National Institute for Health and Care Excellence

Draft for consultation

Neonatal parenteral nutrition

[G] Stopping parenteral nutrition

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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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Contents

Stopping parenteral nutrition in preterm and term babies	6
Review question	6
Introduction	6
Summary of the protocol	6
Clinical evidence	7
Summary of clinical studies included in the evidence review	7
Quality assessment of clinical outcomes included in the evidence review	9
Economic evidence	9
Summary of studies included in the economic evidence review	9
Economic model	10
Evidence statements	10
The committee's discussion of the evidence	14
References	16
Appendices	18
Appendix A – Review protocols	18
Review protocol for review question: What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?	
Appendix B – Literature search strategies	23
Literature search strategies for review question: What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?	
Appendix C – Clinical evidence study selection	
Clinical study selection for: What amount of enteral feed (measured in terms ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?	of
Appendix D – Clinical evidence tables	28
Clinical evidence tables for review question: What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer	20
required? Appendix E – Forest plots	
Appendix F – GRADE tables	
GRADE tables for review question: What amount of enteral feed (measured i	
terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?	
Appendix G – Economic evidence study selection	44
Economic evidence study selection for review question: What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?	44
Appendix H – Economic evidence tables	45

Econo	mic evidence tables for review question: What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?	. 45
Appendix I -	- Economic evidence profiles	. 46
Econo	mic evidence profiles for review question: What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?	. 46
Appendix J -	- Economic analysis	. 47
	mic evidence analysis for review question: What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?	
Appendix K	– Excluded studies	. 48
Exclud	led clinical and economic studies for review question: What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?	. 48
Clinica	ıl studies	. 48
Econo	mic studies	. 54
Appendix L -	- Research recommendations	. 55
Resea	rch recommendations for review question: What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?	. 55

Stopping parenteral nutrition in preterm and term babies

3 Review question

- What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation
- 5 to parenteral nutrition and enteral feeds) indicates that parenteral nutrition is no longer
- 6 required?

7 Introduction

- 8 Determining when parenteral nutrition (PN) can be stopped is important; there is risk of
- 9 malnutrition and associated complications to the baby if parenteral feeding is stopped too
- 10 early. In contrast, the longer PN is continued, the greater the risk of line sepsis and PN
- 11 associated liver disease. Guidance for clinicians on how to determine when it is safe to
- transition from combined parenteral and enteral nutrition to enteral nutrition (EN) alone is
- 13 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	 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	 Volume of PN (ml/kg/day or kcal/kg/day) plus EN (ml/kg/day or kcal/kg/day Ratio of PN (ml/kg/day or kcal/kg/day) and EN (ml/kg/day or kcal/kg/day)
Comparison	 Comparison volume of PN (ml/kg/day or kcal/kg/day) plus EN (ml/kg/day or kcal/kg/day)
	 Different ratio of PN (ml/kg/day or kcal/kg/day) to EN (ml/kg/day or kcal/kg/day)
Outcomes	Critical
	 Total nutritional intake, protein/carbohydrate/lipid/energy (for example, kcal/kg/day as a total of the parental + enteral)
	Growth/Anthropometric measures:
	o Weight gain
	Linear growth
	Head circumference
	 Body composition (lean mass, fat-free mass, fat mass, adipose tissue, nitrogen accretion)
	Important
	Mortality
	Duration of hospital stay
	Necrotising enterocolitis



- Neurodevelopmental outcomes (general cognitive abilities at two years, measured using validated scales only)
- Duration of PN
- Sepsis (central line infections)
- 1 EN: enteral nutrition; PICO: population, intervention, comparison and outcome; PN: parenteral nutrition
- 2 For further details see the review protocol in appendix A.

3 Clinical evidence

4 Included studies

- 5 As limited RCT evidence was available, we also included observational studies. Three
- 6 studies were identified for inclusion in this review (Choi 2016, Dinerstein 2006, and Perrem
- 7 2019).
- 8 One randomised controlled trial (RCT) (n=137) compared stopping PN when EN reached
- 9 100ml/kg/day to stopping PN when EN reached 140ml/kg/day (Perrem 2019).
- 10 One observational study (n=87) compared stopping PN when EN reached 100ml/kg/day to
- 11 stopping PN when EN reached 120ml/kg/day (Choi 2016).
- One observational study (n=182compared stopping PN when EN reached 60kcal/kg/day to
- 13 stopping PN when EN reached 100kcal/kg/day (Dinerstein 2006).
- 14 The included studies are summarised in Table 2.
- 15 See the literature search strategy in appendix B, study selection flow chart in appendix C,
- study evidence tables in appendix D, forest plots in appendix E, and GRADE tables in
- 17 appendix F.

18 Excluded studies

- 19 Studies not included in this review are listed, and reasons for their exclusions are provided,
- in appendix K.

21 Summary of clinical studies included in the evidence review

22 Summaries of the studies included in this review are presented in Table 2.

23 Table 2: Summary of included studies

Study	Population	Intervention	Comparison	Outcomes	Comments
Choi 2016	N=87*	100ml/kg/day (n=50)	120ml/kg/day (n=37)	 Weight gain Length <10th 	Multiple differences
Observational study	VLBW infants (<1500g) born before 34	Nutrition supplements	Nutrition supplements	centile at 36 and 40 weeks post conception	nutrition protocols apart from point at
Korea	weeks' gestation	added and PN discontinued at 100ml/kg/day of	added and PN discontinued at 120ml/kg/day of	 Head circumference <10th centile at 	which PN was discontinued.
	Mean GA: 29.4 weeks (SD 2.5)	EN	EN	36 and 40 weeks post	difficult to conclude if any
		Proteins started on day 1 at	Proteins started on day 1 at	conception	differences are a result of

Study	Population	Intervention	Comparison	Outcomes	Commonto
Study	Population Mean BW: 1126g (SD 235) *124 infants are included in the paper but 37 infants from period 3 were not included in this review question	Intervention 3.0g/kg/day and advanced to 4.0g/kg/day Lipids were started on day 1 at 1.0/kg/day and advanced to 3.5g/kg/day	Comparison 1.5g/kg/day and advanced to 3.5g/kg/day Lipids were started on day 2 at 0.5g/kg/day and advanced to 3.0g/kg/day	Outcomes Duration (days) of hospital stay Necrotising enterocolitis Duration (days) of PN Sepsis	when PN was stopped
Dinerstein 2006 Observational study Argentina	Infants weighing between 750g and 1500g at birth GA range 24 to 36 weeks Median BW 1230g for 60kcal/kg/day cohort and 1245g for 100kcal/kg/day cohort	60kcal/kg/day (n=117) PN was discontinued when infants reached 60kcal/kg/day of EN Fluids started at 80ml/kg/day and advanced to 150ml/kg/day over 7 to 10 days AA started on day 3 at 0.5g/kg/day and advanced to 3g/kg/day Glucose started (day NR) at 5.6mg/kg/minut e and advanced to 8 to 9mg/kg/minute Lipids started on day 3 or 4 at 0.5g/kg/day and advanced to 3g/kg/day	100kcal/kg/day (n=65) PN was discontinued when infants reached 100kcal/kg/day of EN Fluids started at 80ml/kg/day and advanced to 150 to 180ml/kg/day over 7 days AA started on day 1 at 1.5g/kg/day and advanced to 4g/kg/day Glucose started on day 1 at 5.6mg/kg/minute and advanced to 13mg/kg/minute and advanced to 13mg/kg/minute Lipids started at 24 hours at 0.5g/kg/day and advanced to 3.5g/kg/day	Weight gain Length (cm) at 40 weeks postmenstrual age Head circumference (cm) at 40 weeks postmenstrual age Necrotising enterocolitis Late onset sepsis	Multiple differences between nutrition protocols apart from point at which PN was discontinued. Therefore, difficult to conclude if any differences are a result of when PN was stopped
Perrem 2019 RCT Ireland	N=137 VLBW (<1500g), preterm infants with a peripherally inserted central venous catheter (PICC) and less than	100ml/kg/day (n=67) PICC was removed and PN discontinued when infants reached 100ml/kg/day of EN	140ml/kg/day (n=70) PICC was removed and PN discontinued when infants reached 140ml/kg/day of EN	 Weight gain Linear growth Head circumference Mortality Necrotising enterocolitis Central venous catheter- 	There were 12 protocol violations in both arms, where PICC was removed at a different time point than specified by the group allocation. Results of per-

Study Population	Intervention	Comparison	Outcomes	Comments
Mean GA: 28 weeks (SD 2. Median BW 1060g for 100mll/kg/day group and 1070g for 140ml/kg/day group	Fluid started at 60-80ml/kg/day 6 on day 1 using standard bag Standard bag content/100ml:	Fluid started at 60-80ml/kg/day on day 1 using standard bag standard bag content/100ml: amino acids 2.51g, glucose 10%, sodium 2.5mmol, potassium 2mmol, calcium 1mmol, magnesium 0.15mmol, phosphate 1.25mmol Fluids increased up to 150 to 160ml/kg/day	associated late onset sepsis	protocol analysis were not significantly different from results of intention-to- treat analysis for primary outcome.

- AA: amino acids; BW: Birth weight; EN: enteral nutrition; GA: gestational age; NR: not reported; PICC:
- 2 peripherally inserted central venous catheter; PN: Parenteral nutrition; RCT: Randomised controlled trail; SD:
- 3 standard deviation; VLBW: Very low birth weight
- 4 See the full evidence tables in appendix D and the forest plots in appendix E.

5 Quality assessment of clinical outcomes included in the evidence review

- 6 GRADE was conducted to assess the quality of outcomes. Evidence was identified for critical
- 7 and important outcomes. The clinical evidence profiles can be found in appendix F.

8 Economic evidence

9 Included studies

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

14 Excluded studies

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

16 Summary of studies included in the economic evidence review

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

1 Economic model

- 2 This question was a medium priority for economic evaluation. However, the identified clinical
- data was very limited and insufficient to inform useful economic modelling in this area.

4 Evidence statements

5 Clinical evidence statements

6 Stopping PN when EN reached 100ml/kg/day versus stopping PN when EN reached 7 140ml/kg/day

8 Weight gain

- Moderate quality evidence from 1 RCT (n=137) showed no clinically important difference in time to regain birth weight in VLBW and ELBW babies who had PN stopped at 100ml/kg/day of EN compared with 140ml/kg/day of EN. However, there was uncertainty around the effect: Mean difference (MD) 1.50 days (95% CI 0.31 to 2.69).
- Moderate quality evidence from 1 RCT (n=83) showed no clinically important difference in time to regain birth weight in VLBW babies who had PN stopped at 100ml/kg/day of EN compared with 140ml/kg/day of EN. However, there was uncertainty around the effect:
 MD 0.60 days (95% CI -0.80 to 2.00).
- Moderate quality evidence from 1 RCT (n=54) showed a clinically important difference in time to regain birth weight in ELBW babies who had PN stopped at 100ml/kg/day of EN compared with 140ml/kg/day of EN with longer time to regain birthweight associated with stopping at the lower EN. However, there was uncertainty around the effect: MD 2.80 days (95% CI 0.80 to 4.80).
- High quality evidence from 1 RCT (n=137) showed no clinically important difference in weight at discharge in babies who had PN stopped at 100ml/kg/day of EN compared with 140ml/kg/day of EN, MD 13.00g (95% CI -191.05 to 217.05).
- Low quality evidence from 1 RCT (n=137) showed no clinically important difference in the number of babies whose weight was less than the 10th centile at discharge between those who had PN stopped at 100ml/kg/day of EN compared with 140ml/kg/day of EN. However, there was high uncertainty around the effect: Relative risk (RR) 1.09 (95% CI 0.70 to 1.68).

Linear growth

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- High quality evidence from 1 RCT (n=137) showed no clinically important difference in length at discharge in babies who had PN stopped at 100ml/kg/day of EN compared with 140ml/kg/day of EN, MD 0.90cm (95% CI -0.54 to 2.34).
- Low quality evidence from 1 RCT (n=137) showed no clinically important difference in the number of babies whose length was less than the 10th centile at discharge between those who had PN stopped at 100ml/kg/day of EN compared with 140ml/kg/day of EN.
 However, there was high uncertainty around the effect: RR 1.07 (95% CI 0.78 to 1.48).

38 **Head circumference**

High quality evidence from 1 RCT (n=137) showed no clinically important difference in head circumference at discharge in babies who had PN stopped at 100ml/kg/day of EN compared with 140ml/kg/day of EN, MD 0.10cm (95% CI -0.87 to 0.67).

1 • Low quality evidence from 1 RCT (n=137) showed a clinically important difference in the 2 number of babies whose head circumference was less than the 10th centile at discharge 3 between those who had PN stopped at 100ml/kg/day of EN compared with 140ml/kg/day of EN with more babies below the 10th centile in the group of babies stopped at the lower 4 5 EN. However, there was high uncertainty around the effect: RR 1.27 (95% CI 0.68 to 6 2.37).

Mortality

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 Low quality evidence from 1 RCT (n=137) showed no clinically important difference in mortality in babies who had PN stopped at 100ml/kg/day of EN compared with 140ml/kg/day of EN. However, there was high uncertainty around the effect: RR 1.04 (95% CI 0.22 to 5.00).

