



## Donor breast milk versus infant formula for preterm infants: systematic review and meta-analysis

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## ORIGINAL ARTICLE

## Donor breast milk versus infant formula for preterm infants: systematic review and meta-analysis

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A supplementary appendix is available at <http://adc.bmj.com/supplemental>

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**Objectives:** To compare the effect of donor breast milk with infant formula in preterm infants. Separate comparisons with formula were made for donor breast milk that was: (1) given as a sole diet; (2) given as a supplement to mother's own breast milk; and (3) fortified with macronutrients and micronutrients. The main outcomes were death, necrotising enterocolitis (NEC), infection, growth and development.

**Data sources:** Electronic databases—Cochrane, CENTRAL, MEDLINE, EMBASE, CINAHL, and HMIC: DH.

**Review methods:** Systematic review and meta-analysis of trials and observational studies of preterm or low birthweight infants.

**Results:** Seven studies (including five randomised controlled trials), all from the 1970s and 1980s, fulfilled the inclusion criteria. All studies compared the effect of sole donor breast milk with formula (combined  $n = 471$ ). One of these also compared the effect of donor breast milk with formula given as a supplement to mother's own milk ( $n = 343$ ). No studies examined fortified donor breast milk. A meta-analysis based on three studies found a lower risk of NEC in infants receiving donor breast milk compared with formula (combined RR 0.21, 95% CI 0.06 to 0.76). Donor breast milk was associated with slower growth in the early postnatal period, but its long-term effect is unclear.

**Conclusion:** Donor breast milk is associated with a lower risk of NEC and slower growth in the early postnatal period, but the quality of the evidence is limited. Further research is needed to confirm these findings and measure the effect of fortified or supplemented donor breast milk.

Breast milk is the recommended form of enteral nutrition for all infants, including those born preterm.<sup>1</sup> Donor breast milk is an alternative form of milk when the mother's own milk is not available or is in short supply. The advantages of breast milk over infant formulas include:

- the presence of active enzymes that enhance the maturation of the underdeveloped gut<sup>2 3</sup>;
- earlier tolerance of full enteral feeding;
- anti-infective properties which protect the newborn from infection.

Although the nutrient concentrations in preterm breast milk tend to be either the same as or higher than those in term breast milk, there is concern that breast milk, whether maternal or donated, may be inadequate to support the increased nutritional requirements of the preterm or very low birthweight infant.<sup>4 5</sup>

The use of donor breast milk varies across the world. Currently in the UK, 17 human milk banks supplying 50–60 neonatal units (Gillian Weaver, UK Association for Milk Banking, personal communication, 2004). In these banks all donor milk is frozen following Holder pasteurisation (heated to 62.5 °C for 30 min). However, this not only inactivates HIV, cytomegalovirus and other viruses, it also affects the nutritional and immunological properties of breast milk.<sup>3</sup> For example, it is estimated that 34% of the small amount of immunoglobulin G is destroyed, although most of the secretory immunoglobulin A remains (0–30% destroyed).<sup>2</sup> As donor milk is usually provided by women who deliver at term, and is pasteurised, it cannot be presumed that it will have the same effect as mother's own milk.

Although several systematic reviews have compared the effect of breast milk with infant formula in preterm infants,<sup>4–7</sup>

none has specifically focused on pasteurised donor breast milk or separated out the effects of donor breast milk given as a sole diet and that given as a supplement to mother's own milk. The aim of this systematic review, therefore, was to compare the effects of pasteurised donor breast milk and infant formula in preterm infants. We separated the effects of donor milk given as a sole diet, donor milk given as a supplement to mother's own milk, and donor milk that was fortified with macronutrients or micronutrients to encompass the current provision of donor milk in clinical practice. The main outcomes of interest were death, necrotising enterocolitis (NEC), infection, growth and development. We did not investigate whether pasteurisation adequately eliminates microbial contaminants.

## METHODS

### Search strategy

We carried out standard search strategies using the databases Cochrane, CENTRAL, MEDLINE, EMBASE, CINAHL and HMIC: DH from their inception to August 2004. Details of the search strategy can be found in the appendix (see <http://adc.bmj.com/supplemental>). There was no language restriction. There was no systematic attempt to search the "grey" literature although some articles were found in the references for indexed articles.

### Selection

We included studies on the basis of study design, population and comparison groups (box 1). All donor breast milk had to be donated from someone other than the infant's mother and it had to be pasteurised. All clinical outcomes were included.

