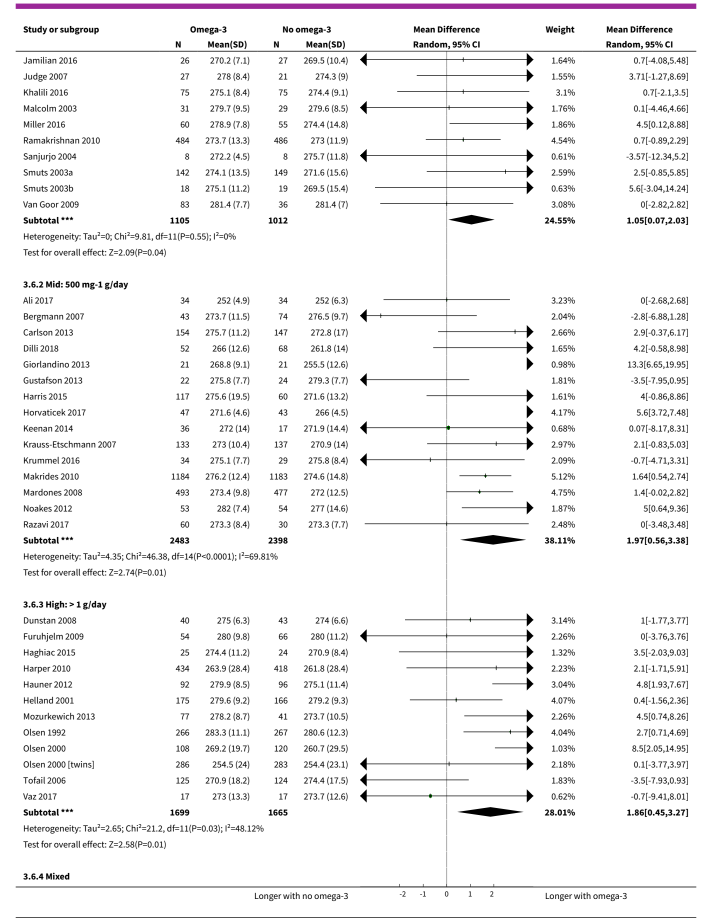




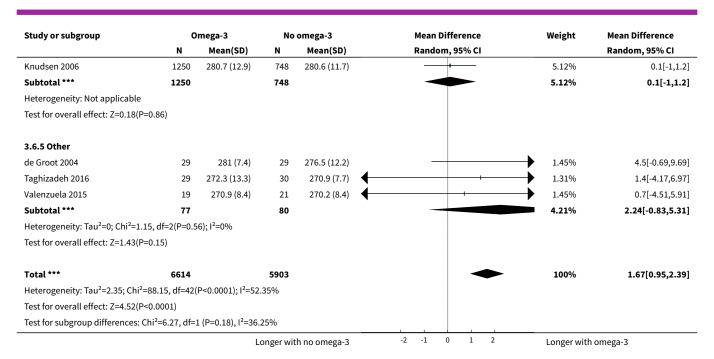
Analysis 3.6. Comparison 3 Dose (DHA/EPA) subgroups, Outcome 6 Length of gestation (days).

Study or subgroup	0	mega-3	No	omega-3		Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Random, 95% CI		Random, 95% CI
3.6.1 Low: < 500 mg/day								
Harris 2015	107	275 (19.5)	61	271.6 (13.2)			1.56%	3.4[-1.56,8.36]
Hurtado 2015	44	275.1 (12.6)	46	277.2 (10.5)	\leftarrow	- 	1.63%	-2.1[-6.9,2.7]
		Lo	onger wit	:h no omega-3		-2 -1 0 1 2	Longer with	omega-3





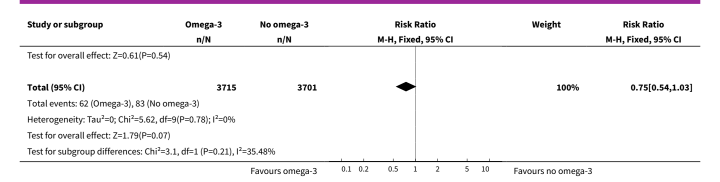




Analysis 3.7. Comparison 3 Dose (DHA/EPA) subgroups, Outcome 7 Perinatal death.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
3.7.1 Low: < 500 mg/day					
Khalili 2016	0/75	1/75	+	1.78%	0.33[0.01,8.05]
Ramakrishnan 2010	6/489	11/488		13.08%	0.54[0.2,1.46]
Subtotal (95% CI)	564	563		14.87%	0.52[0.2,1.33]
Total events: 6 (Omega-3), 12 (No om	nega-3)				
Heterogeneity: Tau ² =0; Chi ² =0.08, df	=1(P=0.77); I ² =0%				
Test for overall effect: Z=1.37(P=0.17))				
3.7.2 Mid: 500 mg-1 g/day					
Ali 2017	2/34	3/34	+	3.56%	0.67[0.12,3.74]
Horvaticek 2017	1/56	0/43	•	0.67%	2.32[0.1,55.48]
Makrides 2010	3/1197	12/1202 —	•	14.23%	0.25[0.07,0.89]
Subtotal (95% CI)	1287	1279		18.46%	0.41[0.16,1.02]
Total events: 6 (Omega-3), 15 (No om	nega-3)				
Heterogeneity: Tau ² =0; Chi ² =2.03, df	=2(P=0.36); I ² =1.41%				
Test for overall effect: Z=1.92(P=0.05))				
3.7.3 High: > 1 g/day					
Bulstra-Ramakers 1994	2/32	3/31	+	3.62%	0.65[0.12,3.61]
Harper 2010	16/434	17/418		20.58%	0.91[0.46,1.77]
Olsen 2000	19/1126	23/1126		27.33%	0.83[0.45,1.51]
Onwude 1995	1/113	2/119	· ·	2.31%	0.53[0.05,5.73]
Tofail 2006	12/159	11/165		12.83%	1.13[0.51,2.49]
Subtotal (95% CI)	1864	1859	*	66.67%	0.89[0.61,1.29]
Total events: 50 (Omega-3), 56 (No or	mega-3)				
Heterogeneity: Tau ² =0; Chi ² =0.74, df	=4(P=0.95); I ² =0%				
		Favours omega-3	0.1 0.2 0.5 1 2 5 10	Favours no omega-3	

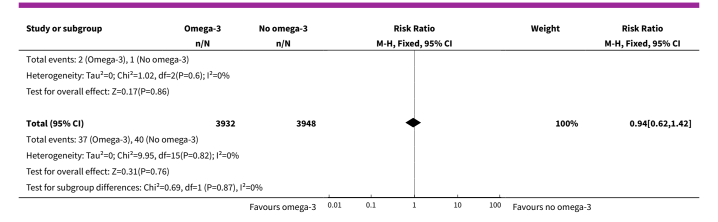




Analysis 3.8. Comparison 3 Dose (DHA/EPA) subgroups, Outcome 8 Stillbirth.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
3.8.1 Low: < 500 mg/day					
Ramakrishnan 2010	2/489	3/488		6.64%	0.67[0.11,3.96]
Subtotal (95% CI)	489	488		6.64%	0.67[0.11,3.96]
Total events: 2 (Omega-3), 3 (No	o omega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.45(P=	=0.65)				
3.8.2 Mid: 500 mg/day-1 g/day	,				
Horvaticek 2017	1/48	0/50		1.08%	3.12[0.13,74.82]
Jamilian 2016	1/27	0/27		1.11%	3[0.13,70.53]
Makrides 2010	1/1197	7/1202 —		15.44%	0.14[0.02,1.16]
Min 2014	2/60	0/57		1.13%	4.75[0.23,96.93]
Min 2016	0/58	1/57 —		3.34%	0.33[0.01,7.88]
Subtotal (95% CI)	1390	1393		22.1%	0.7[0.27,1.83]
Total events: 5 (Omega-3), 8 (No	o omega-3)				
Heterogeneity: Tau ² =0; Chi ² =5.6	64, df=4(P=0.23); I ² =29.09 ⁰	%			
Test for overall effect: Z=0.74(P=	=0.46)				
3.8.3 High: > 1 g/day					
Bulstra-Ramakers 1994	1/32	0/31		1.12%	2.91[0.12,68.81]
Haghiac 2015	1/25	0/25		- 1.11%	3[0.13,70.3]
Helland 2001	1/301	0/289		1.13%	2.88[0.12,70.43]
Olsen 1992	1/266	1/267		2.21%	1[0.06,15.96]
Olsen 2000	16/1056	19/1085		41.42%	0.87[0.45,1.67]
Onwude 1995	0/113	2/119 —		5.38%	0.21[0.01,4.34]
Tofail 2006	8/159	6/165		13.02%	1.38[0.49,3.9]
Subtotal (95% CI)	1952	1981	•	65.38%	1.03[0.62,1.69]
Total events: 28 (Omega-3), 28 (No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =2.8	39, df=6(P=0.82); I ² =0%				
Test for overall effect: Z=0.1(P=0	0.92)				
3.8.4 Other					
de Groot 2004	0/40	1/39 —		3.36%	0.33[0.01,7.75]
Oken 2013	1/31	0/17		1.41%	1.69[0.07,39.3]
	1/30	0/30		- 1.11%	3[0.13,70.83
Taghizadeh 2016	1/30	0/30			



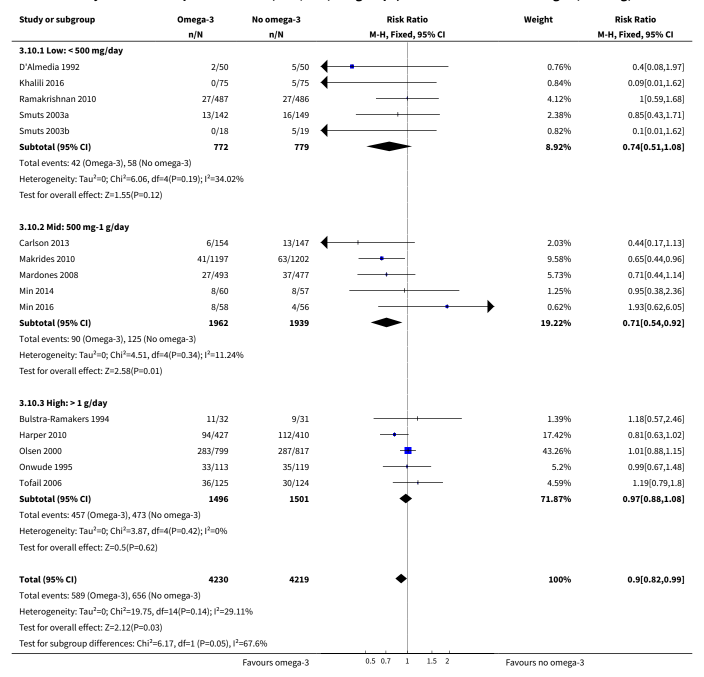


Analysis 3.9. Comparison 3 Dose (DHA/EPA) subgroups, Outcome 9 Neonatal death.

Study or subgroup	Omega-3 No omega-3		Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI	
3.9.1 Low: < 500 mg/day						
Khalili 2016	0/75	1/75 —	+	5.37%	0.33[0.01,8.05]	
Ramakrishnan 2010	4/487	8/486		28.67%	0.5[0.15,1.65]	
Subtotal (95% CI)	562	561		34.04%	0.47[0.15,1.44]	
Total events: 4 (Omega-3), 9 (N	o omega-3)					
Heterogeneity: Tau ² =0; Chi ² =0.	05, df=1(P=0.82); I ² =0%					
Test for overall effect: Z=1.32(P	=0.19)					
3.9.2 Mid: 500 mg/day-1 g/da	у					
Carlson 2013	1/154	1/147		3.66%	0.95[0.06,15.12]	
Makrides 2010	2/1197	5/1202		17.86%	0.4[0.08,2.07]	
Subtotal (95% CI)	1351	1349		21.53%	0.5[0.12,1.98]	
Total events: 3 (Omega-3), 6 (N	o omega-3)					
Heterogeneity: Tau ² =0; Chi ² =0.	28, df=1(P=0.6); l ² =0%					
Test for overall effect: Z=0.99(P	=0.32)					
3.9.3 High: > 1 g/day						
Bisgaard 2016	0/365	0/371			Not estimable	
Bulstra-Ramakers 1994	1/32	3/31		10.91%	0.32[0.04,2.94]	
Olsen 2000	3/1126	4/1144	+	14.21%	0.76[0.17,3.4]	
Onwude 1995	1/113	0/119		1.74%	3.16[0.13,76.73]	
Tofail 2006	4/159	5/165		17.57%	0.83[0.23,3.04]	
Subtotal (95% CI)	1795	1830	*	44.43%	0.78[0.34,1.78]	
Total events: 9 (Omega-3), 12 (No omega-3)					
Heterogeneity: Tau ² =0; Chi ² =1.	36, df=3(P=0.72); I ² =0%					
Test for overall effect: Z=0.6(P=	0.55)					
Total (95% CI)	3708	3740	•	100%	0.61[0.34,1.11]	
Total events: 16 (Omega-3), 27	(No omega-3)					
Heterogeneity: Tau ² =0; Chi ² =2.	24, df=7(P=0.95); I ² =0%					
Test for overall effect: Z=1.61(P	=0.11)					
Test for subgroup differences:	Chi ² =0.6, df=1 (P=0.74), I ² =	:0%				



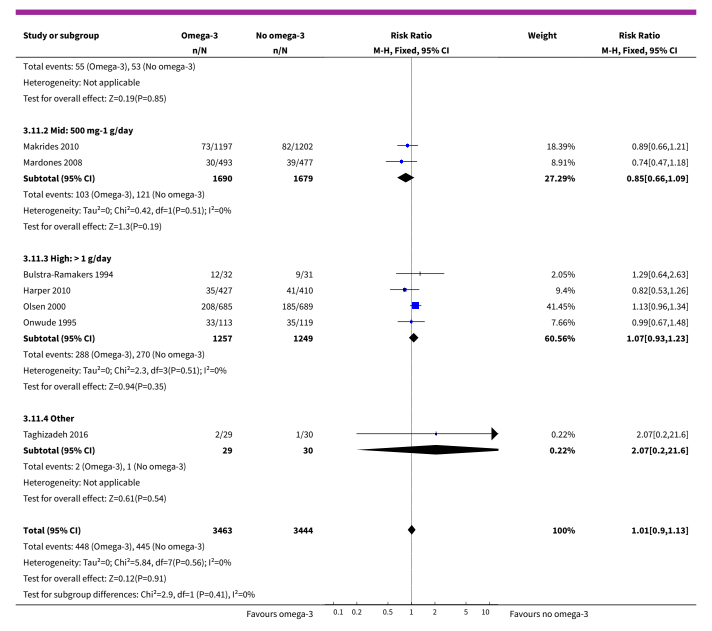
Analysis 3.10. Comparison 3 Dose (DHA/EPA) subgroups, Outcome 10 Low birthweight (< 2500 g).



Analysis 3.11. Comparison 3 Dose (DHA/EPA) subgroups, Outcome 11 Small-for-gestational age/IUGR.

Study or subgroup	Omega-3	No omega-3		Risk	Ratio			Weight	Risk Ratio
	n/N	n/N		M-H, Fix	ed, 95% C	:1			M-H, Fixed, 95% CI
3.11.1 Low: < 500 mg/day									
Ramakrishnan 2010	55/487	53/486		_	+-			11.92%	1.04[0.73,1.48]
Subtotal (95% CI)	487	486		. •				11.92%	1.04[0.73,1.48]
		Favours omega-3	0.1 0.2	0.5	1 2	5	10	Favours no omega-3	

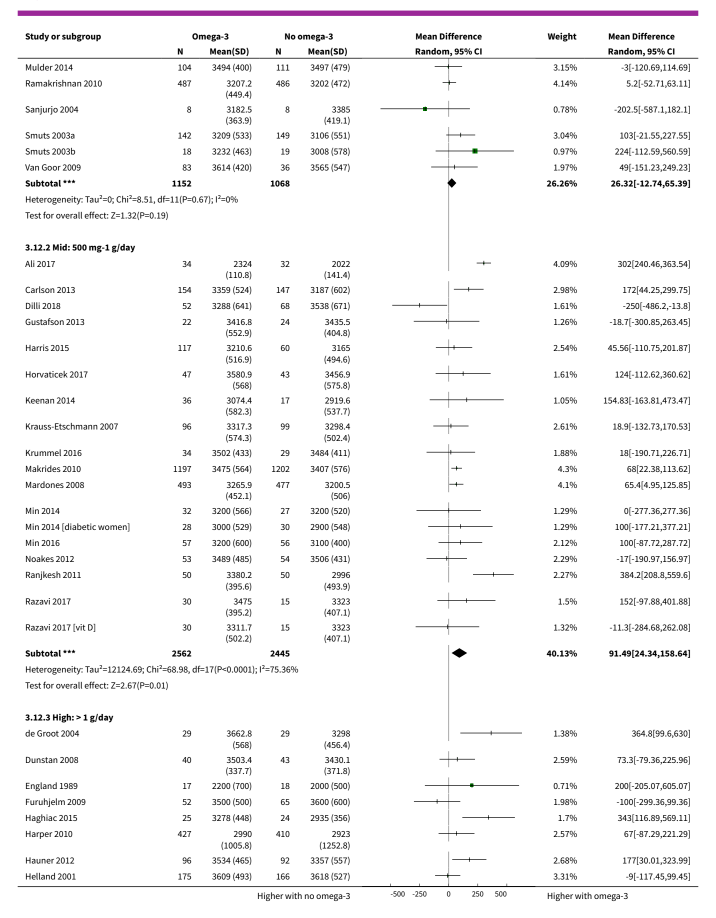




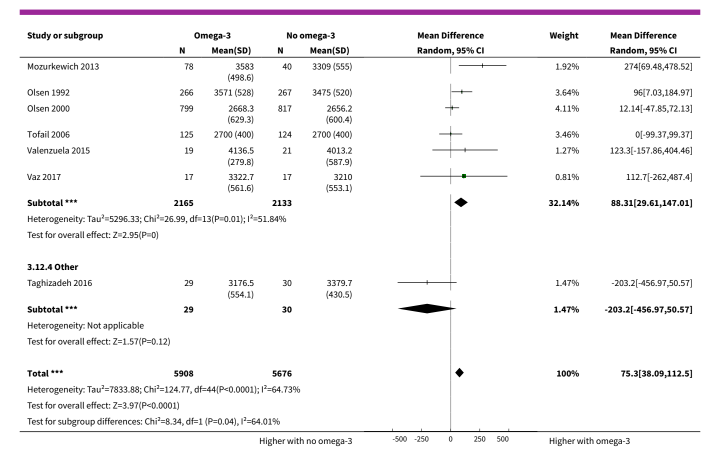
Analysis 3.12. Comparison 3 Dose (DHA/EPA) subgroups, Outcome 12 Birthweight (g).

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
3.12.1 Low: < 500 mg/day							
Harris 2015	107	3220.9 (494.2)	61	3165 (494.6)		2.55%	55.91[-99.57,211.39]
Hurtado 2015	44	3300 (500)	46	3200 (500)		1.9%	100[-106.65,306.65]
Jamilian 2016	26	3418.8 (344.7)	27	3405.2 (465.1)		1.77%	13.6[-206.23,233.43]
Judge 2007	27	3394 (430)	21	3224.6 (431.2)	+	1.53%	169.38[-76.22,414.98]
Khalili 2016	75	3260 (360)	75	3230 (430)		3%	30[-96.92,156.92]
Malcolm 2003	31	3507.7 (500.8)	29	3645.1 (495)		1.48%	-137.4[-389.46,114.66]
		Н	igher wit	h no omega-3	-500 -250 0 250 500	Higher with	n omega-3









Comparison 4. Timing subgroups

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Preterm birth (< 37 weeks)	26	10304	Risk Ratio (M-H, Fixed, 95% CI)	0.89 [0.81, 0.97]
1.1 ≤ 20 weeks GA start	12	6563	Risk Ratio (M-H, Fixed, 95% CI)	0.85 [0.76, 0.95]
1.2 > 20 weeks GA start	13	3693	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.82, 1.23]
1.3 Mixed	1	48	Risk Ratio (M-H, Fixed, 95% CI)	0.11 [0.01, 2.22]
2 Early preterm birth (< 34 weeks)	9	5204	Risk Ratio (M-H, Fixed, 95% CI)	0.58 [0.44, 0.77]
2.1 ≤ 20 weeks GA start	8	5090	Risk Ratio (M-H, Fixed, 95% CI)	0.56 [0.43, 0.75]
2.2 > 20 weeks GA start	1	114	Risk Ratio (M-H, Fixed, 95% CI)	4.83 [0.24, 98.44]
3 Prolonged gestation (> 42 weeks)	6	5141	Risk Ratio (M-H, Fixed, 95% CI)	1.61 [1.11, 2.33]
3.1 ≤ 20 weeks GA start	5	4608	Risk Ratio (M-H, Fixed, 95% CI)	2.35 [1.29, 4.28]

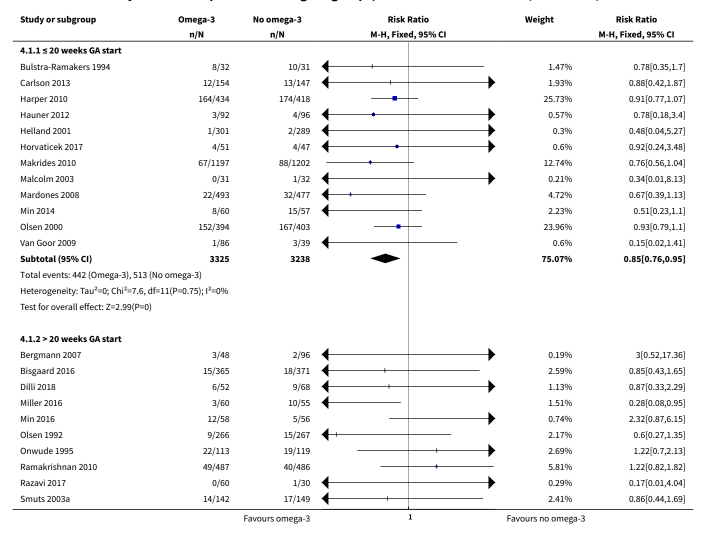


Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size	
3.2 > 20 weeks GA start	1	533	Risk Ratio (M-H, Fixed, 95% CI)	1.19 [0.73, 1.93]	
4 Pre-eclampsia (hyper- tension with proteinuria)	20	8306	Risk Ratio (M-H, Fixed, 95% CI)	0.84 [0.69, 1.01]	
4.1 ≤ 20 weeks GA start	13	6296	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.74, 1.15]	
4.2 > 20 weeks GA start	6	1883	Risk Ratio (M-H, Fixed, 95% CI)	0.79 [0.53, 1.18]	
4.3 Not reported	1	127	Risk Ratio (M-H, Fixed, 95% CI)	0.07 [0.01, 0.54]	
5 Caesarean section	28	8481	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.91, 1.03]	
5.1 ≤ 20 weeks GA start	13	4995	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.88, 1.07]	
5.2 > 20 weeks GA start	14	2617	Risk Ratio (M-H, Fixed, 95% CI)	0.98 [0.87, 1.10]	
5.3 Mixed	1	869	Risk Ratio (M-H, Fixed, 95% CI)	0.95 [0.83, 1.08]	
6 Length of gestation (days)	43	12517	Mean Difference (IV, Random, 95% CI)	1.67 [0.95, 2.39]	
6.1 ≤ 20 weeks GA start	23	9396	Mean Difference (IV, Random, 95% CI)	1.99 [1.08, 2.90]	
6.2 > 20 weeks GA start	20	3121	Mean Difference (IV, Random, 95% CI)	1.18 [-0.05, 2.40]	
7 Perinatal death	10	7416	Risk Ratio (M-H, Fixed, 95% CI)	0.75 [0.54, 1.03]	
7.1 ≤ 20 weeks GA start	6	5815	Risk Ratio (M-H, Fixed, 95% CI)	0.72 [0.49, 1.07]	
7.2 > 20 weeks GA start	4	1601	Risk Ratio (M-H, Fixed, 95% CI)	0.79 [0.46, 1.38]	
8 Stillbirth	16	7880	Risk Ratio (M-H, Fixed, 95% CI)	0.94 [0.62, 1.42]	
8.1 ≤ 20 weeks GA start	8	5537	Risk Ratio (M-H, Fixed, 95% CI)	0.88 [0.52, 1.48]	
8.2 > 20 weeks GA start	7	2295	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.50, 2.07]	
8.3 Mixed	1	48	Risk Ratio (M-H, Fixed, 95% CI)	1.69 [0.07, 39.30]	
9 Neonatal death	9	7448	Risk Ratio (M-H, Fixed, 95% CI)	0.61 [0.34, 1.11]	
9.1 ≤ 20 weeks GA start	6	5415	Risk Ratio (M-H, Fixed, 95% CI)	0.60 [0.26, 1.36]	
9.2 > 20 weeks GA start	3	2033	Risk Ratio (M-H, Fixed, 95% CI)	0.62 [0.26, 1.49]	
10 Low birthweight (< 2500 g)	15	8449	Risk Ratio (M-H, Fixed, 95% CI)	0.90 [0.82, 0.99]	
10.1 ≤ 20 weeks GA start	9	6553	Risk Ratio (M-H, Fixed, 95% CI)	0.88 [0.79, 0.97]	
10.2 > 20 weeks GA start	6	1896	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.81, 1.28]	

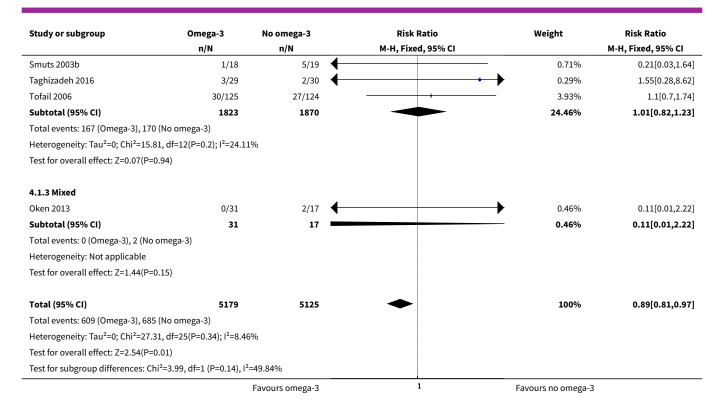


Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
11 Small-for-gestational age/IUGR	8	6907	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.90, 1.13]
11.1 ≤ 20 weeks GA start	5	5643	Risk Ratio (M-H, Fixed, 95% CI)	1.00 [0.88, 1.14]
11.2 > 20 weeks GA start	3	1264	Risk Ratio (M-H, Fixed, 95% CI)	1.03 [0.79, 1.34]
12 Birthweight (g)	43	11584	Mean Difference (IV, Random, 95% CI)	75.69 [37.84, 113.55]
12.1 ≤ 20 weeks GA start	25	7802	Mean Difference (IV, Random, 95% CI)	83.26 [44.09, 122.43]
12.2 > 20 weeks GA start	17	3747	Mean Difference (IV, Random, 95% CI)	42.96 [-34.14, 120.06]
12.3 Not reported	1	35	Mean Difference (IV, Random, 95% CI)	200.0 [-205.07, 605.07]

Analysis 4.1. Comparison 4 Timing subgroups, Outcome 1 Preterm birth (< 37 weeks).



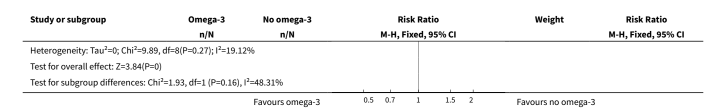




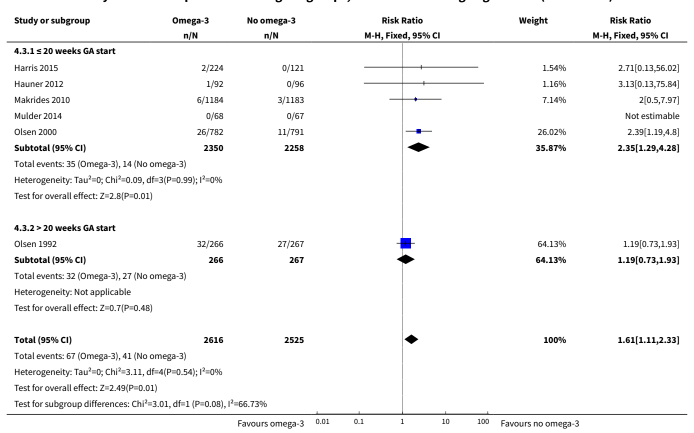
Analysis 4.2. Comparison 4 Timing subgroups, Outcome 2 Early preterm birth (< 34 weeks).