12 **Necrotising enterocolitis**

• Low quality evidence from 1 RCT (n=137) showed a clinically important difference in the number of babies with necrotising enterocolitis between those who had PN stopped at 100ml/kg/day of EN compared with 140ml/kg/day of EN with more babies with necrotising enterocolitis associated with stopping at a higher EN. However, there was high uncertainty around the effect: RR 0.35 (95% CI 0.04 to 3.27).

18 Sepsis

 Low quality evidence from 1 RCT (n=137) showed a clinically important difference in the 19 20 number of babies with central venous catheter-associated late onset sepsis between 21 those who had PN stopped at 100ml/kg/day of EN compared with 140ml/kg/day of EN 22 with more babies with central venous catheter-associated late onset sepsis associated 23 with stopping at a higher EN. However, there was high uncertainty around the effect: RR 24 0.35 (95% CI 0.04 to 3.27).

Stopping PN when EN reached 100ml/kg/day versus stopping PN when EN reached 25 26 120ml/kg/day

27 Weight gain

- Very low quality evidence from 1 observational study (n=87) showed no clinically important difference in time to regain birth weight in babies who had PN stopped at 100ml/kg/day of EN compared with 120ml/kg/day of EN. However, there was uncertainty around the effect: MD -3.40 days (95% CI -6.45 to 0.35).
- 32 • Very low quality evidence from 1 observational study (n=87) showed a clinically important difference in the number of babies whose weight was less than the 10th centile at 36 33 weeks' post conception between those who had PN stopped at 100ml/kg/day of EN 34 35 compared with 120ml/kg/day of EN, with more babies below the 10th centile in the group of babies stopped at the higher EN. However, there was uncertainty around the effect: RR 36 0.72 (95% CI 0.58 to 0.90).
- 38 Very low quality evidence from 1 observational study (n=87) showed a clinically important difference in the number of babies whose weight was less than the 10th centile at 40 39 weeks' post conception between those who had PN stopped at 100ml/kg/day of EN 40 compared with 120ml/kg/day of EN, with more babies below the 10th centile in the group 41 42 of babies stopped at the higher EN. However, there was uncertainty around the effect: RR 43 0.56 (95% CI 0.39 to 0.80).

Linear growth

- Very low quality evidence from 1 observational study (n=87) showed a clinically important difference in the number of babies whose length was less than the 10th centile at 36 weeks' post conception between those who had PN stopped at 100ml/kg/day of EN compared with 120ml/kg/day of EN, with more babies below the 10th centile in the group of babies stopped at the higher EN. However, there is uncertainty around the effect: RR 0.76 (95% CI 0.59 to 1.00).
 - Very low quality evidence from 1 observational study (n=87) showed a clinically important difference in the number of babies whose length was less than the 10th centile at 40 weeks' post conception between those who had PN stopped at 100ml/kg/day of EN compared with 120ml/kg/day of EN, with more babies below the 10th centile in the group of babies stopped at the higher EN. However, there is uncertainty around the effect: RR 0.79 (95% CI 0.63 to 0.99).

Head circumference

- Very low quality evidence from 1 observational study (n=87) showed a clinically important difference in the number of babies whose head circumference was less than the 10th centile at 36 weeks' post conception between those who had PN stopped at 100ml/kg/day of EN compared with 120ml/kg/day of EN, with more babies below the 10th centile in the group of babies stopped at the higher EN: RR 0.54 (95% CI 0.37 to 0.77).
- Very low quality evidence from 1 observational study (n=87) showed a clinically important difference in the number of babies whose head circumference was less than the 10th centile at 40 weeks' post conception between those who had PN stopped at 100ml/kg/day of EN compared with 120ml/kg/day of EN, with more babies below the 10th centile in the group of babies stopped at the higher EN. However, there is uncertainty around the effect: RR 0.60 (95% CI 0.41 to 0.88).

Duration of hospital stay

• Very low quality evidence from 1 observational study (n=87) showed no clinically important difference in duration of hospital stay in babies who had PN stopped at 100ml/kg/day of EN compared with 120ml/kg/day of EN. However, there is uncertainty around the effect: MD -7.90 days (95% CI -19.10 to 3.30).

Necrotising enterocolitis

Very low quality evidence from 1 observational study (n=87) showed a clinically important difference in the number of babies with necrotising enterocolitis between those who had PN stopped at 100ml/kg/day of EN compared with 120ml/kg/day of EN, with more babies with necrotising enterocolitis associated with stopping at a lower EN. However, there is uncertainty around the effect: Peto odds ratio (POR) 5.82 (95% CI 0.35, 97.53).

Duration of PN

Very low quality evidence from 1 observational study (n=87) showed no clinically important difference in duration of PN in babies who had PN stopped at 100ml/kg/day of EN compared with 120ml/kg/day of EN. However, there is uncertainty around the effect: MD -3.50 days (95% CI -9.88 to 2.88).

Sepsis

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Very low quality evidence from 1 observational study (n=87) showed a clinically important difference in the number of babies with sepsis between those who had PN stopped at 100ml/kg/day of EN compared with 120ml/kg/day of EN, with more babies with sepsis associated with stopping at a lower EN. However, there is high uncertainty around the effect: RR 0.79 (95% CI 0.46 to 1.34).

7 Stopping PN when EN reached 60kcal/kg/day versus stopping PN when EN reached 8 100kcal/kg/day

Weight gain

Very low quality evidence from 1 observational study (n=182) showed a clinically important difference in the number of babies with postnatal growth failure at 40 weeks' postmenstrual age between those who had PN stopped at 60kcal/kg/day of EN compared with 100kcal/kg/day of EN, with more babies with growth failure in the group of babies stopped at the lower EN. However, there is uncertainty around the effect: RR 1.45 (95% CI 1.17 to 1.80).

Linear growth

Very low quality evidence from 1 observational study (n=182) showed no clinically important difference in length at 40 weeks' postmenstrual age in babies who had PN stopped at 60kcal/kg/day of EN compared with 100kcal/kg/day of EN. However, there is uncertainty around the effect: MD -1.00cm (95% CI -1.81 to -0.19).

21 Head circumference

Very low quality evidence from 1 observational study (n=182) showed a clinically important difference in head circumference at 40 weeks' postmenstrual age in babies who had PN stopped at 60kcal/kg/day of EN compared with 100kcal/kg/day of EN, with greater head circumference in the group of babies stopped at the higher EN: MD -0.90cm (95% CI -0.96 to -0.84).

27 Necrotising enterocolitis

Very low quality evidence from 1 observational study (n=182) showed a clinically important difference in the number of babies with necrotising enterocolitis between those who had PN stopped at 60kcal/kg/day of EN compared with 100kcal/kg/day of EN, with more babies with necrotising enterocolitis associated with stopping at a lower EN. However, there is high uncertainty around the effect: RR 2.70 (95% CI 0.46 to 15.75).

33 Late onset sepsis

Very low quality evidence from 1 observational study (n=182) showed no clinically important difference in the number of babies with late onset sepsis between those who had PN stopped at 60kcal/kg/day of EN compared with 100kcal/kg/day of EN. However, there is high uncertainty around the effect: RR 0.90 (95% CI 0.52 to 1.55).

38 Economic evidence statements

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

1 The committee's discussion of the evidence

2 Interpreting the evidence

3 The outcomes that matter most

- 4 EN is less energy dense than PN, which is designed to provide an ideal energy composition.
- 5 Additionally, EN must be digested in order to release energy, as opposed to PN which is
- 6 delivered 'ready to use'. Therefore, the committee prioritised total nutritional intake, growth
- 7 and body composition measures as critical outcomes, as nutritional intake has a direct
- 8 impact on growth and body composition.
- 9 Mortality and neurodevelopmental outcomes were selected as important outcomes as these
- are also likely to be affected by differences in nutrition. Sepsis was selected as an important
- 11 outcome as there is an increased risk of sepsis with longer duration of PN due to the
- presence of inserted lines. For this reason, duration of PN was also selected as an important
- outcome to help quantify the risk or benefit associated with stopping PN at different
- 14 thresholds. Finally, duration of hospital stay and necrotising enterocolitis were selected as
- important outcomes as these may be affected by EN, nutritional intake and duration of PN.

16 The quality of the evidence

- 17 The quality of the evidence for this review was assessed using GRADE methodology. The
- observational evidence was very low quality due to risk of bias in the included studies and
- 19 uncertainty around the effects. The RCT evidence ranged from low to high quality and was
- 20 downgraded due to uncertainty around the effects.
- 21 The committee noted that the evidence from the observational studies was confounded due
- to differences in the PN regimens used between arms, which were in addition to the
- 23 difference in the point at which PN was stopped. For example, in one study (Choi 2016), the
- 24 nutritional intake from PN was higher in the arm that stopped PN at a smaller volume of EN
- compared with the arm that stopped at a higher volume of EN. It was therefore often difficult
- 26 to interpretdifferences in the outcomes between groups. The committee were also concerned
- that the duration of PN in one of the observational studies (Choi 2016) was much longer than
- they would expect based on their knowledge, so may not be representative of practice. The
- 29 committee also acknowledged that it was difficult to make comparisons across the studies
- due to the overlap in doses administered to babies in the low and high energy arms of the
- 31 studies. Therefore, the committee agreed that it was not possible to draw firm conclusions
- 32 about the point at which PN should be stopped from the observational evidence.
- 33 There was no RCT evidence available for nutritional intake, body composition or
- 34 neurodevelopmental outcomes and duration of PN and hospital stay were not reported
- 35 sufficiently for analysis.

36 Benefits and harms

- 37 The committee discussed that, based on their expertise, there are a number of factors that
- 38 need to be considered when making the decision to stop PN, beyond the volume of EN that
- 39 has been reached. How well enteral feeds are being tolerated should be considered as
- further PN may be required if babies stop tolerating enteral feeds. Further, the nutritional
- 41 composition of different types of EN is not equivalent and consideration should therefore be
- 42 given to the composition of EN before PN is stopped to avoid nutritional deficits.
- 43 Consideration should also be given to the relative benefit of additional nutritional intake from
- 44 maintaining PN and the risk of venous catheter sepsis, which may be dependent on the

- 1 gestational age and size of the baby. Finally, the committee discussed the need to consider
- the individual baby's circumstances as the decision to stop PN for babies with complex or
- 3 surgical conditions is more complicated. The committee discussed that clinicians may want
- 4 to wait until increasing volumes of EN have been tolerated for a longer period of time before
- 5 stopping PN, and to consider how difficult it may have been to obtain venous access, and
- 6 how many venous access sites are remaining before PN is stopped. However, there was no
- 7 evidence for this group so the committee could not make specific recommendations.
- 8 The RCT evidence showed that it took longer for extremely low birth weight babies to regain
- 9 their birthweight if PN was stopped at a smaller volume of EN (100ml/kg/day) compared with
- 10 a larger volume of EN (140ml/kg/day). There were no clinically important differences based
- on the stopping point of PN for any of the growth outcomes in extremely low and very low
- 12 birth weight babies combined, with the exception of head circumference which was more
- 13 likely to be in the bottom 10th centile when PN was stopped at a smaller volume of EN. There
- were no clinically important differences in mortality rates between groups.
- Low quality evidence showed an increased risk of sepsis and necrotising enterocolitis when
- 16 PN was stopped at a larger volume of EN compared with a smaller volume of EN. The
- 17 committee agreed this was consistent with their expectation of increased risk of sepsis with
- longer duration of PN and that the decision to stop PN requires a balance between
- 19 optimising nutritional intake and minimising the risk of sepsis. However, the committee
- acknowledged that the study was powered to detect differences in time taken to regain
- 21 birthweight, not adverse events.
- The committee agreed that for extremely low birth weight babies, prioritising nutrition, and
- therefore growth, was most critical and that PN should be stopped at a larger volume of EN.
- 24 Whilst the evidence related to extremely low birth weight the committee decided to make
- 25 recommendations based on gestational age (extremely preterm babies <28⁺⁰ weeks'
- gestation) rather than birthweight to be consistent with other recommendations made in this
- 27 guideline and because extremely preterm babies would be more likely to have extremely low
- birth weight than babies born at a later gestational age. The committee noted that EN was
- fortified at 140ml/kg/day in the RCT, which is not common in clinical practice so there was
- 30 concern from the committee that nutrition may be inadequate if PN was stopped at
- 31 140ml/kg/day when EN was not fortified. Therefore, the committee recommended that
- 32 clinicians consider stopping PN when EN reached 140-150ml/kg/day. The upper limit of
- 33 150ml/kg/day was recommended as the committee agreed that this was the threshold at
- which the potential benefits of continuing PN would not outweigh the risks and, therefore, the
- 35 balance favoured discontinuing PN.
- 36 The committee discussed that it is important to maintain nutritional intake when transitioning
- from parenteral to enteral nutrition. However, the committee discussed that clinicians may
- consider stopping PN at a lower volume of enteral feeds for babies born from 28⁺⁰ weeks'
- 39 gestation onwards than for extremely preterm babies, as the balance between the risk from
- 40 nutritional shortfall and the risk of sepsis may be different in this population. The committee
- 41 acknowledged that this balance may be different for growth restricted babies but did not have
- 42 evidence to support a specific recommendation for this group. The committee discussed the
- lower threshold for stopping PN in babies born from 28⁺⁰ weeks' gestation at length. Whilst
- there was little evidence of growth deficits for very low birthweight babies in the RCT when
- 45 PN was stopped at 100ml/kg/day, the committee were concerned that this threshold was too
- low and would result in stopping PN before adequate nutritional intake had been achieved.
- Therefore, the committee recommended by informal consensus that clinicians consider
- stopping PN when EN reached 120-140ml/kg/day for these babies.