**Abbreviation:** NEC, necrotising enterocolitis

**Box 1 Inclusion criteria****Study design**

- Randomised controlled trials (RCTs), quasi-RCTs and observational studies.

**Population**

- Preterm (<37 weeks' gestation) or low birthweight (<2500 g) infants.

**Types of intervention/exposure**

- Sole diet of pasteurised donor milk v sole diet of formula.
- Mother's own milk supplemented with pasteurised donor milk v mother's own milk supplemented with formula.
- Fortified, pasteurised donor milk v fortified formula.

**Outcome measures**

- All clinical outcomes were considered. The primary outcomes were necrotising enterocolitis and infection. All other short-term and long-term health and development outcomes including any data on adverse events were considered as secondary outcomes.
- Studies which only assessed the risk of transmissible diseases from milk banks were excluded.

**Data abstraction**

The articles identified by our search strategy were screened (title and abstract) by two independent reviewers (CB and MQ). Then those articles which potentially met the inclusion criteria were critically reviewed. The decision to include or exclude a specific article was made by consensus of the two reviewers. For two studies, information on whether the milk was pasteurised was not available in the published reports but was supplied by the authors of the primary studies.

**Validity assessment**

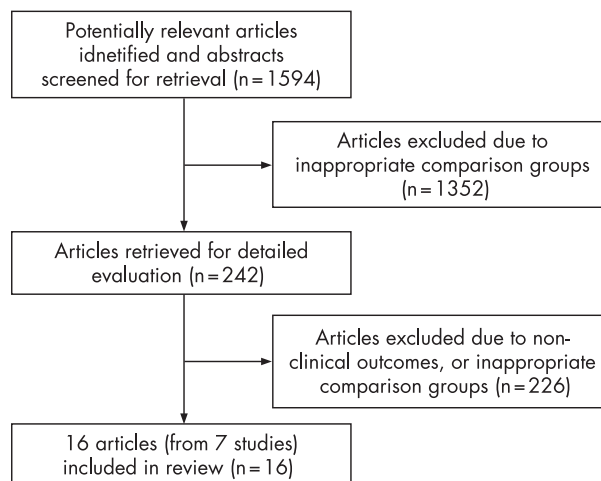
We assessed all studies for methodological quality in terms of sample size estimation, proportion followed up, blinding of outcome assessment, comparison of baseline variables and assessment of confounding. The RCTs were also assessed for method of randomisation, allocation concealment and blinding of intervention.

**Quantitative data synthesis**

Data were extracted and summarised into evidence tables, which included effect measures with 95% CIs. Where outcome measures were given in different units across studies, these were made consistent where possible. Meta-analyses were planned if there were sufficient data, and it was anticipated that a fixed-effects model would be used unless there was evidence of significant heterogeneity ( $p < 0.10$ ). We conducted and reported this review in accordance with the guidelines set out in the Quality Of Reporting Of Meta-analyses (QUOROM) statement.<sup>8</sup>

**RESULTS**

The search produced 1594 articles which were screened (title and abstract). Of these, 242 potentially addressed the research questions but on reading the articles only 16, arising from 7 studies, met the inclusion criteria (fig 1).



**Figure 1** Results of search and selection of articles (QUOROM statement flow diagram<sup>8</sup>)

**Study characteristics**

Table 1 provides details of the characteristics of the seven included studies.<sup>9–24</sup> The combined sample size of the studies which compared the sole diet of donor breast milk with sole formula was 471 infants. Three of these studies started in the 1970s (in Wales,<sup>9</sup> Finland<sup>10</sup> and Hungary<sup>11</sup>) and four in the 1980s (in the USA,<sup>12</sup> England,<sup>13–22</sup> South Africa<sup>23</sup> and France<sup>24</sup>). One of these studies<sup>13–22</sup> (Lucas *et al*<sup>13</sup> will be cited as the primary paper for this study) also compared donor milk with formula, both given as a supplement to mother's own milk. This study was conducted in England in the 1980s, with a sample size of 343.<sup>13</sup> We did not find any other studies comparing donor milk with formula as a supplement to mother's own milk. No studies were found comparing fortified donor milk with formula.

There was considerable variability between the studies in terms of the type of donor milk (preterm v term; drip v expressed) and formulas used, the feeding regimens and the period of intervention. The study population also varied in terms of mean gestation (range 30.2–33.2 weeks) and mean birth weight (range 1310–1954 g) (table 1).

