The Benefits of Expressed Maternal Milk and Donor Breast Milk for Preventing Necrotizing Enterocolitis in Preterm Infants: Systematic Review and Meta-Analysis

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Abstract

Objectives: To examine the protective effects that feeding with expressed maternal milk and donor breast milk compared with formula milk reduces the risk of development of necrotizing enterocolitis in preterm infants.

Data sources: The studies for our systematic review were searched from our library’s electronic databases including PUBMED/MEDLINE, SCIENCEDIRECT (1997-2008), EBSCOHOST (1965-2008), OVID (1993-2008) and Cochrane Library.

Methods: Systematic review and meta-analysis of randomized controlled trials or quasi-randomized controlled trials.

Results: In our systematic review and meta-analysis only five trials fulfilled the prespecified inclusion criteria. Except for one study, all the rest included studies which were initiated nearly three decades ago. None of the individual trials found any statistically significant difference in the incidence of necrotizing enterocolitis. However, meta-analysis found that preterm infants feeding with donor breast milk was associated with a significantly reduced relative risk of necrotizing enterocolitis. Suspected necrotizing enterocolitis was three times less likely (relative risk 0.31; 95% confidence interval 0.12-0.81; p=0.02<0.05) and confirmed necrotizing enterocolitis was four times less likely (relative risk 0.24; 95% confidence interval 0.07-0.76; p=0.02<0.05) in premature infants feeding with donor breast milk compared with formula milk given as a sole diet. No data to date was available to be combined in our meta-analysis to compare expressed maternal milk with formula milk given as a sole diet.

Conclusion: Feeding with donor breast milk is associated with a lower risk of necrotizing enterocolitis in preterm infants, but the protective benefits of donor breast milk are described as of borderline effects and the quality of the evidence is limited. Further trials should be focused on the effects of fortified expressed maternal milk and donor breast milk for preventing necrotizing enterocolitis in premature infants.

Keywords: Preterm infants; Necrotizing enterocolitis; Expressed maternal milk; Donor breast milk; Formula milk; Systematic review; Meta-analysis

Introduction

As advances in neonatology and the modern neonatal intensive care unit (NICU) have improved, much more premature infants could survive after birth. However, necrotizing enterocolitis (NEC) remains a major cause of neonatal morbidity and death. The mortality rate (15%–25%) for affected infants has not changed appreciably in the past 30 years [1]. Necrotizing enterocolitis is primarily a disease of premature infants; >90% of those affected were born prior to 36 weeks gestation [2]. Prematurity is the only risk factor for necrotizing enterocolitis consistently identified in case-control studies. The mechanism of the development of necrotizing enterocolitis is unclear; a leading hypothesis is that the immature intestinal epithelial cells mount an exaggerated inflammatory response to intestinal injury in preterm infants [3].

At present more attention is being focused on the nutritional management of the preterm infants those who are vulnerable to necrotizing enterocolitis. Breast milk is the recommended source of enteral nutrition for all infants including those preterm infants [4]. Compared with formula feeding, a putative benefit of breast milk for feeding preterm infants is that the delivery of immunologic factors to the immature gut mucosa may decrease the risk of necrotizing enterocolitis. However, for some reason mothers those who delivered preterm may be unable to provide directly breast feeding or sufficient breast milk for their premature infants. When directly breast feeding is not available, the preferred alternative is expressed maternal milk (EMM) or donor breast milk (DBM) [5,6].

A theoretical concern is that feeding preterm with expressed maternal milk and donor breast milk do not completely equal to directly breast feeding. Storage and processing of expressed maternal milk and donor breast milk alters some of the immunologic and nutritional properties [7]. Early clinical study suggested that feeding with donor breast milk is associated with lower growth rates in the preterm infants during the short postnatal term [8]. However, it is unclear whether the decrease of non-nutrient components in expressed maternal milk and donor breast milk during storage and processing may confer immunoprotective benefits. The objective of our systematic review and meta-analysis is to determine if enteral feeding with expressed maternal milk and donor breast milk compared with formula milk reduces the incidence of necrotizing enterocolitis in preterm infants.