- 1 The committee agreed that the thresholds specified in these recommendations should be the
- 2 point at which PN is no longer prescribed, but that any existing PN which is still in place
- 3 should be finished. In practical terms, the committee agreed this would mean stopping PN
- 4 within 24 hours of EN volume reaching the specified volumes.

5 Cost effectiveness and resource use

- 6 No economic studies were identified which were applicable to this review question.
- 7 The committee discussed that these recommendations would reduce variation in practice as
- 8 the point at which parental nutrition is stopped is inconsistent across services. For some
- 9 services, these recommendations will result in providing PN for a longer duration, which
- would have increased costs associated with additional PN and nurse time required to
- administer PN. However, for other services, these recommendations would result in
- 12 providing PN for a shorter duration and may produce cost savings.
- 13 The volume of PN is decreased as the volume of EN is increased during the transition from
- 14 PN to EN. Therefore, the committee were concerned that stopping PN at larger volumes of
- 15 EN may result in very small volumes of PN being prescribed, which would not be practical or
- economically viable i.e. due to a high wastage of PN. As a result, the committee discussed
- whether there should be a minimum volume of parental nutrition that is prescribed, such as
- 18 30ml/kg/day. This could mean that PN is stopped at this point, or that this volume is
- maintained, rather than decreased further, until the threshold of EN is reached. However, the
- 20 minimal volume of parental nutrition was outside the scope of this review question, so no
- 21 evidence was reviewed to support a minimal volume.
- The committee noted that there is a need to balance the risk and benefits of PN including
- optimising nutritional intake and minimising the risk of sepsis and necrotising enterocolitis i.e.
- there was evidence of an increased risk of sepsis and necrotising enterocolitis when PN is
- stopped at a larger volume of EN compared with a smaller volume of EN. However, the
- 26 committee questioned the finding for necrotising enterocolitis due to the studies being
- 27 underpowered to detect a difference in this outcome. The recommendations in this area will
- 28 mean that PN is stopped at an optimal time and will potentially result in fewer cases of
- 29 sepsis. Sepsis incurs substantial costs to the NHS. According to the Hospital Episode
- 30 Statistics 2016/17 the mean duration of stay at NICU was 8 days for babies with sepsis. The
- 31 mean cost of stay at NICU is £1,445 per day. Any strategy that reduces the risk of sepsis is
- 32 likely to represent a cost effective use of NHS resources.

33 Other factors the committee took into account

- Whilst the focus of this review question was the point at which PN should be stopped, the
- 35 committee agreed that it was also important to ensure that adequate nutritional intake has
- been reached before weaning from PN is started. However, they could not make
- 37 recommendations in this area as it was outside the scope of the review question.

38 References

- 39 **Choi 2016**
- 40 Choi, A. Y., Lee, Y. W., Chang, M. Y., Modification of nutrition strategy for improvement of
- 41 postnatal growth in very low birth weight infants, Korean Journal of Pediatrics, 59, 165-173,
- 42 2016

DRAFT FOR CONSULTATION

Stopping parenteral nutrition in preterm and term babies

1 Dinerstein 2006

- Dinerstein, A., Nieto, R. M., Solana, C. L., Perez, G. P., Otheguy, L. E., Larguia, A. M., Early
- 3 and aggressive nutritional strategy (parenteral and enteral) decreases postnatal growth
- 4 failure in very low birth weight infants, Journal of Perinatology, 26, 436-42, 2006

5 **Perrem 2019**

- 6 Perrem, L., Semberova, J., O'Sullivan, A., Kieran, E. A., O'Donnel, C. P. F., White, M. J.,
- 7 Miletin, J., Effect of early parenteral nutrition discontinuation on time to regain birth weight in
 - very low birth weight infants: a randomized controlled trial, Journal of Parenteral and Enteral
- 9 Nutrition, 2019

10

8

11

Appendices

2 Appendix A – Review protocols

- 3 Review protocol for review question: What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in
- 4 relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?

5 Table 3: Review protocol – stopping parenteral nutrition in preterm and term babies

Field (based on PRISMA-P	Content
Review question RQ 1.2	What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?
Type of review question	Intervention
Objective of the review	Determining when parenteral nutrition can be stopped is important; there is risk of malnutrition and associated complications to the baby if parenteral feeding is stopped too early. In contrast, the longer parenteral feeding is continued, the greater the risk of line sepsis. Guidance for clinicians on how to determine when it is safe to transition from combined parenteral and enteral nutrition to enteral nutrition alone is required.
Eligibility criteria –	• Babies born preterm, up to 28 days after their due birth date (preterm babies)
population/disease/condition/issue/domain	Babies born at term, up to 28 days after their birth (term babies)
Eligibility criteria – intervention(s)/exposure(s)/prognostic factor(s)	Volume of PN (ml/kg/day or kcal/kg/day) plus EN (ml/kg/day or kcal/kg/day Ratio of PN (ml/kg/day or kcal/kg/day) and EN (ml/kg/day or kcal/kg/day)
Eligibility criteria – comparator(s)/control or reference (gold) standard	Comparison volume of PN (ml/kg/day or kcal/kg/day) plus EN (ml/kg/day or kcal/kg/day) Different ratio of PN (ml/kg/day or kcal/kg/day) to EN (ml/kg/day or kcal/kg/day)
Outcomes and prioritisation	Critical
	 Total nutritional intake, protein/carbohydrate/lipid/energy (for example, kcal/kg/day as a total of the parental + enteral)
	Growth/Anthropometric measures: Weight rain
	Weight gainLinear growth
	Head circumference (mm)
	o Body composition (lean mass, fat-free mass, fat mass, adipose tissue, nitrogen accretion)
	Important

Field (based on PRISMA-P	Content
	Mortality
	Duration of hospital stay
	Necrotising enterocolitis
	 Neurodevelopmental outcomes (general cognitive abilities at two years, measured using validated scales only)
	Duration of PN
	Sepsis (central line infections)
Eligibility criteria – study design	Systematic reviews of RCTs RCTs Comparative cohort studies (only if RCTs unavailable or limited data to inform decision making),
	retrospective or prospective
Other inclusion exclusion criteria	No sample size restriction No date restriction Studies conducted in the community, residential, primary and secondary care will be included
Proposed sensitivity/sub-group analysis, or	Stratified analysis
meta-regression	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)
	Where evidence exists, consideration will be given to the specific needs of population subgroups: 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)
	Birth weight: 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)
	Important confounders (when comparative observational studies are included for interventional reviews): Age of baby (first 2 weeks vs. later)
	Preterm (Very early <28 weeks' GA; 28-31 weeks' GA; 32-36 weeks' GA)
	Birth weight: Low birth weight (< 2500g); very low birth weight (< 1500g) and extremely low birth weight (< 1000g)
Selection process – duplicate	Study Selection
screening/selection/analysis	Studies will be imported to the NGA STAR database for screening by one reviewer. Dual sifting will not be performed for this review question. All full texts identified will be screened for inclusion by two reviewers. All

ield (based on <u>PRISMA-P</u>	Content
	disagreements will be resolved by discussion between the two reviewers. The senior systematic reviewer or guideline lead will act as arbiter where necessary.
Pata management (software)	Data Analysis
	Where data is available, pair-wise meta-analysis using a fixed effects model, will be used to combine results from similar studies, this will be performed using Cochrane Review Manager (RevMan5). Heterogeneity will be considered, and if a random-effects model is considered more appropriate, it will be conducted. Quality Assessment
	Appraisal of methodological quality will be conducted using the appropriate tool:
	ROBIS (systematic reviews and meta-analyses),
	Cochrane risk of bias tool for RCTs (RCT or comparative cohort studies).
	· · · · · · · · · · · · · · · · · · ·
	Cochrane risk of bias tool, ROBINS-I (Non-randomised studies)
	The quality of evidence for each outcome will be assessed using GRADEpro:
	Outcomes will be downgraded if the randomisation and/or concealment methods are unclear or inadequate. Outcomes will also be downgraded if there is considerable missing data (if there is a dropout of more than 20%, or if there is a difference of >20% between groups.
	Heterogeneity will be assessed using the I2, outcomes will be downgraded once if I2 >50%, twice if I2 >80%. Imprecision: Outcomes will be downgraded if the 95% CI is imprecise (i.e. crosses 0.8 or 1.25, (dichotomous) or -0.5 or 0.5 (continuous)). Outcomes will be downgraded two levels depending on how many lines of imprecision are crossed. If the clinical decision threshold is NOT crossed, we will consider whether the criterion for Optimal Information Size is met, if not we will downgrade one level for dichotomous outcomes with less than 300 events, and downgrade one level for continuous outcomes when less than 400 participants are included.
	Clinical effectiveness
	For dichotomous outcomes, minimal important differences will be considered using thresholds of RR >0.80 and <1.25.
	For continuous outcomes, minimal important differences will be considered 0.5 times the SD of the control group
nformation 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.

Field (based on PRISMA-P	Content
Author contacts	Developer: The National Guideline Alliance
	https://www.nice.org.uk/guidance/indevelopment/gid-ng10037
Highlight if amendment to previous protocol	For details please see section 4.5 of <u>Developing NICE guidelines: the manual</u> 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/
Criteria for quantitative synthesis (where suitable)	For details please see section 6.4 of <u>Developing NICE guidelines: the manual 2014</u> .
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 <u>Developing NICE guidelines: the manual 2014</u> .
Assessment of confidence in cumulative evidence	For details please see sections 6.4 and 9.1 of <u>Developing NICE guidelines: the manual 2014</u> .
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 NGA 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

Fie	eld (based on PRISMA-P	Content
Ro	oles of sponsor	NICE funds the National Guideline Alliance (to develop guidelines for those working in the NHS, public health, and social care in England
PR	ROSPERO registration number	The review is not registered with PROSPERO

CDSR: Cochrane Database of Systematic Reviews; CCTR: Cochrane controlled trials register; DARE: Database of Abstracts of Reviews of Effects; EN: enteral nutrition; GRADE: Grading of Recommendations Assessment, Development and Evaluation; HTA: Health Technology Assessment; ICF: International Classification of Functioning, Disability and Health; NGA: National Guideline Alliance; NHS: National health service; NICE: National Institute for Health and Care Excellence; PN: parenteral nutrition; PROSPERO: International prospective register of systematic reviews; RCT: randomised controlled trial; RoB: risk of bias; ROBINS-I: risk of bias in non-randomised studies of interventions; ROBIS; risk of bias in systematic reviews; RR: risk ratio; SD: standard deviation

1 Appendix B – Literature search strategies

- 2 Literature search strategies for review question: What amount of enteral feed
- (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and
- enteral feeds) indicates that parenteral nutrition is no longer required?

5 Databases: Medline; Medline EPub Ahead of Print; and Medline In-Process &

6 Other Non-Indexed Citations

	Non-indexed Gitations
#	Searches
1	INFANT, NEWBORN/
2	(neonat\$ or newborn\$ or new-born\$ or baby or babies).ti,ab.
3	PREMATURE BIRTH/
4	((preterm\$ or pre-term\$ or pre-matur\$ or pre-matur\$) adj5 (birth? or born)).ab,ti.
5	exp INFANT, PREMATURE/
6	((preterm\$ or pre-term\$ or prematur\$ or pre-matur\$) adj5 infan\$).ti,ab.
7	(pre#mie? or premie or premies).ti,ab.
8	exp INFANT, LOW BIRTH WEIGHT/
9	(low adj3 birth adj3 weigh\$ adj5 infan\$).ti,ab.
10	((LBW or VLBW) adj5 infan\$).ti,ab.
11	INTENSIVE CARE, NEONATAL/
12	INTENSIVE CARE UNITS, NEONATAL/
13	NICU?.ti,ab.
14	or/1-13
15	PARENTERAL NUTRITION/
16	PARENTERAL NUTRITION, TOTAL/
17	PARENTERAL NUTRITION SOLUTIONS/
18	ADMINISTRATION, INTRAVENOUS/ and (nutrition\$ or feed\$ or fed\$).ti,ab.
19	INFUSIONS, INTRAVENOUS/ and (nutrition\$ or feed\$ or fed\$).ti,ab.
20	CATHETERIZATION, CENTRAL VENOUS/ and (nutrition\$ or feed\$ or fed\$).ti,ab.
21	exp CATHETERIZATION, PERIPHERAL/ and (nutrition\$ or feed\$).ti,ab.
22	((parenteral\$ or intravenous\$ or intra-venous\$ or IV or venous\$ or infusion?) adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
23	((peripheral\$ or central\$) adj3 line? adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
24	(catheter\$ adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
25	(drip? adj3 (nutrition\$ or feed\$) or fed\$)).ti,ab.
26	or/15-25
27	ENTERAL NUTRITION/
28	INTUBATION, GASTROINTESTINAL/
29	GASTROSTOMY/
30	JEJUNOSTOMY/
31	((enteral\$ or tube? or oral\$ or sip) adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
32	((Nasogastric\$ or gastrointestinal\$) adj3 (tube? or intubate\$ or nutrition\$ or feed\$ or fed\$)).ti,ab.
33	Gastrostom\$.ti,ab.
34	Jejunostom\$.ti,ab.
35	or/27-34
36	((wean\$ or decreas\$ or de-creas\$ or halt\$ or ceas\$ or cessat\$ or suspen\$ or stop\$ or end\$ or discontinu\$ or discontinu\$ or finish\$ or transition\$) adj5 (parenteral\$ or intravenous\$ or intra-venous\$ or IV or venous\$ or infusion?) adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
37	((wean\$ or decreas\$ or de-creas\$ or halt\$ or ceas\$ or cessat\$ or suspen\$ or stop\$ or end\$ or discontinu\$ or discontinu\$ or finish\$ or transition\$) adj5 (PN or SPN or IPN or TPN or STD-PN or IND-PN)).ti,ab.
38	or/36-37
39	((Advanc\$ or Achiev\$ or Establish\$ or Tolera\$ or Success\$ or Full\$) adj3 (enteral\$ or tube? or oral\$ or sip) adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
40	((amount? or volume? or ratio?) adj10 (parenteral\$ or intravenous\$ or intra-venous\$ or IV or venous\$ or infusion?) adj10 (enteral\$ or tube? or oral\$ or sip) adj10 (nutrition\$ or feed\$ or fed\$)).ti,ab.
41	((ml? or milliliter? or kcal? or kilocalor\$) adj3 (kg? or kilogram?) adj3 (d or day)).ti,ab.
42	ENERGY INTAKE/
43	NUTRITIONAL STATUS/
44	((energy or volume? or kcal or kilocalorie? or nutrition\$) adj3 (goal? or target\$)).ti,ab.
45	((optimi\$ or success\$) adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
46	or/42-45