Study or subgroup	Omega-3 No omega-3		Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI	
4.2.1 ≤ 20 weeks GA start						
Bulstra-Ramakers 1994	3/32	6/31	+	4.92%	0.48[0.13,1.77]	
Carlson 2013	1/154	7/147	+	5.78%	0.14[0.02,1.09]	
Harris 2015	4/224	7/121	—	7.34%	0.31[0.09,1.03]	
Horvaticek 2017	1/51	0/47	+	0.42%	2.77[0.12,66.36]	
Makrides 2010	13/1197	27/1202	—	21.74%	0.48[0.25,0.93]	
Mardones 2008	2/493	10/477	—	8.2%	0.19[0.04,0.88]	
Min 2014	4/60	4/57	+	3.31%	0.95[0.25,3.62]	
Olsen 2000	42/394	60/403		47.88%	0.72[0.5,1.04]	
Subtotal (95% CI)	2605	2485		99.59%	0.56[0.43,0.75]	
Total events: 70 (Omega-3), 121 (N	No omega-3)					
Heterogeneity: Tau ² =0; Chi ² =8.08,	df=7(P=0.33); I ² =13.399	%				
Test for overall effect: Z=4.01(P<0.	.0001)					
4.2.2 > 20 weeks GA start						
Min 2016	2/58	0/56	+	0.41%	4.83[0.24,98.44]	
Subtotal (95% CI)	58	56		0.41%	4.83[0.24,98.44]	
Total events: 2 (Omega-3), 0 (No o	mega-3)					
Heterogeneity: Not applicable						
Test for overall effect: Z=1.02(P=0.	31)					
Total (95% CI)	2663	2541	•	100%	0.58[0.44,0.77]	
Total events: 72 (Omega-3), 121 (N	lo omega-3)					





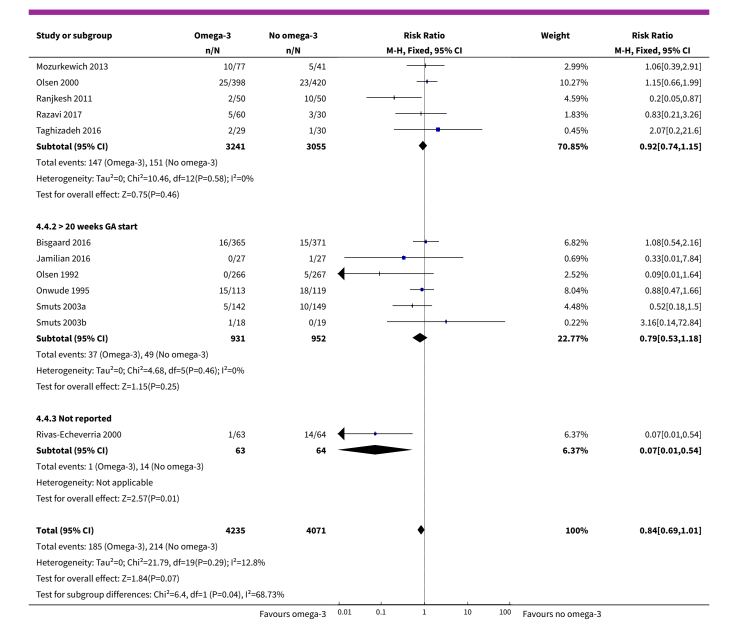
Analysis 4.3. Comparison 4 Timing subgroups, Outcome 3 Prolonged gestation (> 42 weeks).



Analysis 4.4. Comparison 4 Timing subgroups, Outcome 4 Pre-eclampsia (hypertension with proteinuria).

Study or subgroup	Omega-3	No omega-3		Risk Ratio			Weight	Risk Ratio
	n/N	'N n/N		M-H, Fixed, 95% CI				M-H, Fixed, 95% CI
4.4.1 ≤ 20 weeks GA start								
Bulstra-Ramakers 1994	5/32	3/31			-		1.4%	1.61[0.42,6.19]
Carlson 2013	2/154	2/147	-	+	_		0.94%	0.95[0.14,6.69]
D'Almedia 1992	2/50	5/50	_	-+-			2.29%	0.4[0.08,1.97]
Harper 2010	20/434	20/418		-			9.35%	0.96[0.53,1.76]
Harris 2015	2/224	0/121	_	+		_	0.3%	2.71[0.13,56.02]
Horvaticek 2017	4/43	5/38					2.44%	0.71[0.2,2.44]
Makrides 2010	60/1197	58/1202		+			26.55%	1.04[0.73,1.48]
Mardones 2008	8/493	16/477		+			7.46%	0.48[0.21,1.12]
		Favours omega-3	0.01 0.1	1	10	100	Favours no omega-3	

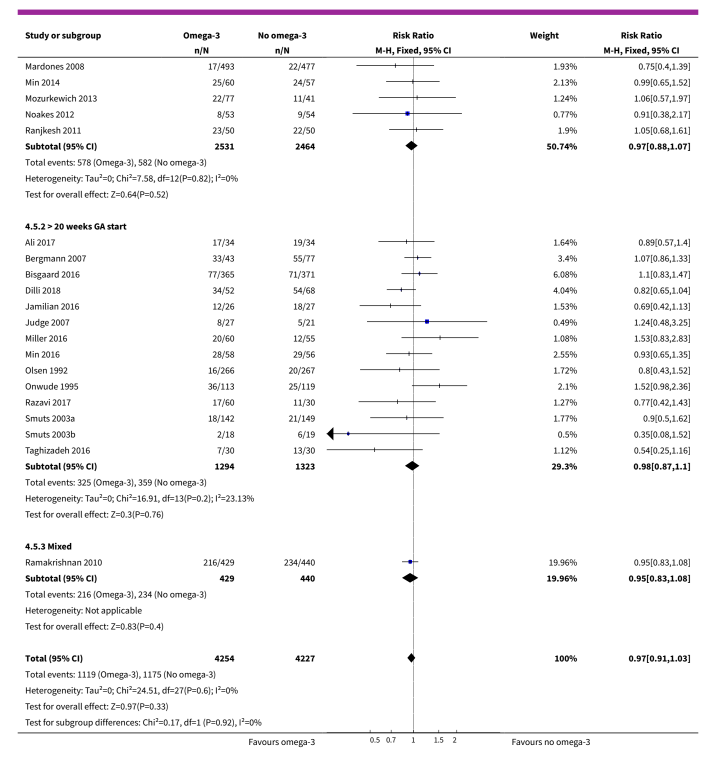




Analysis 4.5. Comparison 4 Timing subgroups, Outcome 5 Caesarean section.

Study or subgroup	r subgroup Omega-3 No omega-3 Risk Ratio n/N n/N M-H, Fixed, 95% CI		Risk Ratio	Weight	Risk Ratio
				M-H, Fixed, 95% CI	
4.5.1 ≤ 20 weeks GA start					
Bulstra-Ramakers 1994	8/32	10/31		0.88%	0.78[0.35,1.7]
Carlson 2013	46/154	44/147	- +	3.89%	1[0.71,1.41]
de Groot 2004	3/29	4/29	+ - -	0.35%	0.75[0.18,3.06]
Dunstan 2008	11/40	8/43		0.67%	1.48[0.66,3.3]
Hauner 2012	31/96	31/92		2.73%	0.96[0.64,1.44]
Helland 2001	28/175	14/166		1.24%	1.9[1.04,3.48]
Khalili 2016	30/75	33/75		2.85%	0.91[0.62,1.33]
Makrides 2010	326/1197	350/1202	, , -= 	30.17%	0.94[0.82,1.06]
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	

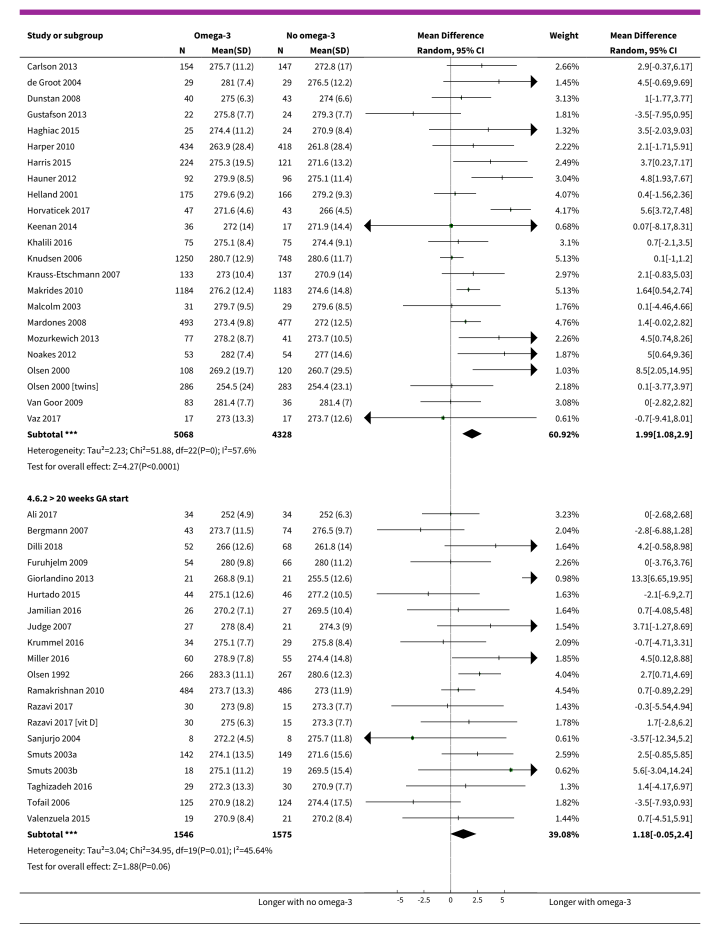




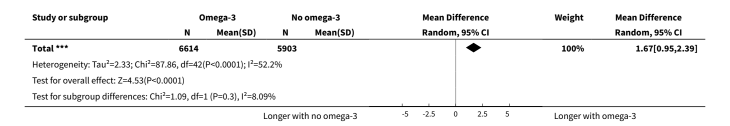
Analysis 4.6. Comparison 4 Timing subgroups, Outcome 6 Length of gestation (days).

Study or subgroup	Oı	Omega-3 No o		omega-3	Mean Difference			Weight	Mean Difference		
	N	Mean(SD)	N	Mean(SD)		Rand	om, 9	5% CI			Random, 95% CI
4.6.1 ≤ 20 weeks GA start											
		Longer with no omega-3		-5	-2.5	0	2.5	5	Longer with	omega-3	

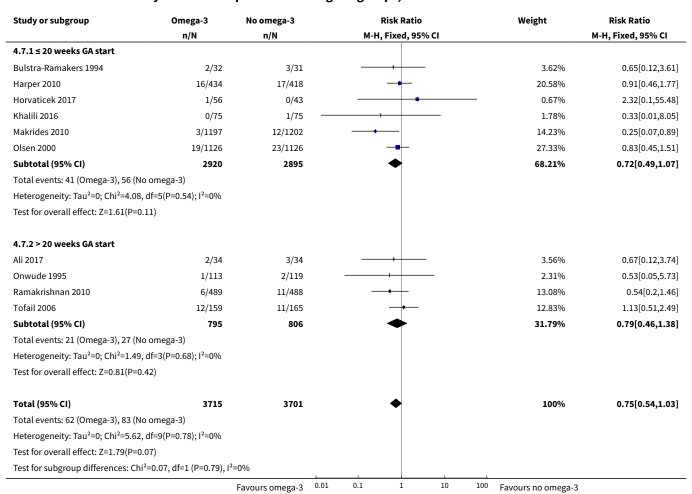








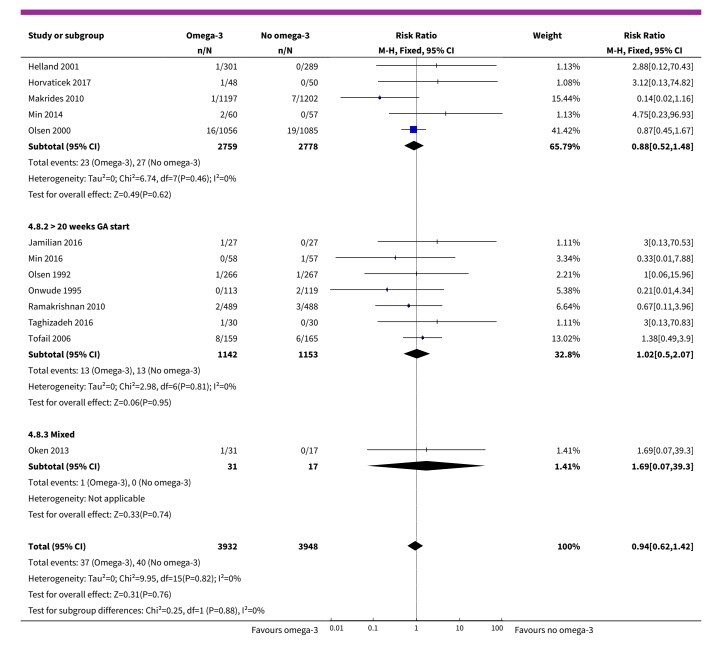
Analysis 4.7. Comparison 4 Timing subgroups, Outcome 7 Perinatal death.



Analysis 4.8. Comparison 4 Timing subgroups, Outcome 8 Stillbirth.

Study or subgroup	Omega-3	No omega-3	R	isk Ratio		Weight	Risk Ratio	
	n/N	n/N	М-Н, І	Fixed, 95% CI			M-H, Fixed, 95% CI	
4.8.1 ≤ 20 weeks GA start								
Bulstra-Ramakers 1994	1/32	0/31		+		1.12%	2.91[0.12,68.81]	
de Groot 2004	0/40	1/39				3.36%	0.33[0.01,7.75]	
Haghiac 2015	1/25	0/25		+ .		1.11%	3[0.13,70.3]	
		Favours omega-3	0.01 0.1	1 10	100	Favours no omega-3		

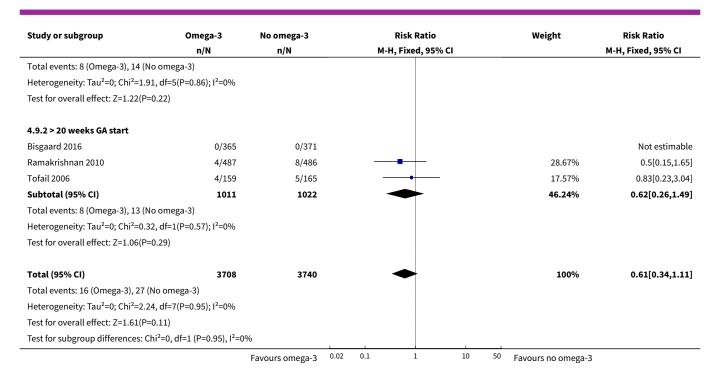




Analysis 4.9. Comparison 4 Timing subgroups, Outcome 9 Neonatal death.

Study or subgroup	Omega-3	No omega-3		Ris	k Ratio			Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI						M-H, Fixed, 95% CI	
4.9.1 ≤ 20 weeks GA start										
Bulstra-Ramakers 1994	1/32	3/31	_	+	-			10.91%	0.32[0.04,2.94]	
Carlson 2013	1/154	1/147			+			3.66%	0.95[0.06,15.12]	
Khalili 2016	0/75	1/75	-	+				5.37%	0.33[0.01,8.05]	
Makrides 2010	2/1197	5/1202						17.86%	0.4[0.08,2.07]	
Olsen 2000	3/1126	4/1144		-	•			14.21%	0.76[0.17,3.4]	
Onwude 1995	1/113	0/119		-			\rightarrow	1.74%	3.16[0.13,76.73]	
Subtotal (95% CI)	2697	2718		_	+			53.76%	0.6[0.26,1.36]	
		Favours omega-3	0.02	0.1	1	10	50	Favours no omega-3		

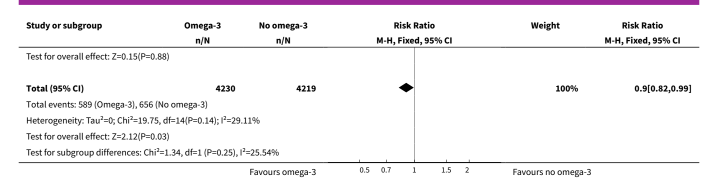




Analysis 4.10. Comparison 4 Timing subgroups, Outcome 10 Low birthweight (< 2500 g).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
4.10.1 ≤ 20 weeks GA start					
Bulstra-Ramakers 1994	11/32	9/31	+	1.39%	1.18[0.57,2.46]
Carlson 2013	6/154	13/147		2.03%	0.44[0.17,1.13]
D'Almedia 1992	2/50	5/50	 • 	0.76%	0.4[0.08,1.97]
Harper 2010	94/427	112/410		17.42%	0.81[0.63,1.02]
Khalili 2016	0/75	5/75		0.84%	0.09[0.01,1.62]
Makrides 2010	41/1197	63/1202		9.58%	0.65[0.44,0.96]
Mardones 2008	27/493	37/477		5.73%	0.71[0.44,1.14]
Min 2014	8/60	8/57		1.25%	0.95[0.38,2.36]
Olsen 2000	283/799	287/817		43.26%	1.01[0.88,1.15]
Subtotal (95% CI)	3287	3266	•	82.27%	0.88[0.79,0.97]
Total events: 472 (Omega-3), 539 ((No omega-3)				
Heterogeneity: Tau²=0; Chi²=13.87	7, df=8(P=0.09); I ² =42.33	3%			
Test for overall effect: Z=2.45(P=0.	.01)				
4.10.2 > 20 weeks GA start					
Min 2016	8/58	4/56		0.62%	1.93[0.62,6.05]
Onwude 1995	33/113	35/119		5.2%	0.99[0.67,1.48]
Ramakrishnan 2010	27/487	27/486		4.12%	1[0.59,1.68]
Smuts 2003a	13/142	16/149	+	2.38%	0.85[0.43,1.71]
Smuts 2003b	0/18	5/19	4	0.82%	0.1[0.01,1.62]
Tofail 2006	36/125	30/124	- +	4.59%	1.19[0.79,1.8]
Subtotal (95% CI)	943	953	*	17.73%	1.02[0.81,1.28]
Total events: 117 (Omega-3), 117 ((No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =4.71,	df=5(P=0.45): I ² =0%		ĺ		





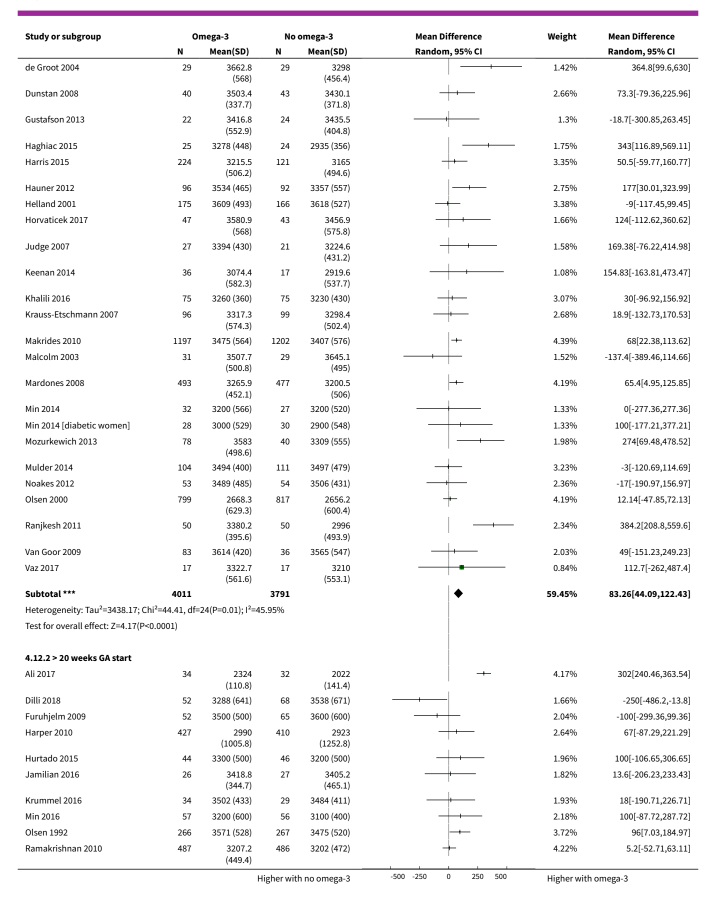
Analysis 4.11. Comparison 4 Timing subgroups, Outcome 11 Small-for-gestational age/IUGR.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio M-H, Fixed, 95% CI
	n/N	n/N	M-H, Fixed, 95% CI		
4.11.1 ≤ 20 weeks GA start					
Bulstra-Ramakers 1994	12/32	9/31		2.05%	1.29[0.64,2.63
Harper 2010	35/427	41/410		9.4%	0.82[0.53,1.26
Makrides 2010	73/1197	82/1202	-+	18.39%	0.89[0.66,1.21
Mardones 2008	30/493	39/477	-+	8.91%	0.74[0.47,1.18
Olsen 2000	208/685	185/689		41.45%	1.13[0.96,1.34
Subtotal (95% CI)	2834	2809	*	80.2%	1[0.88,1.14
Total events: 358 (Omega-3), 356	(No omega-3)				
Heterogeneity: Tau²=0; Chi²=5.49,	, df=4(P=0.24); I ² =27.159	%			
Test for overall effect: Z=0.02(P=0	.98)				
4.11.2 > 20 weeks GA start					
Onwude 1995	33/113	35/119		7.66%	0.99[0.67,1.48
Ramakrishnan 2010	55/487	53/486	-	11.92%	1.04[0.73,1.48
Taghizadeh 2016	2/29	1/30	+	0.22%	2.07[0.2,21.6
Subtotal (95% CI)	629	635	*	19.8%	1.03[0.79,1.34
Total events: 90 (Omega-3), 89 (N	o omega-3)				
Heterogeneity: Tau²=0; Chi²=0.37,	, df=2(P=0.83); I ² =0%				
Test for overall effect: Z=0.22(P=0	.82)				
Total (95% CI)	3463	3444	♦	100%	1.01[0.9,1.13
Total events: 448 (Omega-3), 445	(No omega-3)				
Heterogeneity: Tau²=0; Chi²=5.84,	, df=7(P=0.56); I ² =0%				
Test for overall effect: Z=0.12(P=0	.91)				
Test for subgroup differences: Chi	i ² =0.04 df=1 (P=0.85) I ²	=0%			

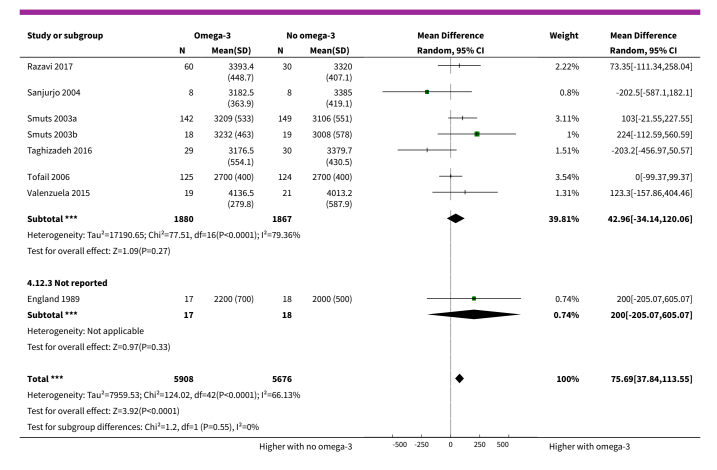
Analysis 4.12. Comparison 4 Timing subgroups, Outcome 12 Birthweight (g).

Study or subgroup	Oı	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
4.12.1 ≤ 20 weeks GA start							
Carlson 2013	154	3359 (524)	147	3187 (602)		3.06%	172[44.25,299.75]
		Н	ligher wit	h no omega-3	-500 -250 0 250 500	Higher with	omega-3









Comparison 5. DHA/mixed subgroups

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Preterm birth (< 37 weeks)	26	10304	Risk Ratio (M-H, Fixed, 95% CI)	0.89 [0.81, 0.97]
1.1 DHA/largely DHA	12	4744	Risk Ratio (M-H, Fixed, 95% CI)	0.84 [0.69, 1.02]
1.2 Mixed DHA/EPA	9	4172	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.83, 1.03]
1.3 Mixed DHA/EPA/other	5	1388	Risk Ratio (M-H, Fixed, 95% CI)	0.71 [0.45, 1.11]
2 Early preterm birth (< 34 weeks)	9	5204	Risk Ratio (M-H, Fixed, 95% CI)	0.58 [0.44, 0.77]
2.1 DHA/largely DHA	5	3260	Risk Ratio (M-H, Fixed, 95% CI)	0.46 [0.28, 0.76]
2.2 Mixed DHA/EPA	2	860	Risk Ratio (M-H, Fixed, 95% CI)	0.69 [0.49, 0.99]
2.3 Mixed DHA/EPA/other	2	1084	Risk Ratio (M-H, Fixed, 95% CI)	0.41 [0.14, 1.25]
3 Prolonged gestation (> 42 weeks)	6	5141	Risk Ratio (M-H, Fixed, 95% CI)	1.61 [1.11, 2.33]



Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
3.1 DHA/largely DHA	3	2847	Risk Ratio (M-H, Fixed, 95% CI)	2.12 [0.60, 7.49]
3.2 Mixed DHA/EPA	2	2106	Risk Ratio (M-H, Fixed, 95% CI)	1.54 [1.04, 2.28]
3.3 Mixed DHA/EPA/other	1	188	Risk Ratio (M-H, Fixed, 95% CI)	3.13 [0.13, 75.84]
4 Pre-eclampsia (hyper- tension with proteinuria)	20	8306	Risk Ratio (M-H, Fixed, 95% CI)	0.84 [0.69, 1.01]
4.1 DHA/largely DHA	6	3454	Risk Ratio (M-H, Fixed, 95% CI)	0.98 [0.71, 1.33]
4.2 Mixed DHA/EPA	9	3506	Risk Ratio (M-H, Fixed, 95% CI)	0.90 [0.69, 1.18]
4.3 Mixed DHA/EPA/other	5	1346	Risk Ratio (M-H, Fixed, 95% CI)	0.40 [0.23, 0.71]
5 Caesarean section	28	8481	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.91, 1.03]
5.1 DHA/largely DHA	9	4327	Risk Ratio (M-H, Fixed, 95% CI)	0.95 [0.87, 1.03]
5.2 Mixed DHA/EPA	10	2433	Risk Ratio (M-H, Fixed, 95% CI)	1.10 [0.95, 1.27]
5.3 Mixed DHA/EPA/other	9	1721	Risk Ratio (M-H, Fixed, 95% CI)	0.88 [0.75, 1.02]
6 Gestational length (days)	43	12517	Mean Difference (IV, Random, 95% CI)	1.67 [0.95, 2.39]
6.1 DHA/largely DHA	14	4791	Mean Difference (IV, Random, 95% CI)	2.44 [0.91, 3.98]
6.2 Mixed DHA/EPA	17	5760	Mean Difference (IV, Random, 95% CI)	1.23 [0.21, 2.24]
6.3 Mixed DHA/EPA/other	12	1966	Mean Difference (IV, Random, 95% CI)	1.42 [0.33, 2.50]
7 Perinatal death	10	7416	Risk Ratio (M-H, Fixed, 95% CI)	0.75 [0.54, 1.03]
7.1 DHA/largely DHA	3	3475	Risk Ratio (M-H, Fixed, 95% CI)	0.44 [0.21, 0.91]
7.2 Mixed DHA/EPA	6	3873	Risk Ratio (M-H, Fixed, 95% CI)	0.88 [0.60, 1.27]
7.3 Mixed DHA/EPA/other	1	68	Risk Ratio (M-H, Fixed, 95% CI)	0.67 [0.12, 3.74]
8 Stillbirth	16	7880	Risk Ratio (M-H, Fixed, 95% CI)	0.94 [0.62, 1.42]
8.1 DHA/largely DHA	5	3639	Risk Ratio (M-H, Fixed, 95% CI)	0.69 [0.28, 1.70]
8.2 Mixed DHA/EPA	8	3987	Risk Ratio (M-H, Fixed, 95% CI)	1.06 [0.65, 1.73]
8.3 Mixed DHA/EPA/other	3	254	Risk Ratio (M-H, Fixed, 95% CI)	0.70 [0.14, 3.51]
9 Neonatal death	9	7448	Risk Ratio (M-H, Fixed, 95% CI)	0.61 [0.34, 1.11]
9.1 DHA/largely DHA	3	3673	Risk Ratio (M-H, Fixed, 95% CI)	0.50 [0.20, 1.23]
9.2 Mixed DHA/EPA	6	3775	Risk Ratio (M-H, Fixed, 95% CI)	0.73 [0.33, 1.62]

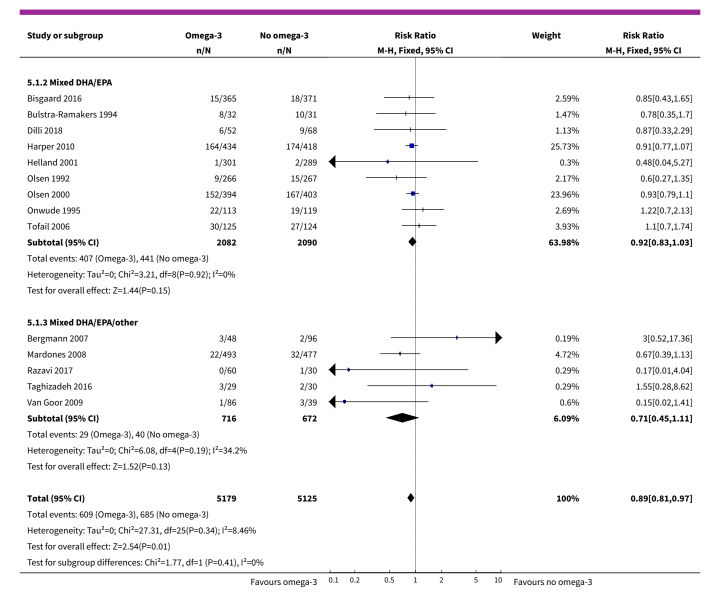


Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
10 Low birthweight (< 2500 g)	15	8449	Risk Ratio (M-H, Fixed, 95% CI)	0.90 [0.82, 0.99]
10.1 DHA/largely DHA	6	4118	Risk Ratio (M-H, Fixed, 95% CI)	0.72 [0.56, 0.93]
10.2 Mixed DHA/EPA	6	3147	Risk Ratio (M-H, Fixed, 95% CI)	0.96 [0.87, 1.07]
10.3 Mixed DHA/EPA/other	3	1184	Risk Ratio (M-H, Fixed, 95% CI)	0.78 [0.51, 1.18]
11 Small-for-gestational age/IUGR	8	6907	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.90, 1.13]
11.1 DHA/largely DHA	2	3372	Risk Ratio (M-H, Fixed, 95% CI)	0.95 [0.75, 1.20]
11.2 Mixed DHA/EPA	4	2506	Risk Ratio (M-H, Fixed, 95% CI)	1.07 [0.93, 1.23]
11.3 Mixed EPA/DHA/other	2	1029	Risk Ratio (M-H, Fixed, 95% CI)	0.78 [0.50, 1.22]
12 Birthweight (g)	43	11584	Mean Difference (IV, Random, 95% CI)	75.69 [37.84, 113.55]
12.1 DHA/largely DHA	17	6121	Mean Difference (IV, Random, 95% CI)	52.60 [26.96, 78.23]
12.2 Mixed DHA/EPA	15	4429	Mean Difference (IV, Random, 95% CI)	72.72 [6.67, 138.78]
12.3 Mixed DHA/EPA/oth- er	11	1034	Mean Difference (IV, Random, 95% CI)	113.65 [12.54, 214.75]

Analysis 5.1. Comparison 5 DHA/mixed subgroups, Outcome 1 Preterm birth (< 37 weeks).