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Methods

Search strategy

The standard search strategy of our systematic review included electronic search and manual search. Electronic search was carried out in our library’s databases including PUBMED/MEDLINE, SCIENCEDIRECT (1997-2008), EBSCOHOST (1965-2008), EMBASE (1974-2008), OVID (1993-2008) and Cochrane Library. There was no language restriction. As supplement, manual search had also been undertaken; references in studies identified as relevant, and in previous reviews and standard textbooks of neonatal medicine and nutrition were examined.

Selection

In this systematic we selected studies on the basis of study design, participant, intervention and outcome. Expressed maternal milk had to be collected from the own mother of each preterm infant. Donor breast milk had to be donated from someone other than the infant’s mother and it had to be pasteurized or purchased from human milk bank. The studies which had severe methodological faults would be excluded; if the data was equivocal it would be excluded before we got the clarification from its authors. The followings were details for the inclusion criteria:

- **Study design**: Randomized controlled trials (RCTs) or quasi-RCTs.
- **Participant**: Preterm infants <37 weeks gestation.
- **Intervention**: Donor breast milk versus formula milk given as a sole diet.
- **Outcome**: Suspected NEC (included confirmed NEC and the cases reported by individual trial only without confirmed evidence).

Valid assessment

We assessed the methodological quality of the included randomized controlled trials. Quality of the trials was evaluated in terms of allocation concealment, blinding of parents or carers and assessors to intervention, and completeness of assessment in all randomized subjects. Additional information was requested from the authors of each trial to clarify methodology and results as necessary. Taken clinical heterogeneity into consideration, we also assessed the baseline variables and confounding factors of the included studies.

Data abstraction

The title and abstract of studies identified via our search strategy were screened by two independent reviewers. The full text of each study which potentially met the inclusion criteria was critically reviewed by both two reviewers. Then the decision to include or exclude a specific article was made by consensus of the two reviewers. If the two reviewers could not get consensus on one article, we would turn to the third reviewer.

Quantitative data synthesis

Data were separately extracted and summarized into evidence tables by each reviewer, compared data, and resolved differences by consensus. If there were sufficient data and no evidence of significant heterogeneity (p>0.10), meta-analysis would be performed by a fixed-effects model. If there was evidence of significant heterogeneity (p<0.10) or no sufficient data was available, meta-analysis would be switched to a random-effects model or non-quantitative systematic review would be performed. Effects were expressed as relative risk (RR) and 95% confidence interval (CI) and risk difference (RD) and 95% CI for a categorical data, p<0.05 was considered statistically significant.

Results

Figure 1 is the QUOROM statement flow diagram of this systematic review. After the first round of screening, thirteen potentially relevant full texts were identified by two independent reviewers. Then those full texts were critically reviewed in accordance with the above inclusion criteria, eight articles [11-18] (seven [11-17] due to no relevant clinical outcome, one [18] due to methodological fault) were excluded on consensus of the two reviewers. At last only five randomized controlled trials were included in our systematic review and meta-analysis.

Description of studies

The summary of characteristics of the five included trials is shown in the Table 1. There were 862 preterm infants had been enrolled in our systematic review. Four studies [19-22] of these included trials compared donor breast milk with formula milk given as a sole diet for preterm infants. Two studies [22,23] compared donor breast milk with formula milk given as a supplement diet to expressed maternal milk. No data to date was available which compared expressed maternal milk with formula milk given as a sole diet for preterm infants.

Table 2 provides assessments of methodological quality of included trials. Except for one study [21] which was quasi-RCT, all the rest of studies are RCTs with allocation concealment. All the studies had not been taking blinded of intervention and none could specify blinding of outcome. Complete follow up was performed in all included studies.

Findings of the included studies

Effects on suspected NEC: None of the included individual studies

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**Figure 1**: Flow diagram of search and selection of included trials.
Sensitivity analysis

Sensitivity analysis is an approach to test how robust the results of our systematic review are relative to key decisions and assumptions that were made in the process of conducting the review. Preterm infants enrolled in the formula milk group were fed with term formula milk in one included study [19]. Term formula milk is different with preterm formula milk which was used in other included studies. To eliminate this potential confounding factor, the data abstracted from the studies in which preterm infants were fed with preterm formula milk, were combined in sensitivity analysis, and we found a statistically significant difference (RD -0.05; 95% CI -0.10-0.00; p=0.03<0.05) (Figure 8).

