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Effects of neonatal nutrition interventions on neonatal mortality and child health and development outcomes: A systematic review

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Abstract

Background: The last two decades have seen a significant decrease in mortality for children <5 years of age in low and middle-income countries (LMICs); however, neonatal (age, 0–28 days) mortality has not decreased at the same rate. We assessed three neonatal nutritional interventions that have the potential of reducing morbidity and mortality during infancy in LMICs.

Objectives: To determine the efficacy and effectiveness of synthetic vitamin A, dextrose oral gel, and probiotic supplementation during the neonatal period.

Search Methods: We conducted electronic searches for relevant studies on the following databases: PubMed, CINAHL, LILACS, SCOPUS, and CENTRAL, Cochrane Central Register for Controlled Trials, up to November 27, 2019.

Selection Criteria: We aimed to include randomized and quasi-experimental studies. The target population was neonates in LMICs. The interventions included synthetic vitamin A supplementation, oral dextrose gel supplementation, and probiotic supplementation during the neonatal period. We included studies from the community and hospital settings irrespective of the gestational age or birth weight of the neonate.

Data Collection and Analysis: Two authors screened the titles and extracted the data from selected studies. The risk of bias (ROB) in the included studies was assessed according to the Cochrane Handbook of Systematic Reviews. The primary outcome was all-cause mortality. The secondary outcomes were neonatal sepsis, necrotizing enterocolitis (NEC), prevention and treatment of neonatal hypoglycaemia, adverse events, and neurodevelopmental outcomes. Data were meta-analyzed by random effect models to obtain relative risk (RR) and 95% confidence interval (CI) for dichotomous outcomes and mean difference with 95% CI for continuous outcomes. The overall rating of evidence was determined by the Grading of Recommendations Assessment, Development, and Evaluation (GRADE) approach.

This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited. © 2021 The Authors. *Campbell Systematic Reviews* published by John Wiley & Sons Ltd on behalf of The Campbell Collaboration. Main Results: Sixteen randomized studies (total participants 169,366) assessed the effect of vitamin A supplementation during the neonatal period. All studies were conducted in low- and middle-income (LMIC) countries. Thirteen studies were conducted in the community setting and three studies were conducted in the hospital setting, specifically in neonatal intensive care units. Studies were conducted in 10 different countries including India (four studies), Guinea-Bissau (three studies), Bangladesh (two studies), and one study each in China, Ghana, Indonesia, Nepal, Pakistan, Tanzania, and Zimbabwe. The overall ROB was low in most of the included studies for neonatal vitamin A supplementation. The pooled results from the community based randomized studies showed that there was no significant difference in all-cause mortality in the vitamin A (intervention) group compared to controls at 1 month (RR, 0.99; 95% CI, 0.90-1.08; six studies with 126,548 participants, statistical heterogeneity l^2 0%, funnel plot symmetrical, grade rating high), 6 months (RR, 0.98; 95% CI, 0.89-1.07; 12 studies with 154,940 participants, statistical heterogeneity l^2 43%, funnel plot symmetrical, GRADE quality high) and 12 months of age (RR, 1.04; 95% CI, 0.94-1.14; eight studies with 118,376 participants, statistical heterogeneity I^2 46%, funnel plot symmetrical, GRADE quality high). Neonatal vitamin A supplementation increased the incidence of bulging fontanelle by 53% compared to control (RR, 1.53; 95% CI, 1.12-2.09; six studies with 100,256 participants, statistical heterogeneity I^2 65%, funnel plot symmetrical, GRADE quality high). We did not identify any experimental study that addressed the use of dextrose gel for the prevention and/or treatment of neonatal hypoglycaemia in LMIC. Thirtythree studies assessed the effect of probiotic supplementation during the neonatal period (total participants 11,595; probiotics: 5854 and controls: 5741). All of the included studies were conducted in LMIC and were randomized. Most of the studies were done in the hospital setting and included participants who were preterm (born < 37 weeks gestation) and/or low birth weight (<2500 g birth weight). Studies were conducted in 13 different countries with 10 studies conducted in India, six studies in Turkey, three studies each in China and Iran, two each in Mexico and South Africa, and one each in Bangladesh, Brazil, Colombia, Indonesia, Nepal, Pakistan, and Thailand. Three studies were at high ROB due to lack of appropriate randomization sequence or allocation concealment. Combined data from 25 studies showed that probiotic supplementation reduced all-cause mortality by 20% compared to controls (RR. 0.80: 95% CI. 0.66–0.96: total number of participants 10.998. number needed to treat 100, statistical heterogeneity l^2 0%, funnel plot symmetrical. GRADE quality high). Twenty-nine studies reported the effect of probiotics on the incidence of NEC, and the combined results showed a relative reduction of 54% in the intervention group compared to controls (RR, 0.46; 95% CI, 0.35–0.59; total number of participants 5574, number needed to treat 17, statistical heterogeneity l^2 24%, funnel plot symmetrical, GRADE quality high). Twenty-one studies assessed the effect of probiotic supplementation during the neonatal period on neonatal sepsis, and the combined results showed a relative reduction of 22% in the intervention group compared to controls (RR, 0.78; 95% CI, 0.70-0.86; total number of