**Methodological quality of studies**

The methodological quality of the studies is summarised in table 2. Five studies were RCTs, however, only one of these trials<sup>13</sup> reported that a sample size estimate had been calculated prior to the trial starting. This study also had adequate methodological quality in terms of allocation concealment and randomisation. None of the remaining four RCTs included a sample size estimate, and two did not specify the method of randomisation.<sup>9–11</sup> One of the studies did not specify whether it was randomised.<sup>24</sup> One study,<sup>23</sup> which was designed as a trial, stopped recruitment in the formula group due to an "outbreak" of NEC, and continued to recruit in the formula group from another hospital. We have only used outcome data from the first hospital.

**Findings of the included studies**

Tables 3–5 present the main results of the seven studies included in the review.

**Effect on mortality**

Mortality was reported in one study<sup>13</sup> for both sole diet and supplementary diet comparisons. This study found a lower risk of mortality at ages 9 months and 18 months in the donor milk group compared with the formula group, for both sole and

**Table 1** Summary of the characteristics of the seven included studies

Study Time period Design	Sample size	DBM group Type o DBM (n)	Formula group Type of formula (n)	Baseline data in study population*		Feeding regimen Route of delivery	Dose	Pre-allocation nutrition	When allocated feed started	Infants fed allocated diet until	Additional nourishment	Post diet regimen	
				Study inclusion criteria Birth weight (g)	Mean gestation (weeks)								Mean birth weight (g)
Davies <sup>9</sup> 1972-1973 RCT	68	Term (34)	PTF (34)	28-36*	32.9	1954	Not stated	50 ml/kg increased to 200 ml/kg or until satisfied	Not stated	2 months old	If diet was not tolerated infants received 10% dextrose IV. All infants received vitamin and iron supplements DBM group were given vitamins A, D and C and ferrous sulphate Not stated	N/A	
Reihe <i>et al</i> <sup>10</sup> 1972-1975 RCT	106	Term (22)	Not stated (84)	28-36	31-33	1680	Smallest infants initially fed by gavage, others teat	Formula: 150 ml/kg/day DBM: 170 ml/kg/day	Not stated	2400 g	Not stated	Not stated	
Schultz <i>et al</i> <sup>11</sup> 1980 RCT	20	"Mature" (10)	Not stated (10)	30-37*	33.2	1698	Not stated	Increased to 170 ml/kg/day by 1 week	None	Before 24 h (usually 6-12 h)	Not stated	Not stated	
Gross <sup>2</sup> 1980-1982 RCT	67	Preterm (20)	PTF (46)	27-33	30.9	1322	Gastric gavage	24 ml/kg/day increased to 180 ml/kg/day	Not stated	Day 1-6 (mean day 3)	1800 g	IV dextrose and electrolytes days 1-5 after birth, and 1 ml Poly Vi Sol from day 7; hypotraumatic babies received NaCl in their milk	All fed formula
Lucas A <i>et al</i> <sup>13,22</sup> 1982-1984 RCT	159	Term (21) "Mainly drip" (83)	PTF (76)	<18.50	31.3	1401	Nasogastric tube	Up to 200 ml/kg/24 h	In 3 centres all infants received DBM for first 24 h	Median 7 days	2000 g or discharge	In total (trials A and B) 90/502 babies required initial parenteral nutrition	Not stated
Lucas B <i>et al</i> <sup>13,22</sup> 1982-1984 RCT	343	"Mainly drip" (170)	PTF (173)	<18.50	31.1	1388	Nasogastric tube	Up to 200 ml/kg/24 h	In 3 centres all infants received DBM for first 24 h	Median 7 days	2000 g or discharge	See above	Not stated
Cooper <i>et al</i> <sup>23</sup> Early 1980s Observational†	39	Preterm (24)	PTF (15)	<36	1200-1500	1359	Nasogastric tube then bottle	Formula: 175 ml/kg/day DBM: 170-200 ml/kg/day	Not stated	When enteral feeding started	1800 g	Daily vitamin D.	N/A
Péret <i>et al</i> <sup>24</sup> 1984 Study design unclear	12	Not stated (6)	PTF (6)	"VLBW"	30.2	1310	Nasogastric gavage or teat	Formula: mean 157 ml/kg/day DBM: mean 168 ml/kg/day	Parenteral glucose and amino acid solution until oral feeding; all infants initially fed DBM within 24-48 hours until full diet had been reached	Study 1: once full oral feeding obtained	Not stated	Not stated	N/A

DBM, donor breast milk; IV, intravenous; N/A, not applicable as outcomes measured whilst on allocated diet; PTF, preterm formula; RCT, randomised controlled study; VLBW, very low birth weight.

\* "Drip" milk: the milk that drips from one breast while the infant feeds at the other breast.