Study or subgroup	Omega-3 No omeg			Risk Ratio	Weight	Risk Ratio	
	n/N	n/N		M-H, Fixed, 95% CI		M-H, Fixed, 95% CI	
5.1.1 DHA/largely DHA							
Carlson 2013	12/154	13/147			1.93%	0.88[0.42,1.87]	
Hauner 2012	3/92	4/96	-	+	0.57%	0.78[0.18,3.4]	
Horvaticek 2017	4/51	4/47			0.6%	0.92[0.24,3.48]	
Makrides 2010	67/1197	88/1202		 	12.74%	0.76[0.56,1.04]	
Malcolm 2003	0/31	1/32	\leftarrow	+	0.21%	0.34[0.01,8.13]	
Miller 2016	3/60	10/55	\leftarrow		1.51%	0.28[0.08,0.95]	
Min 2014	8/60	15/57			2.23%	0.51[0.23,1.1]	
Min 2016	12/58	5/56		-	0.74%	2.32[0.87,6.15]	
Oken 2013	0/31	2/17	-		0.46%	0.11[0.01,2.22]	
Ramakrishnan 2010	49/487	40/486		+	5.81%	1.22[0.82,1.82]	
Smuts 2003a	14/142	17/149			2.41%	0.86[0.44,1.69]	
Smuts 2003b	1/18	5/19	\leftarrow	•	0.71%	0.21[0.03,1.64]	
Subtotal (95% CI)	2381	2363		•	29.94%	0.84[0.69,1.02]	
Total events: 173 (Omega-3), 204 (No	omega-3)						
Heterogeneity: Tau²=0; Chi²=16.52, df	=11(P=0.12); I ² =33.4	11%					
Test for overall effect: Z=1.76(P=0.08)							

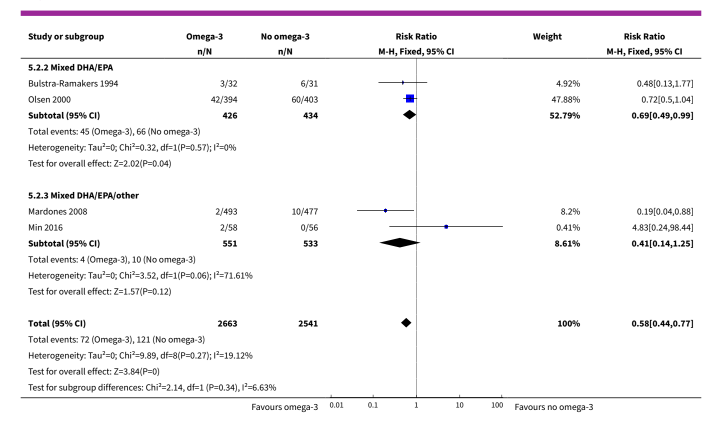




Analysis 5.2. Comparison 5 DHA/mixed subgroups, Outcome 2 Early preterm birth (< 34 weeks).

Study or subgroup	Omega-3	Omega-3 No omega-3		Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fix	ed, 95% CI		M-H, Fixed, 95% CI
5.2.1 DHA/largely DHA						
Carlson 2013	1/154	7/147	+	+	5.78%	0.14[0.02,1.09]
Harris 2015	4/224	7/121		-	7.34%	0.31[0.09,1.03]
Horvaticek 2017	1/51	0/47		+	0.42%	2.77[0.12,66.36]
Makrides 2010	13/1197	27/1202		-	21.74%	0.48[0.25,0.93]
Min 2014	4/60	4/57		+	3.31%	0.95[0.25,3.62]
Subtotal (95% CI)	1686	1574	•		38.59%	0.46[0.28,0.76]
Total events: 23 (Omega-3), 45 (N	lo omega-3)					
Heterogeneity: Tau ² =0; Chi ² =4.1,	df=4(P=0.39); I ² =2.41%					
Test for overall effect: Z=3.07(P=	0)					
		Favours omega-3	0.01 0.1	1 10	100 Favours no omega-3	

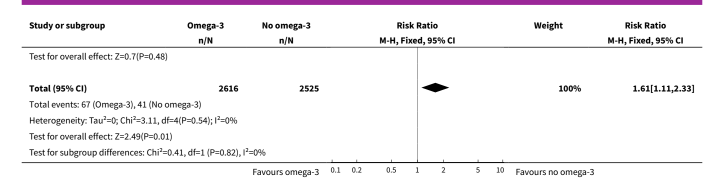




Analysis 5.3. Comparison 5 DHA/mixed subgroups, Outcome 3 Prolonged gestation (> 42 weeks).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
5.3.1 DHA/largely DHA					
Harris 2015	2/224	0/121		1.54%	2.71[0.13,56.02]
Makrides 2010	6/1184	3/1183	- • 	7.14%	2[0.5,7.97]
Mulder 2014	0/68	0/67			Not estimable
Subtotal (95% CI)	1476	1371		8.68%	2.12[0.6,7.49]
Total events: 8 (Omega-3), 3 (No	omega-3)				
Heterogeneity: Tau ² =0; Chi ² =0.03	3, df=1(P=0.86); I ² =0%				
Test for overall effect: Z=1.17(P=0	0.24)				
5.3.2 Mixed DHA/EPA					
Olsen 1992	32/266	27/267	- 	64.13%	1.19[0.73,1.93]
Olsen 2000	26/782	11/791		26.02%	2.39[1.19,4.8]
Subtotal (95% CI)	1048	1058	•	90.15%	1.54[1.04,2.28]
Total events: 58 (Omega-3), 38 (N	lo omega-3)				
Heterogeneity: Tau ² =0; Chi ² =2.62	2, df=1(P=0.11); I ² =61.81	%			
Test for overall effect: Z=2.14(P=0	0.03)				
5.3.3 Mixed DHA/EPA/other					
Hauner 2012	1/92	0/96	+	1.16%	3.13[0.13,75.84]
Subtotal (95% CI)	92	96		1.16%	3.13[0.13,75.84]
Total events: 1 (Omega-3), 0 (No	omega-3)				
Heterogeneity: Not applicable					

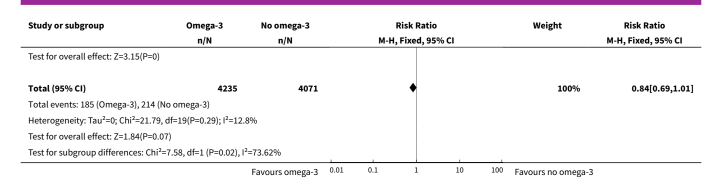




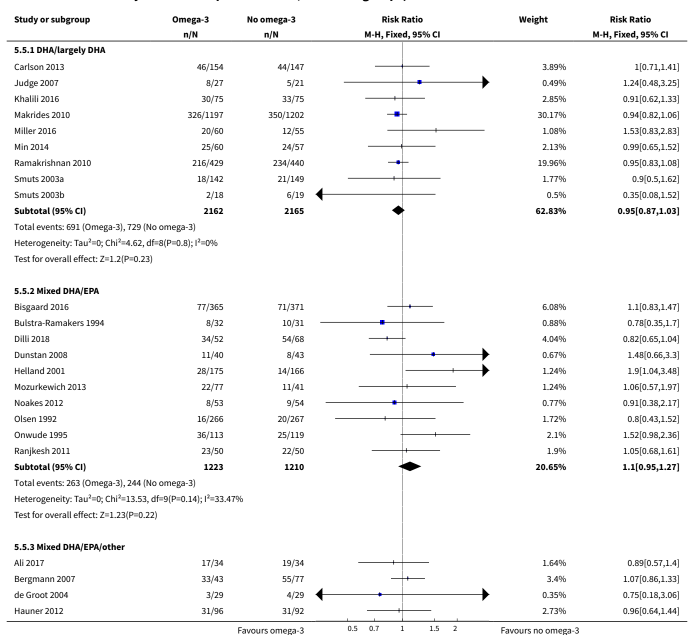
Analysis 5.4. Comparison 5 DHA/mixed subgroups, Outcome 4 Pre-eclampsia (hypertension with proteinuria).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
5.4.1 DHA/largely DHA					
Carlson 2013	2/154	2/147		0.94%	0.95[0.14,6.69]
Harris 2015	2/224	0/121	+	0.3%	2.71[0.13,56.02]
Horvaticek 2017	4/43	5/38		2.44%	0.71[0.2,2.44]
Makrides 2010	60/1197	58/1202	+	26.55%	1.04[0.73,1.48]
Smuts 2003a	5/142	10/149		4.48%	0.52[0.18,1.5]
Smuts 2003b	1/18	0/19		- 0.22%	3.16[0.14,72.84]
Subtotal (95% CI)	1778	1676	•	34.92%	0.98[0.71,1.33]
Total events: 74 (Omega-3), 75 ((No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =2.7	7, df=5(P=0.75); I ² =0%				
Test for overall effect: Z=0.16(P=	=0.88)				
5.4.2 Mixed DHA/EPA					
Bisgaard 2016	16/365	15/371	-	6.82%	1.08[0.54,2.16]
Bulstra-Ramakers 1994	5/32	3/31		1.4%	1.61[0.42,6.19]
Harper 2010	20/434	20/418		9.35%	0.96[0.53,1.76]
Jamilian 2016	0/27	1/27	• · · · · · · · · · · · · · · · · · · ·	0.69%	0.33[0.01,7.84]
Mozurkewich 2013	10/77	5/41		2.99%	1.06[0.39,2.91]
Olsen 1992	0/266	5/267		2.52%	0.09[0.01,1.64]
Olsen 2000	25/398	23/420	- •-	10.27%	1.15[0.66,1.99]
Onwude 1995	15/113	18/119	-	8.04%	0.88[0.47,1.66]
Ranjkesh 2011	2/50	10/50		4.59%	0.2[0.05,0.87]
Subtotal (95% CI)	1762	1744	•	46.67%	0.9[0.69,1.18]
Total events: 93 (Omega-3), 100	(No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =8.7	74, df=8(P=0.36); I ² =8.46%	ı			
Test for overall effect: Z=0.75(P=	=0.45)				
5.4.3 Mixed DHA/EPA/other					
D'Almedia 1992	2/50	5/50		2.29%	0.4[0.08,1.97]
Mardones 2008	8/493	16/477		7.46%	0.48[0.21,1.12]
Razavi 2017	5/60	3/30		1.83%	0.83[0.21,3.26]
Rivas-Echeverria 2000	1/63	14/64		6.37%	0.07[0.01,0.54]
Taghizadeh 2016	2/29	1/30		0.45%	2.07[0.2,21.6]
Subtotal (95% CI)	695	651	◆	18.41%	0.4[0.23,0.71]
Total events: 18 (Omega-3), 39 (No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =5.9	95, df=4(P=0.2); I ² =32.8%				

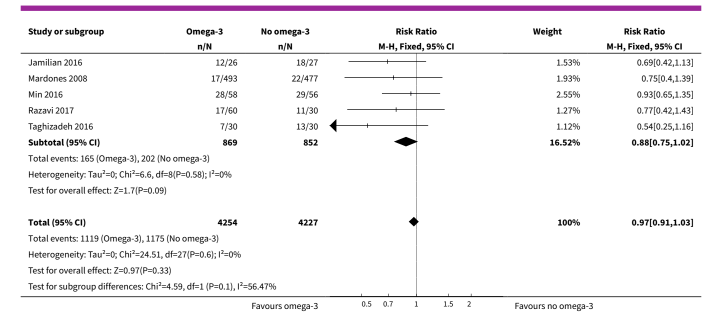




Analysis 5.5. Comparison 5 DHA/mixed subgroups, Outcome 5 Caesarean section.



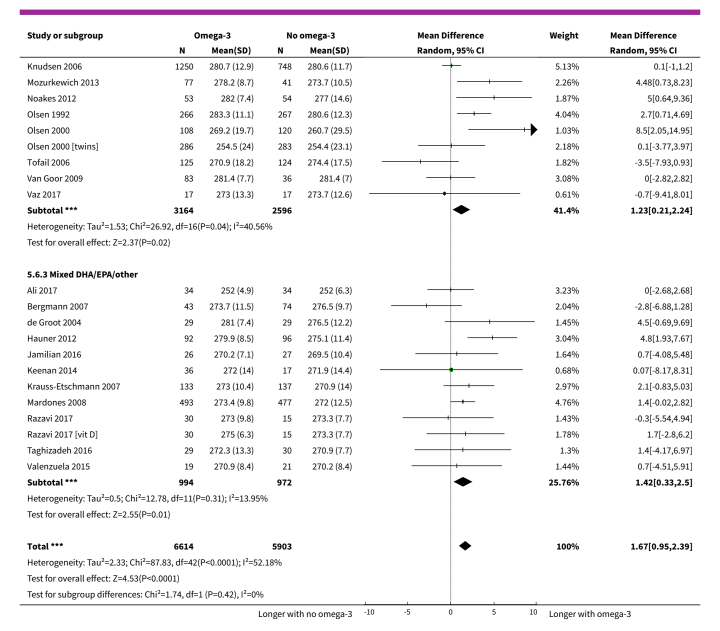




Analysis 5.6. Comparison 5 DHA/mixed subgroups, Outcome 6 Gestational length (days).

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
5.6.1 DHA/largely DHA							
Carlson 2013	154	275.7 (11.2)	147	272.8 (17)	+	2.66%	2.9[-0.37,6.17]
Giorlandino 2013	21	268.8 (9.1)	21	255.5 (12.6)	_	0.98%	13.3[6.65,19.95]
Gustafson 2013	22	275.8 (7.7)	24	279.3 (7.7)		1.81%	-3.5[-7.95,0.95]
Harris 2015	224	275.3 (19.5)	121	271.6 (13.2)		2.49%	3.7[0.23,7.17]
Horvaticek 2017	47	271.6 (4.6)	43	266 (4.5)		4.18%	5.6[3.72,7.48]
Judge 2007	27	278 (8.4)	21	274.3 (9)	-	1.54%	3.71[-1.27,8.69]
Krummel 2016	34	275.1 (7.7)	29	275.8 (8.4)		2.09%	-0.7[-4.71,3.31]
Makrides 2010	1184	276.2 (12.4)	1183	274.6 (14.8)	-	5.13%	1.64[0.54,2.74]
Malcolm 2003	31	279.7 (9.5)	29	279.6 (8.5)		1.75%	0.1[-4.46,4.66]
Miller 2016	60	278.9 (7.8)	55	274.4 (14.8)		1.85%	4.5[0.12,8.88]
Ramakrishnan 2010	484	273.7 (13.3)	486	273 (11.9)	+	4.54%	0.7[-0.89,2.29]
Sanjurjo 2004	8	272.2 (4.5)	8	275.7 (11.8)	<u> </u>	0.61%	-3.57[-12.34,5.2]
Smuts 2003a	142	274.1 (13.5)	149	271.6 (15.6)	+	2.59%	2.5[-0.85,5.85]
Smuts 2003b	18	275.1 (11.2)	19	269.5 (15.4)	+	0.62%	5.6[-3.04,14.24]
Subtotal ***	2456		2335		•	32.84%	2.44[0.91,3.98]
Heterogeneity: Tau ² =4.59; Chi ² =	41.39, df=13(P<0.0001); I ² =68	.59%				
Test for overall effect: Z=3.12(P=	=0)						
5.6.2 Mixed DHA/EPA							
Dilli 2018	52	266 (12.6)	68	261.8 (14)	+	1.64%	4.2[-0.58,8.98]
Dunstan 2008	40	275 (6.3)	43	274 (6.6)		3.13%	1[-1.77,3.77]
Furuhjelm 2009	54	280 (9.8)	66	280 (11.2)		2.26%	0[-3.76,3.76]
Haghiac 2015	25	274.4 (11.2)	24	270.9 (8.4)	-	1.32%	3.5[-2.03,9.03]
Harper 2010	434	263.9 (28.4)	418	261.8 (28.4)		2.22%	2.1[-1.71,5.91]
Helland 2001	175	279.6 (9.2)	166	279.2 (9.3)		4.07%	0.4[-1.56,2.36]
Hurtado 2015	44	275.1 (12.6)	46	277.2 (10.5)		1.63%	-2.1[-6.9,2.7]
Khalili 2016	75	275.1 (8.4)	75	274.4 (9.1)		3.1%	0.7[-2.1,3.5]

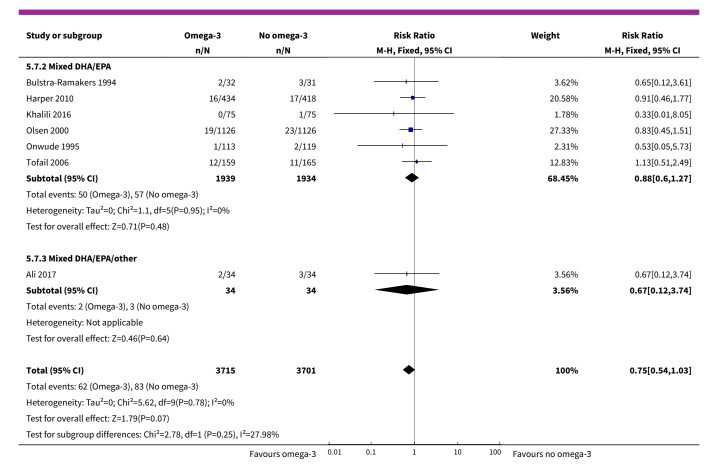




Analysis 5.7. Comparison 5 DHA/mixed subgroups, Outcome 7 Perinatal death.

Study or subgroup	Omega-3	No omega-3		Risk Ratio			Weight	Risk Ratio	
	n/N n/N M-H, Fixed, 95% CI				M-H, Fixed, 95% CI				
5.7.1 DHA/largely DHA									
Horvaticek 2017	1/56	0/43			-			0.67%	2.32[0.1,55.48]
Makrides 2010	3/1197	12/1202						14.23%	0.25[0.07,0.89]
Ramakrishnan 2010	6/489	11/488			+			13.08%	0.54[0.2,1.46]
Subtotal (95% CI)	1742	1733		<	▶			27.98%	0.44[0.21,0.91]
Total events: 10 (Omega-3), 23 (No o	omega-3)								
Heterogeneity: Tau²=0; Chi²=1.99, df	f=2(P=0.37); I ² =0%								
Test for overall effect: Z=2.22(P=0.03	3)								
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3	

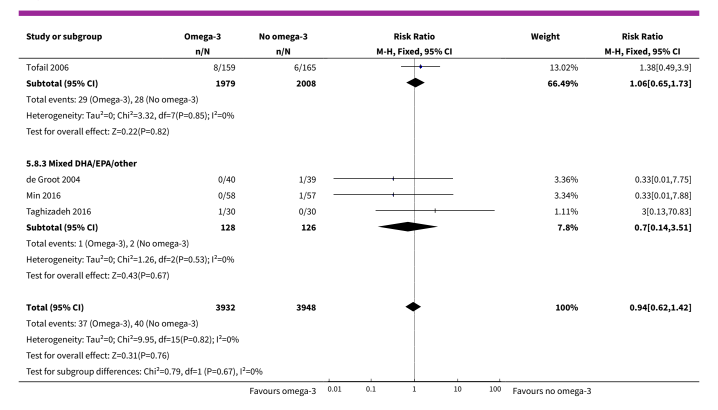




Analysis 5.8. Comparison 5 DHA/mixed subgroups, Outcome 8 Stillbirth.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
5.8.1 DHA/largely DHA					
Horvaticek 2017	1/48	0/50		1.08%	3.12[0.13,74.82]
Makrides 2010	1/1197	7/1202		15.44%	0.14[0.02,1.16]
Min 2014	2/60	0/57		1.13%	4.75[0.23,96.93]
Oken 2013	1/31	0/17		1.41%	1.69[0.07,39.3]
Ramakrishnan 2010	2/489	3/488		6.64%	0.67[0.11,3.96]
Subtotal (95% CI)	1825	1814	*	25.71%	0.69[0.28,1.7]
Total events: 7 (Omega-3), 10 (N	lo omega-3)				
Heterogeneity: Tau²=0; Chi²=4.9	1, df=4(P=0.3); I ² =18.59%)			
Test for overall effect: Z=0.8(P=0	0.42)				
5.8.2 Mixed DHA/EPA					
Bulstra-Ramakers 1994	1/32	0/31		1.12%	2.91[0.12,68.81]
Haghiac 2015	1/25	0/25		1.11%	3[0.13,70.3]
Helland 2001	1/301	0/289		1.13%	2.88[0.12,70.43]
Jamilian 2016	1/27	0/27		1.11%	3[0.13,70.53]
Olsen 1992	1/266	1/267		2.21%	1[0.06,15.96]
Olsen 2000	16/1056	19/1085	-	41.42%	0.87[0.45,1.67]
Onwude 1995	0/113	2/119		5.38%	0.21[0.01,4.34]
		Favours omega-3	0.01 0.1 1 10	100 Favours no omega-3	





Analysis 5.9. Comparison 5 DHA/mixed subgroups, Outcome 9 Neonatal death.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
5.9.1 DHA/largely DHA					
Carlson 2013	1/154	1/147		3.66%	0.95[0.06,15.12]
Makrides 2010	2/1197	5/1202		17.86%	0.4[0.08,2.07]
Ramakrishnan 2010	4/487	8/486		28.67%	0.5[0.15,1.65]
Subtotal (95% CI)	1838	1835		50.2%	0.5[0.2,1.23]
Total events: 7 (Omega-3), 14 (No	omega-3)				
Heterogeneity: Tau ² =0; Chi ² =0.28,	df=2(P=0.87); I ² =0%				
Test for overall effect: Z=1.51(P=0.3	13)				
5.9.2 Mixed DHA/EPA					
Bisgaard 2016	0/365	0/371			Not estimable
Bulstra-Ramakers 1994	1/32	3/31		10.91%	0.32[0.04,2.94]
Khalili 2016	0/75	1/75 —		5.37%	0.33[0.01,8.05]
Olsen 2000	3/1126	4/1144		14.21%	0.76[0.17,3.4]
Onwude 1995	1/113	0/119		— 1.74%	3.16[0.13,76.73]
	•	•	,		
Tofail 2006	4/159	5/165		17.57%	0.83[0.23,3.04]
Subtotal (95% CI)	1870	1905		49.8%	0.73[0.33,1.62]
Total events: 9 (Omega-3), 13 (No o					
Heterogeneity: Tau ² =0; Chi ² =1.61,					
Test for overall effect: Z=0.78(P=0.4	44)				
Total (95% CI)	3708	3740	•	100%	0.61[0.34,1.11]
Total events: 16 (Omega-3), 27 (No	omega-3)				
		Favours omega-3 0.01	0.1 1 10	100 Favours no omega-3	}



Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI				M-H, Fixed, 95% CI		
Heterogeneity: Tau ² =0; Chi ² =2	2.24, df=7(P=0.95); I ² =0%								
Test for overall effect: Z=1.61(P=0.11)								
Test for subgroup differences:	Chi ² =0.38, df=1 (P=0.54),	12=0%							
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3	

Analysis 5.10. Comparison 5 DHA/mixed subgroups, Outcome 10 Low birthweight (< 2500 g).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI	
5.10.1 DHA/largely DHA						
Carlson 2013	6/154	13/147		2.03%	0.44[0.17,1.13]	
Makrides 2010	41/1197	63/1202		9.58%	0.65[0.44,0.96]	
Min 2014	8/60	8/57		1.25%	0.95[0.38,2.36]	
Ramakrishnan 2010	27/487	27/486	+	4.12%	1[0.59,1.68]	
Smuts 2003a	13/142	16/149	 -	2.38%	0.85[0.43,1.71]	
Smuts 2003b	0/18	5/19		0.82%	0.1[0.01,1.62]	
Subtotal (95% CI)	2058	2060	•	20.18%	0.72[0.56,0.93]	
Total events: 95 (Omega-3), 132	(No omega-3)					
Heterogeneity: Tau ² =0; Chi ² =5.3	-					
Test for overall effect: Z=2.51(P=						
5.10.2 Mixed DHA/EPA						
Bulstra-Ramakers 1994	11/32	9/31		1.39%	1.18[0.57,2.46]	
Harper 2010	94/427	112/410	+	17.42%	0.81[0.63,1.02]	
Khalili 2016	0/75	5/75		0.84%	0.09[0.01,1.62]	
Olsen 2000	283/799	287/817	•	43.26%	1.01[0.88,1.15]	
Onwude 1995	33/113	35/119		5.2%	0.99[0.67,1.48]	
Tofail 2006	36/125	30/124	 	4.59%	1.19[0.79,1.8]	
Subtotal (95% CI)	1571	1576		72.7%	0.96[0.87,1.07]	
Total events: 457 (Omega-3), 478					. , .	
Heterogeneity: Tau ² =0; Chi ² =6.52	- ·	%				
Test for overall effect: Z=0.7(P=0						
5.10.3 Mixed DHA/EPA/other						
D'Almedia 1992	2/50	5/50		0.76%	0.4[0.08,1.97]	
Mardones 2008	27/493	37/477		5.73%	0.71[0.44,1.14]	
Min 2016	8/58	4/56		0.62%	1.93[0.62,6.05]	
Subtotal (95% CI)	601	583	•	7.12%	0.78[0.51,1.18]	
Total events: 37 (Omega-3), 46 (N	lo omega-3)				. , .	
Heterogeneity: Tau ² =0; Chi ² =3.26						
Test for overall effect: Z=1.16(P=						
Total (95% CI)	4230	4219	•	100%	0.9[0.82,0.99]	
Total events: 589 (Omega-3), 656	6 (No omega-3)				. ,	
Heterogeneity: Tau ² =0; Chi ² =19.		11%				
Test for overall effect: Z=2.12(P=						
Test for subgroup differences: Ch	•	=58 12%				