Authors' Conclusions: Vitamin A supplementation during the neonatal period does not reduce all-cause neonatal or infant mortality in LMICs in the community setting. However, neonatal vitamin A supplementation increases the risk of Bulging Fontanelle. No experimental or quasi-experimental studies were available from LMICs to assess the effect of dextrose gel supplementation for the prevention or treatment of neonatal hypoglycaemia. Probiotic supplementation during the neonatal period seems to reduce all-cause mortality, NEC, and sepsis in babies born with low birth weight and/or preterm in the hospital setting. There was clinical heterogeneity in the use of probiotics, and we could not recommend any single strain of probiotics for wider use based on these results. There was a lack of studies on probiotic supplementation in the community setting. More research is needed to assess the effect of probiotics administered to neonates in-home/community setting in LMICs.

1 | PLAIN LANGUAGE SUMMARY

1.1 | Neonatal probiotic supplementation can improve infant illness and reduce death, but vitamin A does not, and may have adverse effects

Nutritional support during the 1st month of life is vital for the short- and long-term survival of the newborn. Neonatal nutrition interventions have the potential to decrease death and illness in young infants in LMICs.

1.1.1 | What is this review about?

This review assesses the efficacy of synthetic vitamin A, dextrose, and probiotic supplementation during the neonatal period. These interventions were assessed separately and not in combination with each other.

What is the aim of this review?

This Campbell systematic review assesses the efficacy of three neonatal nutritional interventions that have the potential of reducing morbidity and mortality during infancy in LMICs: synthetic vitamin A, dextrose, and probiotic supplementation.

1.1.4 | What are the main findings of the review?

Combined results from thirteen vitamin A studies conducted in the community settings showed that there was no significant effect of vitamin A supplementation for reduction of death in young infants at

1.1.2 | What studies are included?

Sixteen studies that assessed the effect of vitamin A supplementation during the neonatal period were included. Thirteen of these studies were conducted in the community setting and three studies were conducted in the hospital setting. All the included studies on neonatal vitamin A supplementation were conducted in LMICs. Most of the studies had a low ROB.

No experimental studies were found that evaluated the use of dextrose for the prevention or treatment of low blood sugar during the neonatal period.

Thirty-three studies assessed the use of probiotics during the 1st month of life. All included studies on probiotic supplementation were randomized and conducted in LMICs. Most of the included studies had a low ROB. The probiotics studies mainly included babies born early and/or with low birth weight, and these studies were mostly conducted in hospital settings.