#	Searches
47	PARENTERAL NUTRITION/mt [Methods]
48	ENTERAL NUTRITION/mt [Methods]
49	14 and 35 and 38
50	14 and 26 and 39
51	14 and 40
52	14 and 26 and 35 and 41
53	14 and 26 and 35 and 46
54	14 and 47 and 48
55	or/49-54
56	limit 55 to english language
57	LETTER/
58	EDITORIAL/
59	NEWS/
60	exp HISTORICAL ARTICLE/
61	ANECDOTES AS TOPIC/
62	COMMENT/
63	CASE REPORT/
64	(letter or comment*).ti.
65	or/57-64
66	RANDOMIZED CONTROLLED TRIAL/ or random*.ti,ab.
67	65 not 66
68	ANIMALS/ not HUMANS/
69	exp ANIMALS, LABORATORY/
70	exp ANIMAL EXPERIMENTATION/
71	exp MODELS, ANIMAL/
72	exp RODENTIA/
73	(rat or rats or mouse or mice).ti.
74	or/67-73
75	56 not 74

1 Databases: Embase; and Embase Classic

#	Searches
1	NEWBORN/
2	(neonat\$ or newborn\$ or new-born\$ or baby or babies).ti,ab.
3	PREMATURITY/
4	((preterm\$ or pre-term\$ or prematur\$ or pre-matur\$) adj5 (birth? or born)).ab,ti.
5	((preterm\$ or pre-term\$ or prematur\$ or pre-matur\$) adj5 infan\$).ti,ab.
6	(pre#mie? or premie or premies).ti,ab.
7	exp LOW BIRTH WEIGHT/
8	(low adj3 birth adj3 weigh\$ adj5 infan\$).ti,ab.
9	((LBW or VLBW) adj5 infan\$).ti,ab.
10	NEWBORN INTENSIVE CARE/
11	NEONATAL INTENSIVE CARE UNIT/
12	NICU?.ti,ab.
13	or/1-12
14	PARENTERAL NUTRITION/
15	TOTAL PARENTERAL NUTRITION/
16	PERIPHERAL PARENTERAL NUTRITION/
17	PARENTERAL SOLUTIONS/
18	INTRAVENOUS FEEDING/
19	INTRAVENOUS DRUG ADMINISTRATION/ and (nutrition\$ or feed\$ or fed\$).ti,ab.
20	exp INTRAVENOUS CATHETER/ and (nutrition\$ or feed\$ or fed\$).ti,ab.
21	((parenteral\$ or intravenous\$ or intra-venous\$ or IV or venous\$ or infusion?) adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
22	((peripheral\$ or central\$) adj3 line? adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
23	(catheter\$ adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
24	(drip? adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
25	or/14-24
26	ENTERIC FEEDING/
27	exp DIGESTIVE TRACT INTUBATION/
28	GASTROSTOMY/
29	JEJUNOSTOMY/
30	((enteral\$ or tube? or oral\$ or sip) adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
31	((Nasogastric\$ or gastrointestinal\$) adj3 (tube? or intubate\$ or nutrition\$ or feed\$ or fed\$)).ti,ab.

#	Searches
32	Gastrostom\$.ti,ab.
33	Jejunostom\$.ti,ab.
34	or/26-33
35	((wean\$ or decreas\$ or de-creas\$ or halt\$ or ceas\$ or cessat\$ or suspen\$ or stop\$ or end\$ or discontinu\$ or discontinu\$ or finish\$ or transition\$) adj5 (parenteral\$ or intravenous\$ or intra-venous\$ or IV or venous\$ or infusion?) adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
36	((wean\$ or decreas\$ or de-creas\$ or halt\$ or ceas\$ or cessat\$ or suspen\$ or stop\$ or end\$ or discontinu\$ or discontinu\$ or finish\$ or transition\$) adj5 (PN or SPN or IPN or TPN or STD-PN or IND-PN)).ti,ab.
37	or/35-36
38	((Advanc\$ or Achiev\$ or Establish\$ or Tolera\$ or Success\$ or Full\$) adj3 (enteral\$ or tube? or oral\$ or sip) adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
39	((amount? or volume? or ratio?) adj10 (parenteral\$ or intravenous\$ or intra-venous\$ or IV or venous\$ or infusion?) adj10 (enteral\$ or tube? or oral\$ or sip) adj10 (nutrition\$ or feed\$ or fed\$)).ti,ab.
40	((ml? or milliliter? or kcal? or kilocalor\$) adj3 (kg? or kilogram?) adj3 (d or day)).ti,ab.
41	CALORIC INTAKE/
42	NUTRITIONAL STATUS/
43	((energy or volume? or kcal or kilocalorie? or nutrition\$) adj3 (goal? or target\$)).ti,ab.
44	((optimi\$ or success\$) adj3 (nutrition\$ or feed\$ or fed\$)).ti,ab.
45	or/41-44
46	13 and 34 and 37
47	13 and 25 and 38
48	13 and 39
49	13 and 25 and 34 and 40
50	13 and 25 and 34 and 45
51	or/46-50
52	limit 51 to english language
53	letter.pt. or LETTER/
54	note.pt.
55	editorial.pt.
56	CASE REPORT/ or CASE STUDY/
57	(letter or comment*).ti.
58	or/53-57
59	RANDOMIZED CONTROLLED TRIAL/ or random*.ti,ab.
60	58 not 59
61	ANIMAL/ not HUMAN/
62	NONHUMAN/
63	exp ANIMAL EXPERIMENT/
64	exp EXPERIMENTAL ANIMAL/
65	ANIMAL MODEL/
66	exp RODENT/
67	(rat or rats or mouse or mice).ti.
68	or/60-67
69	52 not 68

1 Databases: Cochrane Central Register of Controlled Trials; Cochrane Database of

2 Systematic Reviews; Database of Abstracts of Reviews of Effects; and Health

3 Technology Assessment

• • • • • • • • • • • • • • • • • • • •	neregy recomment
#	Searches
#1	MeSH descriptor: [INFANT, NEWBORN] this term only
#2	(neonat* or newborn* or new-born* or baby or babies):ti,ab
#3	MeSH descriptor: [PREMATURE BIRTH] this term only
#4	((preterm* or pre-term* or pre-matur* or pre-matur*) near/5 (birth* or born)):ti,ab
#5	MeSH descriptor: [INFANT, PREMATURE] explode all trees
#6	((preterm* or pre-term* or pre-matur* or pre-matur*) near/5 infan*):ti,ab
#7	(pre?mie? or premie or premies):ti,ab
#8	MeSH descriptor: [INFANT, LOW BIRTH WEIGHT] explode all trees
#9	(low near/3 birth near/3 weigh* near/5 infan*):ti,ab
#10	((LBW or VLBW) near/5 infan*):ti,ab
#11	MeSH descriptor: [INTENSIVE CARE, NEONATAL] this term only
#12	MeSH descriptor: [INTENSIVE CARE UNITS, NEONATAL] this term only
#13	NICU?:ti,ab
#14	#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11 or #12 or #13

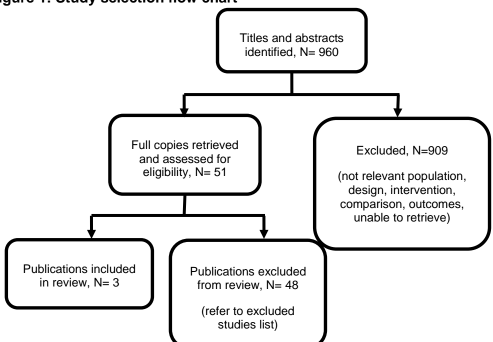
#	Searches
#15	MeSH descriptor: [PARENTERAL NUTRITION] this term only
#16	MeSH descriptor: [PARENTERAL NUTRITION, TOTAL] this term only
#17	MeSH descriptor: [PARENTERAL NUTRITION SOLUTIONS] this term only
#18	MeSH descriptor: [ADMINISTRATION, INTRAVENOUS] this term only
#19	MeSH descriptor: [INFUSIONS, INTRAVENOUS] this term only
#20	MeSH descriptor: [CATHETERIZATION, CENTRAL VENOUS] this term only
#21	MeSH descriptor: [CATHETERIZATION, PERIPHERAL] explode all trees
#22	#18 or #19 or #20 or #21
#23	(nutrition* or feed* or fed*) :ti,ab
#24	#22 and #23
#25	((parenteral* or intravenous* or intra-venous* or IV or venous* or infusion?) near/3 (nutrition* or feed* or fed*)) :ti,ab
#26	((peripheral* or central*) near/3 line? near/3 (nutrition* or feed* or fed*)) :ti,ab
#27	(catheter* near/3 (nutrition* or feed* or fed*)) :ti,ab
#28	(drip? near/3 (nutrition* or feed* or fed*)) :ti,ab
#29	#15 or #16 or #17 or #24 or #25 or #26 or #27 or #28
#30	MeSH descriptor: [ENTERAL NUTRITION] this term only
#31	MeSH descriptor: [INTUBATION, GASTROINTESTINAL] this term only
#32	MeSH descriptor: [GASTROSTOMY] this term only
#33	MeSH descriptor: [JEJUNOSTOMY] this term only
#34	((enteral* or tube? or oral* or sip) near/3 (nutrition* or feed* or fed*)) :ti,ab
#35	((Nasogastric* or gastrointestinal*) near/3 (tube? or intubate* or nutrition* or feed* or fed*)) :ti,ab
#36	Gastrostom*:ti,ab
#37	Jejunostom*:ti,ab
#38	#30 or #31 or #32 or #33 or #34 or #35 or #36 or #37
#39	((wean* or decreas* or de-creas* or halt* or ceas* or cessat* or suspen* or stop* or end* or discontinu* or finish* or transition*) near/5 (parenteral* or intravenous* or intra-venous* or IV or venous* or infusion*) near/3 (nutrition* or feed* or fed*)):ti,ab
#40	((wean* or decreas* or de-creas* or halt* or ceas* or cessat* or suspen* or stop* or end* or discontinu* or dis-continu* or finish* or transition*) near/5 (PN or SPN or IPN or TPN or STD-PN or IND-PN)):ti,ab
#41	#39 or #40
#42	((Advanc* or Achiev* or Establish* or Tolera* or Success* or Full*) near/3 (enteral* or tube* or oral* or sip) near/3 (nutrition* or feed* or fed*)):ti,ab
#43	((amount* or volume* or ratio*) near/10 (parenteral* or intravenous* or intra-venous* or IV or venous* or infusion*) near/10 (enteral* or tube* or oral* or sip) near/10 (nutrition* or feed* or fed*)):ti,ab
#44	((ml* or milliliter* or kcal* or kilocalor*) near/3 (kg* or kilogram*) near/3 (d or day)):ti,ab
#45	MeSH descriptor: [ENERGY INTAKE] this term only
#46	MeSH descriptor: [NUTRITIONAL STATUS] this term only
#47	((energy or volume* or kcal or kilocalorie* or nutrition*) near/3 (goal* or target*)):ti,ab
#48	((optimi* or success*) near/3 (nutrition* or feed* or fed*)):ti,ab
#49	#45 or #46 or #47 or #48
#50	MeSH descriptor: [PARENTERAL NUTRITION] this term only and with qualifier(s): [methods - MT]
#51	MeSH descriptor: [ENTERAL NUTRITION] this term only and with qualifier(s): [methods - MT]
#52	#14 and #38 and #41
#53	#14 and #29 and #42
#54	#14 and #43
#55	#14 and #29 and #38 and #44
#56	#14 and #29 and #38 and #49
#57	#14 and #50 and #51
#58	#52 or #53 or #54 or #55 or #56 or #57

1

1 Appendix C - Clinical evidence study selection

- 2 Clinical study selection for: What amount of enteral feed (measured in terms of
- 3 ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that
- 4 parenteral nutrition is no longer required?