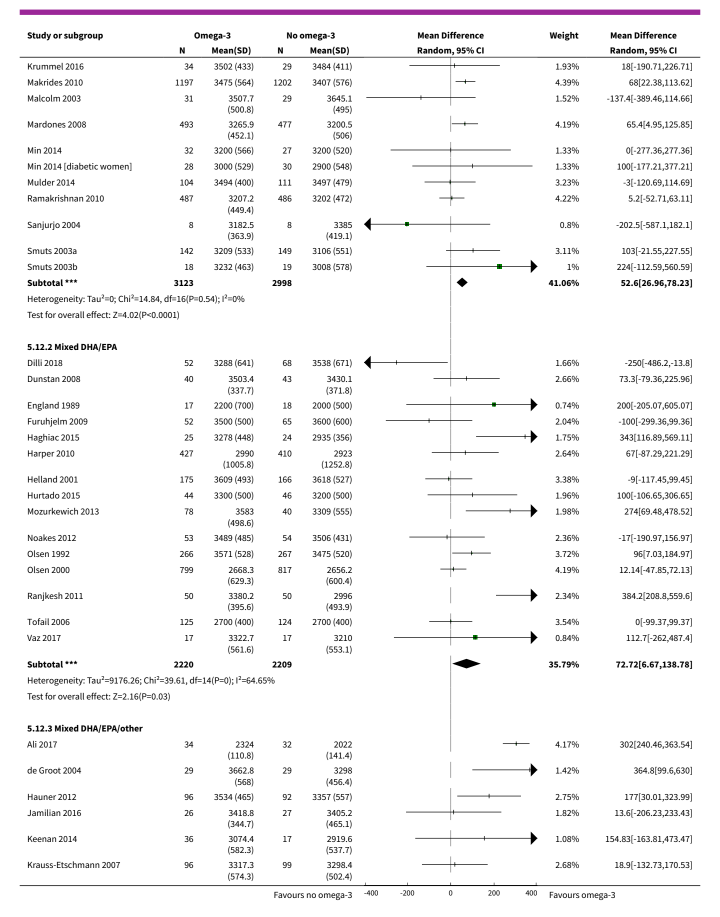
Analysis 5.11. Comparison 5 DHA/mixed subgroups, Outcome 11 Small-for-gestational age/IUGR.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N		M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
5.11.1 DHA/largely DHA					
Makrides 2010	73/1197	82/1202	+	18.39%	0.89[0.66,1.21]
Ramakrishnan 2010	55/487	53/486	+	11.92%	1.04[0.73,1.48]
Subtotal (95% CI)	1684	1688	*	30.31%	0.95[0.75,1.2]
Total events: 128 (Omega-3), 135	(No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =0.38	8, df=1(P=0.54); I ² =0%				
Test for overall effect: Z=0.44(P=	0.66)				
5.11.2 Mixed DHA/EPA					
Bulstra-Ramakers 1994	12/32	9/31	- 	2.05%	1.29[0.64,2.63]
Harper 2010	35/427	41/410		9.4%	0.82[0.53,1.26]
Olsen 2000	208/685	185/689	+	41.45%	1.13[0.96,1.34]
Onwude 1995	33/113	35/119	+	7.66%	0.99[0.67,1.48]
Subtotal (95% CI)	1257	1249	*	60.56%	1.07[0.93,1.23]
Total events: 288 (Omega-3), 270) (No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =2.3,	df=3(P=0.51); I ² =0%				
Test for overall effect: Z=0.94(P=	0.35)				
5.11.3 Mixed EPA/DHA/other					
Mardones 2008	30/493	39/477	-+	8.91%	0.74[0.47,1.18]
Taghizadeh 2016	2/29	1/30		0.22%	2.07[0.2,21.6]
Subtotal (95% CI)	522	507	*	9.13%	0.78[0.5,1.22]
Total events: 32 (Omega-3), 40 (N	No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =0.7,	df=1(P=0.4); I ² =0%				
Test for overall effect: Z=1.11(P=	0.27)				
Total (95% CI)	3463	3444	•	100%	1.01[0.9,1.13]
Total events: 448 (Omega-3), 445	5 (No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =5.8 ⁴	4, df=7(P=0.56); I ² =0%				
Test for overall effect: Z=0.12(P=	0.91)				
Test for subgroup differences: Ch	ni²=2.23, df=1 (P=0.33), I²	=10.21%			

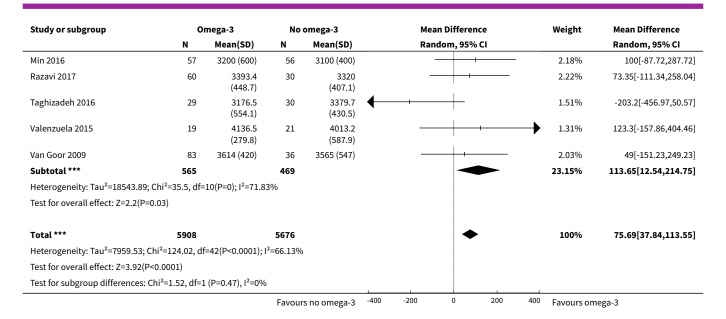
Analysis 5.12. Comparison 5 DHA/mixed subgroups, Outcome 12 Birthweight (g).

Study or subgroup	0	mega-3	No	omega-3		Ме	an Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Ra	ndom, 95% CI			Random, 95% CI
5.12.1 DHA/largely DHA										
Carlson 2013	154	3359 (524)	147	3187 (602)				_	3.06%	172[44.25,299.75]
Gustafson 2013	22	3416.8 (552.9)	24	3435.5 (404.8)					1.3%	-18.7[-300.85,263.45]
Harris 2015	224	3215.5 (506.2)	121	3165 (494.6)			+		3.35%	50.5[-59.77,160.77]
Horvaticek 2017	47	3580.9 (568)	43	3456.9 (575.8)			+		1.66%	124[-112.62,360.62]
Judge 2007	27	3394 (430)	21	3224.6 (431.2)			-	-	1.58%	169.38[-76.22,414.98]
Khalili 2016	75	3260 (360)	75	3230 (430)					3.07%	30[-96.92,156.92]
			Favour	s no omega-3	-400	-200	0 200	400	Favours om	ega-3









Comparison 6. Risk subgroups

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Preterm birth (< 37 weeks)	27	10304	Risk Ratio (M-H, Fixed, 95% CI)	0.89 [0.81, 0.97]
1.1 Increased/high risk	12	3702	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.83, 1.03]
1.2 Low risk	10	3241	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.71, 1.20]
1.3 Any/mixed risk	5	3361	Risk Ratio (M-H, Fixed, 95% CI)	0.71 [0.54, 0.93]
2 Early preterm birth (< 34 weeks)	10	5204	Risk Ratio (M-H, Fixed, 95% CI)	0.58 [0.44, 0.77]
2.1 Increased/high risk	6	2104	Risk Ratio (M-H, Fixed, 95% CI)	0.67 [0.49, 0.93]
2.2 Low risk	3	701	Risk Ratio (M-H, Fixed, 95% CI)	0.31 [0.12, 0.79]
2.3 Any/mixed risk	1	2399	Risk Ratio (M-H, Fixed, 95% CI)	0.48 [0.25, 0.93]
3 Prolonged gestation (> 42 weeks)	6	5141	Risk Ratio (M-H, Fixed, 95% CI)	1.61 [1.11, 2.33]
3.1 Increased/high risk	1	1573	Risk Ratio (M-H, Fixed, 95% CI)	2.39 [1.19, 4.80]
3.2 Low risk	4	1201	Risk Ratio (M-H, Fixed, 95% CI)	1.26 [0.79, 2.01]
3.3 Any/mixed risk	1	2367	Risk Ratio (M-H, Fixed, 95% CI)	2.00 [0.50, 7.97]
4 Pre-eclampsia (hyper- tension with proteinuria)	20	8306	Risk Ratio (M-H, Fixed, 95% CI)	0.84 [0.69, 1.01]

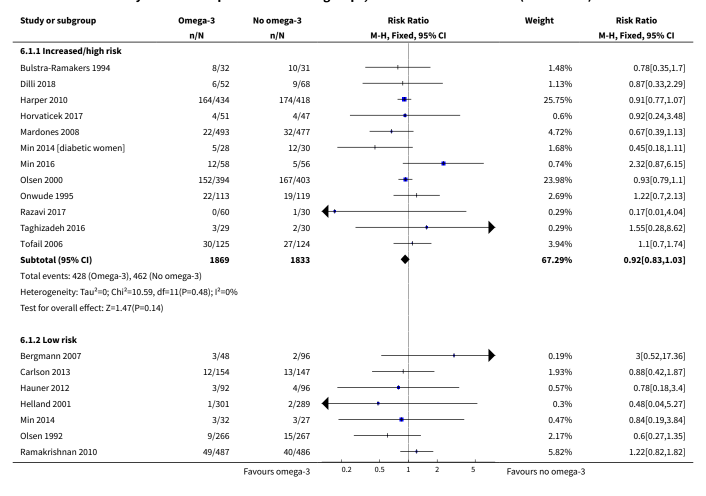


Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
4.1 Increased/high risk	12	3564	Risk Ratio (M-H, Fixed, 95% CI)	0.76 [0.59, 0.99]
4.2 Low risk	5	1507	Risk Ratio (M-H, Fixed, 95% CI)	0.59 [0.28, 1.24]
4.3 Any/mixed risk	3	3235	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.74, 1.37]
5 Caesarean section	29	8481	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.91, 1.03]
5.1 Increased/high risk	12	2046	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.80, 1.05]
5.2 Low risk	14	3185	Risk Ratio (M-H, Fixed, 95% CI)	0.99 [0.89, 1.09]
5.3 Any/mixed risk	3	3250	Risk Ratio (M-H, Fixed, 95% CI)	0.98 [0.87, 1.10]
6 Length of gestation (days)	43	12517	Mean Difference (IV, Random, 95% CI)	1.67 [0.95, 2.39]
6.1 Increased/high risk	18	3707	Mean Difference (IV, Random, 95% CI)	2.17 [0.65, 3.68]
6.2 Low risk	22	4330	Mean Difference (IV, Random, 95% CI)	1.41 [0.52, 2.29]
6.3 Any/mixed group	3	4480	Mean Difference (IV, Random, 95% CI)	1.27 [-0.36, 2.91]
7 Perinatal death	10	7416	Risk Ratio (M-H, Fixed, 95% CI)	0.75 [0.54, 1.03]
7.1 Increased/high risk	6	3566	Risk Ratio (M-H, Fixed, 95% CI)	0.84 [0.56, 1.26]
7.2 Low risk	2	1127	Risk Ratio (M-H, Fixed, 95% CI)	0.52 [0.20, 1.33]
7.3 Any/mixed risk	2	2723	Risk Ratio (M-H, Fixed, 95% CI)	0.67 [0.35, 1.26]
8 Stillbirth	16	7880	Risk Ratio (M-H, Fixed, 95% CI)	0.94 [0.62, 1.42]
8.1 Increased/high risk	9	3137	Risk Ratio (M-H, Fixed, 95% CI)	1.06 [0.65, 1.72]
8.2 Low risk	5	2296	Risk Ratio (M-H, Fixed, 95% CI)	1.13 [0.40, 3.23]
8.3 Any/mixed risk	2	2447	Risk Ratio (M-H, Fixed, 95% CI)	0.27 [0.06, 1.27]
9 Neonatal death	9	7448	Risk Ratio (M-H, Fixed, 95% CI)	0.61 [0.34, 1.11]
9.1 Increased/high risk	4	2889	Risk Ratio (M-H, Fixed, 95% CI)	0.78 [0.34, 1.78]
9.2 Low risk	3	1424	Risk Ratio (M-H, Fixed, 95% CI)	0.52 [0.19, 1.45]
9.3 Any/mixed risk	2	3135	Risk Ratio (M-H, Fixed, 95% CI)	0.40 [0.08, 2.07]
10 Low birthweight (< 2500 g)	15	8449	Risk Ratio (M-H, Fixed, 95% CI)	0.90 [0.82, 0.99]
10.1 Increased/high risk	7	4081	Risk Ratio (M-H, Fixed, 95% CI)	0.96 [0.87, 1.07]
10.2 Low risk	6	1869	Risk Ratio (M-H, Fixed, 95% CI)	0.73 [0.52, 1.02]

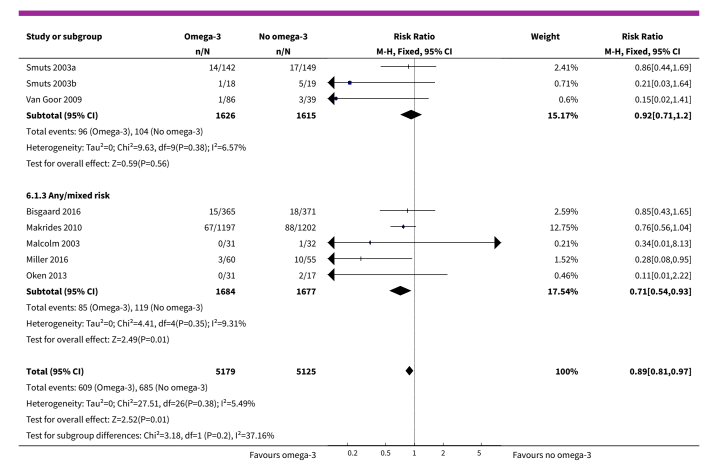


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
10.3 Any/mixed risk	2	2499	Risk Ratio (M-H, Fixed, 95% CI)	0.63 [0.44, 0.92]
11 Small-for-gestational age/IUGR	8	6907	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.90, 1.13]
11.1 Increased/high risk	6	3535	Risk Ratio (M-H, Fixed, 95% CI)	1.03 [0.90, 1.18]
11.2 Low risk	1	973	Risk Ratio (M-H, Fixed, 95% CI)	1.04 [0.73, 1.48]
11.3 Any/mixed risk	1	2399	Risk Ratio (M-H, Fixed, 95% CI)	0.89 [0.66, 1.21]
12 Birthweight (g)	43	11584	Mean Difference (IV, Random, 95% CI)	75.69 [37.84, 113.55]
12.1 Increased/high risk	19	4848	Mean Difference (IV, Random, 95% CI)	105.52 [30.84, 180.21]
12.2 Low risk	23	4337	Mean Difference (IV, Random, 95% CI)	46.63 [13.90, 79.36]
12.3 Any/mixed group	1	2399	Mean Difference (IV, Random, 95% CI)	68.0 [22.38, 113.62]

Analysis 6.1. Comparison 6 Risk subgroups, Outcome 1 Preterm birth (< 37 weeks).



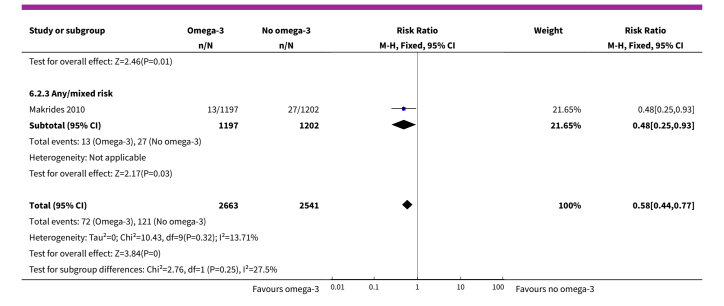




Analysis 6.2. Comparison 6 Risk subgroups, Outcome 2 Early preterm birth (< 34 weeks).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
6.2.1 Increased/high risk					
Bulstra-Ramakers 1994	3/32	6/31		4.9%	0.48[0.13,1.77]
Horvaticek 2017	1/51	0/47		0.42%	2.77[0.12,66.36]
Mardones 2008	2/493	10/477		8.17%	0.19[0.04,0.88]
Min 2014 [diabetic women]	3/32	4/30		3.32%	0.7[0.17,2.88]
Min 2016	2/58	0/56	-	0.41%	4.83[0.24,98.44]
Olsen 2000	42/394	60/403		47.67%	0.72[0.5,1.04]
Subtotal (95% CI)	1060	1044	•	64.88%	0.67[0.49,0.93]
Total events: 53 (Omega-3), 80 (No	o omega-3)				
Heterogeneity: Tau ² =0; Chi ² =5.37,	df=5(P=0.37); I ² =6.94%				
Test for overall effect: Z=2.4(P=0.0	2)				
6.2.2 Low risk					
Carlson 2013	1/154	7/147		5.76%	0.14[0.02,1.09]
Harris 2015	4/224	7/121		7.3%	0.31[0.09,1.03]
Min 2014	1/28	0/27		- 0.41%	2.9[0.12,68.15]
Subtotal (95% CI)	406	295	•	13.47%	0.31[0.12,0.79]
Total events: 6 (Omega-3), 14 (No	omega-3)				
Heterogeneity: Tau ² =0; Chi ² =2.52,	df=2(P=0.28); I ² =20.58 ^c	%			
		Favours omega-3	0.01 0.1 1 10	100 Favours no omega-3	





Analysis 6.3. Comparison 6 Risk subgroups, Outcome 3 Prolonged gestation (> 42 weeks).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
6.3.1 Increased/high risk					
Olsen 2000	26/782	11/791		26.02%	2.39[1.19,4.8]
Subtotal (95% CI)	782	791	•	26.02%	2.39[1.19,4.8]
Total events: 26 (Omega-3), 11 (No	omega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=2.45(P=0.0	01)				
6.3.2 Low risk					
Harris 2015	2/224	0/121		1.54%	2.71[0.13,56.02]
Hauner 2012	1/92	0/96		1.16%	3.13[0.13,75.84]
Mulder 2014	0/68	0/67			Not estimable
Olsen 1992	32/266	27/267		64.13%	1.19[0.73,1.93]
Subtotal (95% CI)	650	551	*	66.83%	1.26[0.79,2.01]
Total events: 35 (Omega-3), 27 (No	omega-3)				
Heterogeneity: Tau ² =0; Chi ² =0.61, o	df=2(P=0.74); I ² =0%				
Test for overall effect: Z=0.96(P=0.3	34)				
6.3.3 Any/mixed risk					
Makrides 2010	6/1184	3/1183	+	7.14%	2[0.5,7.97]
Subtotal (95% CI)	1184	1183		7.14%	2[0.5,7.97]
Total events: 6 (Omega-3), 3 (No or	nega-3)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.98(P=0.3	33)				
Total (95% CI)	2616	2525	•	100%	1.61[1.11,2.33]
Total events: 67 (Omega-3), 41 (No	omega-3)				
Heterogeneity: Tau ² =0; Chi ² =3.11, o	df=4(P=0.54); I ² =0%				
Test for overall effect: Z=2.49(P=0.0	01)				
Test for subgroup differences: Chi ²	=2.36, df=1 (P=0.31), I ²	=15.24%			



Analysis 6.4. Comparison 6 Risk subgroups, Outcome 4 Pre-eclampsia (hypertension with proteinuria).

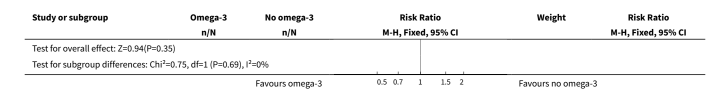
Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
6.4.1 Increased/high risk					
Bulstra-Ramakers 1994	5/32	3/31		1.4%	1.61[0.42,6.19]
Harper 2010	20/434	20/418	-	9.35%	0.96[0.53,1.76]
Horvaticek 2017	4/43	5/38		2.44%	0.71[0.2,2.44]
Jamilian 2016	0/27	1/27 —	•	0.69%	0.33[0.01,7.84]
Mardones 2008	8/493	16/477		7.46%	0.48[0.21,1.12]
Mozurkewich 2013	10/77	5/41		2.99%	1.06[0.39,2.91]
Olsen 2000	25/398	23/420		10.27%	1.15[0.66,1.99]
Onwude 1995	15/113	18/119		8.04%	0.88[0.47,1.66]
Ranjkesh 2011	2/50	10/50		4.59%	0.2[0.05,0.87]
Razavi 2017	5/60	3/30		1.83%	0.83[0.21,3.26]
Rivas-Echeverria 2000	1/63	14/64		6.37%	0.07[0.01,0.54]
Taghizadeh 2016	2/29	1/30		0.45%	2.07[0.2,21.6]
Subtotal (95% CI)	1819	1745	•	55.88%	0.76[0.59,0.99]
Total events: 97 (Omega-3), 119					
Heterogeneity: Tau ² =0; Chi ² =15.1	- ·	7%			
Test for overall effect: Z=2.03(P=					
	,				
6.4.2 Low risk					
Carlson 2013	2/154	2/147		0.94%	0.95[0.14,6.69]
Harris 2015	2/224	0/121		- 0.3%	2.71[0.13,56.02]
Olsen 1992	0/266	5/267		2.52%	0.09[0.01,1.64]
Smuts 2003a	5/142	10/149		4.48%	0.52[0.18,1.5]
Smuts 2003b	1/18	0/19		- 0.22%	3.16[0.14,72.84]
Subtotal (95% CI)	804	703		8.45%	0.59[0.28,1.24]
Total events: 10 (Omega-3), 17 (N					
Heterogeneity: Tau ² =0; Chi ² =3.96	- '				
Test for overall effect: Z=1.39(P=					
	,				
6.4.3 Any/mixed risk					
Bisgaard 2016	16/365	15/371		6.82%	1.08[0.54,2.16]
D'Almedia 1992	2/50	5/50		2.29%	0.4[0.08,1.97]
Makrides 2010	60/1197	58/1202	-	26.55%	1.04[0.73,1.48]
Subtotal (95% CI)	1612	1623	•	35.67%	1.01[0.74,1.37]
Total events: 78 (Omega-3), 78 (N					
Heterogeneity: Tau ² =0; Chi ² =1.37					
Test for overall effect: Z=0.04(P=					
1111101 010101 011001 Z 0104(1 -1	/				
Total (95% CI)	4235	4071	•	100%	0.84[0.69,1.01]
Total events: 185 (Omega-3), 214					- ,
Heterogeneity: Tau ² =0; Chi ² =21.7	· -	%			
Test for overall effect: Z=1.84(P=					
	ni ² =2.72, df=1 (P=0.26), I ² :	-26 420%			



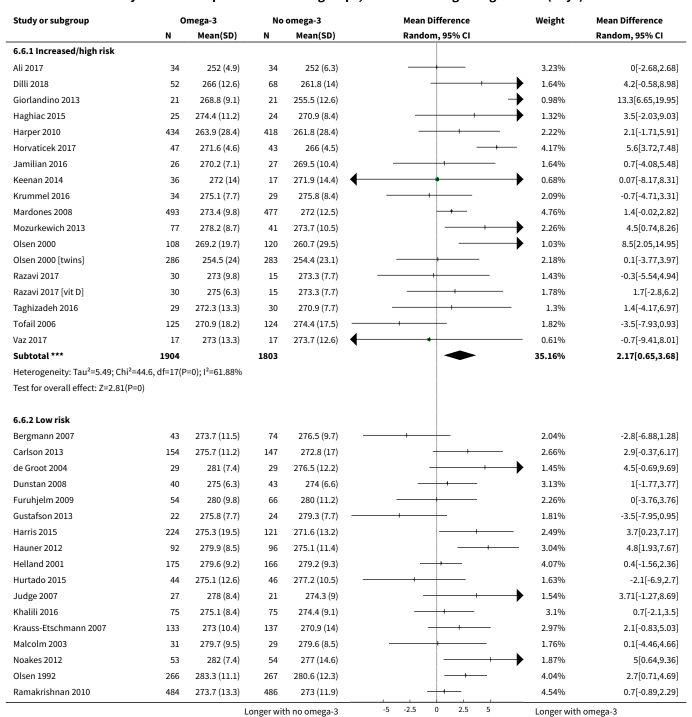
Analysis 6.5. Comparison 6 Risk subgroups, Outcome 5 Caesarean section.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
6.5.1 Increased/high risk	17/24	10/24		1.040/	0.00[0.57.1.4]
Ali 2017	17/34	19/34		1.64%	0.89[0.57,1.4]
Bulstra-Ramakers 1994	8/32	10/31		0.88%	0.78[0.35,1.7]
Dilli 2018	34/52	54/68		4.04%	0.82[0.65,1.04]
Jamilian 2016	12/26	18/27		1.53%	0.69[0.42,1.13]
Mardones 2008	17/493	22/477		1.93%	0.75[0.4,1.39]
Min 2014 [diabetic women]	16/28	17/30		1.42%	1.01[0.64,1.58]
Min 2016	28/58	29/56		2.55%	0.93[0.65,1.35]
Mozurkewich 2013	22/77	11/41	- -	1.24%	1.06[0.57,1.97]
Onwude 1995	36/113	25/119	 	2.1%	1.52[0.98,2.36]
Ranjkesh 2011	23/50	22/50		1.9%	1.05[0.68,1.61]
Razavi 2017	17/60	11/30		1.27%	0.77[0.42,1.43]
Taghizadeh 2016	7/30	13/30 —	+ +	1.12%	0.54[0.25,1.16]
Subtotal (95% CI)	1053	993	•	21.63%	0.92[0.8,1.05]
Total events: 237 (Omega-3), 251	(No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =10.63	1, df=11(P=0.48); l ² =0%				
Test for overall effect: Z=1.23(P=0.	.22)				
6.5.2 Low risk					
Bergmann 2007	33/43	55/77		3.41%	1.07[0.86,1.33]
Carlson 2013	46/154	44/147		3.89%	1[0.71,1.41]
de Groot 2004	3/29	4/29	•	0.35%	0.75[0.18,3.06]
Dunstan 2008	11/40	8/43		0.67%	1.48[0.66,3.3]
Hauner 2012	31/96	31/92		2.74%	0.96[0.64,1.44]
Helland 2001	28/175	14/166		1.24%	1.9[1.04,3.48]
Judge 2007	8/27	5/21		0.49%	1.24[0.48,3.25]
Khalili 2016	30/75	33/75		2.85%	0.91[0.62,1.33]
Min 2014	9/32	7/27		0.66%	1.08[0.47,2.52]
Noakes 2012	8/53	9/54		0.77%	0.91[0.38,2.17]
Olsen 1992	16/266	20/267		1.73%	0.8[0.43,1.52]
Ramakrishnan 2010	216/429	234/440	-	19.97%	0.95[0.83,1.08]
Smuts 2003a	18/142	21/149		1.77%	0.9[0.5,1.62]
Smuts 2003b	2/18	6/19		0.5%	0.35[0.08,1.52]
Subtotal (95% CI)	1579	1606	<u> </u>	41.02%	0.99[0.89,1.09]
Total events: 459 (Omega-3), 491		1000	Ĭ	41.02 /0	0.55[0.05,1.05]
Heterogeneity: Tau ² =0; Chi ² =9.51,					
Test for overall effect: Z=0.29(P=0.					
6.5.3 Any/mixed risk					
Bisgaard 2016	77/365	71/371		6.09%	1.1[0.83,1.47]
Makrides 2010	326/1197	350/1202		30.18%	0.94[0.82,1.06]
Miller 2016	20/60			1.08%	
	•	12/55 1628		1.08% 37.35%	1.53[0.83,2.83]
Subtotal (95% CI)	1622	1628	T	31.35%	0.98[0.87,1.1]
Total events: 423 (Omega-3), 433					
Heterogeneity: Tau ² =0; Chi ² =3.15, Test for overall effect: Z=0.35(P=0.					
Total (95% CI)	4254	4227		100%	0.97[0.91,1.03]
Total events: 1119 (Omega-3), 117		4221	T	100%	0.51[0.51,1.03]
Total events. 1113 (Onnega-3), 117	2 (140 OHIERA-2)				

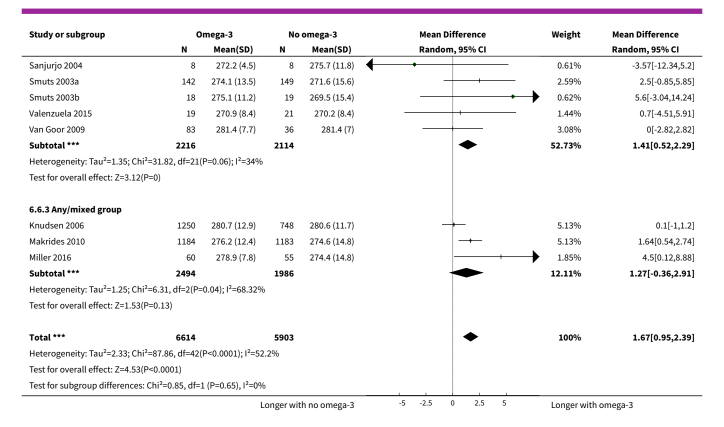




Analysis 6.6. Comparison 6 Risk subgroups, Outcome 6 Length of gestation (days).