1.1.3 | Key results

Combined results from thirteen vitamin A studies conducted in the community settings showed that there was no significant effect of vitamin A supplementation for reduction of death in young infants at 1, 6, or 12 months of age. Neonatal vitamin A supplementation increases the risk of bulging fontanelle by 53%. The pooled data from probiotics studies showed that this intervention reduced the risk of death by 20% compared to controls. Further analysis showed that compared to controls, probiotic supplementation reduced the risk of a severe form of gastrointestinal illness in neonates called NEC by 54%. Probiotic supplementation also reduced the risk of blood infection called sepsis by 22% compared to controls. The quality grade ratings for these outcomes were "high."

1, 6, or 12 months of age. However, neonatal vitamin A supplementation increased the incidence of bulging fontanelle by 53%.

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The pooled data from probiotics studies showed that this intervention reduced the risk of death by 20% compared to controls. Further analysis showed that compared to controls, probiotic supplementation reduced the risk of a severe form of gastrointestinal illness in neonates called NEC by 54%. Probiotic supplementation also reduced the risk of blood infection called sepsis by 22% compared to controls. The quality grade ratings for these outcomes were "high."

1.1.5 | What do the findings of this review mean?

Vitamin A supplementation during the 1st month of life does not reduce the risk of death during the 1st year of life in LMICs. However, neonatal vitamin A supplementation increases the risk of bulging fontanelle, which may cause damage to the brain.

We did not find any experimental studies from LMICs that assessed the use of dextrose gel supplementation during the 1st month of life for the prevention or treatment of low blood sugar.

Probiotic supplementation during the 1st month of life to babies born preterm and/or low birthweight can reduce the risk of death, blood infection and bowel sickness (NEC).

There was clinical heterogeneity in the use of probiotics and we could not recommend any single strain or combination of probiotics for wider use based of these results.

There is a lack of studies on probiotic supplementation in the 1st month of life in community settings. More research is needed to assess the effect of probiotics administered to neonates in home/ community settings in LMICs.

1.1.6 | How up-to-date is this review?

The review authors searched for studies published up to November 2019.

2 | BACKGROUND

The decline in rates of neonatal (age, 0–28 days) mortality has been slower than the decline in child mortality between 1990 and 2016 (Alkema et al., 2014; Bhutta et al., 2015). Neonatal mortality accounted for 46% of child mortality in 2016 compared to 40% of all under-five mortality rates in 1990 (WHO, 2017a). Globally, the percentage of neonatal mortality is the highest in South Asia and Sub-Saharan Africa (Alkema et al., 2014). Optimal nutritional support during the neonatal period is vital to the short and long term survival of the newborn (Bhutta et al., 2013; WHO, 2017b). Poor nutritional status of neonates is a major cause of illness and can lead to poor growth, increased risk of infection, bleeding, and neonatal death (Bhutta et al., 2013; WHO, 2017b). The risk of morbidity and mortality during the neonatal period is higher in LMICs where many

births happen at home and the prevalence of maternal malnutrition and incidence of low birth weight (birth weight <2500 g) and preterm birth (gestational age <37 weeks) is high (Bhutta et al., 2013; Lee et al., 2017; WHO, 2017b). This review focused on three nutritional interventions during neonatal periods that have the potential to reduce illness and death during infancy in LMIC.

2.1 | Description of the condition

The approach to nutritional management of newborn depends on maternal nutritional status, comorbidities during pregnancy (such as gestational diabetes), pregnancy duration (term vs. preterm birth), events at birth (such as birth asphyxia), birth weight (low birth weight vs. normal birth weight) and available resources for postpartum care of the mother and the baby (such as skilled birth attendant, home vs. facility birth, availability of neonatal intensive care) (Bhutta et al., 2013; WHO, 2015, 2017a, 2017b). The most important nutritional intervention after birth is breastfeeding, which is covered in a separate Campbell review of this series. There are a number of other nutritional interventions that have been proposed in addition to breastfeeding. It is beyond the scope of this review to comprehensively evaluate all the possible nutritional interventions during the neonatal period. We limited our review to the following three interventions: neonatal synthetic vitamin A supplementation, oral dextrose gel supplementation, and probiotic supplementation during the neonatal period in LMIC. Below in this section and in the rest of the introduction, we describe the rationale and importance of reviewing these interventions.