Figure 1: Study selection flow chart



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1 Appendix D – Clinical evidence tables

- 2 Clinical evidence tables for review question: What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or
- 3 ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?

4 Table 4: Clinical evidence tables

Study details	Participants	Interventions	Methods	Outcomes and Results	Comments
Full citation	Sample size	Interventions	Details	Results	Limitations
Choi, A. Y., Lee, Y. W., Chang, M. Y., Modification of nutrition strategy for improvement of postnatal growth in very low birth weight infants, Korean Journal of Pediatrics, 59, 165- 173, 2016	N=124; only n=87 of interest for the purpose of this review question (n=50 100ml/kg/day; n=37 120ml/kg/day) n=37 infants from period 3 were not included as differences between nutritional protocols in period 2 and 3 are changes to enteral	Infants admitted between October 2010 and September 2011 formed the first cohort. Infants admitted between October 2011 and December 2012 formed the second cohort (after modifications to PN protocol, and	PN regimen prior to stopping: On day 1 of life, 60ml of fluid was started with glucose infusion rate of 4 to 6mg/kg/minute and increased to 10 to 15mg/kg/minute based on blood glucose level. Proteins were supplied using a 10% amino acid 100ml solution and lipids were supplied using a	Weight gain - time (days) to regain birth weight - mean (SD): 100ml/kg/day (n=50): 14.3 (5.3) 120ml/kg/day (n=37): 17.7 (8.3) Weight gain - weight <10th centile at 36 weeks post conception - n/N:	Quality of study assessed using ROBINS-I Confounding bias: Low risk. Selection of participants' bias: Moderate risk. Retrospective study; start and follow-up of the two cohorts differ. Classification of interventions bias: Low
688689 Country/ies where the	practices, not PN Characteristics	minimal modifications to EN protocol). Period 1 -	20% lipid 250ml solution. During period 1, proteins	100ml/kg/day: 33/50 120ml/kg/day: 34/37	risk. Intervention groups clearly defined.
study was carried out Korea	Gestational age (weeks) - mean (SD) 100ml/kg/day: 29+1	120ml/kg/day: Nutritional supplements were	were started on the first day of life at 1.5g/kg/day and advanced to 3.5g/kg/day; lipids were	Weight gain - weight <10th centile at 40 weeks post	Deviations from intended interventions bias: Unclear risk.
Study type Observational study	(2+2) 120ml/kg/day: 29+5 (2+5)	added and PN discontinued when infants reached 100ml/kg/day of	started on the second day of life at 0.5g/kg/day and advanced to 3.0g/kg/day. During period 2, proteins	conception - n/N: 100ml/kg/day: 22/50 120ml/kg/day: 29/37	Protocol violations, if any occurred, are not reported.
Aim of the study The aim of the study was to: 1) modify parenteral and enteral	Birth weight (g) - mean (SD) 100ml/kg/day: 1120 (222)	enteral feeds Period 2 - 100ml/kg/day:	was started on the first day of life at 3.0g/kg/day and advanced to 4.0g/kg/day; lipids were	Linear growth - length <10th centile at 36	Missing data bias: Low risk. No missing data. Measurement of outcomes bias: Low risk.

Study details	Participants	Interventions	Methods	Outcomes and Results	Comments
nutrition protocols based on recent literature, and 2) evaluate the impact of modified protocols on growth of very low birth weight infants Study dates October 2010 to April 2014 (only interested in October 2012 for the purpose of this review question) Source of funding No sources of funding reported	120ml/kg/day: 1135 (252) Length (cm) at birth - mean (SD) 100ml/kg/day: 36.3 (3.0) 120ml/kg/day: 37.2 (3.0) Head circumference (cm) at birth - mean (SD) 100ml/kg/day: 26.2 (1.8) 120ml/kg/day: 26.5 (1.9) Inclusion criteria Very low birth weight infants (<1,500g) born before 34 weeks' gestation who were admitted to 1 NICU between October 2010 and April 2014. Exclusion criteria Died during hospitalisation; transferred to other hospitals; serious congenital malformation that needed surgical treatment	Nutritional supplements were added and PN discontinued when infants reached 120ml/kg/day of enteral feeds	started on the first day of life at 1.0/kg/day and advanced to 3.5g/kg/day. Enteral regimen: Minimal enteral feeding was started as soon as possible at 10 to 15ml/kg/day and increased by 10 to 30ml/kg/day. Human milk was preferred but preterm formula was used if milk was unavailable. Milk was fortified and supplemented with protein when full enteral feeds were reached. During period 1, the goal for enteral feeding was 3.0 to 3.5g/kg/day of protein and 120kcal/kg/day; protein goals were the same in period 2 but calorie goal was 130kcal/kg/day. Power analysis: Not stated Statistical analysis: Analysis was conducted using SPSS version 22.0. Chi-squared tests and Fisher exact tests were used for categorical variables, Kruskal-Wallis	weeks post conception - n/N: 100ml/kg/day: 31/50 120ml/kg/day: 30/37 Linear growth - length <10th centile at 40 weeks' post conception - n/N: 100ml/kg/day: 34/50 120ml/kg/day: 32/37 Head circumference <10th centile at 36 ' post conception - n/N: 100ml/kg/day: 21/50 120ml/kg/day: 29/37 Head circumference <10th centile at 40 weeks' post conception - n/N: 100ml/kg/day: 29/37 Head circumference <10th centile at 40 weeks' post conception - n/N: 100ml/kg/day: 21/50 120ml/kg/day: 26/37 Duration (days) of hospital stay - mean (SD): 100ml/kg/day (n=50): 68.9 (21.4) 120mlkg/day (n=37): 76.8 (29.5)	Unlikely that outcome assessors were blind to intervention for safety reasons but all outcomes are objective. Selection of the reported results bias: Moderate risk. Insufficient reporting of growth rates and measurements at 40 weeks' post conceptional age for weight, height and head circumference. Other information There are multiple differences between the nutrition protocols followed during the different time periods apart from the point at which PN was discontinued. Therefore, it is difficult to conclude whether any differences observed between groups are a result of when PN was stopped.

Study details	Participants	Interventions	Methods	Outcomes and Results	Comments
Study details	raticipants	interventions	test was used for serial variants and Pearson coefficient was used for correlations.	Necrotising enterocolitis - n/N: 100ml/kg/day: 2/50 120ml/kg/day: 0/37 Duration (days) of PN - mean (SD): 100ml/kg/day (n=50): 25.5 (14.3) 120mlkg/day (n=37): 29.0 (15.5) Sepsis - n/N: 100ml/kg/day: 17/50 120ml/kg/day: 16/37	Comments
Full citation Dinerstein, A., Nieto, R. M., Solana, C. L., Perez, G. P., Otheguy, L. E., Larguia, A. M., Early and aggressive nutritional strategy (parenteral and enteral) decreases postnatal growth failure in very low birth weight infants, Journal of Perinatology, 26, 436- 42, 2006 Ref Id 378253	Sample size N=200 (n=128 100kcal/kg/day; n=72 60kcal/kg/day) N=182 analysed (n=117 100kcal/kg/day [n=6 died; n=5 transferred before 40 weeks postmenstrual age]; n=65 60kcal/kg/day [n=3 died; n=4 transferred before 40 weeks postmenstrual age]) Characteristics Gestational age (weeks) - median (range) 60kcal/kg/day: 30 (24- 36)	Interventions Infants born between August 2001 and July 2003 were treated according to the aggressive nutritional regimen and infants born during the preceding year were treated according to the conservative nutritional regimen. Aggressive regimen - 100kcal/kg/day: PN was discontinued when infants reached 100kcal/kg/day of enteral feeds	Details Aggressive nutritional regimen: Fluids were begun at 80ml/kg/day and advanced to 150 to 180ml/kg/day over 7 days. Amino acids were delivered from day 1 in a 10% solution starting at 1.5g/kg/day and advanced by 0.5g/kg/day to 4g/kg/day, unless there was renal failure or metabolic acidosis. Glucose infusion began on day 1 at a rate of 5.6mg/kg/minute and advanced by	Results Nutritional intake (kcal/kg/week)* - median (min/max): Week 1 100kcal/kg/day: 497 (331/731) 60kcal/kg/day: 351 (195/605) Week 2 100kcal/kg/day: 770 (411/1120) 60kcal/kg/day: 679 (348/917) Week 3 100kcal/kg/day: 875 (333/1371) 60kcal/kg/day: 802	Limitations Quality of study assessed using ROBINS-I Confounding bias: Low risk. Selection of participants' bias: Moderate risk. Start and follow-up of the two cohorts differ. Classification of interventions bias: Low risk. Intervention groups clearly defined. Deviations from intended interventions bias: Unclear risk. Protocol violations, if any

Study details	Participants	Interventions	Methods	Outcomes and Results	Comments
No sources of funding reported			amino acids using a 10% solution at the discretion of neonatal staff on day 3 of life at a rate of 0.5g/kg/day; this was advanced by 0.5g/kg/day up to 3g/kg/day. Glucose was started (day not reported) at a rate of 5.6mg/kg/minute and advanced by 1mg/kg/minute every day to a maximum of 8 to 9mg/kg/minute. Lipids were introduced in a 20% solution at day 3 or 4 of life at a rate of 0.5g/kg/day and increased by 0.5g/kg/day up to 3g/kg/day. Enteral feeds were introduced when the attending physician thought the infant was clinically stable. Power analysis: Not stated Statistical analysis: Analysis was conducted using Stata version 7. Categorical variables were analysed using chisquared tests. Student's t tests were used for	postmenstrual age - mean (SD): 100kcal/kg/day (n=117): 35.2 (0.17) 60kcal/kg/day (n=65): 34.3 (0.21) Necrotising enterocolitis - n/N: 100kcal/kg/day: 2/117 60kcal/kg/day: 3/65 Duration (days) of PN* - median (range): 100kcal/kg/day: 10 (5-36) 60kcal/kg/day: 4 (0-37) Sepsis - late onset sepsis - n/N: 100kcal/kg/day: 30/117 60kcal/kg/day: 15/65 *Not analysed as data is reported as medians	

Study details	Participants	Interventions	Methods	Outcomes and Results	Comments
·			normally distributed continuous variables and Mann-Whitney U-tests were used when data was not normally distributed.		
Full citation Perrem, L., Semberova, J., O'Sullivan, A., Kieran, E. A., O'Donnel, C. P. F., White, M. J., Miletin, J., Effect of early parenteral nutrition discontinuation on time to regain birth weight in very low birth weight in very low birth weight infants: a randomized controlled trial, Journal of Parenteral and Enteral Nutrition, 2019 Ref Id 1009143 Country/ies where the study was carried out Ireland Study type Randomised controlled trial Aim of the study	Sample size N=317 assessed for eligibility (n=104 did not meet inclusion criteria; n=54 reached 100 mL/kg/day of enteral feeds before study enrolment; n=20 declined participation) N=139 randomised (n=69 100ml/kg/day; n=70 140ml/kg/day) N=137 analysed (n=67 100ml/kg/day [n=1 withdrew consent; n=1 died prior to primary outcome]; n=70 140ml/kg/day) Characteristics Gestational age (weeks) - mean (SD) 100ml/kg/day: 28.7 (2.6) 140ml/kg/day: 28.5 (2.1) Birth weight Weight (g) - median (IQR) 100ml/kg/day: 1060 (800-1300)	Interventions Infants were randomised to the below groups when they were clinically stable and had a PICC inserted, but before they reached 100m;/kg/day of enteral feeds. 100ml/kg/day: PICC was removed and PN discontinued when infants reached 100ml/kg/day of enteral feeds 140ml/kg/day: PICC was removed and PN discontinued when infants reached 140ml/kg/day: PICC was removed and PN discontinued when infants reached 140ml/kg/day of enteral feeds	PN regimen prior to stopping: All infants received 60-80ml/kg/day PN within the first 24 hours of life using a standard bag of preterm PN (content/100 mL: amino acids 2.51g, glucose 10%, sodium 2.5mmol, potassium 2mmol, calcium 1mmol, magnesium 0.15mmol, phosphate 1.25mmol). This was increased by 10-20ml/kg/day up to a target volume of 150-160ml/kg/day. Enteral regimen: If infants were stable, 10-20ml/kg/day of enteral feeds were started on the first day of life. Expressed breast milk (EBM), either maternal or donor, was the preferred choice of enteral feeds and was received by all infants ≤1250g and growth-restricted infants >1250g at higher risk of	Results Weight gain - time (days) to regain birth weight - mean (SD): All infants (VLBW & ELBW) 100ml/kg/day (n=67): 10.8 (3.6) 140ml/kg/day (n=70): 9.3 (3.5) VLBW infants 100ml/kg/day (n=41): 10.7 (3.5) 140ml/kg/day (n=42): 10.1 (3.0) ELBW infants 100ml/kg/day (n=26): 10.9 (3.7) 140ml/kg/day (n=28): 8.1 (3.8) Weight gain - weight (g) at discharge - mean (SD): 100ml/kg/day (n=67): 2391 (660) 140ml/kg/day (n=70): 2378 (551)	Limitations Quality of study assessed using Cochrane risk of bias tool Selection bias Random sequence generation: Low risk. Computer-generated in blocks of 4 and 8, stratified by birth weight (<1000g or 1000-1499g) and centre. Allocation concealment: Low risk. Randomisation list prepared by research assistant who had no involvement in the trial and placed in sealed, opaque, sequentially numbered envelopes. Performance bias Blinding of participants and personnel: Low risk. Infants would be unaware of their assignment. Caregivers were not blinded, likely due to safety reasons.