Discussion

In order to get the appropriate conclusion from our systematic review and meta-analysis, the limitations should be carefully considered before we discuss its results. First of all, there were only five included trials in our meta-analysis and no data was available to compare expressed maternal milk versus preterm formula milk given as a sole diet, therefore a noteworthy publication bias examined by ‘funnel-plot’ was inevitable. For another, except for one study [23], the rest included studies which were started nearly three decades ago, and in the past 30 years the composition of the formula milk and feeding practices of preterm infants had changed greatly. Still another, there were several methodological faults in terms of randomization, allocation concealment, blinding of intervention and outcome, and a degree of clinical heterogeneity. These limitations would be considered carefully in our following discussion.

Based on the available data of the included studies, our meta-analysis suggests that donor breast milk has a degree of protective effects for preventing necrotizing enterocolitis in preterm infants. Although none of the included studies individually found any significant difference, the significant differences were shown in our meta-analysis. Furthermore, the relative risk estimates and the risk difference estimate are both statistically significant, the results of sensitivity analysis are consistent with the overall effects shown in

<table>
<thead>
<tr>
<th>Included Study</th>
<th>Participant</th>
<th>Intervention</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gross 1983</td>
<td>Gestation 27-33 weeks Birth weight &lt;1600g. DBM (n=41) vs TFM (n=26) given as sole diet</td>
<td>Suspected NEC: DBM(14/41) ; TFM(3/26) Confirmed NEC: DBM(1/41) ; TFM(3/26)</td>
<td></td>
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<tr>
<td>Tyson 1983</td>
<td>Birth weight &lt;1500g. DBM (n=37) vs PFM (n=44) given as sole diet</td>
<td>Suspected NEC: DBM(0/37) ; PFM(2/44) Confirmed NEC: DBM(0/37) ; PFM(1/44)</td>
<td></td>
</tr>
<tr>
<td>Cooper 1984</td>
<td>Gestation=36 weeks Birth weight 1200-1500g DBM (n=24) vs PFM (n=15) given as sole diet</td>
<td>Suspected NEC: DBM(1/24) ; PFM(3/15) Confirmed NEC: DBM(1/24) ; PFM(3/15)</td>
<td></td>
</tr>
<tr>
<td>Lucas 1990A</td>
<td>159 preterm infants Birth weight &lt;1850g. DBM (n=83) vs PFM (n=76) given as sole diet</td>
<td>Suspected NEC: DBM(3/83) ; PFM(6/76) Confirmed NEC: DBM(1/83) ; PFM(4/76)</td>
<td></td>
</tr>
<tr>
<td>Lucas 1990B</td>
<td>343 preterm infants Birth weight &lt;1850 g. DBM (n=170) vs PFM (n=173) given as the supplement diet for EMM</td>
<td>Suspected NEC: DBM(6/170) ; PFM(6/173) Confirmed NEC: DBM(2/170) ; PFM(5/173)</td>
<td></td>
</tr>
<tr>
<td>Schanler 2005</td>
<td>Gestation 23-29 weeks DBM (n=92) vs PFM (n=81) given as the supplement diet for EMM</td>
<td>Suspected NEC: DBM(5/92) ; PFM(10/81) Confirmed NEC: DBM(5/92) ; PFM(10/81)</td>
<td></td>
</tr>
</tbody>
</table>

Note: Lucas 1990A and Lucas 1990B were parallel studies in one trial. RCT: randomized controlled trial.

Table 2: Methodological quality of the included studies.