2.1.1 | Neonatal vitamin A deficiency (VAD)

Globally, about 190 million children and 19.1 million pregnant women are vitamin A deficient based on serum retinol levels (i.e., serum retinol <0.70 μ mol/L) (WHO, 2009a). VAD is most prevalent in South Asia and Africa (Stevens et al., 2015). VAD is associated with increased risk of blindness, infections, and mortality (Imdad et al., 2017). Most of the newborns are vitamin A deficient and rely on supplementation from maternal breast milk (Haider et al., 2017). High prevalence of maternal VAD in LMICs increases the risk of neonatal VAD. There has been interest in vitamin A supplementation during neonatal period to assess if it reduces risk of illness and death (Haider et al., 2017; WHO, 2009b), as it has been shown to reduce morbidity and mortality in children 6–59 months of age (Imdad et al., 2017).

2.1.2 | Hypoglycemia during the neonatal period

Hypoglycemia (low blood sugar) is common during the immediate neonatal period. The definition of neonatal hypoglycaemia varies. The American Academy of Pediatrics defines neonatal hypoglycaemia as blood glucose below 47 mg/dl (2.61 mmol/L); however, other societies such as the Pediatric Endocrine Society define neonatal hypoglycaemia as blood glucose <50 mg/dl (2.77 mmol/L; Thompson-Branch & Havranek, 2017; Thornton et al., 2015). Recurrent, severe, and/or persistent hypoglycaemia can lead to complications such as death; there is limited evidence to show that blood sugars below a certain level leads to long-term brain damage (Kaiser et al., 2015; McKinlav et al., 2017: Thornton et al., 2015). About 10-15% of otherwise healthy newborns have low blood sugar, and the rate is much higher among infants with additional risk factors such as large for gestational age, small for gestational age, low birth weight, preterm birth, infant of diabetic mother, and newborns with perinatal asphyxia (Thompson-Branch & Havranek, 2017). Additional risk factors for neonatal hypoglycaemia include neonatal sepsis, prolonged labor, and maternal medication use such as use of β-agonists and β-blockers (Thompson-Branch & Havranek, 2017). The recommended initial intervention to treat neonatal hypoglycaemia is to offer feeding in the form of breastfeeding followed by formula feeding if breastfeeding is unsuccessful. Persistent hypoglycaemia may require IV dextrose supplementation and admission to a neonatal intensive care unit (Thompson-Branch & Havranek, 2017; Thornton et al., 2015). In LMIC, where a significant proportion of births happen at home and incidence of low birth weight and preterm birth is high, prevention and treatment of hypoglycaemia encounters additional challenges (Singhal et al., 1991, 1992; WHO, 2017b; Williams, 1997). The instruments to test blood sugar might not be available in lowresource settings; In addition, formula, IV dextrose, and intensive care units might not be available to treat persistent and/or severe hypoglycaemia. Recent studies have tested simple interventions such as oral dextrose gel to treat neonatal hypoglycaemia and to prevent hypoglycaemia in high-risk newborns (Hegarty et al., 2016; Weston et al., 2016).

2.1.3 | Neonatal sepsis and NEC

Neonatal sepsis and NEC are neonatal morbidities that can be fatal (Oza et al., 2015; WHO, 2017b). Neonatal sepsis is the presence of an infection leading to systemic illness. Bacterial sepsis is common in LMIC and is a significant risk factor of morbidity and mortality in these countries (WHO, 2017a). NEC is a condition that occurs in newborns and can lead to intestinal injury and death. The extent of injury may vary from mucosal injury to full thickness intestinal wall injury. NEC happens most commonly in preterm babies and especially in extremely preterm babies (<28 weeks gestational age; AlFaleh & Anabrees, 2014; Patel & Denning, 2015). Multiple factors lead to the development of NEC in preterm infants including altered bacterial gut flora affecting the protective intestinal barrier, decreased intestinal motility, and the increased susceptibility of preterm infants to inflammation and infections (Patel & Denning, 2015). Recent studies have shown that an imbalance between commensal bacteria and pathogenic bacteria (intestinal dysbiosis) makes babies vulnerable to pathogenic bacterial growth in the intestine which then



