

Neonatal parenteral nutrition

[D4] Lipid emulsions

NICE guideline tbc

Evidence reviews

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Draft for Consultation

These evidence reviews were developed by the National Guideline Alliance which is part of the Royal College of Obstetricians and Gynaecologists

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1 Safety and effectiveness of different lipid 2 formulations in parenteral nutrition for 3 preterm and term babies

4 Review question

5 What is the clinical effectiveness, efficacy and safety of lipid formulations from different
6 sources (for example, soya, fish oil, or mixed sources)?

7 Introduction

8 Lipid emulsions are a vital part of neonatal parenteral nutrition (PN). However, soybean oil-
9 based lipid emulsions (S-LE), which have been used conventionally, contain a high amount
10 of polyunsaturated fatty acid (PUFA) and phytosterols which may contribute to adverse
11 events in neonates, including PN associated liver disease (PNALD). There are newer lipid
12 emulsions available from other lipid sources that decrease fatty acid content. Guidance on
13 the effectiveness and safety of different lipid emulsions is required.

14 Summary of the protocol

15 Please see Table 1 for a summary of the Population, Intervention, Comparison and Outcome
16 (PICO) characteristics of this review.

17 **Table 1: Summary of the protocol (PICO table)**

Population	<ul style="list-style-type: none"> • Babies born preterm, up to 28 days after their due birth date (preterm babies) • Babies born at term, up to 28 days after their birth (term babies)
Intervention	Any concentrations of: <ul style="list-style-type: none"> • Soy based (Intralipid) • Specific Multicomponent (Soy, MCT, olive oil, fish oil) SMOFlipid • Olive oil +soy oil (clinoleic) • Fish oil (omegaven)
Comparison	Each other
Outcomes	<p>Critical</p> <ul style="list-style-type: none"> • Growth/anthropometric measures: <ul style="list-style-type: none"> ○ Weight gain ○ Linear growth ○ Head circumference • Neurodevelopmental outcomes (general cognitive abilities at two years corrected age as measured by a validated scale) • Adverse effects of lipids: <ul style="list-style-type: none"> ○ PN related liver disease (abnormal liver function, cholestasis, conjugated Hyperbilirubinaemia, Intrahepatocellular lipid) <p>Important</p> <ul style="list-style-type: none"> • Mortality • Adverse effects of lipids: <ul style="list-style-type: none"> ○ Infection including sepsis ○ Hyperglycaemia (due to high rates of infusion)



- Hypertriglyceridemia
- Hyperlipidaemia
- Duration of hospital stay
- Nutritional intake (g/kg/day, proportion of lipid received or essential fatty acids at 2 days)

1 MCT: medium chain triglycerides; PN: parenteral nutrition; SMOF: Specific multicomponent (soy, MCT, olive oil,
2 fish oil) lipid

3 PICO characteristics were agreed in collaboration with the guideline committee and the Cochrane Neonatal. See
4 the Cochrane reviews for the full review protocols. Additional outcomes were included in the Cochrane reviews
5 that will not be presented in this evidence report, as the guideline committee did not consider them to be critical or
6 important outcomes.

7 Clinical evidence

8 Included studies

9 In a collaboration with Cochrane Neonatal two Cochrane reviews were conducted specifically
10 to address this topic for the guideline to be included in this review (Kapoor 2018a and Kapoor
11 2018b). This was initially planned as an update review by Cochrane Neonatal, but
12 adjustments were made to address the needs of the current guideline (such as additional
13 comparisons and a separate review for late-preterm and term babies).

14 For methodological considerations related to this collaboration see supplementary material
15 C.

16 One review compared the safety and efficacy of different lipid emulsions in preterm (before
17 27 weeks' gestation) babies (Kapoor 2018a).

18 One review compared the safety and efficacy of different lipid emulsions in term and late
19 preterm (between 34 and 36⁺⁶ weeks' gestation) babies (Kapoor 2018b).

20 Studies are summarised in Table 2 (with hyperlinks to the full reviews) and hyperlinks to
21 study evidence tables are in appendix D.

22 See the Cochrane reviews for the literature search strategy, study selection flow chart, forest
23 plots, and GRADE tables.

24 Excluded studies

25 See the Cochrane reviews for list of excluded studies with reasons for their exclusions.

26 Summary of clinical studies included in the evidence review

27 Summaries of the studies that were included in this review are presented Table 2.

28 **Table 2: Summary of included studies**

Study	Population	Comparisons	Outcomes	Comments
Kapoor 2018a Systematic review	<u>Included in review</u> N = 29 studies N = 2037 babies	<ul style="list-style-type: none"> ● Fish oil lipid emulsion versus non-fish oil lipid emulsion ● Fish oil lipid emulsion versus another fish oil lipid emulsion 	<ul style="list-style-type: none"> ● Weight gain ● Linear growth ● Head growth ● Neurodevelopmental outcomes ● Parenteral nutrition associated liver disease (PNALD)/cholestasis 	Two of the 3 studies of babies with surgical conditions of cholestasis were stopped early which may have
	<u>Included in meta-analysis</u> N = 26 studies	<ul style="list-style-type: none"> ● Alternative lipid emulsion versus 		

Study	Population	Comparisons	Outcomes	Comments
	<p>N = 1890 babies</p> <p>Preterm babies born before 37 weeks' gestation</p> <p>Subgroups: preterm babies without surgical conditions of cholestasis; preterm babies with surgical conditions; preterm babies with cholestasis</p>	<p>soybean oil-based lipid emulsion</p> <ul style="list-style-type: none"> Alternative lipid emulsion versus another alternative lipid emulsion 	<ul style="list-style-type: none"> Death before discharge Sepsis Hyperglycaemia Hypertriglyceridemia Duration of hospital stay 	introduced bias.
<p>Kapoor 2018b</p> <p>Systematic review</p>	<p><u>Included in review</u></p> <p>N = 9 studies N = 273 babies</p> <p><u>Included in meta-analysis</u></p> <p>N = 5 studies N = 150 babies</p> <p>Term babies born at 37 weeks' gestation or after; late preterm babies born between 34⁺⁰ and 36⁺⁶ weeks' gestation</p> <p>Subgroups: term or late preterm babies with surgical conditions; term or late preterm babies with cholestasis</p>	<ul style="list-style-type: none"> Fish oil lipid emulsion versus non-fish oil lipid emulsion 	<ul style="list-style-type: none"> Weight gain Head growth Neurodevelopmental outcomes PNALD/cholestasis Death before discharge Sepsis Hyperglycaemia Hypertriglyceridemia Duration of hospital stay 	Three of the 5 studies that contributed data to the review were stopped early which may have introduced bias.