Study details	Participants	Interventions	Methods	Outcomes and Results	Comments
To compare the effect of stopping parenteral nutrition at 2 different volumes of enteral feeding on the time taken for very low birth weight infants to regain birth weight Study dates July 2013 to April 2016 Source of funding No sources of funding reported	140ml/kg/day: 1070 (860-1280) <100g - n (%) 100ml/kg/day: 26 (39) 140ml/kg/day: 28 (40) <2nd centile - n (%) 100ml/kg/day: 7 (10) <10th centile - n (%) 100ml/kg/day: 21 (31) 140ml/kg/day: 19 (27) Head circumference (cm) at birth - mean (SD) 100ml/kg/day: 25.7 (2.4) 140ml/kg/day: 25.6 (3.8) Length (cm) at birth - mean (SD) 100ml/kg/day: 36.0 (10.9) 140ml/kg/day: 37.4 (12.4) Inclusion criteria VLBW (<1500g), preterm infants who had a peripherally inserted central venous catheter (PICC) and were receiving less than 100ml/kg/day of enteral feeds		necrotising enterocolitis (NEC). Preterm formula (80kcal/100ml) was given to appropriately grown infants >1250g if maternal EBM was not available by day 2 of life. If there were no contraindications, enteral feeds were advanced at 20-30ml/kg/day, as tolerated, and fortified when they reached 140ml/kg/day. Enteral feeds were discontinued if NEC was suspected and resumed at the same volume once NEC and infection were excluded. Power analysis: 140 patients (allowing for 10% mortality and 10% withdrawal from the trial) would be needed to detect a 2-day difference (defined as a clinically significant difference by the authors) in time taken to regain birth weight between groups. Statistical analysis: Conducted based on intention-to-treat. Normally distributed	Weight gain - weight <10th centile at discharge - n/N: 100ml/kg/day: 26/67 140ml/kg/day: 25/70 Linear growth - length (cm) at discharge - mean (SD): 100ml/kg/day (n=67): 44.7 (3.9) 140ml/kg/day (n=70): 43.8 (4.7) Linear growth - length <10th centile at discharge - n/N: 100ml/kg/day: 35/70 Head circumference (cm) at discharge - mean (SD): 100ml/kg/day (n=67): 32.0 (2.5) 140ml/kg/day (n=70): 32.1 (2.1) Head circumference <10th centile at discharge - n/N: 100ml/kg/day: 17/67 140ml/kg/day: 14/70	Blinding of outcome assessment: Low risk. Outcome assessors were not blind to treatment allocation but all outcomes are objective. Attrition bias Incomplete outcome data: Low risk. Only 2 infants in the intervention arm were withdrawn from the study; no infants in the intervention arm were withdrawn. Reporting bias Selective reporting: Low risk. All outcomes reported Other bias Other sources of bias: Low risk. None. Other information In both arms, there were 12 protocol violations where PICC was removed at a different time point than specified by the group allocation (median delay of 9.5 hours in intervention group [100ml/kg/day] and median 6.5 hours early in

Study details	Participants	Interventions	Methods	Outcomes and Results	Comments
	Exclusion criteria Major congenital anomaly; likely to need prolonged PN or intravenous antibiotics after achieving 100ml/kg/day of enteral feeds; death or transfer out of NICU likely to occur before birth weight could be regained		continuous variables were analysed using t-tests and dichotomous and categorical variables were analysed with chi-squared tests, Mann-Whitney U tests and Fisher's exact tests.	Mortality - n/N: 100ml/kg/day: 3/67 140ml/kg/day: 3/70 Duration (days) of hospital stay* - median (IQR): 100ml/kg/day: 49.6 (38.2-69.2) 140ml/kg/day: 53.9 (38.9-71.3) p=0.86 Necrotising enterocolitis - n/N: 100ml/kg/day: 1/67 140ml/kg/day: 3/70 Duration (days) of PN* - median (IQR): 100ml/kg/day: 5.6 (4.5-6.9) 140ml/kg/day: 5.6 (4.5-6.9) 140ml/kg/day: 6.1 (5.2-8.1) p=0.04 Sepsis - central venous catheter-associated late onset sepsis - n/N: 100ml/kg/day: 1/67 140ml/kg/day: 1/67 140ml/kg/day: 3/70	control group [140ml/kg/day]); results of per-protocol analysis were not significantly different from results of intention-to-treat analysis for primary outcome (time in days to regain birth weight).

Study details	Participants	Interventions	Methods	Outcomes and Results	Comments
				*Not analysed as data is reported as medians	

EBM: expressed breast milk; EN: enteral nutrition; IQR: interquartile range; NEC: necrotising enterocolitis; NICU: neonatal intensive care unit; N: number; PICC: peripherally inserted central venous catheter; PN: parenteral nutrition; ROBINS-I: risk of bias in non-randomised studies of interventions; SD: standard deviation; VLBW: very low brithweight.

1 Appendix E – Forest plots

- Forest plots for review question: What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?
- 5 No meta-analysis was conducted for this review; therefore there are no forest plots.

1 Appendix F – GRADE tables

- 2 GRADE tables for review question: What amount of enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and enteral feeds) indicates that parenteral nutrition is no longer required?
- 4 Table 5: Clinical evidence profile for stopping PN when EN reached 100ml/kg/day versus stopping PN when EN reached 140ml/kg/day

Quality	assessment						No of patient	s	Effect			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	100ml/kg/day	140ml/kg/day	Relative (95% CI)	Absolute	Quality	Importance
Neight	gain - time (d	days) to ı	regain birth wei	ght - All infant	s (VLBW & EL	BW) (Better ind	icated by lowe	er values)				
1	randomised trials		no serious inconsistency	no serious indirectness	serious ¹	none	67	70	-	MD 1.5 higher (0.31 to 2.69 higher)		CRITICAL
Weight	gain - time (d	days) to i	regain birth wei	ght - VLBW inf	ants (Better i	ndicated by lowe	er values)					
1	randomised trials		no serious inconsistency	no serious indirectness	serious ²	none	41	42	-	MD 0.6 higher (0.8 lower to 2 higher)		CRITICAL
Weight	gain - time (d	days) to i	regain birth wei	ght - ELBW inf	ants (Better i	ndicated by lowe	er values)					
1	randomised trials		no serious inconsistency	no serious indirectness	serious ³	none	26	28	-	MD 2.8 higher (0.8 to 4.8 higher)	⊕⊕⊕O MODERATE	CRITICAL
Weight	gain - weigh	t (g) at di	scharge (Better	indicated by I	nigher values)						
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	67	70	-	MD 13 higher (191.05 lower to 217.05 higher)		CRITICAL

Quality a	assessment						No of patient	S	Effect			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	100ml/kg/day	140ml/kg/day	Relative (95% CI)	Absolute	Quality	Importance
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ⁴	none	26/67 (38.8%)	25/70 (35.7%)	RR 1.09 (0.7 to 1.68)	32 more per 1000 (from 107 fewer to 243 more)	⊕⊕OO LOW	CRITICAL
Linear g	rowth - leng	th (cm) a	nt discharge (Be	tter indicated	by higher val	ues)						
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	67	70	-	MD 0.9 higher (0.54 lower to 2.34 higher)	⊕⊕⊕⊕ HIGH	CRITICAL
Linear g	rowth - leng	th <10th	centile at disch	arge								
1	randomised trials		no serious inconsistency	no serious indirectness	very serious ⁴	none	36/67 (53.7%)	35/70 (50%)	RR 1.07 (0.78 to 1.48)	35 more per 1000 (from 110 fewer to 240 more)	⊕⊕OO LOW	CRITICAL
Head ci	rcumference	(cm) at	discharge (Bette	er indicated by	/ higher value	s)						
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	67	70	-	MD 0.1 lower (0.87 lower to 0.67 higher)		CRITICAL
Head cir	cumference	<10th ce	entile at dischar	ge								
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ⁴	none	17/67 (25.4%)	14/70 (20%)	RR 1.27 (0.68 to 2.37)	54 more per 1000 (from 64 fewer to 274 more)		CRITICAL
Mortalit	у											
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ⁴	none	3/67 (4.5%)	3/70 (4.3%)		2 more per 1000 (from 33 fewer to 171 more)	⊕⊕OO LOW	IMPORTANT
Necrotis	sing enteroc	olitis										

Quality assessment No of patients Effect											
No of studies		Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	100ml/kg/day	Relative (95% CI)	Absolute	Quality	Importance
1		no serious risk of bias		no serious indirectness	very serious ⁴	none	1/67 (1.5%)	RR 0.35 (0.04 to 3.27)	28 fewer per 1000 (from 41 fewer to 97 more)	⊕⊕OO LOW	IMPORTANT
Central	venous cath	eter-asso	ociated late ons	et sepsis							
1		no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ⁴	none	1/67 (1.5%)	RR 0.35 (0.04 to 3.27)	28 fewer per 1000 (from 41 fewer to 97 more)		IMPORTANT

CI: confidence interval; ELBW: extremely low birthweight; MD: mean difference; RR: risk ratio; VLBW: very low birthweight.

Table 6: Clinical evidence profile for stopping PN when EN reached 100ml/kg/day versus stopping PN when EN reached 120ml/kg/day

			•	e e e e e e e e e e e e e e e e e e e			angrady re	.,				
Quality assessment							No of patients Eff		Effect			
No of studies	Design	Risk of bias	Inconsistency	Indirectness		Other considerations	100ml/kg/day	120ml/ka/day	Relative (95% CI)	Absolute	Quality	Importance
Weight (gain - time (da	ys) to re	gain birth weigh	t (Better indic	ated by lower	values)						
1	observational studies		no serious inconsistency	no serious indirectness	serious ²	none	50	37		MD 3.4 lower (6.45 to 0.35 lower)	⊕OOO VERY LOW	CRITICAL
Weight 9	gain - weight <	<10th cer	ntile - At 36 weel	ks' post conce	ption							
1	observational studies		no serious inconsistency	no serious indirectness	serious ³	none	33/50 (66%)		RR 0.72 (0.58 to 0.9)	257 fewer per 1000 (from 92	0000	CRITICAL

¹ Evidence was downgraded by 1 due to serious imprecision, 95% confidence interval crosses one default MID for continuous outcomes, calculated as 0.5 x SD control at baseline (3.5).

² Evidence was downgraded by 1 due to serious imprecision, 95% confidence interval crosses one default MID for continuous outcomes, calculated as 0.5 x SD control at baseline (3.0).

³ Evidence was downgraded by 1 due to serious imprecision, 95% confidence interval crosses one default MID for continuous outcomes, calculated as 0.5 x SD control at baseline (3.8).

⁴ Evidence was downgraded by 2 due to very serious imprecision, 95% confidence interval crosses two default MID for dichotomous outcomes (0.8 and 1.25).

										fewer to 386 fewer)		
Weight	gain - weight <	10th cer	ntile - At 40 wee	ks' post conce	eption							
1	observational studies	serious ¹	no serious inconsistency	no serious indirectness	serious ³	none	22/50 (44%)	29/37 (78.4%)	RR 0.56 (0.39 to 0.8)	345 fewer per 1000 (from 157 fewer to 478 fewer)	⊕OOO VERY LOW	CRITICAL
Linear	growth - length	<10th c	entile - At 36 we	eks' post con	ception							
1	observational studies		inconsistency	no serious indirectness	serious ³	none	31/50 (62%)	30/37 (81.1%)	RR 0.76 (0.59 to 1)	195 fewer per 1000 (from 332 fewer to 0 more)	VERY	CRITICAL
Linear	growth - length	<10th c	entile - At 40 we	eeks' post con	ception							
1	observational studies	serious ¹	no serious inconsistency	no serious indirectness	serious ³	none	34/50 (68%)	32/37 (86.5%)	RR 0.79 (0.63 to 0.99)	182 fewer per 1000 (from 9 fewer to 320 fewer)	⊕OOO VERY LOW	CRITICAL
Head ci	ircumference <	10th cen	tile - At 36 wee	ks' post conce	ption							
1	observational studies	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	21/50 (42%)	29/37 (78.4%)	RR 0.54 (0.37 to 0.77)	361 fewer per 1000 (from 180 fewer to 494 fewer)	⊕OOO VERY LOW	CRITICAL
Head ci	ircumference <	10th cen	tile - At 40 weel	ks' post conce	ption							
1	observational studies	serious ¹	no serious inconsistency	no serious indirectness	serious ³	none	21/50 (42%)	26/37 (70.3%)	RR 0.6 (0.41 to 0.88)	281 fewer per 1000 (from 84 fewer to 415 fewer)		CRITICAL
Duratio	n (days) of hos	pital sta	y (Better indica	ted by lower v	alues)							
1	observational studies	serious ¹	no serious inconsistency	no serious indirectness	serious ⁴	none	50	37	-	MD 7.9 lower (19.1 lower to 3.3 higher)		IMPORTANT
Necroti	sing enterocoli	tis										
1	observational studies	serious ¹	no serious inconsistency	no serious indirectness	serious ⁶	none	2/50 (4%)	0/37 (0%)	Peto OR 5.82 (0.35 to 97.53)	-	⊕OOO VERY LOW	IMPORTANT
Duratio	n (days) of PN	(Better i	ndicated by low	ver values)								

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1	observational studies	serious ¹	no serious indirectness	serious ⁶	none	50	37	-	MD 3.5 lower (9.88 lower to 2.88 higher)	0000	IMPORTANT
Sepsis											
1	observational studies	serious ¹	no serious indirectness	very serious ⁵	none	17/50 (34%)	16/37 (43.2%)	(0.46 to	91 fewer per 1000 (from 234 fewer to 147 more)	VERY	IMPORTANT

CI: confidence interval; MD: mean difference; OR: odds ratio; RR: risk ratio.