causes inflammation that may contribute to neonatal sepsis and/or NEC (Arrieta et al., 2014; Deshmukh et al., 2014; Gewolb et al., 1999; Panigrahi et al., 2017). There is an increasing interest in correction of intestinal dysbiosis by probiotics to prevent NEC and neonatal sepsis. Data from early studies on probiotic use in neonates from LMIC is encouraging (AlFaleh & Anabrees, 2014; Rao et al., 2016).

2.2 | Description of the intervention

2.2.1 | Neonatal vitamin A supplementation

Vitamin A is a term used for a subclass of the family of fat soluble compounds named retinoic acids. Vitamin A is found in nature in two forms, provitamin A carotenoids and preformed vitamin A, which is essential to human bodily function. Plant-based foods are the main source of provitamin A carotenoids, of which β -carotene is the most commonly known. Animal-based foods are the main source of preformed vitamin A (Bates, 1995; Haider & Bhutta, 2011). Vitamin A from animal sources (retinol, retinal, retinoic acid, and retinyl esters) is the most active form, and synthetic vitamin A retinol has been used in most intervention trials in the past (Haider & Bhutta, 2011; Imdad et al., 2017). Plant-based foods may not be an adequate source of vitamin A, as the gastrointestinal conversion ratio from carotenoidto-retinol varies from 6:1 to 26:1 (US Institute of Medicine, Food and Nutrition Board). VAD may, therefore, exist in areas even when there is high consumption of plant-based foods such as in South Asia and Africa (Imdad et al., 2017; Stevens et al., 2015).

2.2.2 | Oral dextrose gel supplementation during neonatal period

Dextrose gel is a thickened aqueous solution that contains the concentrated simple carbohydrate dextrose. It can be administered by direct application to oral, buccal, or sublingual mucosa and can increase blood sugars rapidly by absorption through the highly vascularized and thin mucus membranes of the oral mucosa (Hegarty et al., 2016). Detxrose gel is a low cost nonproprietary intervention, and the gel can be prepared in hospital pharmacies. The typical ingredients include water, glucose, a gelling agent, and preservatives (Hegarty et al., 2016). The decision to use dextrose gel in a neonate should be taken on individual basis and should be avoided in neonates with compromised neurological or respiratory status (Hegarty et al., 2016; Weston et al., 2016).

2.2.3 | Probiotic supplementation during neonatal period

Prebiotics are supplements that promote the growth of commensal bacteria (AlFaleh & Anabrees, 2014; Panigrahi et al., 2017). Probiotics contain live bacteria that enrich pools of commensal bacteria

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(AlFaleh & Anabrees, 2014; Millar et al., 2003; Panigrahi et al., 2017). Synbiotics are a combination of prebiotics and probiotics and might have synergistic effect (Johnson-Henry et al., 2016, Nandhini et al., 2016; Panigrahi et al., 2017). These supplements are meant to optimise gut health and their hypothesized mechanisms of actions include enhanced gut barrier function, inhibition of gut colonization with pathogenic bacteria, improvement in colonization with healthy commensal bacteria that protect the infant from enteropathogenic infection through production of acetate, enhanced innate immunity, and increased maturation of the enteric nervous system (Rao et al., 2016). Recent data have shown that probiotic supplements can prevent the incidence of NEC in preterm babies (AlFaleh & Anabrees, 2014: Millar et al., 2003; Patel & Denning, 2015; van den Akker et al., 2018). There is also promising data on use of probiotocs/synbiotics for prevention of neonatal sepsis (Rao et al., 2016; Panigrahi et al., 2017). The most commonly used strains of probiotics include Lactobacillus and Bifidobacterium (Rao et al., 2016).