1 *PNALD: parenteral nutrition associated liver disease*

2 See appendix D for the full evidence tables.

3 Quality assessment of clinical outcomes included in the evidence review

4 GRADE was conducted to assess the quality of outcomes. Evidence was identified for critical
5 and important outcomes. The clinical evidence profiles can be found in the Cochrane
6 reviews.

1 Economic evidence

2 Included studies

3 A systematic review of the economic literature was conducted but no economic studies were
4 identified which were applicable to this review question. A single economic search was
5 undertaken for all topics included in the scope of this guideline. Please see supplementary
6 material D for details.

7 Excluded studies

8 No studies were identified which were applicable to this review question.

9 Economic evidence statements

10 No studies were identified which were applicable to this review question.

11 Summary of studies included in the economic evidence review

12 No economic evaluations were identified which were applicable to this review question.

13 Economic model

14 This question was medium priority for economic evaluation. However, the identified clinical
15 data was insufficient to inform de-novo economic modelling in this area.

16 Evidence statements

17 Clinical evidence statements

18 Evidence statements were based on the GRADE analysis carried out by the authors of the
19 Cochrane reviews.

20 Fish oil lipid emulsion (MOFS-LE) compared with non-fish oil lipid emulsion for preterm 21 babies

22 *Weight gain*

23 **Rate of weight gain**

24 • Low quality evidence from 5 randomised controlled trials (RCTs) (n=347) showed no
25 clinically important difference in rate of weight gain in babies who received MOFS-LE
26 compared with S-LE. However, there was uncertainty around the effect: Mean difference
27 (MD) 0.71g/kg/day (95% CI -0.17 to 1.60).

28 *PNALD/cholestasis*

29 **Direct bilirubin \geq 2mg/dl (equivalent to 34.2mmol/L)**

30 • Low quality evidence from 4 RCTs (n=328) showed a clinically important difference in the
31 rate of direct bilirubin \geq 2mg/dl in babies who received fish oil lipid emulsions compared
32 with non-fish oil lipid emulsions. Fewer babies receiving fish oil lipid emulsions had
33 PNALD/cholestasis. However, there was high uncertainty around the effect: Relative risk
34 (RR) 0.61 (95% CI 0.24 to 1.56).

1 Any definition

- 2 • Very low quality evidence from 11 RCTs (n=1154) showed a clinically important difference
3 in the rate of PNALD/cholestasis (using any definition) in babies who received fish oil lipid
4 emulsions compared with non-fish oil lipid emulsions. Fewer babies receiving fish oil lipid
5 emulsions had PNALD/cholestasis. However, there was uncertainty around the effect: RR
6 0.63 (95% CI 0.43 to 0.91).

7 Death before discharge

- 8 • Low quality evidence from 9 RCTs (n=855) showed no clinically important difference in
9 rate of death before discharge in babies who received MOFS-LE compared with S-LE.
10 However, there was uncertainty around the effect: RR 1.24 (95% CI 0.81 to 1.90).

11 Sepsis**12 Culture positive**

- 13 • Low quality evidence from 7 RCT (n=774) showed no clinically important difference in the
14 rate of cultures positive for sepsis in babies who received fish oil lipid emulsions
15 compared with non-fish oil lipid emulsions. However, there was uncertainty around the
16 effect: RR 1.16 (95% CI 0.91 to 1.48).

**17 Fish oil lipid emulsion (MOFS-LE) compared with another fish oil lipid emulsion (MFS-LE)
18 for preterm babies****19 Weight gain****20 Rate of weight gain**

- 21 • Low quality evidence from 1 RCT (n=55) showed no clinically important difference in rate
22 of weight gain in babies who received MOFS-LE compared with MFS-LE. However, there
23 was uncertainty around the effect: MD 4g/kg/day (95% CI -2.03 to 10.03).

24 PNALD/cholestasis

- 25 • Low quality evidence from 1 RCT (n=55) showed no clinically important difference in the
26 rate of direct bilirubin \geq 2mg/dl in babies who received MOFS-LE compared with MFS-LE.
27 However, there was high uncertainty around the effect: RR 0.96 (95% CI 0.06 to 14.65).

28 Death before discharge

- 29 • Low quality evidence from 1 RCT (n=60) showed no clinically important difference in the
30 rate of death before discharge in babies who received MOFS-LE compared with MFS-LE.
31 However, there was high uncertainty around the effect: RR 1.00 (95% CI 0.15 to 6.64).

32 Sepsis

- 33 • Low quality evidence from 1 RCT (n=55) showed a clinically important difference in the
34 rate of sepsis in babies who received MOFS-LE compared with MFS-LE. More babies
35 receiving MOFS-LE had sepsis. However, there was high uncertainty around the effect:
36 RR 1.69 (95% CI 0.56 to 5.11).

1 **Alternative lipid emulsion compared with Soybean based emulsion (S-LE) for preterm**
2 **babies**

3 ***Weight gain***

4 **Rate of weight gain**

- 5 • Low quality evidence from 1 RCT (n=60) showed no clinically important difference in rate
6 of weight gain in babies who received MS-LE compared with S-LE. However, there was
7 uncertainty around the effect: MD -2.67g/kg/day (95% CI -8.20 to 2.86).
- 8 • Low quality evidence from 2 RCTs (n=123) showed no clinically important difference in
9 rate of weight gain in babies who received OS-LE compared with S-LE. However, there
10 was uncertainty around the effect: MD -0.42 (95% CI -5.15 to 4.30).

11 ***Death before discharge***

- 12 • Low quality evidence from 3 RCTs (n=224) showed no clinically important difference in
13 the rate of death before discharge in babies who received OS-LE compared with S-LE.
14 However, there was high uncertainty around the effect: RR 1.00 (95% CI 0.21 to 4.82).