<table>
<thead>
<tr>
<th>Included Study</th>
<th>Study Design</th>
<th>Allocation Concealment</th>
<th>Blinding of Intervention</th>
<th>Blinding of Outcome</th>
<th>Complete Follow up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gross 1983</td>
<td>RCT</td>
<td>Yes</td>
<td>No</td>
<td>Unclear</td>
<td>Yes</td>
</tr>
<tr>
<td>Tyson 1983</td>
<td>RCT</td>
<td>Yes</td>
<td>No</td>
<td>Unclear</td>
<td>Yes</td>
</tr>
<tr>
<td>Cooper 1984</td>
<td>quasi-RCT</td>
<td>Unclear</td>
<td>No</td>
<td>Unclear</td>
<td>Yes</td>
</tr>
<tr>
<td>Lucas 1990 A</td>
<td>RCT</td>
<td>Yes</td>
<td>No</td>
<td>Unclear</td>
<td>Yes</td>
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<tr>
<td>Lucas 1990 B</td>
<td>RCT</td>
<td>Yes</td>
<td>No</td>
<td>Unclear</td>
<td>Yes</td>
</tr>
<tr>
<td>Schanler 2005</td>
<td>RCT</td>
<td>Yes</td>
<td>No</td>
<td>Unclear</td>
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</tbody>
</table>

Note: Lucas 1990A and Lucas 1990B were parallel studies in one trial. RCT: randomized controlled trial.

Note: Relative risk of Suspected NEC in preterm infants fed with donor breast milk vs formula milk (RR: 0.54; 95%CI: 0.31-0.95; p<0.03), the overall effect favors donor breast milk.

**Figure 2:** Relative risk of Suspected NEC in preterm infants fed with donor breast milk vs formula milk.

Note: Relative risk of Confirmed NEC in preterm infants fed with donor breast milk vs formula milk (RR: 0.34; 95%CI: 0.17-0.68; p<0.002), the overall effect favors donor breast milk.

**Figure 3:** Relative risk of Confirmed NEC in preterm infants fed with donor breast milk vs formula milk.

Note: Relative risk of Suspected NEC in preterm infants fed with donor breast milk vs formula milk given as a sole diet (RR: 0.31;95%CI: 0.12-0.81; p<0.02), the overall effect favors donor breast milk.

**Figure 4:** Relative risk of Suspected NEC in preterm infants fed with donor breast milk vs formula milk given as a sole diet.

Note: Relative risk of Confirmed NEC in preterm infants fed with donor breast milk vs formula milk given as a sole diet (RR: 0.32; 95%CI: 0.22-0.45; p<0.001), the overall effect favors donor breast milk.

**Figure 5:** Relative risk of Confirmed NEC in preterm infants fed with donor breast milk vs formula milk given as a sole diet.

Note: Risk difference of Confirmed NEC in preterm infants fed with donor breast milk vs formula milk given as a sole diet (RR: -0.65; 95%CI: -0.10- -0.00; p<0.03), the overall effect favors donor breast milk.

**Figure 6:** Risk difference of Suspected NEC in preterm infants fed with donor breast milk vs formula milk given as a sole diet (RD: 0.05; 95%CI: 0.00-0.10; p=0.04), the overall effect favors donor breast milk.

**Figure 7:** Risk difference of Suspected NEC in preterm infants fed with donor breast milk vs formula milk given as a sole diet (RD: 0.05; 95%CI: 0.00-0.10; p=0.04), the overall effect favors donor breast milk.

**Figure 8:** Risk difference of Suspected NEC in preterm infants fed with donor breast milk vs formula milk given as a sole diet.

Note: Risk difference of Confirmed NEC in preterm infants fed with donor breast milk vs formula milk given as a sole diet (RD: 0.05; 95%CI: 0.00-0.10; p=0.04), the overall effect favors donor breast milk.

meta-analysis. Suspected NEC is three times less likely and Confirmed NEC is four times less likely in premature infants feeding with donor breast milk compared with formula milk given as a sole diet. Subgroup analysis was performed in this review due to clinical heterogeneity. In subgroup analysis we compared donor breast milk with preterm formula milk given as a supplement diet to expressed maternal milk for preterm infants, and no significant difference was found. No data was available to compared expressed maternal milk with preterm formula milk or donor breast milk. Only one individual study [23] reported the incidence of necrotizing enterocolitis in preterm infants fed with expressed maternal milk given as a sole diet, and no benefit was found. The potential benefits of expressed maternal milk for preventing necrotizing enterocolitis may be estimated in the future systematic review when sufficient data are available.