2.3 | How the intervention might work

2.3.1 | Neonatal vitamin A supplementation

Vitamin A has an effect on cell differentiation and helps maintain normal functioning of epithelial cells (Bates, 1995; Bhutta et al., 2013; Haider & Bhutta, 2011). It is considered anti-infective because it helps to maintain the protective epithelial barrier of the skin and mucosa, which protects the body from infections. Vitamin A helps in the regeneration of the epithelium and therefore maintains the integrity of the body's first line of defence. These mechanisms may help prevent infections in newborns (McCullough et al., 1999; Wolbach, 1933). Synthetic vitamin A supplementation has been shown to reduce morbidity and mortality in children 6-59 months of age (Imdad et al., 2017). The potential side effects of synthetic vitamin A supplementation include vomiting and bulging fontanelle (Imdad et al., 2016, 2017; Haider & Bhutta, 2011; Haider et al., 2017). Excess vitamin A supplementation can cause toxicity that presents in the form of a bulging fontanelle in children under 1 year, headaches, vomiting, diarrhea, loss of appetite, and irritability (Haider et al., 2017; Imdad et al., 2017).

2.3.2 | Oral dextrose gel supplementation during neonatal period

The absorption of dextrose gel through the oral mucosa leads to entry of glucose into lingual veins and into the internal jugular vein. This pathway provides almost immediate delivery of glucose to the systemic circulation and bypasses first pass liver metabolism through the portal circulation. If proven effective in preventing and treating hypoglycaemia, dextrose gel can avoid the need of intravenous glucose and reduce separation of baby from mother (Hegarty et al., 2016; Weston et al., 2016). The intervention is simple enough that it does not require special skills (such as IV placement) and can be administered by community, lay health workers, or the caregiver herself. Potential adverse effects include vomiting, choking, gagging, respiratory distress, and delay of treatment for severe hypoglycaemia (Hegarty et al., 2016; Weston et al., 2016).

2.4 | Probiotics supplementation during neonatal period

Newborn, and especially preterm, babies have immature intestines free of normal commensal bacteria that would normally protect them from developing NEC and sepsis by inhibiting the growth of pathogenic bacteria in the intestines (AlFaleh & Anabrees, 2014; Patel & Denning, 2015; Rao et al., 2016). Probiotics are used to proactively colonize the intestines with beneficial bacteria such as Lactobacillus species (Millar et al., 2003; Patel & Denning, 2015). Probiotics therefore reduce the growth of pathogenic bacteria which would otherwise increase the risk of NEC and sepsis. Also probiotics promote gut immunity by increasing IgA levels and contributing to improved mucosal barrier function (Patel & Denning, 2015). These protective mechanisms reduce intestinal permeability by producing a protective mucosal barrier against bacteria and increase the production of anti-inflammatory cytokines (Deshpande et al., 2017; Millar et al., 2003). Probiotics are especially protective in preterm babies with immature intestinal microbiomes and neonates on antibiotics; antibiotics may reduce bacterial diversity in the intestine and thus also dispose to colonization by pathogenic bacteria causing NEC. Prebiotics and probiotics can be given together in the form of a synbiotic to improve the gut flora and can potentially reduce allcause neonatal mortality (Johnson-Henry et al., 2016; Panigrahi et al., 2017). Probiotics are considered safe; however, there are concerns regarding probiotic supplementation in extremely premature or immunocompromised neonates. A few cases of neonatal sepsis have been reported that were thought to be caused by probiotics (Dani et al., 2016).