† These were the characteristics of the study sample rather than specific inclusion criteria.

‡ Lucas A and Lucas B refer to the Lucas study trial A (sole diet comparison) and trial B (supplementary diet comparison), respectively.

§ Initially a quasi-RCT that became observational due to a change in selection criteria and population.

**Table 2** Methodological quality of the studies included in the review

Study	Sample size estimate	Method of randomisation	Allocation Concealment	Blinding of intervention	Proportion (%) included in analysis*	Blinding of outcome	Baseline variables compared	Assessment of confounding
Davies <sup>9</sup>	No	Randomised but method not stated	Not stated	Not stated	100%	Not stated	Yes	Stratification by gestational age
Raiha <i>et al</i> <sup>10</sup>	No	Blocked randomisation	Coding of bottles	Not stated	95%	Not stated	Yes	Stratification by gestational age
Schultz <i>et al</i> <sup>11</sup>	No	Randomised but method not stated	Not stated	Not stated	100%	Not stated	Yes	No
Gross <sup>12</sup>	No	Random number table	Not stated	Not stated	100%	Not stated	Yes	No
Lucas <i>et al</i> <sup>13-22</sup>	Yes	Blocked randomisation	Opaque, sealed, numbered envelopes	No	100% for short term outcomes 69% at 9 months 86% at 18 months 82% at 8 years	For some outcomes	Yes	For some outcomes
Cooper <i>et al</i> <sup>23</sup>	No	Not random (alternating assignment)	Not stated	Not stated	100%	Not stated	Yes	No
Putet <i>et al</i> <sup>24</sup>	No	Not stated (may not be randomised)	Not stated	Not stated	92%	Not stated	Yes	No

\*This is the percentage included in the analysis after excluding losses to follow-up (for which there were very few for the short-term outcomes) and those with missing values for some outcomes.

supplementary diet comparisons, but this effect was not statistically significant (table 3).

### Effect on NEC

In the sole diet comparison, NEC was reported in three studies.<sup>12 13 23</sup> In all of these studies, there was a lower risk of NEC in the donor milk group than the formula group but the effect was not statistically significant (table 3). For confirmed cases of NEC, the three studies yielded almost identical measures of effect (risk ratio (RR) 0.21–0.22) with similar 95% CIs (the largest study had a much lower overall risk of NEC and hence had a similar level of precision to the smallest study). In view of the homogeneity of these RRs, a fixed-effects meta-analysis and a random-effects meta-analysis gave identical results (combined RR 0.21, 95% CI 0.06 to 0.76,  $p = 0.017$ ). The combined evidence from these studies suggests that donor milk reduces the risk of NEC by about 79% (95% CI 24% to 94%) (fig 2).

The risk of confirmed NEC in the formula group in these three studies was 5.3%,<sup>13</sup> 11.5%<sup>12</sup> and 20.0%.<sup>23</sup> Each study yielded different estimates of the risk difference (RD) 4.1%, 95% CI  $-1.4\%$  to  $9.6\%$ <sup>13</sup>; 9.2% (95% CI  $-4.1\%$  to  $22.3\%$ )<sup>12</sup>; 15.5% (95% CI  $-5.9\%$  to  $37.6\%$ )<sup>23</sup>. The combined estimate of the RD, estimated from a random effects meta-analysis, was 5.4%

(95% CI 0.5% to 10.3%,  $p = 0.032$ ). This suggests that in settings in which the risk of NEC in the formula group is about 5–20%, approximately 18.5 preterm infants (95% CI 9.7 to 200) would need to be given donor milk to prevent one case of NEC.

In the supplementary diet comparison,<sup>13</sup> there were more suspected cases of NEC but fewer confirmed cases in the donor milk group than in the formula group but neither effect reached statistical significance (table 3).

### Effect on other perinatal outcomes

Four studies in the sole diet comparison reported other perinatal morbidity events. Three studies found significantly fewer episodes of feeding intolerance (including NEC) and diarrhoea in the donor milk group compared with the formula group (table 3); in one,<sup>13</sup> infants in the donor milk group were found to tolerate full enteral feeds earlier, another study<sup>12</sup> reported significantly fewer withdrawals due to feeding intolerance (within 9–31 days of starting the intervention) in the donor milk group, and yet another<sup>11</sup> found infants fed donor milk had significantly fewer episodes of mild diarrhoea in the first two weeks after birth. One study<sup>23</sup> reported more withdrawals due to respiratory symptoms within the first few days of starting the intervention in the donor milk group compared with the formula group (12.5% *v* 6.7%), but this