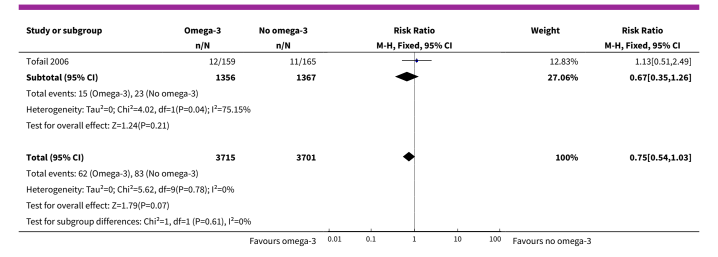




Analysis 6.7. Comparison 6 Risk subgroups, Outcome 7 Perinatal death.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
6.7.1 Increased/high risk					
Ali 2017	2/34	3/34		3.56%	0.67[0.12,3.74]
Bulstra-Ramakers 1994	2/32	3/31		3.62%	0.65[0.12,3.61]
Harper 2010	16/434	17/418	-	20.58%	0.91[0.46,1.77]
Horvaticek 2017	1/56	0/43		- 0.67%	2.32[0.1,55.48]
Olsen 2000	19/1126	23/1126	-	27.33%	0.83[0.45,1.51]
Onwude 1995	1/113	2/119		2.31%	0.53[0.05,5.73]
Subtotal (95% CI)	1795	1771	•	58.08%	0.84[0.56,1.26]
Total events: 41 (Omega-3), 48 (No om	nega-3)				
Heterogeneity: Tau ² =0; Chi ² =0.75, df=	5(P=0.98); I ² =0%				
Test for overall effect: Z=0.84(P=0.4)					
6.7.2 Low risk					
Khalili 2016	0/75	1/75 —		1.78%	0.33[0.01,8.05]
Ramakrishnan 2010	6/489	11/488		13.08%	0.54[0.2,1.46]
Subtotal (95% CI)	564	563		14.87%	0.52[0.2,1.33]
Total events: 6 (Omega-3), 12 (No ome	ega-3)				
Heterogeneity: Tau ² =0; Chi ² =0.08, df=	1(P=0.77); I ² =0%				
Test for overall effect: Z=1.37(P=0.17)					
6.7.3 Any/mixed risk					
Makrides 2010	3/1197	12/1202	 	14.23%	0.25[0.07,0.89]
		Favours omega-3 0.01	0.1 1 10	100 Favours no omega-3	

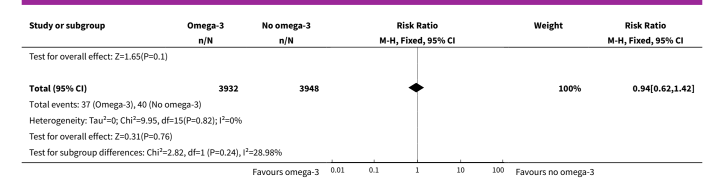




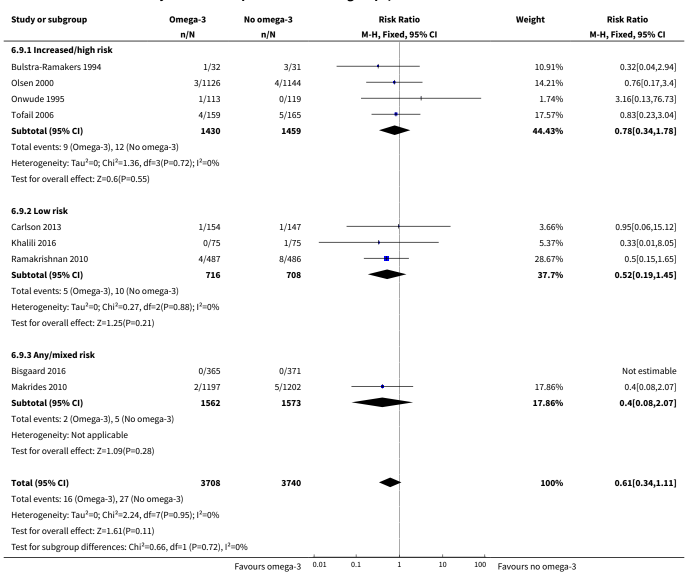
Analysis 6.8. Comparison 6 Risk subgroups, Outcome 8 Stillbirth.

	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
6.8.1 Increased/high risk					
Bulstra-Ramakers 1994	1/32	0/31		1.12%	2.91[0.12,68.81]
Haghiac 2015	1/25	0/25		1.11%	3[0.13,70.3]
Horvaticek 2017	1/48	0/50	+	1.08%	3.12[0.13,74.82]
Jamilian 2016	1/27	0/27		1.11%	3[0.13,70.53]
Min 2016	0/58	1/57 —	+	3.34%	0.33[0.01,7.88]
Olsen 2000	16/1056	19/1085	- ■	41.42%	0.87[0.45,1.67]
Onwude 1995	0/113	2/119 —	+	5.38%	0.21[0.01,4.34]
Taghizadeh 2016	1/30	0/30		- 1.11%	3[0.13,70.83]
Tofail 2006	8/159	6/165		13.02%	1.38[0.49,3.9]
Subtotal (95% CI)	1548	1589	*	68.69%	1.06[0.65,1.72]
Total events: 29 (Omega-3), 28 ((No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =4.3	32, df=8(P=0.83); I ² =0%				
Test for overall effect: Z=0.23(P=	=0.82)				
6.8.2 Low risk					
de Groot 2004	0/40	1/39 —	+	3.36%	0.33[0.01,7.75]
de Groot 2004 Helland 2001	0/40 1/301	1/39 — 0/289		3.36% - 1.13%	0.33[0.01,7.75] 2.88[0.12,70.43]
	·	•			
Helland 2001	1/301	0/289		1.13%	2.88[0.12,70.43]
Helland 2001 Min 2014	1/301 2/60	0/289 0/57		- 1.13% - 1.13%	2.88[0.12,70.43] 4.75[0.23,96.93]
Helland 2001 Min 2014 Olsen 1992	1/301 2/60 1/266	0/289 0/57 1/267		- 1.13% 1.13% 2.21%	2.88[0.12,70.43] 4.75[0.23,96.93] 1[0.06,15.96]
Helland 2001 Min 2014 Olsen 1992 Ramakrishnan 2010	1/301 2/60 1/266 2/489 1156	0/289 0/57 1/267 3/488		- 1.13% - 1.13% 2.21% 6.64%	2.88[0.12,70.43] 4.75[0.23,96.93] 1[0.06,15.96] 0.67[0.11,3.96]
Helland 2001 Min 2014 Olsen 1992 Ramakrishnan 2010 Subtotal (95% CI)	1/301 2/60 1/266 2/489 1156 o omega-3)	0/289 0/57 1/267 3/488		- 1.13% - 1.13% 2.21% 6.64%	2.88[0.12,70.43] 4.75[0.23,96.93] 1[0.06,15.96] 0.67[0.11,3.96]
Helland 2001 Min 2014 Olsen 1992 Ramakrishnan 2010 Subtotal (95% CI) Total events: 6 (Omega-3), 5 (Nother of the companion	1/301 2/60 1/266 2/489 1156 comega-3) 14, df=4(P=0.71); l ² =0%	0/289 0/57 1/267 3/488		- 1.13% - 1.13% 2.21% 6.64%	2.88[0.12,70.43] 4.75[0.23,96.93] 1[0.06,15.96] 0.67[0.11,3.96]
Helland 2001 Min 2014 Olsen 1992 Ramakrishnan 2010 Subtotal (95% CI) Total events: 6 (Omega-3), 5 (No	1/301 2/60 1/266 2/489 1156 comega-3) 14, df=4(P=0.71); l ² =0%	0/289 0/57 1/267 3/488		- 1.13% - 1.13% 2.21% 6.64%	2.88[0.12,70.43] 4.75[0.23,96.93] 1[0.06,15.96] 0.67[0.11,3.96]
Helland 2001 Min 2014 Olsen 1992 Ramakrishnan 2010 Subtotal (95% CI) Total events: 6 (Omega-3), 5 (Nother the companion of	1/301 2/60 1/266 2/489 1156 comega-3) 14, df=4(P=0.71); l ² =0%	0/289 0/57 1/267 3/488		- 1.13% - 1.13% 2.21% 6.64%	2.88[0.12,70.43] 4.75[0.23,96.93] 1[0.06,15.96] 0.67[0.11,3.96] 1.13[0.4,3.23]
Helland 2001 Min 2014 Olsen 1992 Ramakrishnan 2010 Subtotal (95% CI) Total events: 6 (Omega-3), 5 (Notheterogeneity: Tau²=0; Chi²=2.1 Test for overall effect: Z=0.23(P=0.8.3 Any/mixed risk Makrides 2010	1/301 2/60 1/266 2/489 1156 0 omega-3) 14, df=4(P=0.71); I ² =0% =0.82)	0/289 0/57 1/267 3/488 1140		- 1.13% - 1.13% - 2.21% - 6.64% - 14.46%	2.88[0.12,70.43 4.75[0.23,96.93 1[0.06,15.96 0.67[0.11,3.96 1.13[0.4,3.23]
Helland 2001 Min 2014 Olsen 1992 Ramakrishnan 2010 Subtotal (95% CI) Total events: 6 (Omega-3), 5 (Notero and the second and t	1/301 2/60 1/266 2/489 1156 0 omega-3) 1.4, df=4(P=0.71); I ² =0% =0.82)	0/289 0/57 1/267 3/488 1140		- 1.13% - 1.13% 2.21% 6.64% 14.46%	2.88[0.12,70.43 4.75[0.23,96.93 1[0.06,15.96 0.67[0.11,3.96 1.13[0.4,3.23] 0.14[0.02,1.16 1.69[0.07,39.3
Helland 2001 Min 2014 Olsen 1992 Ramakrishnan 2010 Subtotal (95% CI) Total events: 6 (Omega-3), 5 (Nother of the companion of	1/301 2/60 1/266 2/489 1156 0 omega-3) 14, df=4(P=0.71); l ² =0% =0.82) 1/1197 1/31 1228	0/289 0/57 1/267 3/488 1140		- 1.13% - 1.13% 2.21% 6.64% 14.46% 15.44% 1.41%	2.88[0.12,70.43] 4.75[0.23,96.93] 1[0.06,15.96] 0.67[0.11,3.96]



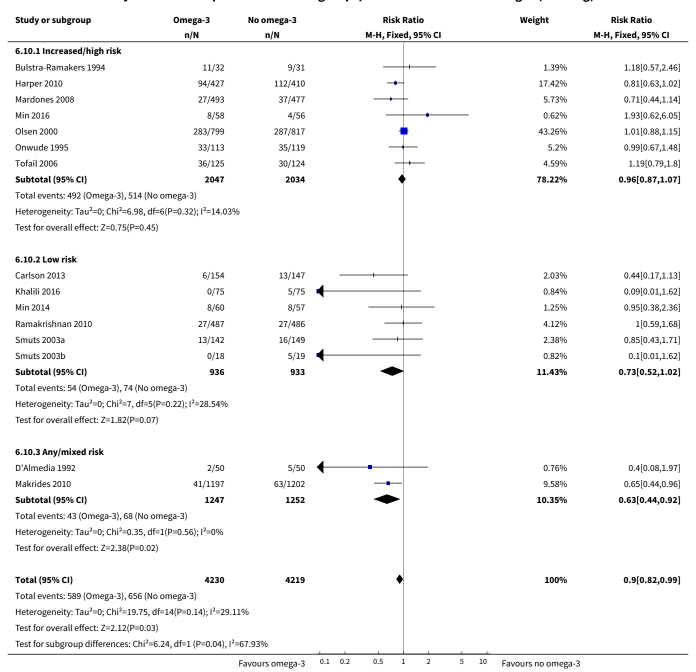


Analysis 6.9. Comparison 6 Risk subgroups, Outcome 9 Neonatal death.





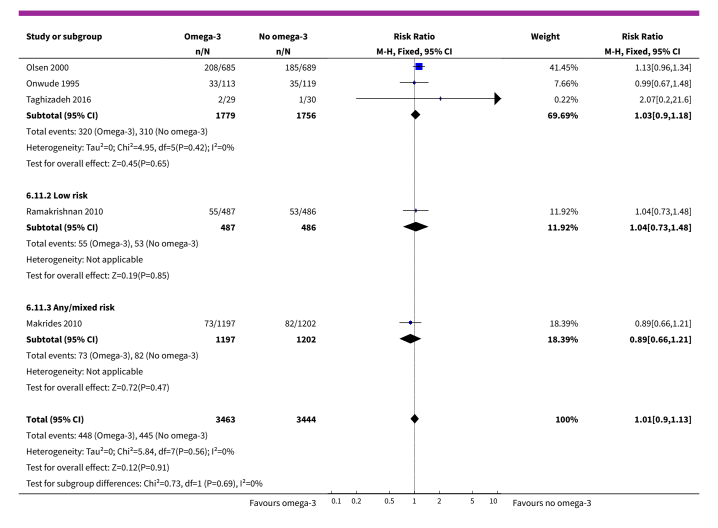
Analysis 6.10. Comparison 6 Risk subgroups, Outcome 10 Low birthweight (< 2500 g).



Analysis 6.11. Comparison 6 Risk subgroups, Outcome 11 Small-for-gestational age/IUGR.

Study or subgroup	Omega-3	No omega-3		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		M-H, Fixed	i, 95% CI				M-H, Fixed, 95% CI
6.11.1 Increased/high risk									
Bulstra-Ramakers 1994	12/32	9/31			+			2.05%	1.29[0.64,2.63]
Harper 2010	35/427	41/410			_			9.4%	0.82[0.53,1.26]
Mardones 2008	30/493	39/477		. +	-			8.91%	0.74[0.47,1.18]
		Favours omega-3	0.1 0.2	0.5 1	2	5	10	Favours no omega-3	

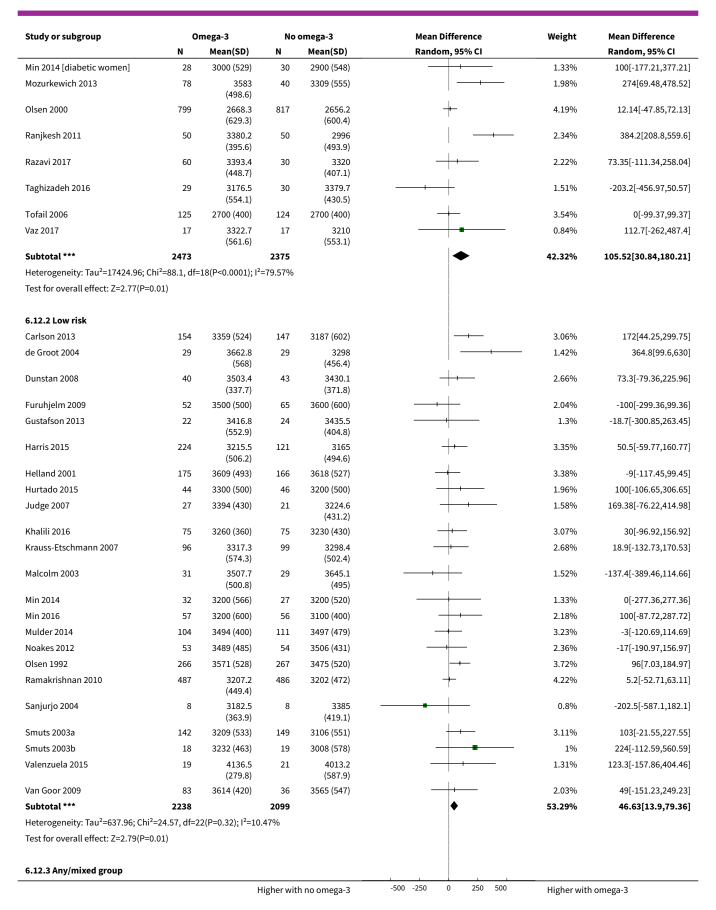




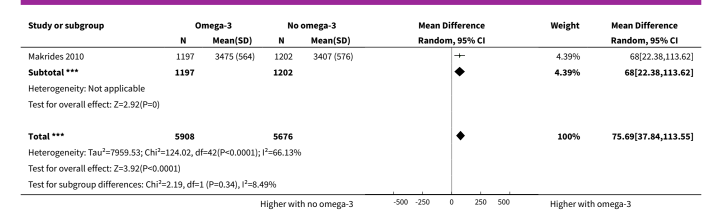
Analysis 6.12. Comparison 6 Risk subgroups, Outcome 12 Birthweight (g).

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
6.12.1 Increased/high risk							
Ali 2017	34	2324 (110.8)	32	2022 (141.4)	+	4.17%	302[240.46,363.54]
Dilli 2018	52	3288 (641)	68	3538 (671)		1.66%	-250[-486.2,-13.8]
England 1989	17	2200 (700)	18	2000 (500)	-	0.74%	200[-205.07,605.07]
Haghiac 2015	25	3278 (448)	24	2935 (356)		1.75%	343[116.89,569.11]
Harper 2010	427	2990 (1005.8)	410	2923 (1252.8)		2.64%	67[-87.29,221.29]
Hauner 2012	96	3534 (465)	92	3357 (557)		2.75%	177[30.01,323.99]
Horvaticek 2017	47	3580.9 (568)	43	3456.9 (575.8)	++-	1.66%	124[-112.62,360.62]
Jamilian 2016	26	3418.8 (344.7)	27	3405.2 (465.1)	- 	1.82%	13.6[-206.23,233.43]
Keenan 2014	36	3074.4 (582.3)	17	2919.6 (537.7)		1.08%	154.83[-163.81,473.47]
Krummel 2016	34	3502 (433)	29	3484 (411)	 +	1.93%	18[-190.71,226.71]
Mardones 2008	493	3265.9 (452.1)	477	3200.5 (506)	 	4.19%	65.4[4.95,125.85]
		Н	ligher wit	h no omega-3	-500 -250 0 250 500	Higher with	n omega-3









Comparison 7. Omega-3 doses: direct comparisons

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Early preterm birth < 34 weeks	1	224	Risk Ratio (M-H, Fixed, 95% CI)	0.91 [0.13, 6.38]
2 Prolonged gestation > 42 weeks	1	224	Risk Ratio (M-H, Fixed, 95% CI)	0.91 [0.06, 14.44]
3 Pre-eclampsia	1	224	Risk Ratio (M-H, Fixed, 95% CI)	0.91 [0.06, 14.44]
4 Induction (post-term)	1	224	Risk Ratio (M-H, Fixed, 95% CI)	0.10 [0.01, 1.87]
5 PROM	1	224	Risk Ratio (M-H, Fixed, 95% CI)	0.30 [0.03, 2.89]
6 PPROM	1	224	Risk Ratio (M-H, Fixed, 95% CI)	1.22 [0.28, 5.32]
7 Length of gestation	2	1474	Mean Difference (IV, Fixed, 95% CI)	0.24 [-1.16, 1.64]
8 Birthweight (g)	1	224	Mean Difference (IV, Fixed, 95% CI)	-110.35 [-242.80, 22.10]
9 Length at birth (cm)	1	224	Mean Difference (IV, Fixed, 95% CI)	0.05 [-0.80, 0.90]
10 Head circumference at birth (cm)	1	224	Mean Difference (IV, Fixed, 95% CI)	-0.24 [-0.87, 0.39]

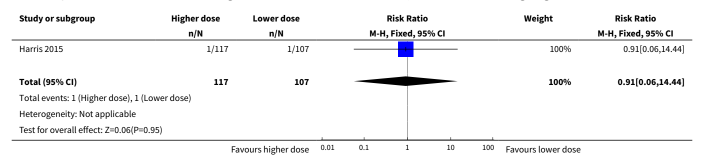
Analysis 7.1. Comparison 7 Omega-3 doses: direct comparisons, Outcome 1 Early preterm birth < 34 weeks.

Study or subgroup	Higher dose	Lower dose			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		М-Н	Fixed, 95	% CI			M-H, Fixed, 95% CI
Harris 2015	2/117	2/107			-	_		100%	0.91[0.13,6.38]
Total (95% CI)	117	107				_		100%	0.91[0.13,6.38]
Total events: 2 (Higher dose),	2 (Lower dose)								
	Fa	vours higher dose	0.01	0.1	1	10	100	Favours lower dose	

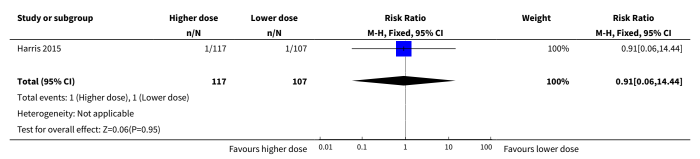


Study or subgroup	Higher dose Lower dose		Risk Ratio					Weight	Risk Ratio
	n/N	n/N		М-Н	, Fixed, 95	% CI			M-H, Fixed, 95% CI
Heterogeneity: Not applicable									
Test for overall effect: Z=0.09(P=0.93)			1			1			
	ļ	Favours higher dose	0.01	0.1	1	10	100	Favours lower dose	

Analysis 7.2. Comparison 7 Omega-3 doses: direct comparisons, Outcome 2 Prolonged gestation > 42 weeks.



Analysis 7.3. Comparison 7 Omega-3 doses: direct comparisons, Outcome 3 Pre-eclampsia.



Analysis 7.4. Comparison 7 Omega-3 doses: direct comparisons, Outcome 4 Induction (post-term).

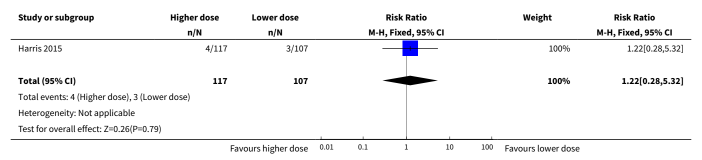
Study or subgroup	Higher dose	Lower dose		Risk Ratio M-H, Fixed, 95% CI			Weight	Risk Ratio	
	n/N	n/N						M-H, Fixed, 95% CI	
Harris 2015	0/117	4/107		1				100%	0.1[0.01,1.87]
Total (95% CI)	117	107	-	-	-			100%	0.1[0.01,1.87]
Total events: 0 (Higher dose), 4 (L	ower dose)								
Heterogeneity: Not applicable									
Test for overall effect: Z=1.54(P=0	.12)								
	Fa	vours higher dose	0.002	0.1	1	10	500	Favours lower dose	



Analysis 7.5. Comparison 7 Omega-3 doses: direct comparisons, Outcome 5 PROM.

Study or subgroup	Higher dose	Lower dose		Risk Ratio			Weight	Risk Ratio		
	n/N	n/N		М-Н,	Fixed, 95%	CI			M-H, Fixed, 95% CI	
Harris 2015	1/117	3/107	-	1				100%	0.3[0.03,2.89]	
Total (95% CI)	117	107	-					100%	0.3[0.03,2.89]	
Total events: 1 (Higher dose),	3 (Lower dose)									
Heterogeneity: Not applicable	2									
Test for overall effect: Z=1.04(P=0.3)									
	Fa	vours higher dose	0.01	0.1	1	10	100	Favours lower dose		

Analysis 7.6. Comparison 7 Omega-3 doses: direct comparisons, Outcome 6 PPROM.



Analysis 7.7. Comparison 7 Omega-3 doses: direct comparisons, Outcome 7 Length of gestation.

Study or subgroup	High	Higher omega-3		Lower omega-3		Me	an Differen	ce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		F	ixed, 95% C	I			Fixed, 95% CI
Harris 2015	117	275.6 (19.5)	107	275 (19.5)			+			7.54%	0.6[-4.5,5.7]
Knudsen 2006	453	280.6 (12.3)	797	280.4 (13.2)						92.46%	0.21[-1.25,1.67]
Total ***	570		904				•			100%	0.24[-1.16,1.64]
Heterogeneity: Tau ² =0; Chi ² =	0.02, df=1(P=0.8	9); I ² =0%									
Test for overall effect: Z=0.33	(P=0.74)										
			Favou	rs higher dose	-10	-5	0	5	10	Favours low	ver dose

Analysis 7.8. Comparison 7 Omega-3 doses: direct comparisons, Outcome 8 Birthweight (g).

Study or subgroup	Hig	her dose	Lov	ver dose	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
Harris 2015	117	3210.6 (516.9)	107	3320.9 (494.2)		100%	-110.35[-242.8,22.1]
Total ***	117		107			100%	-110.35[-242.8,22.1]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.63(P=0.1)							
			Favou	rs lower dose	-200 -100 0 100 200	Favours hig	her dose



Analysis 7.9. Comparison 7 Omega-3 doses: direct comparisons, Outcome 9 Length at birth (cm).

Study or subgroup	Hig	her dose	Lov	ver dose		Mea	an Differen	ce		Weight M	lean Difference
	N	Mean(SD)	N	Mean(SD)		Fi	xed, 95% C	1			Fixed, 95% CI
Harris 2015	117	50 (3.2)	107	49.9 (3.2)			+			100%	0.05[-0.8,0.9]
Total ***	117		107				•			100%	0.05[-0.8,0.9]
Heterogeneity: Tau ² =0; Chi ² =	0, df=0(P<0.0001	.); I ² =100%									
Test for overall effect: Z=0.12	(P=0.91)										
			Favou	rs lower dose	-20	-10	0	10	20	Favours higher de	ose

Analysis 7.10. Comparison 7 Omega-3 doses: direct comparisons, Outcome 10 Head circumference at birth (cm).

Study or subgroup	Hig	her dose	Lov	ver dose	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
Harris 2015	117	34 (2.2)	107	34.2 (2.6)	+	100%	-0.24[-0.87,0.39]
Total ***	117		107		•	100%	-0.24[-0.87,0.39]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.75(P=0.45)							
			Favou	rs lower dose	-10 -5 0 5 10	Favours hig	her dose

Comparison 8. Omega-3 type: direct comparisons

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Gestational diabetes	2		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
1.1 DHA versus EPA	1	77	Risk Ratio (M-H, Fixed, 95% CI)	0.15 [0.02, 1.14]
1.2 DHA versus DHA/AA	1	86	Risk Ratio (M-H, Fixed, 95% CI)	0.33 [0.01, 7.96]
2 Caesarean section	1	77	Risk Ratio (M-H, Fixed, 95% CI)	1.23 [0.61, 2.51]
2.1 DHA versus EPA	1	77	Risk Ratio (M-H, Fixed, 95% CI)	1.23 [0.61, 2.51]
3 Adverse events: cessation	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
3.1 DHA versus EPA	1	77	Risk Ratio (M-H, Fixed, 95% CI)	0.82 [0.24, 2.83]
4 Pre-eclampsia	1	77	Risk Ratio (M-H, Fixed, 95% CI)	0.26 [0.06, 1.13]
4.1 DHA versus EPA	1	77	Risk Ratio (M-H, Fixed, 95% CI)	0.26 [0.06, 1.13]
5 Blood loss at birth (mL)	1	77	Mean Difference (IV, Fixed, 95% CI)	1.0 [-181.94, 183.94]
5.1 DHA versus EPA	1	77	Mean Difference (IV, Fixed, 95% CI)	1.0 [-181.94, 183.94]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
6 Depressive symptoms post- partum: thresholds	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
6.1 Major depressive disorder at 6-8 weeks	1	77	Risk Ratio (M-H, Fixed, 95% CI)	0.68 [0.12, 3.87]
7 Depressive symptoms post- partum: scores	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
7.1 BDI: 6-8 weeks postpartum	1	77	Mean Difference (IV, Fixed, 95% CI)	-1.40 [-3.75, 0.95]
8 Length of gestation (days)	3		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
.1 DHA versus EPA 1 77		Mean Difference (IV, Fixed, 95% CI)	9.10 [5.24, 12.96]	
8.2 EPA/DHA vs ALA	1	1250	Mean Difference (IV, Fixed, 95% CI)	-0.29 [-2.33, 1.75]
8.3 DHA versus DHA/AA	1	83	Mean Difference (IV, Fixed, 95% CI)	0.0 [-3.31, 3.31]
9 Baby admitted to neonatal care	1	78	Risk Ratio (M-H, Fixed, 95% CI)	0.35 [0.08, 1.63]
9.1 DHA versus EPA	1	78	Risk Ratio (M-H, Fixed, 95% CI)	0.35 [0.08, 1.63]
10 Birthweight (g)	2		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
10.1 DHA versus EPA	1	78	Mean Difference (IV, Fixed, 95% CI)	372.0 [151.90, 592.10]
10.2 DHA versus DHA/AA	1	83	Mean Difference (IV, Fixed, 95% CI)	-79.0 [-260.22, 102.22]
11 Infant weight (kg)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
11.1 DHA versus DHA/AA	1	80	Mean Difference (IV, Fixed, 95% CI)	-0.20 [-0.79, 0.39]
12 Infant height (cm)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
12.1 DHA versus DHA/AA	1	80	Mean Difference (IV, Fixed, 95% CI)	-0.80 [-2.50, 0.90]
13 Infant head circumference (cm)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
13.1 At 18 months	1	80	Mean Difference (IV, Fixed, 95% CI)	0.10 [-0.45, 0.65]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
14 Cognition: Scores	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
14.1 DHA versus DHA/AA: BSID II	1	80	Mean Difference (IV, Fixed, 95% CI)	0.90 [-4.71, 6.51]
15 Motor: Scores	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
15.1 DHA versus DHA/AA: BSID II	1	79	Mean Difference (IV, Fixed, 95% CI)	3.40 [-1.07, 7.87]
16 Neurodevelopment	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
16.1 DHA versus DHA/AA: neonatal neurological classi- fication: mildly/definitely ab- normal at 2 weeks	1	67	Risk Ratio (M-H, Fixed, 95% CI)	0.73 [0.28, 1.87]
16.2 DHA versus DHA/AA: general movement quality: mildly/definitely abnormal at 2 weeks	1	67	Risk Ratio (M-H, Fixed, 95% CI)	1.08 [0.68, 1.72]
16.3 DHA versus DHA/AA: general movement quality: mildly/definitely abnormal at 12 weeks	1	83	Risk Ratio (M-H, Fixed, 95% CI)	1.81 [1.11, 2.95]
17 Cerebral palsy	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
17.1 DHA versus DHA/AA	1	80	Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]

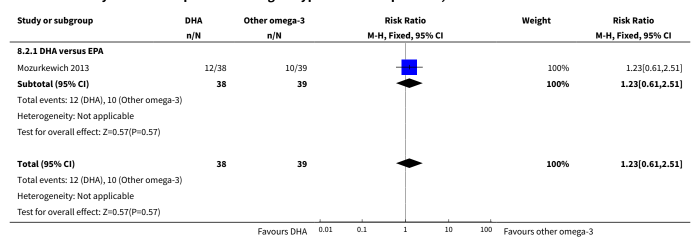
Analysis 8.1. Comparison 8 Omega-3 type: direct comparisons, Outcome 1 Gestational diabetes.