15 ***Sepsis***

16 **Culture positive**

- 17 • Low quality evidence from 2 RCTs (n=164) showed no clinically important difference in
18 the rate of cultures positive for sepsis in babies who received OS-LE compared with S-LE.
19 However, there was high uncertainty around the effect: RR 1.22 (95% CI 0.54 to 2.78).

20 **Alternative lipid emulsion compared with another alternative lipid emulsion for preterm**
21 **babies**

22 ***Weight gain***

23 **Rate of weight gain**

- 24 • Low quality evidence from 1 RCT (n=59) showed no clinically important difference in rate
25 of weight gain in babies who received MS-LE compared with OS-LE. However, there was
26 uncertainty around the effect: MD -1.33g/kg/day (95% CI -7.36 to 4.70).

27 ***PNALD/cholestasis***

- 28 • Low quality evidence from 1 RCT (n=59) showed a clinically important difference in the
29 rate of direct bilirubin \geq 2mg/dl in babies who received MS-LE compared with OS-LE.
30 More babies receiving MS-LE had PNALD/cholestasis. However, there was high
31 uncertainty around the effect: RR 2.90 (95% CI 0.12 to 68.5).

32 ***Sepsis***

- 33 • Low quality evidence from 1 RCT (n=59) showed a clinically important difference in the
34 rate of sepsis in babies who received MS-LE compared with OS-LE. More babies
35 receiving MS-LE had sepsis. However, there was high uncertainty around the effect: RR
36 1.93 (95% CI 0.65 to 5.73).

37 **Fish oil lipid emulsion compared with non-fish oil lipid emulsion for preterm babies with**
38 **surgical conditions**

39 ***PNALD/cholestasis***

- 40 • Very low quality evidence from 1 RCT (n=19) showed no clinically important difference in
41 the rate of direct bilirubin \geq 2mg/dl in babies who received pure fish oil compared with S-

1 LE. However, there was high uncertainty around the effect: RR 1.11 (95% CI 0.08 to
2 15.28).

3 **Sepsis**

4 • Very low quality evidence from 1 RCT (n=19) showed no clinically important difference in
5 the rate of cultures positive for sepsis in babies who received pure fish oil compared with
6 S-LE. However, there was high uncertainty around the effect: RR 1.11 (95% CI 0.39 to
7 3.19).

8 **Fish oil lipid emulsion compared with non-fish oil lipid emulsion for preterm babies with** 9 **cholestasis**

10 **Weight gain**

11 • Very low quality evidence from 1 RCT (n=16) showed a clinically important difference in
12 rate of weight gain in babies who received pure fish oil compared with S-LE. Weight gain
13 was greater in babies who received pure fish oil. However, there was uncertainty around
14 the effect: MD 45g/week (95% CI 15 to 75).

15 **Head growth**

16 • Very low quality evidence from 1 RCT (n=16) showed a clinically important difference in
17 rate of head growth in babies who received pure fish oil compared with S-LE. Head
18 growth was greater in babies who received pure fish oil. However, there was uncertainty
19 around the effect: MD 0.16cm/week (95% CI -0.01 to 0.33).

20 **PNALD/cholestasis**

21 • Very low quality evidence from 2 RCTs (n=40) showed a clinically important difference in
22 the rate of PNALD/cholestasis (using any definition) in babies who received fish oil lipid
23 emulsions compared with non-fish oil lipid emulsions. Fewer babies receiving fish oil had
24 PNALD/cholestasis. However, there was uncertainty around the effect: RR 0.54 (95% CI
25 0.32 to 0.91).

26 • Very low quality evidence from 1 RCT (n=16) showed a clinically important difference in
27 the rate of resolution of PNALD/cholestasis (direct bilirubin < 2 mg/dl) in babies who
28 received pure fish oil compared with Intralipid. More babies receiving pure-fish oil had
29 resolution of PNALD/cholestasis. However, there was high uncertainty around the effect:
30 RR 5.60 (95% CI 0.34 to 93.35).

31 **Death before discharge**

32 • Very low quality evidence from 2 RCTs (n=40) showed a clinically important difference in
33 the rate of death before discharge in babies who received fish oil lipid emulsions
34 compared with non-fish oil lipid emulsions. Fewer babies receiving fish oil died before
35 discharge. However, there was high uncertainty around the effect: RR 0.24 (95% CI 0.03
36 to 1.87).

37 **Sepsis**

38 • Very low quality evidence from 2 RCTs (n=40) showed no clinically important difference in
39 the rate of sepsis in babies who received fish oil lipid emulsions compared with non-fish oil
40 lipid emulsions. However, there was high uncertainty around the effect: RR 1.21 (95% CI
41 0.50 to 2.92).

1 **Fish oil lipid emulsion compared with non-fish oil lipid emulsion for term and late**
2 **preterm babies with surgical conditions**

3 ***PNALD/cholestasis***

- 4 • Very low quality evidence from 1 RCT (n=19) showed no clinically important difference in
5 the rate of direct bilirubin \geq 2mg/dl in babies who received pure fish oil compared with S-
6 LE. However, there was high uncertainty around the effect: RR 1.11 (95% CI 0.08 to
7 15.28).
- 8 • Low quality evidence from 2 RCTs (n=68) showed no clinically important difference in the
9 rate of PNALD/cholestasis (using any definition) in babies who received fish oil lipid
10 emulsions compared with non-fish oil lipid emulsions. However, there was high
11 uncertainty around the effect: RR 1.20 (95% CI 0.38 to 3.76).

12 ***Sepsis***

13 ***Culture positive***

- 14 • Very low quality evidence from 2 RCTs (n=51) showed no clinically important difference in
15 the rate of cultures positive for sepsis in babies who received fish oil lipid emulsions
16 compared with non-fish oil lipid emulsions. However, there was high uncertainty around
17 the effect: RR 1.05 (95% CI 0.47 to 2.34).