**Table 3** Comparisons of mortality, NEC, and other perinatal outcomes between the donor breast milk and formula groups

Outcome and studies	Sole diet comparison				RR (95% CI)	p value	Supplementary diet comparison				RR (95% CI)	p value
	Donor breast milk		Formula				Donor breast milk		Formula			
	No.	n (%)	No.	n (%)			No.	n (%)	No.	n (%)		
Mortality up to 9 months Lucas <i>et al</i> <sup>17</sup>	83	7 (8.4)	76	9 (11.8)	0.71 (0.28 to 1.82)	$p = 0.48$	170	12 (7.1)	173	15 (8.7)	0.81 (0.39 to 1.69)	$p = 0.58$
Mortality up to 18 months Lucas <i>et al</i> <sup>17 19</sup>	83	7 (8.4)	76	9 (11.8)	0.71 (0.28 to 1.82)	$p = 0.48$	170	13 (7.6)	173	16 (9.2)	0.83 (0.41 to 1.67)	$p = 0.59$
Confirmed NEC Cooper <i>et al</i> <sup>23</sup>	24	1 (4.2)	15	3 (20.0)	0.21 (0.02 to 1.82)	$p = 0.113$						
Gross <sup>12</sup>	41	1 (2.4)	26	3 (11.5)	0.21 (0.02 to 1.92)	$p = 0.126$						
Lucas and Cole <sup>18</sup>	86	1 (1.2)	76	4 (5.3)	0.22 (0.03 to 1.93)	$p = 0.13$	167	2 (1.2)	173	5 (2.9)	0.41 (0.08 to 2.11)	$p = 0.27$
Suspected NEC Lucas and Cole <sup>18</sup>	86	3 (3.5)	76	6 (7.9)	0.44 (0.11 to 1.71)	$p = 0.22$	167	8 (4.8)	173	6 (3.5)	1.38 (0.49 to 3.90)	$p = 0.54$
Respiratory symptoms Cooper <i>et al</i> <sup>23</sup>	24	3 (12.5)	15	1 (6.7)	1.87 (0.21 to 16.41)	$p = 0.56$						
Mild diarrhoea (culture negative) Schultz <i>et al</i> <sup>11</sup>	10	0 (0)	10	6 (60)	0	$p = 0.011$						
Feeding intolerance, including NEC Gross <sup>12</sup>	41	1 (2.4)	26	6 (23.1)	0.11 (0.01 to 0.83)	$p = 0.007$						
>2 weeks to tolerate full enteral feeds Lucas <sup>16</sup>	83	9 (11)	76	25 (33)	0.33 (0.16 to 0.66)	$p = 0.0007$						
>3 weeks to tolerate full enteral feeds Lucas <sup>16</sup>	83	4 (5)	76	13 (17)	0.28 (0.10 to 0.83)	$p = 0.012$						

NEC, necrotising ulcerative colitis; No., total; n, number of cases.  
RR < 1 favours donor breast milk.

**Table 4** Comparison of growth between the donor breast milk and formula groups

Outcome and studies	Sole diet comparison						p value	Supplementary diet comparison						p value		
	Donor breast milk			Formula				Donor breast milk			Formula					
	No.	Mean	SD	No.	Mean	SD	MD (95% CI)		No.	Mean	SD	No.	Mean	SD	MD (95% CI)	
Days taken to regain birth weight																
Lucas <i>et al</i> <sup>3</sup>	29	16	–	33	10*	–	6*	p=0.004	67	15*	–	65	13*	–	2*	p=0.14
Gross <sup>12</sup>	40	3.58	0.8	20	10.3	3.58	–6.72 (–7.90 to –5.54)	p<0.001								
Raiha <i>et al</i> <sup>10</sup>	22	16.27	6.30	84	13.56	5.29	2.71 (0.09 to 5.33)	p=0.042								
Schultz <i>et al</i> <sup>1</sup>	10	10.5	–	10	18.9	–	–8.4	“p value NS”								
Weight change: birth to 1 month (g/week)																
Davies <sup>9</sup>	34	135.3	59.8	34	171.8	61.6	–36.5 (–65.9 to –7.1)	p=0.016								
Weight change: 1 to 2 months (g/week)																
Davies <sup>9</sup>	34	225.3	85.8	34	242.4	69.9	–17.1 (–55.0 to 20.8)	p=0.37								
Change in head circumference (mm/week)																
Gross <sup>12</sup>	40	8.4	1.3	20	8.8	2.2	–0.4 (–1.3 to 0.5)	p=0.38								
>2 weeks after regaining birth weight																
Lucas <i>et al</i> <sup>3,19</sup>	83	8.96	3.22	76	10.43	4.9	–1.47 (–2.73 to –0.21)	p=0.026	142	9.1	2.1	144	10.5	3.5	–1.4 (–2.1 to –0.7)	
p=0.0001																
From birth to 1 month																
Davies <sup>9</sup>	34	6.7	1.6	34	7.0	1.4	–0.3 (–1.0 to 0.4)	p=0.41								
From 1 to 2 months																
Davies <sup>9</sup>	34	6.9	2.2	34	7.7	1.8	–0.8 (–1.8 to 0.2)	p=0.106								
Between start of enteral feeds and 36 weeks' gestation																
Pulet <i>et al</i> <sup>4</sup>	6	10	2	6	11	1	–1 (–3.0 to 1.0)	p=0.30								