Table 7: Clinical evidence profile for stopping PN when EN reached 60kcal/kg/day versus stopping PN when EN reached 100kcal/kg/day

	TOOKCair	itg, aa j							1			
Quality assessment No of pat							No of patien	nts Effect				
No of studies		Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	60kcal/kg/day	100kcal/kg/day	Relative (95% CI)	Absolute	Quality	Importance
Weight 9	gain - postnata	al growth	failure at 40 wo	eeks' postmen	strual age							
1	observational studies		no serious inconsistency	no serious indirectness	serious ²	none	50/65 (76.9%)	62/117 (53%)	RR 1.45 (1.17 to 1.8)	238 more per 1000 (from 90 more to 424 more)	0000	CRITICAL
Linear g	rowth - length	(cm) at	40 weeks' postr	menstrual age	(Better indica	ated by higher v	alues)					
1	observational studies		no serious inconsistency	no serious indirectness	serious ³	none	65	117	-	MD 1 lower (1.81 to 0.19 lower)	⊕OOO VERY LOW	CRITICAL
Head cir	Head circumference (cm) at 40 weeks' postmenstrual age (Better indicated by higher values)											

¹ Evidence downgraded by 1 due to moderate risk of selection bias (as timeframe for the 2 cohorts differs) and moderate risk of reporting bias as some outcomes are not reported sufficiently to be included in analysis.

² Evidence was downgraded by 1 due to serious imprecision, 95% confidence interval crosses one default MID for continuous outcomes, calculated as 0.5 x SD control at baseline (8.3).

³ Evidence was downgraded by 1 due to serious imprecision, 95% confidence interval crosses one default MID for dichotomous outcomes (0.8 or 1.25).

⁴ Evidence was downgraded by 1 due to serious imprecision, 95% confidence interval crosses one default MID for continuous outcomes, calculated as 0.5 x SD control at baseline (29.5).

⁵ Evidence was downgraded by 2 due to very serious imprecision, 95% confidence interval crosses two default MID for dichotomous outcomes (0.8 and 1.25).⁶ Evidence was downgraded by 1 due to serious imprecision, 95% confidence interval crosses 1 default MID for continuous outcomes, calculated as 0.5 x SD control at baseline (15.5).

⁶ Evidence was downgraded for risk of imprecision due to low event rate

Quality	Quality assessment							No of patients				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	60kcal/kg/day	100kcal/kg/day	Relative (95% CI)	Absolute	Quality	Importance
1	observational studies	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	65	117	-	MD 0.9 lower (0.96 to 0.84 lower)	⊕OOO VERY LOW	CRITICAL
Necrotiz	ing enterocol	itis										
1	observational studies	serious ¹	no serious inconsistency	no serious indirectness	very serious ⁴	none	3/65 (4.6%)	2/117 (1.7%)	RR 2.7 (0.46 to 15.75)	29 more per 1000 (from 9 fewer to 252 more)	⊕OOO VERY LOW	IMPORTANT
Late ons	set sepsis											
1	observational studies	serious ¹	no serious inconsistency	no serious indirectness	very serious ⁴	none	15/65 (23.1%)	30/117 (25.6%)	RR 0.9 (0.52 to 1.55)	26 fewer per 1000 (from 123 fewer to 141 more)	⊕OOO VERY LOW	IMPORTANT

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

¹ Evidence downgraded by 1 due to moderate risk of selection bias (as timeframe for the 2 cohorts differs).

² Evidence was downgraded by 1 due to serious imprecision, 95% confidence interval crosses one default MID for dichotomous outcomes (0.8 or 1.25).

³ Evidence was downgraded by 1 due to serious imprecision, 95% confidence interval crosses one default MID for continuous outcomes, calculated as 0.5 x SD control at baseline (1.3).

⁴ Evidence was downgraded by 2 due to very serious imprecision, 95% confidence interval crosses both default MID for dichotomous outcomes (0.8 and 1.25).

1 Appendix G - Economic evidence study selection

- 2 Economic evidence study selection for review question: What amount of enteral
- 3 feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and
- 4 enteral feeds) indicates that parenteral nutrition is no longer required?
- 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 tables for review question: What amount of enteral feed
- 3 (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and
- 4 enteral feeds) indicates that parenteral nutrition is no longer required?
- 5 No economic studies were identified which was applicable to this review question.

1 Appendix I – Economic evidence profiles

- 2 Economic evidence profiles for review question: What amount of enteral feed
- 3 (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and
- 4 enteral feeds) indicates that parenteral nutrition is no longer required?
- 5 No economic studies were identified which was applicable to this review question.

1 Appendix J - Economic analysis

- 2 Economic evidence analysis for review question: What amount of enteral feed
- 3 (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and
- 4 enteral feeds) indicates that parenteral nutrition is no longer required?
- 5 No economic analysis was undertaken for this review question.

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1 Appendix K - Excluded studies

- 2 Excluded clinical and economic studies for review question: What amount of
- 3 enteral feed (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to
- 4 PN and enteral feeds) indicates that parenteral nutrition is no longer required?

5 Clinical studies

6 Table 8: Excluded studies and reasons for their exclusion

	INTO IN CACTURE OF THE PROPERTY OF THE PROPERT
Study	Reason for Exclusion
Als, Heidelise, Gilkerson, Linda, Duffy, Frank H., McAnulty, Gloria B., Buehler, Deborah M., Vandenberg, Kathleen, Sweet, Nancy, Sell, Elsa, Parad, Richard B., Ringer, Steven A., Butler, Samantha C., Blickman, Johan G., Jones, Kenneth J., A three-center, randomized, controlled trial of individualized developmental care for very low birth weight preterm infants: medical, neurodevelopmental, parenting, and caregiving effects, Journal of developmental and behavioral pediatrics: JDBP, 24, 399-408, 2003	Intervention does not meet inclusion criteria - observing and classifying neonate behaviour
Bergsten, G., Aziz, K., Lau, G., Brown, K., Brunet, K., Larsen, B., Energy and protein intakes during the transition of parenteral to enteral nutrition, Paediatrics and Child Health, 16, 40A-41A, 2011	Conference abstract - insufficient information reported
Braudis, Nancy J., Curley, Martha A. Q., Beaupre, Karen, Thomas, Kristi C., Hardiman, Gina, Laussen, Peter, Gauvreau, Kimberlee, Thiagarajan, Ravi R., Enteral feeding algorithm for infants with hypoplastic left heart syndrome poststage I palliation, Pediatric critical care medicine: a journal of the Society of Critical Care Medicine and the World Federation of Pediatric Intensive and Critical Care Societies, 10, 460-6, 2009	Intervention does not meet inclusion criteria - feeding algorithm does not specify the volume or ratio of enteral feeds at which PN should be stopped
Brennan, A. M., Fenton, S., Murphy, B. P., Kiely, M. E., Transition Phase Nutrition Recommendations: A Missing Link in the Nutrition Management of Preterm Infants, Journal of Parenteral and Enteral Nutrition, 42, 343-351, 2018	Intervention does not meet inclusion criteria - classification of nutritional requirements according to the nutritional phase of the neonate compared with chronological age
Butler, T. J., Szekely, L. J., Grow, J. L., A standardized nutrition approach for very low birth weight neonates improves outcomes, reduces cost and is not associated with increased rates of necrotizing enterocolitis, sepsis or mortality, Journal of Perinatology, 33, 851-7, 2013	Comparison does not meet inclusion criteria - stopping point for PN was at the discretion of neonatologist (and is not reported)
Camacho, J. E., Santos Oren, M., DuPont, T., Stefanescu, A., Stefanescu, B., Evaluation of a nutrition protocol on infectious and growth	Conference abstract - insufficient information reported

Charles	December Fredrick
Study	Reason for Exclusion
outcomes, Journal of Investigative Medicine, 66, 176-177, 2018	
Chetry, S., Kler, N., Saluja, S., Soni, A., Feasibilty of early total enteral feeds in very low birthweight infants, Journal of Paediatrics and Child Health, 48, 48, 2012	Conference abstract - insufficient information reported
Christmann, V., Visser, R., Engelkes, M., De Grauw, A. M., Van Goudoever, J. B., Van Heijst, A. F. J., The enigma to achieve normal postnatal growth in preterm infants - Using parenteral or enteral nutrition?, Acta Paediatrica, International Journal of Paediatrics, 102, 471-479, 2013	Intervention does not meet inclusion criteria - nutrition protocol does not specify the volume or ratio of enteral feeds at which PN should be stopped
Collins, Carmel T., Chua, Mei Chien, Rajadurai, Victor S., McPhee, Andrew J., Miller, Lisa N., Gibson, Robert A., Makrides, Maria, Higher protein and energy intake is associated with increased weight gain in pre-term infants, Journal of Paediatrics and Child Health, 46, 96-102, 2010	Intervention does not meet inclusion criteria - feeding approach does not specify the volume or ratio of enteral feeds at which PN should be stopped
Cordero Gonzalez, G., Maynez Gonzalez, C. G., Echaniz-Aviles, M. O., Carrera Muinos, S., Yllescas Medrano, E., Corral Kassian, E., Fernandez Carrocera, L. A., Aggressive parenteral nutrition and mean growth rate in newborns less than 1500 g in a 3rd level hospital in Mexico City, Perinatologia y Reproduccion Humana, 32, 54-59, 2018	Full text not written in English
Culpepper, Christine, Hendrickson, Kendra, Marshall, Susan, Benes, Jessica, Grover, Theresa R., Implementation of Feeding Guidelines Hastens the Time to Initiation of Enteral Feeds and Improves Growth Velocity in Very Low Birth-Weight Infants, Advances in neonatal care: official journal of the National Association of Neonatal Nurses, 17, 139-145, 2017	Intervention does not meet inclusion criteria - feeding guideline does not specify the volume or ratio of enteral feeds at which PN should be stopped
Donovan, Ramona, Puppala, Bhagya, Angst, Denise, Coyle, Bryan W., Outcomes of early nutrition support in extremely low-birth-weight infants, Nutrition in clinical practice: official publication of the American Society for Parenteral and Enteral Nutrition, 21, 395-400, 2006	Comparison does not meet inclusion criteria - does not specify the volume or ratio of enteral feeds at which PN should be stopped
Flidel-Rimon, O., Raz, M., Balla, U., Hofi, L., Juster-Reicher, A., Shinwell, E. S., Early, rapidly progressive enteral nutrition promotes growth of very low birth weight (VLBW) infants, Journal of Maternal-Fetal and Neonatal Medicine, 30, 1227-1231, 2017	Intervention does not meet inclusion criteria - quality improvement project does not specify the volume or ratio of enteral feeds at which PN should be stopped
Garratt, J., Norman, S., Wong, H. L., Goh, S. H., Lam, A., Simmer, K., Nathan, L., McLeod, G., An audit to assess the efficacy of changes to nutritional practice on nutrition and growth	Conference abstract - insufficient information reported

Childre	December Evolucion
Study outcomes in an Australian tertiary neonatal unit,	Reason for Exclusion
Journal of Paediatrics and Child Health, 51, 15, 2015	
Hanson, Corrine, Sundermeier, Julie, Dugick, Laura, Lyden, Elizabeth, Anderson-Berry, Ann L., Implementation, process, and outcomes of nutrition best practices for infants <1500 g, Nutrition in clinical practice: official publication of the American Society for Parenteral and Enteral Nutrition, 26, 614-24, 2011	Intervention does not meet inclusion criteria - nutrition regimen does not specify the volume or ratio of enteral feeds at which PN should be stopped (differences between arms are rate of advancement of PN and initiation and fortification of enteral feeds)
Hwang, J. Y., Shastry, P., Cayabyab, R., Ramanathan, R., Bhopal, N. S., Morley, E., Dang, T., Chang, M., Mihalek, A., Lin, E., Chavez, T., Luu, M., Garingo, A., Nair, S., Chin, S., Lin, T., Evaluation of a standardised feeding advancement guideline in preterm neonates less than 32 weeks' gestation, Journal of Investigative Medicine, 66 (1), 177, 2018	Conference abstract - insufficient information
Izquierdo, Montserrat, Martinez-Monseny, Antonio Federico, Pociello, Neus, Gonzalez, Paloma, Del Rio, Ruth, Iriondo, Martin, Iglesias- Platas, Isabel, Changes in Parenteral Nutrition During the First Week of Life Influence Early but Not Late Postnatal Growth in Very Low-Birth- Weight Infants, Nutrition in clinical practice: official publication of the American Society for Parenteral and Enteral Nutrition, 31, 666-72, 2016	Intervention does not meet inclusion criteria - nutrition protocol does not specify the volume or ratio of enteral feeds at which PN should be stopped
Johnson, Mark J., Leaf, Alison A., Pearson, Freya, Clark, Howard W., Dimitrov, Borislav D., Pope, Catherine, May, Carl R., Successfully implementing and embedding guidelines to improve the nutrition and growth of preterm infants in neonatal intensive care: a prospective interventional study, BMJ open, 7, e017727, 2017	Comparison does not meet inclusion criteria - does not specify the volume or ratio of enteral feeds at which PN should be stopped
Khanam, Siraj, Khan, Jafar, Sharma, Deepak, Chawla, Deepak, Murki, Srinivas, Nutritional bundle to improve growth outcomes among very low birth weight infants, The journal of maternal-fetal & neonatal medicine: the official journal of the European Association of Perinatal Medicine, the Federation of Asia and Oceania Perinatal Societies, the International Society of Perinatal Obstetricians, 28, 1851-5, 2015	Intervention does not meet inclusion criteria - PN was stopped at the same volume of enteral feeds in both the intervention and control arms (differences between arms are initiation of PN, quantity of amino acids and lipids, and fortification of enteral feeds)
Kuzma-O'Reilly, Barbara, Duenas, Maria L., Greecher, Coleen, Kimberlin, Lois, Mujsce, Dennis, Miller, Debra, Walker, Donna Jean, Evaluation, development, and implementation of potentially better practices in neonatal intensive care nutrition, Pediatrics, 111, e461-70, 2003	Intervention does not meet inclusion criteria - does not specify the volume or ratio of enteral feeds at which PN should be stopped. Further, description of baseline nutrition practices is insufficient
Kwok, T. C., Dorling, Jon, Ojha, Shalini, Multicentre prospective observational study of	Study design does not meet inclusion criteria - non-comparative