18 **Fish oil lipid emulsion compared with non-fish oil lipid emulsion for term and late**
19 **preterm babies with cholestasis**

20 ***Weight gain***

- 21 • Very low quality evidence from 1 RCT (n=16) showed a clinically important difference in
22 rate of weight gain in babies who received pure fish oil compared with S-LE. Weight gain
23 was greater in babies who received pure fish oil. However, there was uncertainty around
24 the effect: MD 45g/week (95% CI 15 to 75).

25 ***PNALD/cholestasis***

- 26 • Very low quality evidence from 2 RCTs (n=40) showed a clinically important difference in
27 the rate of PNALD/cholestasis (using any definition) in babies who received fish oil lipid
28 emulsions compared with non-fish oil lipid emulsions. Fewer babies receiving fish oil had
29 PNALD/cholestasis. However, there was uncertainty around the effect: RR 0.54 (95% CI
30 0.32 to 0.91).
- 31 • Very low quality evidence from 1 RCT (n=16) showed a clinically important difference in
32 the rate of resolution of PNALD/cholestasis (direct bilirubin $<$ 2 mg/dl) in babies who
33 received pure fish oil compared with Intralipid. More babies receiving pure-fish oil had
34 resolution of PNALD/cholestasis. However, there was high uncertainty around the effect:
35 RR 5.60 (95% CI 0.34 to 93.35).

36 ***Death before discharge***

- 37 • Very low quality evidence from 2 RCTs (n=40) showed a clinically important difference in
38 the rate of death before discharge in babies who received fish oil lipid emulsions
39 compared with non-fish oil lipid emulsions. Fewer babies receiving fish oil died before
40 discharge. However, there was high uncertainty around the effect: RR 0.24 (95% CI 0.03
41 to 1.87).

42 ***Sepsis***

- 43 • Very low quality evidence from 2 RCTs (n=40) showed no clinically important difference in
44 the rate of sepsis in babies who received fish oil lipid emulsions compared with non-fish oil
45 lipid emulsions. However, there was high uncertainty around the effect: RR 1.21 (95% CI
46 0.50 to 2.92).

1 **Hypertriglyceridemia**

- 2 • Very low quality evidence from 1 RCT (n=24) showed a clinically important difference in
3 the rate of hypertriglyceridemia in babies who received MOFS-LE compared with S-LE.
4 Fewer babies receiving MOFS-LE had hypertriglyceridemia. However, there was high
5 uncertainty around the effect: RR 0.79 (95% CI 0.30 to 2.09).

6 **Economic evidence statements**

- 7 No economic evidence was identified which was applicable to this review question.

8 **The committee's discussion of the evidence**

9 **Interpreting the evidence**

10 ***The outcomes that matter most***

11 Lipids are an essential part of PN; therefore, growth and neurodevelopmental outcomes were
12 prioritised as critical outcomes by the committee. PN associated liver disease (PNALD) was
13 also selected as a critical outcome as different lipid emulsions contain varying amounts of
14 polyunsaturated fatty acid and phytosterols which may contribute to PNALD.

15 Other adverse effects of intravenous lipid emulsions (infection including sepsis,
16 hyperglycaemia, hypertriglyceridemia and hyperlipidaemia) were selected as important
17 outcomes as they may vary according to the type of lipid emulsion used. Duration of hospital
18 stay and mortality were selected as important outcomes as they may be affected by both the
19 type of lipid emulsion used and the overall clinical condition of the baby. Nutritional intake
20 was also selected as an important outcome as nutritional composition varies across different
21 lipid emulsions.

22 ***The quality of the evidence***

23 The quality of the Cochrane systematic reviews was assessed using the ROBIS tool. Both
24 reviews were rated as having a low risk of bias. The Cochrane authors performed a GRADE
25 analysis of the outcomes. The evidence was all very low and low quality and was
26 downgraded due to risk of bias in included studies, small sample sizes, small number of
27 events, and uncertainty around effects.

28 There was no evidence for neurodevelopmental outcomes, hyperglycaemia, hyperlipidaemia,
29 duration of hospital stay or nutritional intake.

30 There was no evidence for late preterm or term babies that did not have pre-existing PNALD
31 or surgical conditions.

32 ***Benefits and harms***

33 The evidence on preterm and term babies did not show any clear clinical difference in the
34 range of outcomes when comparing the different lipid emulsions; therefore the committee
35 decided by informal consensus not to make a recommendation on any specific lipid emulsion
36 to administer. However, there was limited evidence of greater weight gain and head growth,
37 greater resolution of PNALD and less mortality and hypertriglyceridemia when babies with
38 PNALD were given fish oil containing lipid emulsions compared with lipid emulsions that did
39 not contain fish oil. The evidence was from preterm and late preterm babies; however, the
40 committee agreed that late preterm and term babies are often treated the same in clinical
41 practice and that term babies with PNALD would be likely to also benefit from fish oil
42 containing lipid emulsions. Therefore, the committee recommended, based on informal
43 consensus, that therapeutic use of fish oil containing lipid emulsions are considered for
44 preterm and term babies with PNALD.

1 For babies with surgical conditions, there was no evidence of advantage of any specific lipid
2 formulation. Therefore, the committee agreed they could not make a recommendation for this
3 population. They discussed that for babies with surgical conditions who are likely to be on
4 long term PN, such as those with little or no remaining bowel, the duration of parental
5 nutrition will be much longer than is covered by the scope of this guideline and these babies
6 may require management by a multidisciplinary team, including a gastroenterologist.

7 There was some evidence of a benefit of fish oil containing lipid emulsions at reducing the
8 rate of PNALD in preterm babies without surgical conditions or pre-existing PNALD. The
9 committee discussed whether or not fish oil containing lipid emulsions should be considered
10 in these babies and could not reach agreement as it was felt that there was no conclusive
11 evidence of benefit, particularly as benefits of fish oil containing lipid emulsions were not
12 seen for outcomes beyond PNALD. There was also no evidence comparing different lipid
13 emulsions in late preterm or term babies without surgical conditions or pre-existing PNALD.
14 The committee discussed that it might be more beneficial to use fish-oil containing lipid
15 emulsions in babies that are likely to be on PN for a longer duration, as there will be a
16 greater risk of developing PNALD; however, the committee did not think there was sufficient
17 evidence to support this as a recommendation. Further, in the absence of surgical conditions,
18 it might not always be possible to know how long babies are likely to need PN.