MD, mean difference; No., total.  
MD<0 favours donor breast milk.  
\*Median not mean.

effect was not statistically significant. One study<sup>15</sup> found significantly higher levels of plasma bilirubin in the donor milk group on day 4–7 and day 25 in the sole and supplementary diet comparison (data not shown).

### Effect on early postnatal growth

Many of the outcome measures reported for growth were not standardised across studies. Therefore it was not possible to undertake meta-analysis on these outcomes. The results for days to regain birth weight, weight change from birth to 2 months, and the change in head circumference are given in table 4. The remaining growth outcomes (other measures of weight, length and skinfold thickness) are available online (see appendix tables B and C at <http://adc.bmj.com/supplemental>). For the sole diet comparison, 9/13 comparisons of early postnatal weight gain (in six studies) were found to be significantly in favour of formula (ie faster or greater weight gain in the formula group), one<sup>12</sup> was significantly in favour of donor milk and in three there was no significant difference (two in favour of formula). There was also a tendency towards smaller growth in head circumference in the sole donor milk group compared with the sole formula group (five comparisons in four

studies) but this difference was only statistically significant in one study. Of the seven comparisons of length gain made (in five studies), five were significantly in favour of formula and in two there was no significant difference (one in favour of formula). There were significantly smaller gains in triceps and subscapular skinfold thickness in the donor milk group than in the formula group (two comparisons in one study). For the supplementary diet comparison there was also significantly slower growth in the donor milk group compared with the formula group in terms of weight gain, increased head circumference and increased skinfold thickness, but no difference in length gain (table 4 and appendix tables B and C at <http://adc.bmj.com/supplemental>).

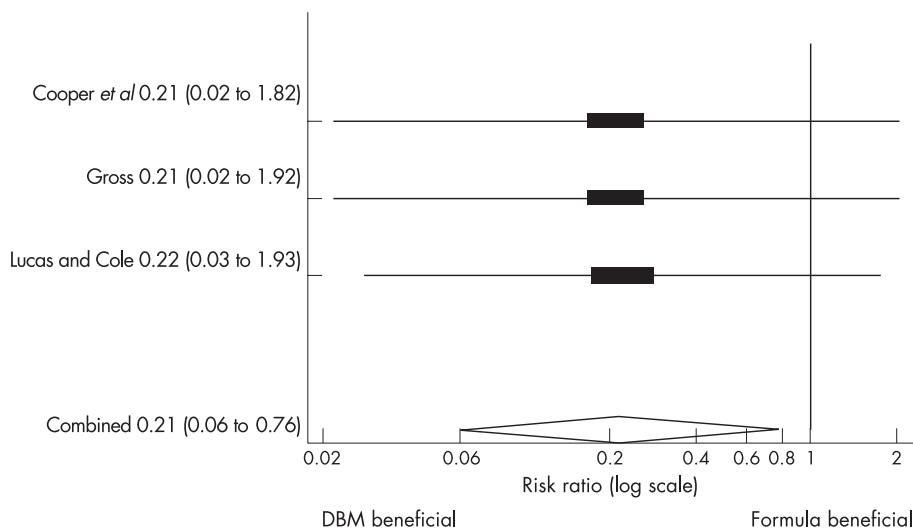
### Effect on growth measured at ages 9 months, 18 months and 7.5–8 years

One study reported several measures of growth at ages 9 months, 18 months and 7.5–8 years<sup>13 22</sup> (see appendix tables D and E at <http://adc.bmj.com/supplemental>). The donor milk group and formula group were similar (with no significant differences) at all ages with respect to weight, height, skinfold thickness, body mass index and (at age 7.5–8 years) waist to