Study or subgroup	DHA	Other omega-3	Risk	Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixe	ed, 95% CI		M-H, Fixed, 95% CI
8.1.1 DHA versus EPA						
Mozurkewich 2013	1/38	7/39	- I	<u> </u> 	100%	0.15[0.02,1.14]
Subtotal (95% CI)	38	39		<u> </u> 	100%	0.15[0.02,1.14]
Total events: 1 (DHA), 7 (Other omega-3)						
Heterogeneity: Not applicable						
Test for overall effect: Z=1.84(P=0.07)						
8.1.2 DHA versus DHA/AA						
Van Goor 2009	0/43	1/43	-		100%	0.33[0.01,7.96]
Subtotal (95% CI)	43	43			100%	0.33[0.01,7.96]
Total events: 0 (DHA), 1 (Other omega-3)						
Heterogeneity: Not applicable						
Test for overall effect: Z=0.68(P=0.5)						
		Favours DHA	0.01 0.1	1 10	100 Favours other omega-3	3

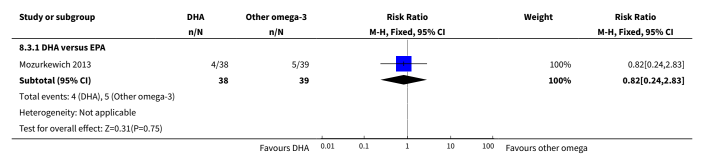


Study or subgroup	DHA n/N	Other omega-3 n/N			Risk Ratio , Fixed, 95			Weight	Risk Ratio M-H, Fixed, 95% CI
Test for subgroup differences: C	hi ² =0.18, df=1 (P=0.67)), I ² =0%							
		Favours DHA	0.01	0.1	1	10	100	Favours other omega	-3

Analysis 8.2. Comparison 8 Omega-3 type: direct comparisons, Outcome 2 Caesarean section.



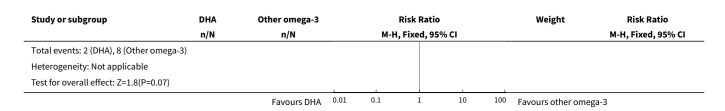
Analysis 8.3. Comparison 8 Omega-3 type: direct comparisons, Outcome 3 Adverse events: cessation.



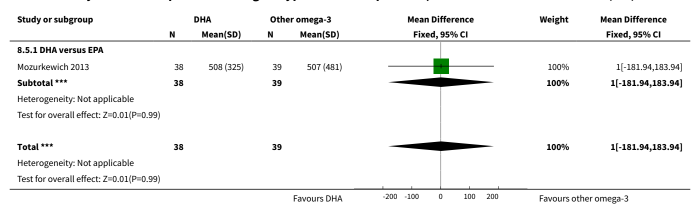
Analysis 8.4. Comparison 8 Omega-3 type: direct comparisons, Outcome 4 Pre-eclampsia.

Study or subgroup	DHA	Other omega-3		Risk	Ratio		Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI				M-H, Fixed, 95% CI	
8.4.1 DHA versus EPA								
Mozurkewich 2013	2/38	8/39		-	<u> </u>		100%	0.26[0.06,1.13]
Subtotal (95% CI)	38	39			<u>[</u>		100%	0.26[0.06,1.13]
Total events: 2 (DHA), 8 (Other omega-3)								
Heterogeneity: Not applicable								
Test for overall effect: Z=1.8(P=0.07)								
Total (95% CI)	38	39		-			100%	0.26[0.06,1.13]
		Favours DHA	0.01	0.1	1 10	100	Favours other omega-3	1

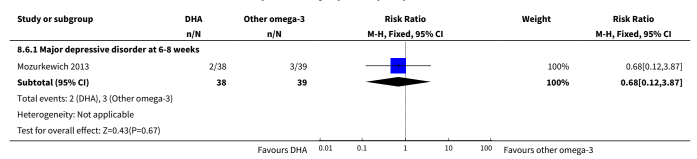




Analysis 8.5. Comparison 8 Omega-3 type: direct comparisons, Outcome 5 Blood loss at birth (mL).



Analysis 8.6. Comparison 8 Omega-3 type: direct comparisons, Outcome 6 Depressive symptoms postpartum: thresholds.



Analysis 8.7. Comparison 8 Omega-3 type: direct comparisons, Outcome 7 Depressive symptoms postpartum: scores.

Study or subgroup		DHA	Othe	r omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
8.7.1 BDI: 6-8 weeks postpartum							
Mozurkewich 2013	38	5.2 (4.8)	39	6.6 (5.7)		100%	-1.4[-3.75,0.95]
Subtotal ***	38		39			100%	-1.4[-3.75,0.95]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.17(P=0.24)							
				Favours DHA	-5 -2.5 0 2.5 5	Favours oth	er omega-3



Analysis 8.8. Comparison 8 Omega-3 type: direct comparisons, Outcome 8 Length of gestation (days).

Study or subgroup		DHA	Othe	er omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
8.8.1 DHA versus EPA							
Mozurkewich 2013	38	282.8 (6.3)	39	273.7 (10.5)		100%	9.1[5.24,12.96]
Subtotal ***	38		39			100%	9.1[5.24,12.96]
Heterogeneity: Not applicable							
Test for overall effect: Z=4.62(P<0.0	0001)						
8.8.2 EPA/DHA vs ALA							
Knudsen 2006	1074	280.4 (12.9)	176	280.7 (12.8)		100%	-0.29[-2.33,1.75]
Subtotal ***	1074		176			100%	-0.29[-2.33,1.75]
Heterogeneity: Tau ² =0; Chi ² =0, df=	0(P<0.000	1); I²=100%					
Test for overall effect: Z=0.28(P=0.7	78)						
8.8.3 DHA versus DHA/AA							
Van Goor 2009	42	281.4 (7.7)	41	281.4 (7.7)		100%	0[-3.31,3.31]
Subtotal ***	42		41			100%	0[-3.31,3.31]
Heterogeneity: Not applicable							
Test for overall effect: Not applicab	ole						
Test for subgroup differences: Chi ²	=18.63, df=	=1 (P<0.0001), I ² =	89.26%				
				Favours DHA	-5 -2.5 0 2.5 5	Favours oth	er omega-3

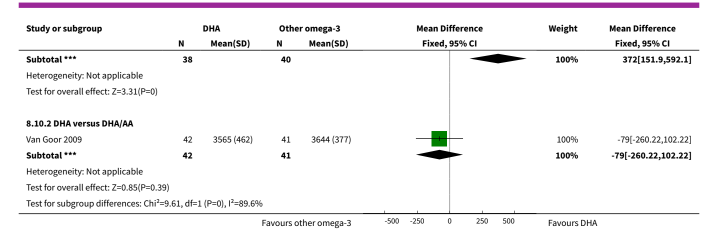
Analysis 8.9. Comparison 8 Omega-3 type: direct comparisons, Outcome 9 Baby admitted to neonatal care.

Study or subgroup	DHA	Other omega-3			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		М-Н	Fixed, 95	% CI			M-H, Fixed, 95% CI
8.9.1 DHA versus EPA									
Mozurkewich 2013	2/38	6/40		-	-			100%	0.35[0.08,1.63]
Subtotal (95% CI)	38	40						100%	0.35[0.08,1.63]
Total events: 2 (DHA), 6 (Other omega-3)					İ				
Heterogeneity: Not applicable					İ				
Test for overall effect: Z=1.34(P=0.18)									
Total (95% CI)	38	40						100%	0.35[0.08,1.63]
Total events: 2 (DHA), 6 (Other omega-3)					İ				
Heterogeneity: Not applicable					İ				
Test for overall effect: Z=1.34(P=0.18)						1			
		Favours DHA	0.01	0.1	1	10	100	Favours other omega-3	

Analysis 8.10. Comparison 8 Omega-3 type: direct comparisons, Outcome 10 Birthweight (g).

Study or subgroup		DHA		r omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
8.10.1 DHA versus EPA							
Mozurkewich 2013	38	3774 (438)	40	3402 (550)		100%	372[151.9,592.1]
			Favours o	ther omega-3	-500 -250 0 250 500	Favours DHA	

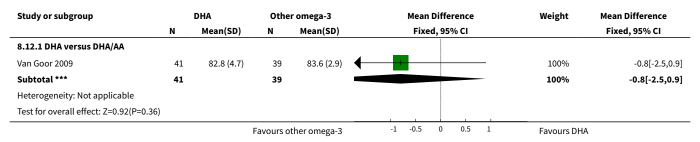




Analysis 8.11. Comparison 8 Omega-3 type: direct comparisons, Outcome 11 Infant weight (kg).

Study or subgroup		DHA	Othe	r omega-3		Mea	n Differ	ence		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fix	ed, 95%	CI			Fixed, 95% CI
8.11.1 DHA versus DHA/AA											
Van Goor 2009	41	11.3 (1.4)	39	11.5 (1.3)						100%	-0.2[-0.79,0.39]
Subtotal ***	41		39				*			100%	-0.2[-0.79,0.39]
Heterogeneity: Not applicable											
Test for overall effect: Z=0.66(P=0.51)											
			Favours o	ther omega-3	-5	-2.5	0	2.5	5	Favours DHA	

Analysis 8.12. Comparison 8 Omega-3 type: direct comparisons, Outcome 12 Infant height (cm).



Analysis 8.13. Comparison 8 Omega-3 type: direct comparisons, Outcome 13 Infant head circumference (cm).

Study or subgroup		DHA	Othe	r omega-3		Mea	n Differen	ce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fix	ced, 95% C	1			Fixed, 95% CI
8.13.1 At 18 months											
Van Goor 2009	41	47.6 (1.1)	39	47.5 (1.4)			-			100%	0.1[-0.45,0.65]
Subtotal ***	41		39				*			100%	0.1[-0.45,0.65]
Heterogeneity: Not applicable											
Test for overall effect: Z=0.35(P=0.72)											
			Favours o	ther omega-3	-5	-2.5	0	2.5	5	Favours DHA	



Analysis 8.14. Comparison 8 Omega-3 type: direct comparisons, Outcome 14 Cognition: Scores.

Study or subgroup		DHA	Othe	r omega-3		Me	an Differen	ce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		F	ixed, 95% C	I			Fixed, 95% CI
8.14.1 DHA versus DHA/AA: BSID II											
Van Goor 2009	41	113.7 (13)	39	112.8 (12.6)			+			100%	0.9[-4.71,6.51]
Subtotal ***	41		39				*			100%	0.9[-4.71,6.51]
Heterogeneity: Not applicable											
Test for overall effect: Z=0.31(P=0.75)											
			Favours o	ther omega-3	-100	-50	0	50	100	Favours DHA	

Analysis 8.15. Comparison 8 Omega-3 type: direct comparisons, Outcome 15 Motor: Scores.

Study or subgroup		DHA		Other omega-3		Mean Difference				Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fi	ixed, 95% CI				Fixed, 95% CI
8.15.1 DHA versus DHA/AA: BSI	DII										
Van Goor 2009	41	95.8 (11.4)	38	92.4 (8.8)						100%	3.4[-1.07,7.87]
Subtotal ***	41		38				•			100%	3.4[-1.07,7.87]
Heterogeneity: Tau ² =0; Chi ² =0, d	f=0(P<0.0001	.); I ² =100%									
Test for overall effect: Z=1.49(P=0	0.14)										
			Favours o	ther omega-3	-50	-25	0	25	50	Favours DHA	

Analysis 8.16. Comparison 8 Omega-3 type: direct comparisons, Outcome 16 Neurodevelopment.

Study or subgroup	DHA	Other omega-3		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		M-H, Fixed, 95% CI			M-H, Fixed, 95% CI
8.16.1 DHA versus DHA/AA: neonatal ne ly/definitely abnormal at 2 weeks	urological cla	assification: mild-					
Van Goor 2009	6/34	8/33				100%	0.73[0.28,1.87]
Subtotal (95% CI)	34	33				100%	0.73[0.28,1.87]
Total events: 6 (DHA), 8 (Other omega-3)							
Heterogeneity: Not applicable							
Test for overall effect: Z=0.66(P=0.51)							
8.16.2 DHA versus DHA/AA: general mov nitely abnormal at 2 weeks	vement quali	ty: mildly/defi-					
Van Goor 2009	20/37	15/30		-		100%	1.08[0.68,1.72]
Subtotal (95% CI)	37	30		*		100%	1.08[0.68,1.72]
Total events: 20 (DHA), 15 (Other omega-3	3)						
Heterogeneity: Not applicable							
Test for overall effect: Z=0.33(P=0.74)							
8.16.3 DHA versus DHA/AA: general mov nitely abnormal at 12 weeks	vement quali	ty: mildly/defi-					
Van Goor 2009	26/42	14/41				100%	1.81[1.11,2.95]
Subtotal (95% CI)	42	41		•		100%	1.81[1.11,2.95]
Total events: 26 (DHA), 14 (Other omega-	3)						
Heterogeneity: Not applicable							
		Favours DHA	0.01	0.1 1 10	100	Favours other omega	-3



Study or subgroup	DHA n/N	Other omega-3 n/N			Risk Ratio , Fixed, 95			Weight	Risk Ratio M-H, Fixed, 95% CI
Test for overall effect: Z=2.4(P=0.02)						1			
		Favours DHA	0.01	0.1	1	10	100	Favours other omega-3	3

Analysis 8.17. Comparison 8 Omega-3 type: direct comparisons, Outcome 17 Cerebral palsy.

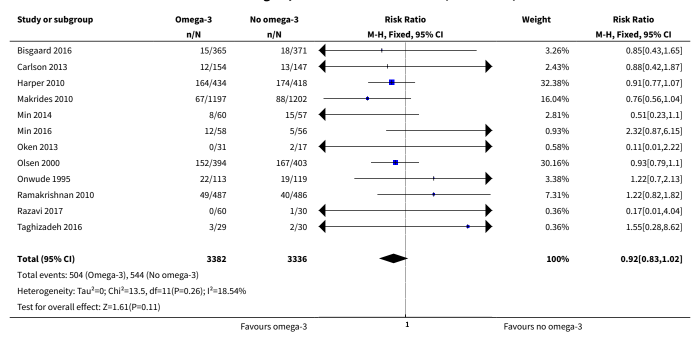
Study or subgroup	DHA	Other omega-3			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		M-H	l, Fixed, 95%	% CI			M-H, Fixed, 95% CI
8.17.1 DHA versus DHA/AA									
Van Goor 2009	0/41	0/39							Not estimable
Subtotal (95% CI)	41	39							Not estimable
Total events: 0 (DHA), 0 (Other omega-3)									
Heterogeneity: Not applicable									
Test for overall effect: Not applicable									
		Favours DHA	0.01	0.1	1	10	100	Favours other omega-3	3

Comparison 9. Sensitivity analysis: omega-3 versus no omega-3

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Preterm birth (< 37 weeks)	12	6718	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.83, 1.02]
2 Early preterm birth (< 34 weeks)	6	4073	Risk Ratio (M-H, Fixed, 95% CI)	0.61 [0.46, 0.82]
3 Prolonged gestation (> 42 weeks)	3	4285	Risk Ratio (M-H, Fixed, 95% CI)	2.32 [1.26, 4.28]
4 Pre-eclampsia (hypertension with proteinuria)	12	6104	Risk Ratio (M-H, Fixed, 95% CI)	1.00 [0.81, 1.25]
5 Caesarean section	12	5239	Risk Ratio (M-H, Fixed, 95% CI)	0.96 [0.89, 1.04]
6 Length of gestation (days)	16	6313	Mean Difference (IV, Fixed, 95% CI)	1.42 [0.73, 2.11]
7 Perinatal death	5	4610	Risk Ratio (M-H, Fixed, 95% CI)	0.60 [0.37, 0.97]
8 Stillbirth	10	6193	Risk Ratio (M-H, Fixed, 95% CI)	0.80 [0.49, 1.31]
9 Neonatal death	6	4791	Risk Ratio (M-H, Fixed, 95% CI)	0.56 [0.25, 1.27]
10 Low birthweight (< 2500 g)	10	6839	Risk Ratio (M-H, Random, 95% CI)	0.87 [0.73, 1.04]
11 Small-for-gestational age/ IUGR	6	5874	Risk Ratio (M-H, Fixed, 95% CI)	1.03 [0.91, 1.16]
12 Birthweight (g)	18	7382	Mean Difference (IV, Fixed, 95% CI)	48.84 [22.93, 74.76]



Analysis 9.1. Comparison 9 Sensitivity analysis: omega-3 versus no omega-3, Outcome 1 Preterm birth (< 37 weeks).



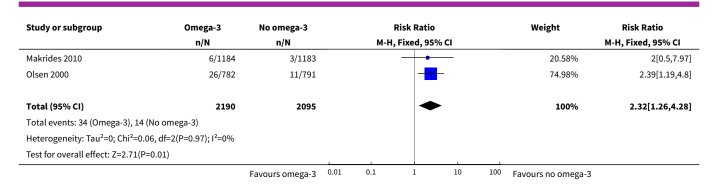
Analysis 9.2. Comparison 9 Sensitivity analysis: omega-3 versus no omega-3, Outcome 2 Early preterm birth (< 34 weeks).

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
Carlson 2013	1/154	7/147	—	6.69%	0.14[0.02,1.09]
Harris 2015	4/224	7/121	—	8.48%	0.31[0.09,1.03]
Makrides 2010	13/1197	27/1202	—	25.15%	0.48[0.25,0.93]
Min 2014	4/60	4/57	+ • • • • • • • • • • • • • • • • • • •	3.83%	0.95[0.25,3.62]
Min 2016	2/58	0/56	+	0.47%	4.83[0.24,98.44]
Olsen 2000	42/394	60/403	-	55.37%	0.72[0.5,1.04]
Total (95% CI)	2087	1986	•	100%	0.61[0.46,0.82]
Total events: 66 (Omega-3), 10	5 (No omega-3)				
Heterogeneity: Tau ² =0; Chi ² =6.	63, df=5(P=0.25); I ² =24.649	6			
Test for overall effect: Z=3.28(P	P=0)				
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	

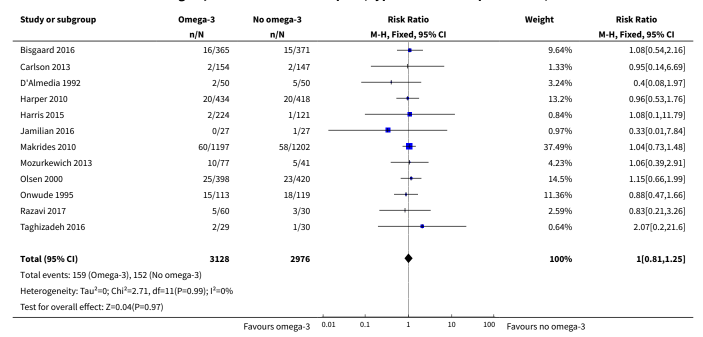
Analysis 9.3. Comparison 9 Sensitivity analysis: omega-3 versus no omega-3, Outcome 3 Prolonged gestation (> 42 weeks).

Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		М-Н,	Fixed, 95	% CI			M-H, Fixed, 95% CI
Harris 2015	2/224	0/121			+			4.45%	2.71[0.13,56.02]
		Favours omega-3	0.01	0.1	1	10	100	Favours no omega-3	





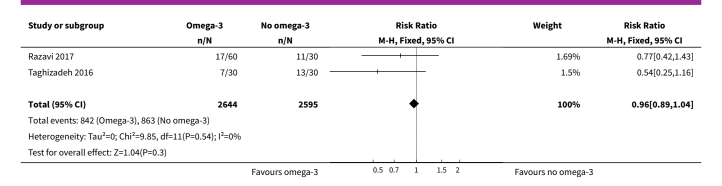
Analysis 9.4. Comparison 9 Sensitivity analysis: omega-3 versus no omega-3, Outcome 4 Pre-eclampsia (hypertension with proteinuria).



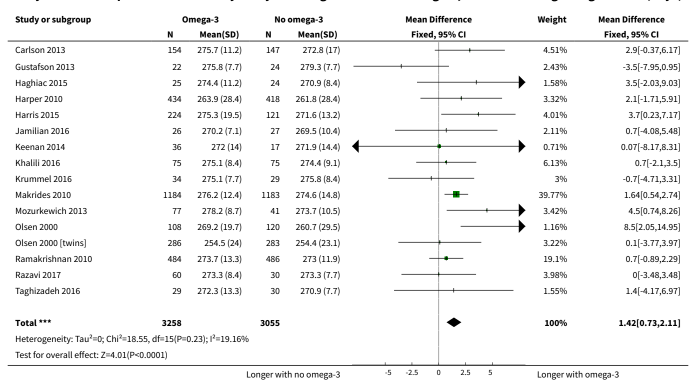
Analysis 9.5. Comparison 9 Sensitivity analysis: omega-3 versus no omega-3, Outcome 5 Caesarean section.

Study or subgroup	Omega-3	No omega-3	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
Bisgaard 2016	77/365	71/371		8.12%	1.1[0.83,1.47]
Carlson 2013	46/154	44/147		5.19%	1[0.71,1.41]
Jamilian 2016	12/26	18/27		2.04%	0.69[0.42,1.13]
Khalili 2016	30/75	33/75		3.81%	0.91[0.62,1.33]
Makrides 2010	326/1197	350/1202	-	40.29%	0.94[0.82,1.06]
Min 2014	25/60	24/57		2.84%	0.99[0.65,1.52]
Min 2016	28/58	29/56		3.4%	0.93[0.65,1.35]
Mozurkewich 2013	22/77	11/41		1.66%	1.06[0.57,1.97]
Onwude 1995	36/113	25/119	 	2.81%	1.52[0.98,2.36]
Ramakrishnan 2010	216/429	234/440	<u> </u>	26.65%	0.95[0.83,1.08]
		Favours omega-3	0.5 0.7 1 1.5 2	Favours no omega-3	





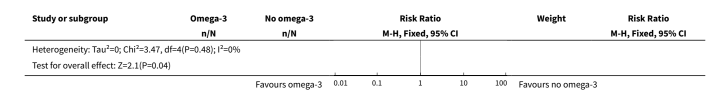
Analysis 9.6. Comparison 9 Sensitivity analysis: omega-3 versus no omega-3, Outcome 6 Length of gestation (days).



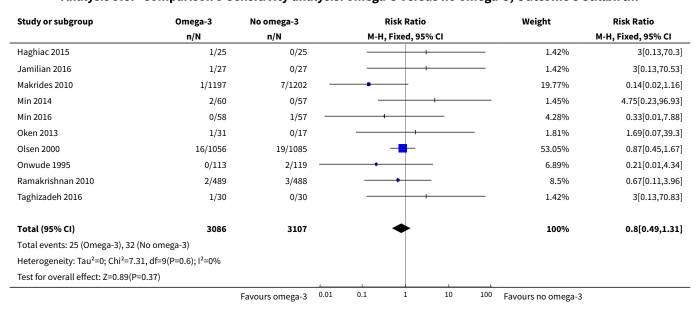
Analysis 9.7. Comparison 9 Sensitivity analysis: omega-3 versus no omega-3, Outcome 7 Perinatal death.

Study or subgroup	Omega-3	No omega-3		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		M-H, Fixed, 95%	CI		M-H, Fixed, 95% CI
Harper 2010	16/434	17/418		-		39.58%	0.91[0.46,1.77]
Khalili 2016	0/75	1/75		+		3.43%	0.33[0.01,8.05]
Makrides 2010	3/1197	12/1202	-			27.37%	0.25[0.07,0.89]
Onwude 1995	1/113	2/119		+	_	4.45%	0.53[0.05,5.73]
Ramakrishnan 2010	6/489	11/488				25.17%	0.54[0.2,1.46]
Total (95% CI)	2308	2302		•		100%	0.6[0.37,0.97]
Total events: 26 (Omega-3), 43 (No ome	ega-3)						
		Favours omega-3	0.01).1 1	10 1	.00 Favours no omega-3	





Analysis 9.8. Comparison 9 Sensitivity analysis: omega-3 versus no omega-3, Outcome 8 Stillbirth.

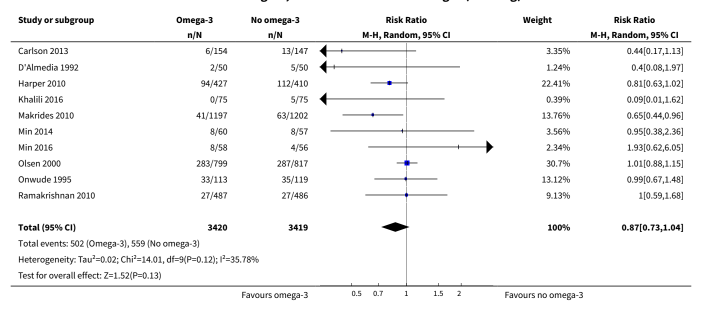


Analysis 9.9. Comparison 9 Sensitivity analysis: omega-3 versus no omega-3, Outcome 9 Neonatal death.

Study or subgroup	Omega-3	No omega-3			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		ı	M-H, Fixed, 95%	CI			M-H, Fixed, 95% CI
Bisgaard 2016	0/365	0/371							Not estimable
Carlson 2013	1/154	1/147			+			6.39%	0.95[0.06,15.12]
Khalili 2016	0/75	1/75	\leftarrow		+			9.37%	0.33[0.01,8.05]
Makrides 2010	2/1197	5/1202			-			31.17%	0.4[0.08,2.07]
Onwude 1995	1/113	0/119		_	+		\rightarrow	3.04%	3.16[0.13,76.73]
Ramakrishnan 2010	4/487	8/486		_	-			50.03%	0.5[0.15,1.65]
Total (95% CI)	2391	2400						100%	0.56[0.25,1.27]
Total events: 8 (Omega-3), 15 (No	omega-3)								
Heterogeneity: Tau ² =0; Chi ² =1.57,	df=4(P=0.81); I ² =0%								
Test for overall effect: Z=1.39(P=0.	.17)								
		Favours omega-3	0.02	0.1	1	10	50	Favours no omega-3	



Analysis 9.10. Comparison 9 Sensitivity analysis: omega-3 versus no omega-3, Outcome 10 Low birthweight (< 2500 g).



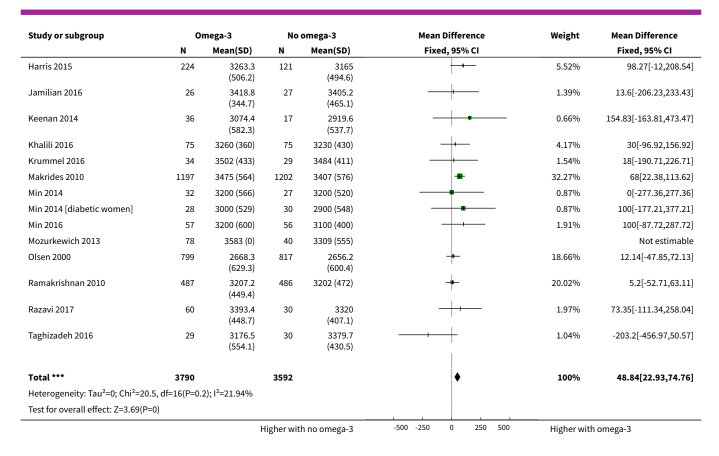
Analysis 9.11. Comparison 9 Sensitivity analysis: omega-3 versus no omega-3, Outcome 11 Small-for-gestational age/IUGR.