Charles	December Evaluation
Study	Reason for Exclusion
feeding practices in 30-33 weeks preterm infants, BMJ paediatrics open, 1, e000040, 2017	
Kwok, T. C., Ojha, S., Dorling, J., Early enteral feeding practice in 29+0 to 32+6 weeks preterm infants in two tertiary neonatal units in United Kingdom, European Journal of Pediatrics, 175, 1642, 2016	Conference abstract - full text identified
Lapointe, M., Barrington, K. J., Savaria, M., Janvier, A., Preventing postnatal growth restriction in infants with birthweight less than 1300 g, Acta Paediatrica, 105, e54-9, 2016	Intervention does not meet inclusion criteria - protocol does not specify the volume or ratio of enteral feeds at which PN should be stopped
Low, C. S., Ho, J. J., Nallusamy, R., Impact on growth of a nutrition policy to improve early oral and parenteral nutrition for preterm infants, Journal of Paediatrics and Child Health, 50, 88, 2014	Conference abstract - full text identified
Low, Chuen Siang, Ho, Jacqueline J., Nallusamy, Revathy, Impact of a new aggressive nutrition policy incorporating early introduction of parenteral nutrition and mother's own milk on growth of preterm infants, World journal of pediatrics: WJP, 12, 450-454, 2016	Intervention does not meet inclusion criteria - policy does not specify the volume or ratio of enteral feeds at which PN should be stopped
McCallie, K. R., Lee, H. C., Mayer, O., Cohen, R. S., Hintz, S. R., Rhine, W. D., Improved outcomes with a standardized feeding protocol for very low birth weight infants, Journal of perinatology: official journal of the California Perinatal Association, 31 Suppl 1, S61-7, 2011	Intervention does not meet inclusion criteria - protocol does not specify the volume or ratio of enteral feeds at which PN should be stopped
Miller, Malki, Donda, Keyur, Bhutada, Alok, Rastogi, Deepa, Rastogi, Shantanu, Transitioning Preterm Infants from Parenteral Nutrition: A Comparison of 2 Protocols, JPEN. Journal of parenteral and enteral nutrition, 41, 1371-1379, 2017	Intervention does not meet inclusion criteria - PN was stopped at the same volume of enteral feeds in both the intervention and control arms. Although the nutritional intake differs between arms as the PN is concentrated in the intervention arm, this is not clearly expressed in terms of a stopping point(differences between arms are volume of PN used during transition, use of dextrose solutions and calorie content of formula if breast milk was unavailable)
Nangia, S., Bishnoi, A., Goel, A., Manda, P., Tiwari, S., Saili, A., Early total enteral feeding in stable very low birth weight infants: A before and after study, Journal of Tropical Pediatrics, 64, 24-30, 2018	Intervention does not meet inclusion criteria - no PN was given in the intervention arm (total enteral feeding started on day 1 of life with no PN)
Narogan, M., Ryumina, I., Grosheva, E., Feeding of very preterm infants: The results application of modern standardized approaches in the practices, Journal of Pediatric Gastroenterology and Nutrition, 62, 859-860, 2016	Conference abstract - insufficient information reported
Olsen, Irene E., Richardson, Douglas K., Schmid, Christopher H., Ausman, Lynne M., Dwyer, Johanna T., Intersite differences in	Insufficient information reported about when PN was stopped at different study sites

Study	Reason for Exclusion
weight growth velocity of extremely premature infants, Pediatrics, 110, 1125-32, 2002	
Olsen, Steven L., Park, Nesha D., Tracy, Kelly, Younger, Darian, Anderson, Betsi, Implementing Standardized Feeding Guidelines, Challenges, and Results, Neonatal network: NN, 37, 218-223, 2018	Comparison does not meet inclusion criteria - stopping point for PN before the standardised guideline was introduced is not specified
Passaro, R. Colby, Savoie, Kate B., Huang, Eunice Y., Use of a Gastroschisis Feeding Guideline to Improve Standardization of Care and Patient Outcomes at an Urban Children's Hospital, Nutrition in clinical practice: official publication of the American Society for Parenteral and Enteral Nutrition, 33, 545-552, 2018	Intervention does not meet inclusion criteria- guideline does not specify stopping point for PN
Perrem, L. M., Semberova, J., O'Sullivan, A., Kieran, E. A., O'Donnell, C. P. F., White, M. J., Miletin, J., Randomised controlled trial comparing PICC removal and parenteral nutrition discontinuation at 100mls/kg/day versus 140mls/kg/day enteral feeding in preterm infants less than 1500g, European Journal of Pediatrics, 175, 1517, 2016	Abstract only - full text has been included
Robinson, J., Gupta, R., Surcouf, J., Knecht, M., Improving preterm infant clinical outcomes through implementation of a feeding protocol, Journal of Investigative Medicine, 67, 529-530, 2019	Abstract only - insufficient information available about feeding protocol.
Rochow, Niels, Fusch, Gerhard, Muhlinghaus, Alexandra, Niesytto, Christian, Straube, Sebastian, Utzig, Norbert, Fusch, Christoph, A nutritional program to improve outcome of very low birth weight infants, Clinical nutrition (Edinburgh, Scotland), 31, 124-31, 2012	Intervention does not meet inclusion criteria - nutrition program does not specify the volume or ratio of enteral feeds at which PN should be stopped
Savoie, Kate B., Bachier-Rodriguez, Marielena, Jones, Tamekia L., Jeffreys, Kristen, Papraniku, Dita, Sevilla, Wednesday Marie A., Tillman, Emma, Huang, Eunice Y., Standardization of Feeding Advancement After Neonatal Gastrointestinal Surgery: Does It Improve Outcomes?, Nutrition in clinical practice: official publication of the American Society for Parenteral and Enteral Nutrition, 31, 810-818, 2016	Intervention does not meet inclusion criteria - feeding advancement strategy does not specify the volume or ratio of enteral feeds at which PN should be stopped
Shim, S. Y., Ahn, H. M., Cho, S. J., Park, E. A., Early aggressive nutrition enhances language development in very low-birthweight infants, Pediatrics International, 56, 845-850, 2014	Intervention does not meet inclusion criteria - PN was stopped at same volume of enteral feeds in both the intervention and control arms
Shinnick, Julia K., Wang, Elizabeth, Hulbert, Cheryl, McCracken, Courtney, Sarson, Gail Yvonne, Piazza, Anthony, Karpen, Heidi, Durham, Megan M., Effects of a Breast Milk Diet on Enteral Feeding Outcomes of Neonates with	Intervention does not meet inclusion criteria - comparison between babies who received 100% of diet from breast milk, between 50 and 100% of diet from breast milk and less than 50% of diet from breast milk. It is unclear if the

Study	Reason for Exclusion
Gastrointestinal Disorders, Breastfeeding medicine: the official journal of the Academy of Breastfeeding Medicine, 2016	remaining percentage only refers to enteral feeds (e.g., formula) or includes parental nutrition
Shores, D. R., Bullard, J. E., Aucott, S. W., Stewart, F. D., Haney, C., Tymann, H., Miller, M. R., Nonyane, B. A. S., Schwarz, K. B., Implementation of feeding guidelines in infants at risk of intestinal failure, Journal of perinatology: official journal of the California Perinatal Association, 35, 941-8, 2015	Intervention does not meet inclusion criteria - feeding guideline does not specify the volume or ratio of enteral feeds at which PN should be stopped
Snyder, R., Allykas, S., Mennonna, A., Rogido, M. R., Nutrition of the extremely low birth weight (ELBW) infants: Are we making a difference?, Archives of Disease in Childhood, 97, A398, 2012	Conference abstract - insufficient information reported
Stefanescu, Beatrice M., Gillam-Krakauer, Maria, Stefanescu, Andrei R., Markham, Melinda, Kosinski, Jennifer L., Very low birth weight infant care: adherence to a new nutrition protocol improves growth outcomes and reduces infectious risk, Early Human Development, 94, 25-30, 2016	Comparison does not meet inclusion criteria - does not specify the volume or ratio of enteral feeds at which PN should be stopped
Stevens, Timothy P., Shields, Eileen, Campbell, Deborah, Combs, Adriann, Horgan, Michael, La Gamma, Edmund F., Xiong, KuangNan, Kacica, Marilyn, Variation in Enteral Feeding Practices and Growth Outcomes among Very Premature Infants: A Report from the New York State Perinatal Quality Collaborative, American Journal of Perinatology, 33, 9-19, 2016	Insufficient presentation of results - not reported separately based on stopping PN at different volumes or ratios of enteral feeds
Street, Jennifer L., Montgomery, Dianne, Alder, Stephen C., Lambert, Diane K., Gerstmann, Dale R., Christensen, Robert D., Implementing feeding guidelines for NICU patients<2000 g results in less variability in nutrition outcomes, JPEN. Journal of parenteral and enteral nutrition, 30, 515-8, 2006	Intervention does not meet inclusion criteria - feeding guideline does not specify the volume or ratio of enteral feeds at which PN should be stopped
Thoene, Melissa K., Lyden, Elizabeth, Anderson-Berry, Ann, Improving Nutrition Outcomes for Infants < 1500 Grams with a Progressive, Evidenced-Based Enteral Feeding Protocol, Nutrition in clinical practice: official publication of the American Society for Parenteral and Enteral Nutrition, 2018	Intervention does not meet inclusion criteria - feeding protocol does not specify the volume or ratio of enteral feeds at which PN should be stopped
Tillman, Emma M., Norman, Johanna L., Huang, Eunice Y., Lazar, Linda F., Crill, Catherine M., Evaluation of parenteral nutrition-associated liver disease in infants with necrotizing enterocolitis before and after the implementation of feeding guidelines, Nutrition in clinical practice: official publication of the American Society for Parenteral and Enteral Nutrition, 29, 234-7, 2014	Intervention does not meet inclusion criteria - feeding guideline does not specify the volume or ratio of enteral feeds at which PN should be stopped

Study	Reason for Exclusion
Tottman, A. C., Bloomfield, F. H., Cormack, B. E., Harding, J. E., Mohd Slim, M. A., Weston, A. F., Alsweiler, J. M., Relationships between Early Nutrition and Blood Glucose Concentrations in Very Preterm Infants, Journal of Pediatric Gastroenterology and Nutrition, 66, 960-966, 2018	Intervention does not meet inclusion criteria - nutrition protocol does not specify the volume or ratio of enteral feeds at which PN should be stopped
Westin, Vera, Klevebro, Susanna, Domellof, Magnus, Vanpee, Mireille, Hallberg, Boubou, Stoltz Sjostrom, Elisabeth, Improved nutrition for extremely preterm infants - A population based observational study, Clinical nutrition ESPEN, 23, 245-251, 2018	Intervention does not meet inclusion criteria - specifies the ratio of enteral feeds to parental nutrition at which lipid infusion should be stopped, but does not specify the volume or ratio of enteral feeds at which all PN should be stopped
Wilson, D. C., Cairns, P., Halliday, H. L., Reid, M., McClure, G., Dodge, J. A., Randomised controlled trial of an aggressive nutritional regimen in sick very low birthweight infants, Archives of disease in childhood. Fetal and neonatal edition, 77, F4-11, 1997	Intervention does not meet inclusion criteria - does not specify the volume or ratio of enteral feeds at which PN should be stopped. The control arm specifies the ratio of enteral feeds to parental nutrition at which PN amino acids and lipids should be stopped, but not when all PN should be stopped

1 Economic studies

- 2 No economic evidence was identified for this review. See supplementary material D for
- 3 further information.

4

1 Appendix L - Research recommendations

- 2 Research recommendations for review question: What amount of enteral feed
- 3 (measured in terms of ml/kg/d, or kcal/kg/d, or ratios in relation to PN and
- 4 enteral feeds) indicates that parenteral nutrition is no longer required?
- 5 No research recommendations were made for this review question.