**Table 5** Comparison of development between the donor breast milk and formula groups

Outcome and studies	Sole diet comparison						RR (95% CI)	p value	Supplementary diet comparison						p value	
	Donor breast milk			Formula					Donor breast milk			Formula				
	No.	n (%)		No.	n (%)			No.	n (%)		No.	n (%)				
Neurological status impaired at 9 months																
Lucas <i>et al</i> <sup>7</sup>	66	4 (6.1%)		56	9 (16.1%)		0.38 (0.12 to 1.16)	p=0.074	140	15 (10.7%)		138	15 (10.9%)	0.99 (0.50 to 1.94)	p=0.97	
Neurological status impaired at 18 months																
Lucas <i>et al</i> <sup>7</sup>	66	4 (6.1%)		56	7 (12.5%)		0.48 (0.15 to 1.57)	p=0.22	140	11 (7.9%)		138	10 (7.2%)	1.08 (0.48 to 2.47)	p=0.85	
	N	Mean	SD	N	Mean	SD	MD		N	Mean	SD	N	Mean	SD	MD	
Development quotient at 9 months (Knobloch)																
Lucas <i>et al</i> <sup>7</sup>	62	97.2	8.5	48	98.2	11.0	–1.0 (–4.7 to 2.7)	p=0.59	133	98.2	10.1	126	101.2	10.5	–3.0 (–5.52 to –0.48)	p=0.020
Mental developmental index at 18 months (Bayley)																
Lucas <i>et al</i> <sup>9</sup>	62	94.8	16.5	52	95.3	19.5	–0.5 (–7.1 to 6.2)	p=0.88	134	102.2	19.7	139	103.8	20.0	–1.6 (–6.33 to 3.13)	p=0.51
Psychomotor developmental index at 18 months (Bayley)																
Lucas <i>et al</i> <sup>9</sup>	62	93.0	14.2	52	94.2	15.9	–1.2 (–6.8 to 4.4)	p=0.67	134	95.5	15.0	139	94.5	16.5	1.0 (–2.76 to 4.76)	p=0.60

MD, mean difference; No., total, n, number of cases.  
RR<1 favours donor breast milk; MD<0 favours donor breast milk.



**Figure 2** Individual and combined estimates of risk ratio of confirmed NEC for sole DBM versus sole formula milk.

hip ratio. Importantly, this study found significantly slower growth in the donor milk group compared with the formula group during the early postnatal period.

Similarly, in the supplementary diet comparison, the slower growth in the donor milk group compared with the formula group during the early postnatal period had no long-term effect on growth in this study; the two groups were broadly similar with the exception of body mass index being significantly higher in the donor milk group at age 18 months.

#### Effect on developmental status

Child development at age 9 months and 18 months was assessed in one study<sup>13</sup> (table 5). In the sole diet comparison there were fewer neurologically impaired children at age 9 months in the donor milk group compared with the formula group but this effect was not statistically significant (RR = 0.38, 95% CI 0.12 to 1.16), and was even weaker when assessed at age 18 months. There were no significant differences between the two feeding groups with respect to mean developmental quotient at age 9 months (Knobloch index) or 18 months (Bayley index) (table 5).

In the supplementary diet comparison, the proportion of neurologically impaired children at age 9 months and 18 months was similar in the donor milk group compared with the formula group. The donor milk group had significantly lower mean developmental scores than the formula group at age 9 months (Knobloch) but not when assessed at age 18 months (Bayley) (table 5).

#### Effect on later outcomes

One study measured systolic and diastolic blood pressure at age 7.5–8 years and found no significant differences between the donor milk and formula groups in both the sole and supplementary comparisons (see appendix table F at <http://adc.bmj.com/supplemental>).<sup>20</sup>

## DISCUSSION

Before discussing the results of this review, a number of potential limitations should be considered. First, only seven studies were included and the total sample size for most outcomes was small. Second, these studies started over 20 years ago, and they may no longer be clinically relevant to contemporary practice as survival of preterm and low birth-weight infants has greatly improved<sup>25</sup> and feeding practices have changed. Third, the methodological quality of most of the

studies was poor in terms of randomisation, blinding of carers and assessors, and assessment of confounding which may have introduced bias, in particular, for the observational data included. Lastly, substantial heterogeneity between the studies makes it difficult to pool evidence across studies. The studies had different inclusion criteria and varied in terms of the type of donor and formula milk used, and the method, timing, and volume of feeding (table 1). These differences may account for some of the observed differences in effects between the studies.