19 **Cost effectiveness and resource use**

20 No economic studies were identified which were applicable to this review question.

21 The committee note that in preterm and term babies with PNALD the use of fish oil
22 containing lipid emulsions may result in avoiding the costs associated with poorer growth,
23 worsening of PNALD and increased risk of hypertriglyceridemia. These are severe long-
24 lasting complications which result in substantial costs to the NHS and also have a
25 detrimental impact on the health-related quality of life of babies and also their parents or
26 carers. Combined with an increased risk of mortality the use of lipid emulsions that do not
27 contain fish oil would result in substantial losses in quality adjusted life years (QALYs). The
28 committee explained that the use of fish oil containing lipid emulsions will have negligible, if
29 any, impact on the unit cost of PN. As a result, the committee was of a view that the use of
30 fish oil containing lipid emulsions would represent a cost effective use of NHS resources in
31 pre-term and term babies with PNALD.

32 **References**

33 **Kapoor 2018a**

34 Kapoor, V., Malviya, M. N., Soll, R., Lipid emulsions for parenterally fed preterm infants.
35 Cochrane Database of Systematic Reviews 2018, Issue 11

36 **Kapoor 2018b**

37 Kapoor, V., Malviya, M. N., Soll, R., Lipid emulsions for parenterally-fed term and late
38 preterm infants. Cochrane Database of Systematic Reviews 2018, Issue 11

1 Appendices

2 Appendix A – Review protocols

3 Review protocol for review question: What is the clinical effectiveness, efficacy and safety of lipid formulations from 4 different sources (for example, soya, fish oil, or mixed sources)?

5 The protocol drafted by the committee is displayed below and formed the basis of discussion with Cochrane Neonatal. The two Cochrane
6 protocols were then adapted in line with this protocol.

7 See Cochrane reviews for the review protocols:

8 Cochrane protocol for preterm babies: <https://www.cochranelibrary.com/cdsr/doi/10.1002/14651858.CD013163/full>

9 Cochrane protocol for term and late- preterm babies: <https://www.cochranelibrary.com/cdsr/doi/10.1002/14651858.CD013171/full>

10 Table 3: Original review protocol presented to the committee and discussed with Cochrane to develop the Cochrane protocols–

Field (based on PRISMA-P)	Content
Review question	What is the clinical effectiveness, efficacy and safety of lipid formulations from different sources (for example, soya, fish oil, or mixed sources)?
Type of review question	Intervention
Objective of the review	Inadequate amounts of calcium and phosphate delivered via PN may contribute to bone disease in preterm and term babies. Delivery of calcium and phosphate should be adequate to achieve retention of amounts which match those in utero, but at a concentration that does not result in adverse events. The aim of this review is to determine the optimal ratio of phosphate to amino acids in preterm and term babies who are receiving PN
Eligibility criteria – population/disease/condition/issue/domain	<ul style="list-style-type: none"> • Babies born preterm, up to 28 days after their due birth date (preterm babies) • Babies born at term, up to 28 days after their birth (term babies).
Eligibility criteria – intervention(s)/exposure(s)/prognostic factor(s)	Any concentrations of: <ul style="list-style-type: none"> • Soy based (Intralipid) • Specific Multicomponent (Soy, MCT, olive oil, fish oil) SMOFlipid • Olive oil +soy oil (clinoleic) • Fish oil (omegaven)
Eligibility criteria – comparator(s)/control or reference (gold) standard	<ul style="list-style-type: none"> • Each other

Field (based on <u>PRISMA-P</u>)	Content
Outcomes and prioritisation	<p>Critical</p> <ul style="list-style-type: none"> • Growth/Anthropometric measures: <ul style="list-style-type: none"> ○ Weight gain (g/kg/d) ○ Linear growth ○ Head circumference (mm) • Neurodevelopmental outcomes • Adverse events of lipids (PN related liver disease, abnormal liver function, cholestasis, conjugated hyperbilirubinaemia, intrahepatocellular lipid) <p>Important</p> <ul style="list-style-type: none"> • Mortality • Adverse events (including sepsis, hyperglycaemia, hypertriglyceridemia, hyperlipidaemia) • Duration of hospital stay • Nutritional intake (g/kg/day – proportion of lipid received or essential fatty acids at 2 days)
Eligibility criteria – study design	<p>Only published full text papers:</p> <p>Systematic reviews of RCTs RCTs Comparative cohort studies (only if RCTs unavailable or limited data to inform decision making)</p> <p>Conference abstracts will only be considered if related to RCTs</p>
Other inclusion exclusion criteria	<p>No sample size restriction No date restriction</p>
Proposed sensitivity/sub-group analysis, or meta-regression	<p>Subgroup analysis:</p> <p>Population subgroups:</p> <p>Age of baby (first 2 weeks vs later) Preterm (extremely preterm <28 weeks' GA; very preterm: 28-31 weeks' GA; moderately preterm: 32-36 weeks' GA) Birthweight: Low birth weight (< 2500g); very low birth weight (< 1500g) and extremely low birth weight (< 1000g) Critically ill babies or those requiring surgery (for example, inotropic support, therapeutic hypothermia or fluid restriction)</p>