Up to now the pathophysiology of necrotizing enterocolitis remains poorly delineated, evidence supports prematurity is an important risk factor and the beneficial effects of breast milk reduce the risk for necrotizing enterocolitis in premature infants [24]. Feeding preterm infants with breast milk would deliver an amount of immunoprotective factors and growth factors to the immature gut mucosa that may decrease the risk of necrotizing enterocolitis. According to basic research, one of the constituents of breast milk that may prove to be therapeutic is epidermal growth factor (EGF), a trophic substance for intestinal growth [25]. A preliminary study of epidermal growth factor...
in neonates diagnosed with necrotizing enterocolitis has shown that epidermal growth factor promotes the repair of intestinal epithelium, and supplementation of epidermal growth factor in animal models has decreased the incidence of necrotizing enterocolitis [26]. Donor breast milk is generally obtained from women who deliver term infants later in their lactation and a loss of the immunologic and nutritional properties during storage and processing in the human milk bank, so the milk has a lower content of protein and host defence protein [7,27]. Although the immunoprotective factors delivered to preterm infants is a small amount by donor breast milk, it may account for the borderline protective effect for preventing necrotizing enterocolitis in our meta-analysis.

Expressed maternal milk is obtained from their own mothers for preterm infants, so it is a form of preterm milk. The results of studies compared preterm milk with term milk suggested that there were much more immunologic and nutritional properties in preterm milk [28]. Theoretically expressed maternal milk is more appropriate than donor breast milk for feeding premature infants, but why no significant difference was found in meta-analysis when donor breast milk versus preterm formula milk were given as a supplement diet to expressed maternal milk? Recently one study enrolled in 1272 preterm infants was performed to determine the association between breast milk intake and risk of necrotizing enterocolitis or death among infants 401 to 1000 g birth weight [17]. This study suggested that a dose-related association of breast milk feeding with a reduction of risk of necrotizing enterocolitis or death after the first two weeks of life among extremely low birth weight infants [17]. These findings may help us elucidate the puzzled results of meta-analysis. When donor breast milk (contains a small amount of immunoprotective factors) versus preterm formula milk (contains no immunoprotective factor) were given as a supplement diet to expressed maternal milk (contains much more immunoprotective factors), the difference of the dose-related effects between the two groups may become relatively less significant.

Are these findings in our systematic review of clinical significance? In meta-analysis with several limitations, we found that donor breast milk was associated with a borderline effect for preventing necrotizing enterocolitis in premature infants. However, the number needed to treat (NNT=1/1RD) was 20 (that meant one case of Confirmed NEC averted if 20 preterm infants received donor breast milk). Additional, donor breast milk given as a sole diet for preterm infants is associated with slower growth at least in the early postnatal period, the long-term effect is unclear [8,29]. The nutrient concentrations in expressed maternal milk and donor breast milk may be inadequate for preterm infants, who have increased nutritional requirements [30]. The nutrient deficits that arise from feeding unfortified donor breast milk may be corrected with nutrient supplementation, and in many clinical guidelines fortifiers were recommended when necessary [5,6]. However, a theoretical concern with human milk fortification is that the added nutrients may affect the intrinsic host defense system of the milk, thereby may increase the risk of development of necrotizing enterocolitis in preterm infants. Although a meta-analysis comparing infants fed unfortified and fortified human milk did not identify any significant difference in necrotizing enterocolitis [31], further research in the safety of human milk fortification is still needed.

In summary, donor breast milk given as a sole diet is associated with lower risk of the development of necrotizing enterocolitis in preterm infants. The clinical applicability of these findings should be considered with caution due to the limitations of our meta-analysis itself and the potential adverse effects on growth found in other studies. On our systematic review itself, the inspiration significance is the most important rather than its findings’ clinical applicability. Consequently, further high quality randomized controlled trials are still needed, and should be focused on the protective effects of fortified expressed maternal milk and donor breast milk for preterm infants, especially the dose-related effects for preventing necrotizing enterocolitis in the extremely low birth weight infants.

References