Since the search was conducted, another trial has been published.<sup>26</sup> This trial, in extremely preterm babies (<30 weeks' gestation), compared fortified donor breast milk with formula, both given as a supplement to mother's own milk. The trial found no significant difference between the two groups in terms of infection-related events or death. For NEC, there was some protection, but the 95% CI was wide (RR 0.56, 0.20 to 1.58). With regard to growth, the trial revealed no effect on length or head circumference gain, but found significantly poorer weight gain in the donor milk group. This resulted in 21% of the infants randomised to donor milk being given

#### What is already known on this topic

- Breast milk is the recommended form of enteral nutrition for all infants.
- Pasteurised donor breast milk is an alternative when mother's own milk is not available.
- Pasteurisation affects the nutritional and immunological properties of breast milk.
- The evidence for the benefits of donor milk compared with formula for preterm infants is limited.

#### What this study adds

- Donor milk given as a sole diet appears to be associated with a lower risk of NEC but slower growth in the early postnatal period when compared with infant formula.
- The long-term effect of donor milk as compared with formula is unclear.

formula; this may have diluted the effect of donor milk on outcomes such as NEC.

Our findings, based on 13 cases of NEC in 268 infants, suggest that donor milk given as a sole diet is associated with a lower risk of NEC compared with formula. Although the observed effects were remarkably similar between the three studies (RR 0.21–0.22), true heterogeneity of effect cannot be ruled out because of differences between the studies in terms of the incidence of NEC (5–20%), the type of donor milk and the timing of feeding initiation. Furthermore, one study<sup>23</sup> was not randomised, and none of the studies considered blinding of the intervention or outcome. These methodological weaknesses may have biased the observed effects, particularly in the study<sup>23</sup> which did not state how NEC was diagnosed; bias due to subjective diagnosis cannot be ruled out. The homogeneous effects found in the three studies reduces the likelihood of a chance finding, and these alternative explanations are unlikely to account wholly for the large effects observed. Moreover, in the study which looked at both sole and supplemented diets,<sup>13</sup> the risk of NEC increased as the amount of formula increased from 1.2% (2/167) in infants who received donor milk and mother's own milk and 1.2% (1/86) in infants who received donor milk only to 2.9% (5/173) in infants who received formula and mother's own milk and to 5.3% (4/76) in infants who received formula only ( $p = 0.053$  for linear trend). The effect of donor milk on NEC seemed weaker in the study in which donor milk and formula were given as a supplement to mother's own milk (RR 0.41, 95% CI 0.08 to 2.11), although the study may not have had sufficient power to detect a modest effect for such a rare outcome (incidence of NEC in formula group was 2.9%). In Schanler and colleagues' recent trial,<sup>26</sup> there was a similar suggestion of some protection of fortified donor milk given as a supplement to mother's own milk.

Breast milk is known to have immune properties as it includes specific immunoglobulin A, lysozyme and lactoferrin.<sup>2</sup> Therefore breast milk is thought to protect the preterm infant from bacterial and viral infections. Holder pasteurisation, however, reduces some of the anti-infective properties of breast milk. Infection was not reported as an outcome in any of the studies included in this review, but it was an outcome in the recent Schanler trial,<sup>26</sup> although no effect was observed.

The nutrient concentrations in human milk may be inadequate for preterm or very low birthweight infants, who have increased nutritional requirements.<sup>2</sup> This may be particularly so for donor milk, as Holder pasteurisation reduces some of the nutritive properties,<sup>2</sup> and the composition of donor milk is not the same as that received by a mother's own infant. This is especially true for drip milk, which was used in one study,<sup>13</sup> and term breast milk, which was used in four studies.<sup>9–12</sup> Inadequate nutrition in the early postnatal period is thought to affect growth, bone mineralisation and neurodevelopment. One trial<sup>12</sup> excluded infants who developed feed intolerance and NEC after randomisation, which may have exaggerated the increase in short-term growth among infants randomised to formula. Although the evidence suggests that donor milk is associated with slower growth in the early postnatal period, only one study included long-term follow-up.<sup>13</sup> Interestingly, this study found no difference between the groups in several growth indices and neurodevelopment at ages 9 months and 18 months, and growth and blood pressure at 7.5–8 years despite observing slower growth in the donor milk group in the short term.

Donor milk given as a sole diet is associated with a lower risk of NEC but slower growth in the early postnatal period. The long-term effect of donor milk as compared with formula is unclear because only one study followed participants into

childhood and adulthood. Further research is needed to confirm our findings and measure the effect of donor breast milk that is fortified or given as a supplement to mother's own milk.

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