Field (based on <u>PRISMA-P</u>)	Content
	First week of life and after first week of life?
Selection process – duplicate screening/selection/analysis	Sifting, data extraction, appraisal of methodological quality and GRADE assessment will be performed by the systematic reviewer. Quality control will be performed by the senior systematic reviewer. A random sample of the references identified in the search will be sifted by a second reviewer. This sample size will be 10% of the total, or 100 studies if the search identifies fewer than 1000 studies. All disagreements in study inclusion will be discussed and resolved between the two reviewers. The senior systematic reviewer or guideline lead will be involved if discrepancies cannot be resolved between the two reviewers.
Data management (software)	Pairwise meta-analyses, if possible, will be performed using Cochrane Review Manager (RevMan5). 'GRADEpro' will be used to assess the quality of evidence for each outcome. NGA STAR software will be used for generating bibliographies/citations, study sifting, data extraction and recording quality assessment using checklists (ROBIS (systematic reviews and meta-analyses); Cochrane risk of bias tool (RCTs or comparative cohort studies); Cochrane risk of bias tool (Non-randomised studies); Newcastle-Ottawa scale (Non-comparative studies)).
Information sources – databases and dates	Sources to be searched: Medline, Medline In-Process, CCTR, CDSR, DARE, HTA, Embase. Limits (e.g. date, study design): All study designs. Apply standard animal/non-English language filters. No date limit. Supplementary search techniques: No supplementary search techniques were used. See appendix B for full strategies.
Identify if an update	This is not an update
Author contacts	Developer: The National Guideline Alliance Guideline website: https://www.nice.org.uk/guidance/indevelopment/gid-ng10037 .
Highlight if amendment to previous protocol	For details please see section 4.5 of Developing NICE guidelines: the manual 2014.
Search strategy – for one database	For details please see appendix B.
Data collection process – forms/duplicate	A standardised evidence table format will be used, and published as appendix D (clinical evidence tables) or H (economic evidence tables).
Data items – define all variables to be collected	For details please see appendix B.
Methods for assessing bias at outcome/study level	Standard study checklists were used to critically appraise individual studies. For details please see section 6.2 of Developing NICE guidelines: the manual 2014. The risk of bias across all available evidence was evaluated for each outcome using an adaptation of the 'Grading of Recommendations Assessment, Development and Evaluation (GRADE) toolbox' developed by the international GRADE working group http://www.gradeworkinggroup.org/

Field (based on <u>PRISMA-P</u>)	Content
Criteria for quantitative synthesis (where suitable)	For details please see section 6.4 of Developing NICE guidelines: the manual 2014.
Methods for analysis – combining studies and exploring (in)consistency	For details of the methods please see supplementary material C.
Meta-bias assessment – publication bias, selective reporting bias	For details please see section 6.2 of Developing NICE guidelines: the manual 2014. If sufficient relevant RCT evidence is available, publication bias will be explored using RevMan software to examine funnel plots. Trial registries will be examined to identify missing evidence: Clinical trials.gov, NIHR Clinical Trials Gateway.
Assessment of confidence in cumulative evidence	For details please see sections 6.4 and 9.1 of Developing NICE guidelines: the manual 2014.
Rationale/context – Current management	For details please see the introduction to the evidence review.
Describe contributions of authors and guarantor	A multidisciplinary committee developed the guideline. The committee was convened by The National Guideline Alliance and chaired by Joe Fawke (Consultant Neonatologist and Honorary Senior Lecturer, University Hospitals Leicester NHS Trust), in line with section 3 of Developing NICE guidelines: the manual 2014. Staff from The National Guideline Alliance, undertook systematic literature searches, appraised the evidence, conducted meta-analysis and cost-effectiveness analysis where appropriate, and drafted the guideline in collaboration with the committee. For details of the methods please see supplementary material C.
Sources of funding/support	The National Guideline Alliance is funded by NICE and hosted by The Royal College of Obstetricians and Gynaecologists.
Name of sponsor	The National Guideline Alliance is funded by NICE and hosted by The Royal College of Obstetricians and Gynaecologists.
Roles of sponsor	NICE funds the National Guideline Alliance to develop guidelines for those working in the NHS, public health, and social care in England.
PROSPERO registration number	This review is not registered with PROSPERO.

- 1 CDSR: Cochrane Database of Systematic Reviews; CCTR: Cochrane Controlled Trials Register; DARE: Database of Abstracts of Reviews of Effects; GA: gestational age;
2 GRADE: Grading of Recommendations Assessment, Development and Evaluation; HTA: Health Technology Assessment; ICF: International Classification of Functioning,
3 Disability and Health; MID: minimally important difference; NGA: National Guideline Alliance; NIHR: National Institute for Health Research; NHS: National health service; NICE:
4 National Institute for Health and Care Excellence; PRISMA-P: preferred reporting items for systematic review and meta-analysis protocols; RCT: randomised controlled trial;
5 RoB: risk of bias; ROBIS: risk of bias in systematic reviews; SD: standard deviation

1 Appendix B – Literature search strategies

2 Literature search strategy for review question: What is the clinical effectiveness,
3 efficacy and safety of lipid formulations from different sources (for example,
4 soya, fish oil, or mixed sources)?

5 See the Cochrane reviews for the literature search strategy:

6 [https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-](https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-preterm-infants)
7 [intravenous-nutrition-preterm-infants](https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-preterm-infants)

8 [https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-](https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-term-and-late-preterm-infants)
9 [intravenous-nutrition-term-and-late-preterm-infants](https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-term-and-late-preterm-infants)

1 **Appendix C – Clinical evidence study selection**

2 **Clinical study selection for review question: What is the clinical effectiveness,**
3 **efficacy and safety of lipid formulations from different sources (for example,**
4 **soya, fish oil, or mixed sources)?**

5 See the Cochrane reviews for the study selection flow charts:

6 [https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-](https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-preterm-infants)
7 [intravenous-nutrition-preterm-infants](https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-preterm-infants)

8 [https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-](https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-term-and-late-preterm-infants)
9 [intravenous-nutrition-term-and-late-preterm-infants](https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-term-and-late-preterm-infants)

1 Appendix D – Clinical evidence tables

2 Clinical evidence tables for review question: What is the clinical effectiveness, efficacy and safety of lipid formulations from different sources (for example, soya, fish oil, or mixed sources)?