Study or subgroup	Omega-3	No omega-3		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		M-H, Fixed, 95% CI			M-H, Fixed, 95% CI
Harper 2010	35/427	41/410		+		10.56%	0.82[0.53,1.26]
Makrides 2010	73/1197	82/1202		+		20.65%	0.89[0.66,1.21]
Olsen 2000	208/685	185/689		•		46.55%	1.13[0.96,1.34]
Onwude 1995	33/113	35/119		+		8.6%	0.99[0.67,1.48]
Ramakrishnan 2010	55/487	53/486		+		13.39%	1.04[0.73,1.48]
Taghizadeh 2016	2/29	1/30			_	0.25%	2.07[0.2,21.6]
Total (95% CI)	2938	2936		,		100%	1.03[0.91,1.16]
Total events: 406 (Omega-3), 3	897 (No omega-3)						
Heterogeneity: Tau ² =0; Chi ² =3	.49, df=5(P=0.62); I ² =0%						
Test for overall effect: Z=0.42(F	P=0.68)				1		
		Favours omega-3	0.01	0.1 1 10	100	Favours no omega-3	

Analysis 9.12. Comparison 9 Sensitivity analysis: omega-3 versus no omega-3, Outcome 12 Birthweight (g).

Study or subgroup	0	mega-3	No	omega-3	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
Carlson 2013	154	3359 (524)	147	3187 (602)		4.12%	172[44.25,299.75]
Gustafson 2013	22	3416.8 (552.9)	24	3435.5 (404.8)		0.84%	-18.7[-300.85,263.45]
Haghiac 2015	25	3278 (448)	24	2935 (356)		1.31%	343[116.89,569.11]
Harper 2010	427	2990 (1005.8)	410	2923 (1252.8)	+-	2.82%	67[-87.29,221.29]
		Н	igher wit	h no omega-3	-500 -250 0 250 500	Higher with	omega-3





ADDITIONAL TABLES

Table 1. Maternal age (years)

Ali 2017 27 (4.3) 27 (4.8) Bergmann 2007 30.9 (4.6) for DHA/FOS group 30.0 (4.62) in vitamin/mineral group; 31 for FOS group Bisgaard 2016; 32.3 (4.3) 32.2 (4.5) Boris 2004 "The three study groups were similar in baseline characteristics with regard to maternal age at deligible (data not shown)". Bosaeus 2015 31.4 (3.9) 31.2 (4.0) Bulstra-Ramakers 1994 Not reported Carlson 2013 25.3 (4.9) 24.8 (4.7) Chase 2015 Not reported	Study ID	Omega-3 (mean (SD)unless otherwise reported)	No omega-3 (mean (SD)unless otherwise reported)
Bisgaard 2016; 32.3 (4.3) 32.2 (4.5) Boris 2004 "The three study groups were similar in baseline characteristics with regard to maternal age at delir (data not shown)". Bosaeus 2015 31.4 (3.9) 31.2 (4.0) Bulstra-Ramakers 1994 Not reported Carlson 2013 25.3 (4.9) 24.8 (4.7)	Ali 2017	27 (4.3)	27 (4.8)
Boris 2004 "The three study groups were similar in baseline characteristics with regard to maternal age at delir (data not shown)". Bosaeus 2015 31.4 (3.9) 31.2 (4.0) Bulstra-Ramakers 1994 Not reported Carlson 2013 25.3 (4.9) 24.8 (4.7)	Bergmann 2007	30.9 (4.6) for DHA/FOS group	30.0 (4.62) in vitamin/mineral group; 31 (4.71) for FOS group
(data not shown)". Bosaeus 2015 31.4 (3.9) 31.2 (4.0) Bulstra-Ramakers 1994 Not reported Carlson 2013 25.3 (4.9) 24.8 (4.7)	Bisgaard 2016;	32.3 (4.3)	32.2 (4.5)
Bulstra-Ramakers 1994 Not reported Carlson 2013 25.3 (4.9) 24.8 (4.7)	Boris 2004		cteristics with regard to maternal age at delivery
Carlson 2013 25.3 (4.9) 24.8 (4.7)	Bosaeus 2015	31.4 (3.9)	31.2 (4.0)
	Bulstra-Ramakers 1994	Not reported	
Chase 2015 Not reported	Carlson 2013	25.3 (4.9)	24.8 (4.7)
	Chase 2015	Not reported	
D'Almedia 1992 "Ages ranged from 14-40 years"	D'Almedia 1992	"Ages ranged from 14-40 years"	



able 1. Maternal ag		
de Groot 2004	30.0 (3.3)	29.2 (3.8)
Dilli 2018	30.9 (5.3)	32.7 (5.9)
Dunstan 2008	30.9 (3.7)	32.6 (3.6)
England 1989	Not reported	
Freeman 2008	31.0 (5.8)	29.7 (6.2)
Furuhjelm 2009	31.1 (4.1)	31.7 (3.9)
Giorlandino 2013	32.6 (4.6)	32.2 (4.8)
Gustafson 2013	25.5 (4.3)	25.6 (4.8)
Haghiac 2015	27 (5)	27 (5)
Harper 2010	Median (interquartile range): 28 (23 - 32)	Median (interquartile range): 27 (24-32)
Harris 2015	In high-dose group 24.5 (12.72);	27.0 (9.05)
	In low-dose group 24.3 (12.72)	
Hauner 2012	31.9 (4.9)	31.6 (4.5)
Helland 2001	28.6 (3.4)	27.6 (3.2)
Horvaticek 2017	29.8 (5.5)	29.6 (4.8)
Hurtado 2015	30.5 (4.8)	29.9 (4.7)
Ismail 2016	27.17 (6.34)	26.71 (5.66)
Jamilian 2016	30.1 (5.3)	30.0 (5.5)
Jamilian 2017	30.7 (3.5) for omega-3 group	30.7 (4.1) for placebo group
	31.2 (4.3) for omega-3 + vitamin D group	31.5 (7.0) for vitamin D group
Judge 2007	23.9 (4.3)	24.7 (4.8)
Judge 2014	Not reported	
Kaviani 2014	26.33 (4.2)	25.15 (4.2)
Keenan 2014	Not reported	
Khalili 2016	25.9 (4.8)	26.9 (4.5)
Knudsen 2006	28.4 for 0.1 g/day EPA + DHA group	28.5 for no treatment group
	28.7 for 0.3 g/day EPA + DHA group	
	28.4 for 0.7 g/day EPA + DHA group	
	28.9 for 1.4 g/day EPA + DHA group	



Table 1.	Maternal age (years) (Continued)
	28.8 for 2.8 g/day EPA + DHA group

	28.8 for 2.2g/day ALA group	
Krauss-Etschmann 2007	Median (range): 30.6 (20.1 - 41.1) for DHA/EPA group	Median (range): 31.1 (18.8 - 40.8) for folate
	Median (range): 31.1 (21.5 - 40.1) for DHA/EPA+folate group	group Median (range): 31.1 (18.4 - 40.3) for no treatment (placebo) group
Krummel 2016	27.9 (4.6)	26.3 (5.0)
Laivuori 1993	Median (IQR): 30.3 (24-40)	Median (IQR): 30.2 (26-32) in placebo group; 32.0 (23-40) in primrose oil group
Makrides 2010	28.9 (5.7)	28.9 (5.6)
Malcolm 2003	Not reported	
Mardones 2008	25.06 (5.73)	25.11 (7.45)
Martin-Alvarez 2012	Not reported	
Miller 2016	31.7 (4.4)	31.2 (4.4)
Min 2014	Median (range): 29 (18 - 42)	Median (range): 29 (18 - 44)
Min 2014 [diabetic women]	Median (range): 34 (20 - 45)	Median (range): 37 (27-45)
Min 2016	Median (range): 31.0 (21.0 - 41.0)	Median (range): 32.0 (21.0 - 44.0)
Mozurkewich 2013	30.6 (4.5) in DHA rich fish oil group; 29.9 (5.0) in EPA rich fish oil group	30.4 (5.9)
Mulder 2014	32.6 (4.04)	33.4 (3.61)
Noakes 2012	29.5 (3.94)	28.4 (4.69)
Ogundipe 2016	Not reported	
Oken 2013	Median (IQR): 32.6 (27.9 - 35.9) advice group;	Median (IQR): 32.4 (27.7 to 34.3)
	27.6 (24.5 - 32.0) advice + gift card group	
Olsen 1992	29.4 (4.4)	olive oil group 29.7 (4.3); placebo/no oil group 29.1 (4.1)
Olsen 2000	Prophylactic trials	Prophylactic trials
	PD trial 29.3 (4.87)	PD trial 30.0 (6.22)
	IUGR trial 30 (4.64)	IUGR trial 29.0 (3.93)
	PIH trial 30.3 (7.01)	PIH trial 28.9 (5.32)
	Twins trial 30.2 (6.18)	Twins trial 30.2 (6.35)
	Therapeutic trials	Therapeutic trials



	Threat-PE trial 32.1 (11.7)	Threat-PE trial 32.9 (14.6)
	Susp-IUGR trial 29.3 (7.88)	Susp-IUGR trial 29.8 (10.3)
Olsen 2000 [twins]	see Olsen 2000	
Onwude 1995	Mean (range): 26.6 (18-39)	Mean (range): 26.1 (16-40)
Otto 2000	30.3 (5.2)	28.3 (4.85)
Pietrantoni 2014	30.86 (4.18)	29.92 (4.80)
Ramakrishnan 2010	26.2 (4.6)	26.3 (4.8)
Ranjkesh 2011	30.06 (7.59)	28.96 (6.40)
Razavi 2017	29.7 (3.6) for omega-3 group	29.2 (3.4) for placebo group
	29.9 (4.0) for omega-3 + vitamin D group	29.9 (5.0) for vitamin D group
Rees 2008	31.2 (4.4)	34.5 (3.8)
Ribeiro 2012	Not reported	
Rivas-Echeverria 2000	Not reported	
Samimi 2015	Median (range): 26.8 (18-39)	Median (range): 26.1 (16-40)
Sanjurjo 2004	34.5 (7.41)	31.25 (5.18)
Smuts 2003a	21.7 (4.3)	21.6 (4.2)
Smuts 2003b	High DHA egg group 19.9 (4.1)	Ordinary egg group 24.8 (7.8)
Su 2008	30.9 (3.9)	31.3 (5.7)
Taghizadeh 2016	28.6 (6.3)	29.4 (4.4)
Tofail 2006	22.1 (4.2)	23.4 (4.5)
Valenzuela 2015	29 (4.7)	28.3 (6.7)
Van Goor 2009	Median (range): 32.3 (22.3 - 43.3) in DHA group;	Median (range): 33.5 (26.0 - 40.3)
	31.5 (24.8 - 41.4) in DHA + AA group	
Van Winden 2017	Not reported	
Vaz 2017	Median (IQR): 25.5 (22.0-34.5)	Median (IQR): 27.0 (21.0 - 31.0)

Abbreviations: IQR (interquartile range)

Table 2. Maternal parity

Study ID Omega-3 No omega-3	
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Ali 2017	Mean (SD): 2.9 (4.8)	Mean (SD): 2.8 (1.6)
Bergmann 2007	> 1: 22 (45.8%) in DHA/FOS group	> 1: 28 (57.1%) in vitamin/mineral group
		24 (51.1%) in FOS group
Bisgaard 2016;	1: 155 (44.8%)	1: 166 (47.6%)
Boris 2004	Not reported	
Bosaeus 2015	Median (IQR): 0.5 (0,1) Median (IQR): 0 (0,1)	
Bulstra-Ramakers 1994	Not reported	
Carlson 2013	Prior pregnancies, N	Prior pregnancies, N
	Mean (SD): 1.2 (1.3)	Mean (SD): 1.3 (1.4)
Chase 2015	Not reported	
D'Almedia 1992	Not reported	
de Groot 2004	0: 11 (38%)	0: 12 (41%)
	1: 15 (52%)	1: 11 (38%)
	2: 3 (10%)	2: 5 (17%)
	3: 0 (0%)	3: 1 (3%)
Dunstan 2008	≥ 1: 15 (45.5%)	≥ 1: 21 (53.8%)
England 1989	Not reported	
Freeman 2008	Primiparous: 24 (77.4%)	Primiparous: 22 (78.6%)
Furuhjelm 2009	Not reported	
Giorlandino 2013	Not reported	
Gustafson 2013	Not reported	
Haghiac 2015	0: 7 (28%)	0: 5 (21%)
	1:18 (72%)	1: 19 (79%)
Harper 2010	Not reported	
Harris 2015	Not reported	
Hauner 2012	Primiparous: 55.8%	Primiparous: 61.2%
Helland 2001	Mean (SD): 0.3 (0.5)	Mean (SD): 0.3 (0.5)
Horvaticek 2017	Nulliparous: 25 (53%)	Nulliparous: 26 (60%)
	Primiparous: 22 (47%)	Primiparous: 17 (40%)



Hurtado 2015	Multiparous: 35.6%	Multiparous: 31.8%		
Ismail 2016	Mean (SD): 1.38 (1.67)	Mean (SD): 1.53 (1.55)		
Jamilian 2016	Not reported			
Jamilian 2017	Not reported			
Judge 2007	Mean (SD): 1.5 (0.8)	Mean (SD): 1.8 (0.8)		
Judge 2014	Not reported			
Kaviani 2014	Not reported			
Keenan 2014	Not reported			
Khalili 2016	1: 38 (50.7%)	1: 37 (49.3%)		
	2: 28 (37.3%)	2: 27 (36%)		
	≥ 3: 9 (12.0%)	≥ 3: 11 (14.7%)		
Knudsen 2006	Primiparous women	Primiparous women		
	0.1 g/day EPA + DHA group: 257 (66.2%)	No treatment group: 513 (66.4%)		
	0.3 g/day EPA + DHA group: 267 (69.5%)			
	0.7 g/day EPA + DHA group: 244 (63.5%)			
	1.4 g/day EPA + DHA group: 247 (64.7%)			
	2.8 g/day EPA + DHA group: 246 (62.9%)			
	2.2 g/day ALA group: 258 (66.3%)			
Krauss-Etschmann 2007	< 2: 56 (86%) for DHA/EPA group; 56 (88%) for DHA/EPA +folate group	< 2: 65 (90%) for folate group; 61 (88%) for placebo group		
	2: 7 (11%) for DHA/EPA group; 6 (9%) for DHA/EPA+folate group	2: 7 (10%).for folate group; 7 (10%) for placebo group		
	> 2: 2 (3%) for DHA/EPA group; 2 (3%) for DHA/EPA+folate group	> 2: 0 (0) for folate group; 1 (1%) for placebo group		
Krummel 2016	Not reported			
Laivuori 1993	Nulliparous: 2 (66%) in fish oil group	Nulliparous: 1 (25%) in primrose oil group; 3		
	Primiparous: 1 in (33%) fish oil group	(75%) in placebo group		
		Primiparous: 3 (60%) in primrose oil group; 2 (40%) in placebo group		
Makrides 2010	Primiparous: 471 (39.3%)	Primiparous: 474 (39.4%)		
	Not reported			
Malcolm 2003	Mean (SD): 1.68 (0.90) Mean (SD): 1.74 (0.91)			
Mardones 2008	Mean (SD): 1.68 (0.90)	Mean (SD): 1.74 (0.91)		



Table 2. Maternal pa	Arity (Continued) Not reported				
Min 2014	0: 18 (40%)	0: 14 (35.0%)			
	1-3: 26 (57.8%)	1-3: 23 (57.5%)			
	> 4: 1 (2.2%)	> 4: 2 (5.0%)			
Min 2014 [diabetic	0: 10 (24%)	0: 7 (14.9%)			
women]	1-3: 27 (65.9%)	1-3: 32 (68.1%)			
	> 4: 3 (7.3%)	> 4: 6 (12.8%)			
Min 2016	0: 33 (50%)	0: 24 (35%)			
	1-3: 27 (41%)	1-3: 40 (57%)			
	≥ 4: 6 (9%)	≥ 4: 5 (7%)			
Mozurkewich 2013	Mean (SD):	Mean (SD): 0.85 (1.2)			
	0.87 (0.83) for EPA rich fish oil group;				
	1.08 (0.94) for DHA rich fish oil group				
Mulder 2014	1: 60.6%	1: 47.7%			
	2: 30.8%	2: 36.7%			
	> 2: 8.6%	> 2: 15.6%			
Noakes 2012	Not reported				
Ogundipe 2016	Not reported				
Oken 2013	Primiparous:	Primiparous:			
	6 (35%) in advice group;	6 (30%) in control group			
	4 (24%) in advice + gift card group				
Olsen 1992	Primiparous:	Primiparous:			
	Fish oil group: 56%	Olive oil group: 61%			
		No oil group: 60%			
Olsen 2000	Prophylactic trials: no nulliparous women except for:	Prophylactic trials: no nulliparous women except for:			
	Twins trial: 52.5% nulliparous	Twins trial: 52.5% nulliparous			
	Therapeutic trials	Therapeutic trials			
	Threat-PE trial: 71.4% nulliparous	Threat-PE trial: 65.6% nulliparous			
	Susp-IUGR trial: 52.0% nulliparous	Susp-IUGR trial: 51.9% nulliparous			
Onwude 1995	Included primiparous and multiparous women				
Otto 2000	Primiparous: 8 (67%)	Primiparous: 5 (42%)			



Pietrantoni 2014	0: 46 (36%)	0: 50 (40%)	
	1: 83 (64%)	1: 76 (60%)	
Ramakrishnan 2010	Not reported		
Ranjkesh 2011	Mean (SD): 0.46 (0.50) Mean (SD): 0.40 (0.49)		
Razavi 2017	Not reported		
Rees 2008	Mean (SD): 1.4 (0.9) Mean (SD): 1.6 (1.2)		
Ribeiro 2012	Not reported		
Rivas-Echeverria 2000	Excluded nulliparous women		
Samimi 2015	Not reported		
Sanjurjo 2004	Mean (SD): 1.63 (0.74) Mean (SD): 1.38 (0.52)		
Smuts 2003a	Nulliparous before study: Nulliparous before study:		
	68%	58%	
Smuts 2003b	Women were excluded if they had more than 4 previous pregnancies	Mean (SD): 2.3 (1.9)	
	Mean (SD): 1.9 (1.1)		
Su 2008	Mean (SD): 1.7 (1.1) Mean (SD): 1.8 (1.1)		
Taghizadeh 2016	Not reported		
Tofail 2006	Women with > 2 children: 16.8%	Women with > 2 children: 31.5%	
Valenzuela 2015	Included women with 1-4 prior births		
Van Goor 2009	Included women with a first or second pregnancy		
Van Winden 2017	Not reported		
Vaz 2017	0-1: 26 (81.2%)	0-1: 18 (64.3%)	
	≥ 2: 6 (18.8%)	≥ 2: 10 (35.7%)	

Table 3. Maternal omega-3 intake criteria

Carlson 2013 Excluded women taking ≥ 300 mg DHA a day Chase 2015 Excluded women planning to take DHA during pregnancy	у	Eligibility criteria
Chase 2015 Excluded women planning to take DHA during pregnancy	on 2013	Excluded women taking ≥ 300 mg DHA a day
	e 2015	Excluded women planning to take DHA during pregnancy
de Groot 2004 Excluded women consuming fish more than twice a week	root 2004	Excluded women consuming fish more than twice a week



Table 3. Maternal omega-3 intake criter

Dunstan 2008	Excluded women consuming fish more than twice a week		
Freeman 2008	Excluded women with a previous intolerance to omega-3 fatty acids		
Furuhjelm 2009	Excluded women with an allergy to fish or undergoing treatment with omega-3 fatty acid supplements		
Giorlandino 2013	Excluded women with an allergy to fish or regular intake of fish oil		
Gustafson 2013	Excluded women taking more than 200 mg DHA a day		
Haghiac 2015	Excluded women with an allergy to fish or fish products; women who do not eat any fish; and women with a regular intake of fish oil (> 500 mg/week in the previous 4 weeks)		
Harper 2010	Excluded women with an allergy to fish or fish products; and women with a regular intake of fish oil supplements (> 500 mg/week at any time during the preceding month)		
Harris 2015	Excluded women with allergies to fish or consumption of salmon, mackerel, rainbow trout or sar- dines at least weekly		
Hauner 2012	Excluded women taking omega-3 supplementation before randomisation		
Helland 2001	Excluded women already taking DHA		
Hurtado 2015	Did not include women taking DHA supplements in pregnancy		
Jamilian 2017	Excluded women taking omega-3 fatty acid supplements		
Kaviani 2014	Excluded women consuming fish more than twice a week		
Keenan 2014	Excluded women consuming ≥ 2 servings of sea fish a week		
Khalili 2016	Excluded women with an allergy to fish oil or fish products; and women consuming fish more than twice a week		
Knudsen 2006	Included women with only limited fish intake and who did not use fish oil capsules during pregnancy		
Krauss-Etschmann 2007	Excluded women who had used fish oil supplements since the beginning of their pregnancy		
Krummel 2016	Excluded women who consumed > 1 fish meal/week or who used DHA-fortified foods or supplements		
Makrides 2010	Excluded women who were already taking DHA supplements		
Malcolm 2003	Excluded women with an allergy to fish products		
Miller 2016	Excluded women with an allergy to seafood or fish oils		
Min 2016	Excluded women taking fish oil supplements		
Mozurkewich 2013	Excluded women taking omega-3 fatty acid supplements and women consuming > 2 fish meals a week		
Mulder 2014	Excluded women taking any lipid or fatty acid supplementation		



Table 3. Matern	ıl omega-3 inta	ke criteria (Continued)
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included women with a diet low in oily fish (excluding canned tuna) ≤ twice per month		
Excluded women with an allergy to fish and fish oil and women previously regularly taking a pre conception fish oil supplement		
Excluded women consuming fish > 3 times a month; or with no contraindications to fish consumption such as allergy, or self-restrictions such as a vegetarian diet		
Excluded women with a fish allergy or regular intake of fish oil		
Excluded women with a fish allergy or regular intake of fish oil		
Only included women who consumed fish at least twice a week (equivalent to 600 g fish a week)		
Excluded women regularly taking fish oil or DHA supplements		
Excluded women taking omega-3 fatty acid supplements		
Excluded women taking fish oil supplements or eating more than 3 oily fish portions per week; no showing any signs of intolerance or allergy to fish		
Excluded women with any signs of intolerance or allergy to fish or using dietary supplements containing omega-3 and omega-6 PUFA		
Excluded women with a diet including polyunsaturated fatty acids (PUFA, ALA supplements) or LCPUFA (EPA and or DHA supplements)		
Excluded women who were vegetarians or vegans		
Excluded women taking any oil supplementation (such as fish oil, flaxseed oil or cod liver oil)		

Table 4. Maternal socioeconomic status

Study ID	omega-3	no omega-3	
Ali 2017	Not reported		
Bergmann 2007 Employed: 31 (77.5%) in DHA/folate group 13 years of schooling: 32 (66.7%) in DHA/folate group		Employed: 35 (85.4%) in Vit/Min group; 30 (78.9%) in folate group 13 years of schooling: 28 (57.1%) in Vit/Min group; 32 (68.1%) in folate group	
Bisgaard 2016;	Household annual income: Low: 33 (9.6%) Medium: 179 (51.9%) High: 133 (38.6%)	Household annual income: Low: 34 (9.7%) Medium: 187 (53.6%) High: 128 (36.7%)	
Boris 2004	Not reported		
Bosaeus 2015	15 or more years of education: 17 (94.4%)	15 or more years of education: 15 (88.2%)	



Table 4.	Materna	l socioeconom	ic status	(Continued)
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Bulstra-Ramakers 1994	Not reported	
Carlson 2013	Maternal education:	Maternal education:
	Mean (SD): 13.69 years (2.67)	Mean (SD): 13.36 years (2.72)
Chase 2015	Not reported	
D'Almedia 1992	"Sixty-nine percent were employed; ninety-four per	cent of their husbands were employed".
de Groot 2004	Education measured on an 8-point scale:	Education measured on an 8-point scale:
	Mean (SD): 4.3 (1.4)	Mean (SD): 3.9 (1.5)
Dunstan 2008	Maternal education:	Maternal education:
	10-12 years: 10 (30.3%)	10-12 years: 9 (23.1%)
	> 12 years: 23 (69.7%)	> 12 years: 30 (76.9%)
England 1989	Not reported	
Freeman 2008	Maternal employment: 61.3% employed	Maternal employment: 60.7% employed
	Maternal education: Mean (SD): 15.5 years ((2.1)	Maternal education, Mean (SD): 14.6 years (2.2)
Furuhjelm 2009	Not reported	
Giorlandino 2013	Not reported	
Gustafson 2013	Maternal education:	Maternal education:
	Mean (SD): 14.0 years (3.1)	Mean (SD): 13.9 years (2.7)
Haghiac 2015	Not reported	
Harper 2010	Maternal education:	Maternal education:
	Median (IQR): 13 years (12-16)	Median (IQR): 13 years (12-16)
Harris 2015	Not reported	
Hauner 2012	Maternal education:	Maternal education:
	63.8% attended ≥ 12 years of school	69.9% attended ≥ 12 years of school
Helland 2001	Maternal education:	Maternal education:
	< 10 years: 2.9%	< 10 years: 1.8%
	10-12 years: 21.4%	10-12 years: 31.1%
	> 12 years: 75.7%	> 12 years: 67.1%
Horvaticek 2017	Not reported	
Hurtado 2015	Not reported	
Ismail 2016	Not reported	



Jamilian 2016	oeconomic status (Continued) Not reported	
Jamilian 2017	Not reported	
Judge 2007	Maternal education:	Maternal education;
	Mean (SD): 12.8 years (2.2)	Mean (SD): 12.2 years (1.5)
Judge 2014	Not reported	
Kaviani 2014	Maternal education:	Maternal education:
	< 6 years: 7.5%	< 6 years: 7.5 %
	6 to 9 years: 12.5%	6 to 9 years: 15%
	9 to 12 years: 20%	9 to 12 years: 10%
Keenan 2014	Not reported	
Khalili 2016	Maternal education:	Maternal education:
	Primary school (1-5 years): 14 (18.7%)	Primary school (1-5 years): 15 (20.0%)
	Seconday school (6-8 years): 23 (30.7%)	Seconday school (6-8 years): 14 (18.7%)
	High school (9-12 years): 33 (44.0%)	High school (9-12 years): 39 (52.0%)
	University (> 12 years): 5 (6.7%)	University (> 12 years): 7 (9.3%)
	Family income:	Family income;
	Adequate: 15 (20%)	Adequate: 13 (17.3%)
	Relatively adequate: 44 (58.7%)	Relatively adequate: 50 (66.7%)
	Non adequate: 16 (21.3%)	Non adequate: 12 (16.0%)
Knudsen 2006	Not reported	
Krauss-Etschmann 2007	Job training of father:	Job training of father:
	None: 29 (45%) for DHA/EPA group; 17 (27%) for DHA/EPA+folate group	None: 33 (47%) for folate group; 27 (40%) for place- bo group
	Apprenticeship: 14 (22%) for DHA/EPA group; 19 (31%) for DHA/EPA+folate group	Apprenticeship: 10 (14%) for folate group; 14 (21%) for placebo group
	University degree: 15 (23%) for DHA/EPA group; 21 (34%) for DHA/EPA+folate group	University degree: 24 (34%) for folate group; 20 (29%) for placebo group
Krummel 2016	Education:	Education:
	Mean (SD): 14.8 years (2.1)	Mean (SD): 14.9 years (3.2)
Laivuori 1993	Not reported	
Makrides 2010	Mother completed secondary education: 755 (63.1%)	Mother completed secondary education: 760 (63.2%)
	Mother completed further education: 816 (68.2%)	Mother completed further education: 824 (68.6%)



able 4. Maternal so	MSSI score: median 28.5, IQR (25.0 - 31.0)	MSSI score: median 29.0, IQR (25.0 - 31.0)
Malcolm 2003	Not reported	
Mardones 2008	Education:	Education:
	> 8 years: 82.1%	> 8 years: 80.7%
	ESOMAR classification:	ESOMAR classification:
	AB (high level): 0.5%	AB (high level): 0.3%
	CA (medium level): 4.4%	CA (medium level): 4.2%
	CB (medium level): 34.9%	CB (medium level): 33.4%
	D (medium - low level): 40.4%	D (medium - low level): 44.6%
	E (low level): 19.8%	E (low level): 17.5%
Martin-Alvarez 2012	Not reported	
Miller 2016	Not reported	
Min 2014	Not reported	
Min 2014 [diabetic women]	Not reported	
Min 2016	Not reported	
Mozurkewich 2013	Not reported	
Mulder 2014	Not reported	
Noakes 2012	Not reported	
Ogundipe 2016	Not reported	
Oken 2013	Working full time: 6 (35%) for advice to eat fish group;	Working full time: 7 (35%) for control group
	9 (50%) for advice to eat fish + gift card group	
Olsen 1992	Not reported	
Olsen 2000	Not reported	
Olsen 2000 [twins]	see Olsen 2000	
Onwude 1995	Not reported	
Otto 2000	Not reported	
Pietrantoni 2014	High school or university degree: 129 (100%)	High school or university degree: 126 (100%)
	Average socioeconomic status (not defined): 129 (100%)	<u>Average socioeconomic status</u> (not defined): 126 (100%)