4 Table 4: Clinical evidence tables for included studies

Study details	Participants	Interventions	Methods	Outcomes and Results	Comments
<p>Full citation Kapoor, V., Malviya, M. N., Soll, R., Lipid emulsions for parenterally fed preterm infants. Cochrane Database of Systematic Reviews 2018a, Issue 11</p> <p>Ref Id N/A</p> <p>Country/ies where the study was carried out Australia, Oman, USA</p> <p>Study type</p>	<p>Preterm babies born before 37 weeks' gestation</p> <p>See Cochrane review for full details.</p>	<ul style="list-style-type: none"> Fish oil lipid emulsion versus non-fish oil lipid emulsion Fish oil lipid emulsion versus another fish oil lipid emulsion Alternative lipid emulsion versus soybean oil-based lipid emulsion Alternative lipid emulsion versus another alternative lipid emulsion 	<p>See Cochrane review for details.</p> <p>https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-preterm-infants</p>	<p>See Cochrane review for details.</p> <p>https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-preterm-infants</p>	<p>Limitations</p> <p>Methodological quality was assessed using the ROBIS tool: <u>Study eligibility criteria - Low concern</u></p> <p>Did the review adhere to pre-defined objectives and eligibility criteria? Yes</p> <p>Were the eligibility criteria appropriate for the review question? Yes</p> <p>Were eligibility criteria unambiguous? Yes</p> <p>Were all restrictions in eligibility criteria based on study characteristics appropriate? Yes</p> <p>Were any restrictions in eligibility criteria based on sources of information appropriate? Yes</p> <p><u>Identification and selection of studies - Low concern</u></p> <p>Did the search include an appropriate range of databases/electronic sources for published and unpublished reports? Yes</p> <p>Were methods additional to database searching used to identify relevant reports? Yes</p> <p>Were the terms and structure of the search strategy likely to retrieve as many eligible studies as possible? Yes</p> <p>Were restrictions based on date, publication format, or language appropriate? Yes</p> <p>Were efforts made to minimise errors in selection of studies? Yes</p>

Study details	Participants	Interventions	Methods	Outcomes and Results	Comments
<p>Systematic review of randomised or quasi-randomised controlled studies</p> <p>Aim of the study To compare the safety and effectiveness of different lipid emulsions for parenteral nutrition in preterm infants.</p> <p>Study dates Articles published up to 18th July 2018</p> <p>Source of funding NIHR Cochrane Programme Grant (16/114/03); Vermont Oxford Network;</p>		<p>See Cochrane review for full details.</p> <p>https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-preterm-infants</p>			<p><u>Data collection and study appraisal - Low concern</u></p> <p>Were efforts made to minimise error in data collection? Yes</p> <p>Were sufficient study characteristics available for both review authors and readers to be able to interpret the results? Yes</p> <p>Were all relevant study results collected for use in the synthesis? Yes</p> <p>Was risk of bias (or methodological quality) formally assessed using appropriate criteria? Yes</p> <p>Were efforts made to minimise errors in risk or bias assessment? Yes</p> <p><u>Synthesis and findings - High concern</u></p> <p>Did the syntheses include all studies that it should? Unclear – The majority of included studies were rated as unclear risk of selection bias as protocols were not available. Therefore, it is unclear if all results were available to the reviewers.</p> <p>Were all predefined analyses followed or departures explained? Yes</p> <p>Was the synthesis appropriate given the nature and similarity in the research questions, study designs and outcomes across included studies? Yes</p> <p>Was between-studies variation (heterogeneity) minimal or addressed in the synthesis? Yes</p> <p>Were the findings robust, e.g., as demonstrated through funnel plot or sensitivity analyses? Probably No – Funnel plots were used where possible to examine publication bias, which was not found. However, for one outcome exclusion of a high risk study reduced the magnitude of the effect and introduced more uncertainty around the estimate. For a number of comparisons and outcomes, only single small studies were available.</p> <p>Were biases in primary studies minimal or addressed in the synthesis? Yes</p> <p><u>Overall risk of bias - Low risk of bias</u></p>

Study details	Participants	Interventions	Methods	Outcomes and Results	Comments
Cochrane Review Incentive Scheme, reference number 17/62/30					<p>Did the interpretation of findings address all of the concerns identified in the phase 2 assessment? Probably Yes – The authors discuss the paucity of large randomised trials and make tentative conclusions based on the results.</p> <p>Was the relevance of identified studies to the review's research question appropriately considered? Yes</p> <p>Did the reviewers avoid emphasising results on the basis of their statistical significance? Yes</p> <p>Other information</p> <p>Two of the 3 studies of cholestatic or surgical preterm babies were stopped early which may have introduced bias.</p>
<p>Full citation Kapoor, V., Malviya, M. N., Soll, R., Lipid emulsions for parenterally-fed term and late preterm infants. Cochrane Database of Systematic Reviews 2018b, Issue 11</p> <p>Ref Id N/A</p> <p>Country/ies where the study was carried out</p>	<p>Term babies born at 37 weeks' gestation or after; late preterm babies born between 34⁺⁰ and 36⁺⁶ weeks' gestation</p> <p>See Cochrane review for full details.</p>	<ul style="list-style-type: none"> Fish oil lipid emulsion versus non-fish oil lipid emulsion <p>See Cochrane review for full details.</p> <p>https://www.cochrane.org/CD013171/NEONATA_systematic-review-lipid-emulsions-intravenous-nutrition-term-and-late-preterm-infants</p>	<p>See Cochrane review for details. https://www.cochrane.org/CD013171/NEONATA_systematic-review-lipid-emulsions-intravenous-nutrition-term-and-late-preterm-infants</p>	<p>See Cochrane review for details.</p> <p>https://www.cochrane.org/CD013171/NEONATA_systematic-review-lipid-emulsions-intravenous-nutrition-term-and-late-preterm-infants</p>	<p>Limitations</p> <p>Methodological quality was assessed using the ROBIS tool: <u>Study eligibility criteria - Low concern</u></p> <p>Did the review adhere to pre-defined objectives and eligibility criteria? Yes</p> <p>Were the eligibility criteria appropriate for the review question? Yes</p> <p>Were eligibility criteria unambiguous? Yes</p> <p>Were all restrictions in eligibility criteria based on study characteristics appropriate? Yes</p> <p>Were any restrictions in eligibility criteria based on sources of information appropriate? Yes</p> <p><u>Identification and selection of studies - Low concern</u></p> <p>Did the search include an appropriate range of databases/electronic sources for published and unpublished reports? Yes</p> <p>Were methods additional to database searching used to identify relevant reports? Yes</p> <p>Were the terms and structure of the search strategy likely to retrieve as many eligible studies as possible? Yes</p>

Study details	Participants	Interventions	Methods	Outcomes and Results	Comments
<p>Australia, Oman, USA</p> <p>Study type Systematic review of randomised or quasi-randomised controlled studies</p> <p>Aim of the study To compare the safety and effectiveness of different lipid emulsions for parenteral nutrition in term and late preterm infants.</p> <p>Study dates Articles published up to 18th June 2018</p> <p>Source of funding NIHR Cochrane Programme</p>					<p>Were restrictions based on date, publication format, or language appropriate? Yes</p> <p>Were efforts made to minimise errors in selection of studies? Probably Yes – It is unclear whether both title and abstracts and full texts were independently assessed but authors report that two authors independently search the database. More information is provided in Kapoor 2018a and it is likely the same methods were used for this review.</p> <p><u>Data collection and study appraisal - Low concern</u></p> <p>Were efforts made to minimise error in data collection? Yes</p> <p>Were sufficient study characteristics available for both review authors and readers to be able to interpret the results? Yes</p> <p>Were all relevant study results collected for use in the synthesis? Yes</p> <p>Was risk of bias (or methodological quality) formally assessed using appropriate criteria? Yes</p> <p>Were efforts made to minimise errors in risk or bias assessment? Yes</p> <p><u>Synthesis and findings - High concern</u></p> <p>Did the syntheses include all studies that it should? Unclear – A number of included studies were rated as unclear risk of selection bias as protocols were not available. Therefore, it is unclear if all results were available to the reviewers.</p> <p>Were all predefined analyses followed or departures explained? Yes</p> <p>Was the synthesis appropriate given the nature and similarity in the research questions, study designs and outcomes across included studies? Yes</p> <p>Was between-studies variation (heterogeneity) minimal or addressed in the synthesis? Yes</p> <p>Were the findings robust, e.g., as demonstrated through funnel plot or sensitivity analyses? Probably No – Authors were unable to use funnel plots due to the number of studies for each outcome.</p>

Study details	Participants	Interventions	Methods	Outcomes and Results	Comments
Grant (16/114/03); Vermont Oxford Network; Cochrane Review Incentive Scheme, reference number 17/62/30					<p>For a number of comparisons and outcomes, only one or two small studies were available.</p> <p>Were biases in primary studies minimal or addressed in the synthesis? Yes</p> <p><u>Overall risk of bias - Low risk of bias</u></p> <p>Did the interpretation of findings address all of the concerns identified in the phase 2 assessment? Probably Yes – The authors discuss the paucity of large randomised trials and make tentative conclusions based on the results.</p> <p>Was the relevance of identified studies to the review's research question appropriately considered? Yes</p> <p>Did the reviewers avoid emphasising results on the basis of their statistical significance? Yes</p> <p>Other information</p> <p>Three of the 5 studies that contributed data to the review were stopped early which may have introduced bias.</p>

1 NIHR: National Institute of Health Research; ROBIS: risk of bias in systematic reviews; USA: United States of America.

1 Appendix E – Forest plots

2 **Forest plots for review question: What is the clinical effectiveness, efficacy and**
3 **safety of lipid formulations from different sources (for example, soya, fish oil,**
4 **or mixed sources)?**

5 See the Cochrane reviews for forest plots:

6 [https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-](https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-preterm-infants)
7 [intravenous-nutrition-preterm-infants](https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-preterm-infants)

8 [https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-](https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-term-and-late-preterm-infants)
9 [intravenous-nutrition-term-and-late-preterm-infants](https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-term-and-late-preterm-infants)

10 Appendix F – GRADE tables

11 **GRADE tables for review question: What is the clinical effectiveness, efficacy and**
12 **safety of lipid formulations from different sources (for example, soya, fish oil,**
13 **or mixed sources)?**

14 See the Cochrane reviews for GRADE tables:

15 [https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-](https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-preterm-infants)
16 [intravenous-nutrition-preterm-infants](https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-preterm-infants)

17 [https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-](https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-term-and-late-preterm-infants)
18 [intravenous-nutrition-term-and-late-preterm-infants](https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-term-and-late-preterm-infants)

1 **Appendix G – Economic evidence study selection**

2 **Economic evidence study selection for review question: What is the clinical** 3 **effectiveness, efficacy and safety of lipid formulations from different sources** 4 **(for example, soya, fish oil, or mixed sources)?**

5 One global search was conducted for all review questions. See supplementary material D for
6 further information.

1 **Appendix H – Economic evidence tables**

2 **Economic evidence study selection for review question: What is the clinical**
3 **effectiveness, efficacy and safety of lipid formulations from different sources**
4 **(for example, soya, fish oil, or mixed sources)?**

5 No economic studies were identified which were applicable to this review question.

6

1 **Appendix I – Economic evidence profiles**

- 2 **Economic evidence profiles for review question: What is the clinical effectiveness,**
- 3 **efficacy and safety of lipid formulations from different sources (for example, soya, fish**
- 4 **oil, or mixed sources)?**
- 5 No economic studies were identified which were applicable to this review question.

1 **Appendix J – Economic analysis**

2 **Economic analysis for review question: What is the clinical effectiveness, efficacy**
3 **and safety of lipid formulations from different sources (for example, soya, fish oil, or**
4 **mixed sources)?**

5 No economic analysis was undertaken for this review question.

6

1 **Appendix K – Excluded studies**

2 **Excluded studies for review question: What is the clinical effectiveness, efficacy**
3 **and safety of lipid formulations from different sources (for example, soya, fish**
4 **oil, or mixed sources)?**

5 **Clinical studies**

6 See the Cochrane reviews for excluded studies:

7 [https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-](https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-preterm-infants)
8 [intravenous-nutrition-preterm-infants](https://www.cochrane.org/CD013163/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-preterm-infants)

9 [https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-](https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-term-and-late-preterm-infants)
10 [intravenous-nutrition-term-and-late-preterm-infants](https://www.cochrane.org/CD013171/NEONATAL_systematic-review-lipid-emulsions-intravenous-nutrition-term-and-late-preterm-infants)

11 **Economic studies**

12 No economic evidence was identified for this review question. See supplementary material D
13 for further information.

14

1 **Appendix L – Research recommendations**

- 2 **Research recommendations for review question: What is the clinical**
- 3 **effectiveness, efficacy and safety of lipid formulations from different sources**
- 4 **(for example, soya, fish oil, or mixed sources)?**
- 5 No research recommendations were made for this review question.