Ramakrishnan 2010	High school education or above: 56.6%	High school education or above: 59.5%	
Ranjkesh 2011	Not reported		
Razavi 2017	Not reported		
Rees 2008	Maternal education:	Maternal education:	
	Mean (SD): 14.5 years (3.5)	Mean (SD): 15.3 (2.9)	
Ribeiro 2012	Not reported		
Rivas-Echeverria 2000	Not reported		
Samimi 2015	Not reported		
Sanjurjo 2004	Not reported		
Smuts 2003a	"Most subjects received government assistance for n	nedical care"	
Smuts 2003b	Not reported		
Su 2008	Not reported		
Taghizadeh 2016	Not reported		
Tofail 2006	Mostly low-income participants	Mostly low-income participants	
	Mothers with > 5 years of schooling: 36.8%	Mothers with > 5 years of schooling: 32.3%	
	Working mothers: 16.0	Working mothers: 12.1%	
	Fathers with stable job: 65.6	Fathers with stable job: 65.3%	
	Family income (taka/month, 1 USD = 59 taka): 64.0	Family income (taka/month, 1 USD = 59 taka): 54.0	
Valenzuela 2015	SES assessed using the ESOMAR criteria:	SES assessed using the ESOMAR criteria:	
	High: 5.3%	High: 19.0%	
	Medium: 73.7%	Medium: 66.7%	
	Low: 21.1%	Low: 14.3%	
Van Goor 2009	Not reported		
Van Winden 2017	Not reported		
Vaz 2017	Family income, not further defined:	Family income (US \$) not further defined:	
	US \$263.2 (181.9-383.0)	US \$304.1 (180.7 - 379.8)	
	Maternal education:	Maternal education:	
	Median (IQR): 11.0 years (7.0 - 11)	Median (IQR): 8.0 years (7.5 - 10.5)	

Abbreviations: DHA: docosahexaenoic acid; EPA: eicosapentaenoic acid; ESOMAR: European Society for Opinion and Marketing Research; IQR: interquartile range; MSSI: maternal social support index; SD: standard deviation; SES: socioeconomic status



Table 5. Maternal ethnicity

Study ID	Omega-3	No omega-3
Ali 2017	Not reported (study conducted in Egypt)	
Bergmann 2007	"Caucasian women"	
Bisgaard 2016	Caucasian:	Caucasian:
	333 (96.2%)	332 (95.1%)
Boris 2004	Not reported (conducted in Denmark)	
Bosaeus 2015	Women of European descent	
Bulstra-Ramakers 1994	Not reported (study conducted in the Netherlands)	
Carlson 2013	Hispanic: 8%	Hispanic: 8%
	Not Hispanic: 92%	Not Hispanic 92%
	African-American: 38%	
Chase 2015	Maternal ethnicity not reported;	Maternal ethnicity not reported;
	reported that 98% of included infants were white	reported that 93% of included infants were white
D'Almedia 1992	Not reported (conducted in Angola)	
de Groot 2004	"White women"	
Dunstan 2008	Caucasian women	
England 1989	Not reported (conducted in South Africa)	
Freeman 2008	Not reported (conducted in USA)	
Furuhjelm 2009	Not reported (conducted in Sweden)	
Giorlandino 2013	Not reported (conducted in Italy)	
Gustafson 2013	28% African-American (conducted in USA)	
Haghiac 2015	African American: 11 (44%)	African American: 6 (25%)
	Caucasian: 10 (40%)	Caucasian: 11 (46%)
	Other (e.g. Hispanic or Asian): 4 (16%)	Other (e.g. Hispanic or Asian): 7 (29%)
Harper 2010	African American: 148 (34.1%)	African American: 145 (34.9%)
	White: 245 (56.5%)	White: 240 (57.7%)
	Asian: 13 (3.0%)	Asian: 5 (1.2%)
	Other: 28 (6.5%)	Other: 26 (6.3%)
	Hispanic/Latina ethnicity: 64 (14.7%)	Hispanic/Latina ethnicity: 57 (13.6%)



Table 5. Maternal ethr		
Harris 2015	Not reported (conducted in USA)	
Hauner 2012	Not reported (conducted in Germany)	
Helland 2001	Not reported (conducted in Norway)	
Horvaticek 2017	Not reported (conducted in Croatia)	
Hurtado 2015	Not reported (conducted in Spain)	
Ismail 2016	Not reported (conducted in Egypt)	
Jamilian 2016	Not reported (conducted in Iran)	
Jamilian 2017	Not reported (conducted in Iran)	
Judge 2007	Not reported (conducted in USA)	
Judge 2014	Not reported (conducted in USA)	
Kaviani 2014	Not reported (conducted in Iran)	
Keenan 2014	African American women	
Khalili 2016	Not reported (conducted in Iran)	
Knudsen 2006	Not reported (conducted in Denmark)	
Krauss-Etschmann 2007	Not reported (conducted in Spain, Germany or Hungary)	
Krummel 2016	African American: 12 (37.5%)	African American: 15 (53.6%)
	White: 20 (62.5%)	White: 13 (46.4%)
Laivuori 1993	Not reported (conducted in Finland)	
Makrides 2010	Not reported (conducted in Australia)	
Malcolm 2003	Not reported (conducted in UK)	
Mardones 2008	"mainly ethnically mixed (American and Hispanic)"	
Martin-Alvarez 2012	Not reported (conducted in Spain)	
Miller 2016	African American: 1 (1.7%)	African American: 0 (0%)
	Caucasian: 55 (92%)	Caucasian: 52 (95%)
	Hispanic: 2 (3%)	Hispanic: 2 (3%)
	Asian: 1 (1.67%)	Asian: 1 (2%)
	Other: 1 (1.67%)	Other: 0 (0%)
Min 2014	Asian: 16 (35.6%)	Asian: 18 (45.0%)
	African/Afro-Caribbean: 10 (22.2%)	African/Afro-Caribbean: 14 (35.0%)
	Caucasian: 13 (28.9%)	Caucasian: 6 (15.0%)



	Others: 6 (13.3%)	Others: 2 (5.0%)
Min 2014 [diabetic	Asian: 18 (43.9%)	Asian: 27 (57.5%)
women]	African/Afro-Caribbean: 15 (36.6%)	African/Afro-Caribbean: 10 (21.3%)
	Caucasian: 5 (12.2%)	Caucasian: 5 (10.6%)
	Others: 3 (7.3%)	Others: 5 (10.6%)
Min 2016	Asian: 40 (60%)	Asian: 44 (62%)
	African/Afro-Caribbean: 18 (27%)	African/Afro-Caribbean: 18 (25%)
	Caucasian: 5 (7%)	Caucasian: 5 (7%)
	Others: 4 (7%)	Others: 4 (6%)
Mozurkewich 2013	White: 33 (85%) for EPA-rich group; 29 (76%) for DHA-rich	White: 34 (83%)
	group	African-American: 2 (5%)
	African-American: 4 (10%) for EPA-rich group; 4 (11%) for DHA-rich group	Hispanic-Latina: 3 (7%)
	Hispanic-Latina: 0 (0%) for EPA-rich group; 4 (11%) for	Asian: 1 (2%)
	DHA-rich group	American Indian or Alaska Native: 1 (2%)
	Asian: 1 (3%) for EPA-rich group; 1 (3%) for DHA-rich group	Native Hawaiian or other Pacific ethnicity: (0%)
	American Indian or Alaska Native: 0 (0%) for EPA-rich group; 0 (0) for DHA-rich group	
	Native Hawaiian or other Pacific ethnicity: 1 (3) for EPArich group; 0 (0%) for DHA-rich group	
Mulder 2014	White: 73.1%	White: 73.9%
	Non-white: 26.9%	Non-white: 26.1%
Noakes 2012	Not reported (conducted in UK)	
Ogundipe 2016	Not reported (conducted in UK)	
Oken 2013	White: 9 (50%) advice only group; 9 (53%) advice+vouch-	White: 9 (45%)
	er group	Black: 2 (10%)
	Black: 2 (11%) advice only group; 2 (12%) advice+voucher group	Asian: 3 (15%)
	Asian: 2 (11%) advice only group; 1 (6%) advice+voucher group	Hispanic/other: 6 (30%)
	Hispanic/other: 5 (28%) advice only group; 5 (29%) advice+voucher group	
Olsen 1992	Not reported (conducted in Denmark)	
Olsen 2000	Not reported (conducted in Denmark, Scotland, Sweden, England, Italy, Netherlands, Norway, Belgium and Russia)	
Olsen 2000 [twins]	See Olsen 2000	



Table 5.	Materna	l ethnicity	(Continued)
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Onwude 1995	Not reported (conducted in UK)	
Otto 2000	Not reported (conducted in the Netherlands)	
Pietrantoni 2014	Caucasians	
Ramakrishnan 2010	Not reported (conducted in Mexico)	
Ranjkesh 2011	Not reported (conducted in Iran)	
Razavi 2017	Not reported (conducted in Iran)	
Rees 2008	Not reported (conducted in Australia)	
Ribeiro 2012	Not reported (conducted in Brazil)	
Rivas-Echeverria 2000	Not reported (conducted in Venezuela)	
Samimi 2015	Not reported (conducted in Iran)	
Sanjurjo 2004	Not reported (conducted in Spain)	
Smuts 2003a	African:104 (73%)	African: 109 (73%)
	Other: 38 (27%)	Other: 40 (27%)
Smuts 2003b	African: 15 (83%)	African: 15 (78%)
	Other: 3 (17%)	Other: 4 (22%)
Su 2008	Not reported (conducted in Taiwan)	
Taghizadeh 2016	Not reported (conducted in Iran)	
Tofail 2006	Not reported (conducted in India)	
Valenzuela 2015	Hispanic: 19 (100%)	Hispanic: 21 (100%)
Van Goor 2009	Not reported (conducted in the Netherlands)	
Van Winden 2017	Neither ethnicity, race or country where study conducted reported	
Vaz 2017	White: 13 (40.6%) White: 5 (17.9%)	
	Non-white: 19 (59.4%)	Non-white: 23 (82.1%)

Table 6. Maternal smoking status

Study ID	Omega-3	No omega-3
Ali 2017	Smokers were excluded	
Bergmann 2007	Smokers were excluded	



Bisgaard 2016	Smoking during pregnancy: 21 (6.1%)	Smoking during pregnancy: 33 (9.5%)	
Boris 2004	"The three study groups were similar in bas percentage of smokers (data not shown)".	eline characteristics with regard to	
Bosaeus 2015	Not reported		
Bulstra-Ramakers 1994	Not reported		
Carlson 2013	History of smoking: 41%	History of smoking: 45%	
	Smoking during pregnancy: 30%	Smoking during pregnancy: 38%	
Chase 2015	Not reported		
D'Almedia 1992	Not reported		
de Groot 2004	Smoking at 14 weeks GA:	Smoking at 14 weeks GA:	
	Yes: 4 (14%)	Yes: 10 (34%)	
Dilli 2018	15 (28%)	24 (35%)	
Dunstan 2008	Smokers were excluded		
England 1989	Not reported		
Freeman 2008	Not reported		
Furuhjelm 2009	Exposure to smoke: (at least 1 of immediate family a smoker)	Exposure to smoke: (at least 1 of immediate family a smoker)	
	9 (17%)	11 (17%)	
Giorlandino 2013	Maternal smoking at baseline: 50%	Maternal smoking at baseline: 48%	
Gustafson 2013	Not reported		
Haghiac 2015	Not reported		
Harper 2010	Smoking during pregnancy: 64 (15%)	Smoking during pregnancy: 72 (17%)	
Harris 2015	Not reported		
Hauner 2012	Smoking before pregnancy: 16%	Smoking before pregnancy: 24%	
Helland 2001	Smoking: 16%	Smoking: 22%	
Horvaticek 2017	Not reported		
Hurtado 2015	Not reported		
Ismail 2016	Not reported		
Jamilian 2016	Smokers were excluded		



Table 6. Maternal smol	king status (Continued)	
Jamilian 2017	Smokers were excluded	
Judge 2007	Smokers were excluded	
Judge 2014	Not reported	
Kaviani 2014	Smokers were excluded	
Keenan 2014	Regular smokers were excluded	
Khalili 2016	Not reported	
Knudsen 2006	Smoked during pregnancy	Smoked during pregnancy
	0.1 g/day EPA + DHA group: 79 (20.3%)	160 (20.7%)
	0.3 g/day EPA + DHA group: 78 (20.3%)	
	0.7 g/day EPA + DHA group: 78 (20.3%)	
	1.4 g/day EPA + DHA group: 79 (20.6%)	
	2.8 g/day EPA + DHA group: 78 (19.9%)	
	2.2g/day ALA group: 79 (20.3%)	
Krauss-Etschmann 2007	Smoking at study entry	Smoking at study entry
	Yes: 8 (12%) for DHA/EPA group; 9 (14%) for DHA/EPA + Folate group	Yes: 5 (7%) for Folate group; 9 (13%) for placebo group
Krummel 2016	"Current or previous use of tobacco" an excl	usion criteria
Laivuori 1993	Not reported	
Makrides 2010	Smoking at trial entry or leading up to pregnancy	Smoking at trial entry or leading up to pregnancy
	358 (29.9%)	407 (33.9%)
Malcolm 2003	Not reported	
Mardones 2008	Not reported	
Martin-Alvarez 2012	Not reported	
Miller 2016	Not reported	
Min 2014	Smoker	<u>Smoker</u>
	6 (13%)	0 (0%)
Min 2014 [diabetic	<u>Smoker</u>	Smoker
women]	2 (4%)	0 (0%)
Min 2016	<u>Smoker</u>	<u>Smoker</u>
	2 (3%)	0 (0%)



Table 6. Maternal smo	king status (Continued)		
Mozurkewich 2013	Not reported		
Mulder 2014	Not reported		
Noakes 2012	Not reported		
Ogundipe 2016	Not reported		
Oken 2013	<u>Never smoker</u>	<u>Never smoker</u>	
	14 (78%) in advice group; 12 (71%) in advice+gift card group	14 (70%) in control group	
Olsen 1992	<u>Smokers</u>	Smokers	
	Fish oil group: 33%	Olive oil group: 29%	
		No oil group: 33%	
Olsen 2000	<u>Smoker</u>	Smoker	
	Prophylactic trials	Prophylactic trials	
	Earl-PD trial 45%	Earl-PD trial 41%	
	Earl-IUGR trial 52%	Earl-IUGR 52%	
	Earl-PIH trial 19%	Earl-PIH trial 24%	
	Twins trial 33%	Twins trial 29%	
	Therapeutic trials	Therapeutic trials	
	Threat-PE trial 18%	Threat-PE trial 21%	
	Susp-IUGR trial 31%	Susp-IUGR trial 30%	
Onwude 1995	Current smoker	<u>Current smoker</u>	
	42 (37%)	32 (27%)	
Otto 2000	Not reported		
Pietrantoni 2014	Smokers were excluded		
Ramakrishnan 2010	Not reported		
Ranjkesh 2011	Not reported		
Razavi 2017	Smokers were excluded		
Rees 2008	<u>Smoker</u>	Smoker	
	0 (0%)	3 (23%)	
Ribeiro 2012	Not reported		
Rivas-Echeverria 2000	Not reported		
Samimi 2015	Smokers were excluded		



Tabl	le 6	. Ma	ternal	smol	king	status	(Continued)
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Sanjurjo 2004	Smoker	Smoker	
	1 (13%)	2 (25%)	
Smuts 2003a	Smoker before pregnancy: 46.8%	Smoker before pregnancy: 38.2%	
	Smoker during pregnancy: 27.0%	Smoker during pregnancy: 21.5%	
Smuts 2003b	Not reported		
Su 2008	Not reported		
Taghizadeh 2016	Smokers were excluded		
Tofail 2006	Not reported		
Valenzuela 2015	Not reported		
Van Goor 2009	Not reported		
Van Winden 2017	Not reported		
Vaz 2017	Not reported		

Table 7. Maternal risk

Study ID	All women included in the study	
Ali 2017	Increased/high-risk (pregnancy complicated with asymmetrical IUGR)	
Bergmann 2007	Low-risk (healthy women)	
Bisgaard 2016	Any/mixed risk (not reported)	
Boris 2004	Low-risk (healthy women)	
Bosaeus 2015	Low-risk (healthy women)	
Bulstra-Ramakers 1994	Increased/high-risk (women with a history of IUGR with or without PIH in the previous pregnancy)	
Carlson 2013	Low-risk (healthy women)	
Chase 2015	Increased/high-risk (Infants at risk of T1D (e.g. mothers with T1D)	
D'Almedia 1992	Mixed risk (21% of all included women had a history of PIH, and 4% a history of preterm birth)	
de Groot 2004	Low-risk (healthy women)	
Dilli 2018	Increased/high risk (women with GDM)	
Dunstan 2008	Low-risk (history of physician-diagnosed allergic rhinitis and/or asthma and 1 or more positive skin prick test to common allergens, but who were otherwise healthy)	
England 1989	Increased/high-risk (women with severe gestational proteinuric hypertension	



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Iani	1 0 7	Maternal	rick	(Continued)

Freeman 2008	Increased/high-risk (pregnant and postpartum women with a major depressive order)
Furuhjelm 2009	Low-risk (pregnant women affected by allergy themselves, of having a husband or previous child with allergies, otherwise healthy)
Giorlandino 2013	Increased/high-risk (pregnancy women with a history of IUGR, fetal demise or pre-eclampsia)
Gustafson 2013	Low-risk (healthy women)
Haghiac 2015	Increased/high-risk: (overweight or obese (BMI ≥ 25)
Harper 2010	Increased/high-risk (women with at least 1 prior spontaneous preterm birth)
Harris 2015	Low-risk (healthy women)
Hauner 2012	Low-risk (healthy women)
Helland 2001	Low-risk (healthy women)
Horvaticek 2017	Increased/high-risk (pregnant women with T1D)
Hurtado 2015	Low-risk (healthy women)
Ismail 2016	Increased/high-risk (oligohydramnios at 30-34 weeks GA)
Jamilian 2016	Increased/high-risk (women with GDM)
Jamilian 2017	Increased/high-risk (women with GDM)
Judge 2007	Low-risk (healthy women)
Judge 2014	Low-risk (healthy women)
Kaviani 2014	Increased/high-risk (women diagnosed with mild depression)
Keenan 2014	Increased/high-risk (women living in urban low-income environments)
Khalili 2016	Low-risk (healthy women)
Knudsen 2006	Any/mixed risk (not reported)
Krauss-Etschmann 2007	Low-risk (healthy women)
Krummel 2016	Increased/high-risk (all women overweight or obese)
Laivuori 1993	Increased/high-risk (women with pre-eclampsia)
Makrides 2010	Any/mixed risk
Malcolm 2003	Low-risk (healthy women) for final outcomes (any/mixed risk for preterm birth outcome)
Mardones 2008	Increased/high-risk (all included women underweight (BMI ≤ 21.2kg/m ² at 10 weeks GA))
Martin-Alvarez 2012	Any/mixed risk (not reported)
Miller 2016	Any/mixed risk



Table 7. Maternal risk (Continued)

Min 2014	Low-risk (healthy women)	
Min 2014 [diabetic women]	Increased/high-risk (women diagnosed with Type 2 diabetes)	
Min 2016	Increased/high-risk (women with GDM)	
Mozurkewich 2013	Increased/high-risk (women with a history of depression)	
Mulder 2014	Low-risk (healthy women)	
Noakes 2012	Low-risk (women with a history of allergy, atopy or asthma)	
Ogundipe 2016	Increased/high-risk: (women at risk of developing pre-eclampsia, fetal growth restriction, gestational diabetes)	
Oken 2013	Any/mixed risk	
Olsen 1992	Low-risk (healthy women)	
Olsen 2000	Increased/high-risk (previous preterm birth or IUGR in previous pregnancy or pregnancy-induced hypertension or twins in current pregnancy; threatening pre-eclampsia or ultrasonically estimated fetal weight below the 10th centile)	
Olsen 2000 [twins]	See Olsen 2000	
Onwude 1995	Increased/high-risk (primigravida with abnormal Doppler blood flow, previous birthweight < 3rd centile, PIH, previous unexplained stillbirth)	
Otto 2000	Low-risk (healthy women)	
Pietrantoni 2014	Low-risk (healthy women)	
Ramakrishnan 2010	Low-risk (healthy women)	
Ranjkesh 2011	Increased/high-risk (women at high risk for pre-eclampsia)	
Razavi 2017	Increased/high-risk (women diagnosed with GDM)	
Rees 2008	Increased/high-risk (current episode of major depression or dysthymia)	
Ribeiro 2012	Any/mixed (not reported)	
Rivas-Echeverria 2000	Increased/high-risk (women at risk of pre-eclampsia)	
Samimi 2015	Increased/high-risk (women with GDM)	
Sanjurjo 2004	Low-risk (healthy women)	
Smuts 2003a	Low-risk (healthy women)	
Smuts 2003b	Low-risk (healthy women)	
Su 2008	Increased/high-risk (women diagnosed with major depressive disorder between 16 weeks and 32 weeks GA)	
Taghizadeh 2016	Increased/high-risk (women with GDM)	



Table 7. Maternal risk (Continued)

Tofail 2006	Increased/high-risk (low income; 28% women undernourished)	
Valenzuela 2015	Low-risk ("women free from any known diseases that could affect fetal growth")	
Van Goor 2009	Low-risk (healthy women)	
Van Winden 2017	Increased/high-risk (women with GDM)	
Vaz 2017	Increased/high-risk (pregnant women classified at risk for postpartum depression)	

Abbreviations: ADA: American Diabetes Association; BMI: body mass index; GA: gestational age; GDM: gestational diabetes mellitus; IUGR: intrauterine growth restriction; OGTT: oral glucose tolerance test; PIH: pregnancy-induced hypertension; PPD: postpartum depression

APPENDICES

Appendix 1. Search terms for ICTRP and ClinicalTrials.gov

ICTRP

omega AND pregnancy

fish oil* AND pregnancy

marine oil* AND pregnancy

EPA AND pregnancy

DHA AND pregnancy

docosahex(a)enoic AND pregnancy

eicosapent(a)enoic AND pregnancy

n3 AND pregnancy

n-3 AND pregnancy

alpha-linoleic acid AND pregnancy

(each line was run separately and manually de-duplicated)

ClinicalTrials.gov

Advanced search

Intervention/treatment = alpha-linoleic acid OR n3 OR n-3 OR omega OR DHA OR EPA OR marine OR docosahexaenoic OR eicosapentaenoic Condition = pregnancy

WHAT'S NEW

Date	Event	Description
16 August 2018	New citation required and conclusions have changed	Preterm and early preterm birth now clearly reduced with omega-3 LCPUFA.
16 August 2018	New search has been performed	Search updated and 64 new trials added



HISTORY

Protocol first published: Issue 4, 2001 Review first published: Issue 3, 2006

Date	Event	Description
11 September 2012	Amended	Contact details updated.
17 September 2008	Amended	Converted to new review format.

CONTRIBUTIONS OF AUTHORS

In the first version of the review (Makrides 2006), Maria Makrides (MM) and Sjurdur Olsen (SO) assessed the trials for inclusion and extracted the data. MM wrote the protocol and review with contributions from Lelia Duley and SO.

For the 2018 update, Philippa Middleton (PM), Emily Shepherd (ES), Jacqueline Gould (JGou) and Judith Gomersall (JGom) joined the review author team. PM, ES and JGom assessed the trials for inclusion, extracted the data and assessed the risk of bias for the new trials. PM and JGom wrote the text, and prepared the graphs and 'Summary of findings' tables. All authors reviewed the text and contributed to editing of the review.

DECLARATIONS OF INTEREST

Philippa Middleton: none known.

Judith C Gomersall: none known.

Emily Shepherd: none known.

Sjurdur F Olsen: Sjurdur Olsen's institution has received grant funding from the Innovation Fund Denmark (grant no. 09-067124, Centre for Fetal Programming) and from the March of Dimes Foundation (6-FY-96-0240, 6-FY97-0553, 6-FY97-0521, 6-FY00-407). Sjurdur Olsen was the principal investigator of three trials included in the review: Olsen 1992, Olsen 2000 and Knudsen 2006.

Maria Makrides: Maria Makrides' institution has received grant funding (NHMRC Fellowship 1061704 Improving the outcomes of mothers and babies through nutritional interventions and NHMRC 1146806 Efficacy and safety of omega-3 DHA supplementation in preterm infants: childhood follow-up of the N3RO trial); and honorarium money from Fonterra (for advice relating to maternal and child nutrition) to fund ECR and MCR conference travel. Professor Makrides is the principal investigator of the DOMInO trial (included) and the ORIP (ongoing) trial trials. She was on the Scientific Advisory Board of Fonterra (to September 2018); and she is President Elect of the International Society for the Study of Fatty Acids and Lipids (ISSFAL).

Jacqueline Gould is an investigator on some of the DOMInO follow-up studies. She has received honoraria from the Nestle Nutrition Institute - these have been paid to Dr Gould's institution to support conference travel.

Dr Gould is an investigator in a clinical trial of high-dose DHA supplementation during pregnancy that was eligible for inclusion in this review. Dr Gould's institute has received project grant funding and a fellowship from the NHMRC and WCH Foundation for work unrelated to this review. She has received honoraria from the Nestle Nutrition Institute and Fonterra - these have been paid to Dr Gould's institution to support conference travel.

Sjurdur Olsen, Maria Makrides and Jacqueline Gould were not involved in any decisions relating to the above studies and assessment for inclusion/exclusions, risk of bias, data extraction, etc. was carried out by other members of the review team who were not directly involved in these studies.

SOURCES OF SUPPORT

Internal sources

- SAHMRI, Australia.
- Women's & Children's Hospital, Australia.



External sources

- Department for International Development, UK.
- · Medical Research Council, UK.
- National Health and Medical Research Council, Australia.
- Danish National Research Foundation, Denmark.

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

In the 2006 version (Makrides 2006), the title was updated from 'Fish oil and other prostaglandin precursor supplementation during pregnancy for reducing pre-eclampsia, preterm birth, low birthweight and intrauterine growth restriction' to 'Marine oil, and other prostaglandin precursor, supplementation for pregnancy uncomplicated by pre-eclampsia and intrauterine growth restriction' to more clearly identify the types of interventions and types of participants included.

For this 2018 update, we have expanded the scope of the review, with the following objective: "To assess the effects of omega-3 long-chain polyunsaturated fatty acids (LCPUFA), as supplements or as dietary additions, during pregnancy on maternal, perinatal, and neonatal outcomes and longer term outcomes for mother and child." We have also expanded the types of participants by including women with pre-eclampsia and other risk factors.

This change in focus has led to a change in outcomes, with outcomes arranged into primary and secondary and more specific outcomes added (e.g. types of morbidities, specific infant development outcomes), with flow-on effects to specification of subgroup analyses. We have also added large-for-gestational age as an outcome.

With changes to methods over the 12-year period, we have used the newer methods for assessing risk of bias and specified outcomes to be used in GRADE and 'Summary of findings' tables.

We have added four new authors, Philippa Middleton, Judith Gomersall, Jacqueline Gould, and Emily Shepherd; and Lelia Duley has stepped down as an author.

INDEX TERMS

Medical Subject Headings (MeSH)

*Infant, Small for Gestational Age; Dietary Supplements; Fatty Acids, Omega-3 [*administration & dosage]; Fetal Death [prevention & control]; Fetal Growth Retardation [*prevention & control]; Fish Oils [administration & dosage]; Gestational Age; Infant, Low Birth Weight; Infant, Postmature; Pre-Eclampsia [*prevention & control]; Pregnancy, High-Risk; Premature Birth [*prevention & control]; Randomized Controlled Trials as Topic; Seafood

MeSH check words

Female; Humans; Infant, Newborn; Pregnancy