2.5 | Why it is important to do this review

2.5.1 | Neonatal vitamin A supplementation

Randomized trials on neonatal vitamin A supplementation have produced conflicting results with some studies (mostly from South Asia) showing a mortality benefit while no major benefit in other studies (mostly from Africa) (Haider et al., 2017) and some studies showing even an increased risk of infant mortality in certain populations (Smith et al., 2016).The exact reason for this difference in results is not clear. Previous reviews (Haider et al., 2017; Gogia & Sachdev, 2009) and a WHO technical consultation (WHO, 2009b) have hypothesized on what factors may explain these varied results. Our group has previously published a Cochrane review on the evidence on neonatal vitamin A supplementation (Haider & Bhutta, 2011), and we wanted to update the previous review. The previous review included studies conducted in the community setting. In this review, we considered studies conducted in both the community and the hospital setting in LMIC. We also included neurodevelopment outcomes for this review that were not covered in the previous Cochrane review.

2.5.2 | Oral dextrose gel supplementation during neonatal period

Oral dextrose gel has been studied in the prevention and treatment of neonatal hypoglycaemia in high-income countries (Hegarty et al., 2017; Weston et al., 2016); however, it was not clear if similar studies were available from LMICs. Our objective was to consider both randomized and nonrandomized observational studies with a control arm. We hypothesized that the use of dextrose may be more beneficial in LMIC than in high-income countries, as the incidence of neonatal hypoglycaemia might be higher in these countries due to an increased rate of preterm and low birth weight birth.

2.5.3 | Probiotics supplementation during neonatal period

The effect of probiotic supplementation for the prevention of NEC and neonatal sepsis has been assessed in previous reviews (AlFaleh & Anabrees, 2014; Rao et al., 2016; van den Akker et al., 2018). Most of these reviews included studies from both high- and LMICs. Deshpande et al. (2017) reviewed studies from LMIC where neonates were supplemented with probiotics. More studies (Amini et al., 2017; Chowdhury et al., 2016; Guney-Varal et al., 2017; Hernández-Enríquez et al., 2016; Hussain et al., 2016) have been published since the publication of Deshpande et al.'s (2017) review. Overall, our objective was to assess the current evidence for the effect of probiotic supplementation during the neonatal period in the hospital and community setting in LMIC.

3 | OBJECTIVES

3.1 | Primary objectives

To determine the efficacy of the following interventions on neonatal morbidity and mortality:

- 1. Synthetic vitamin A supplementation,
- 2. Oral dextrose gel supplementation, and
- 3. Oral probiotic supplementation.

A detailed description of background and methods for this review was published in the form of a protocol as Imdad et al. (2019).

4 | METHODS

4.1 | Criteria for considering studies for this review

4.1.1 | Types of studies

We included the following study designs:

- Randomized controlled trials (RCTs), where participants were randomly assigned either individually or in clusters to intervention and comparison groups. Cross-over designs were also eligible for inclusion.
- Quasi-experimental designs, which include:
- Natural experiments: studies where nonrandom assignment was determined by factors that were out of the control of the investigator. One common type includes allocation based on exogenous geographical variation.
- b. Controlled before-after studies (CBA), in which measures were taken of an experimental group and a comparable control group both before and after the intervention. We also require that appropriate methods were used to control for confounding, such as statistical matching (e.g., propensity score matching or covariate matching) or regression adjustment (e.g., difference-in-differences, instrumental variables).
- c. Regression discontinuity designs; here, allocation to intervention/ control was based upon a cut-off score.
- d. Interrupted time series studies, in which outcomes were measured in the intervention group at at least three time points before the intervention and after the intervention.

4.1.2 | Types of participants

Participants for this review included neonates (aged 0–28 days) from LMICs. We included neonates regardless of their health status. This includes low birth weight and preterm babies. However, studies that focused on neonates with congenital anomalies were excluded. We considered studies that included older age population groups in addition to neonates only if we could disaggregate relevant data for the neonatal population. For example, a study might include infants up to 6 months of age. We included such a study if the disaggregated data were available for neonates (0–28 days). Even though we planned to assess later childhood outcomes, we did not plan to include studies that recruited participants after the neonatal period.

4.1.3 | Types of interventions

The following interventions were included in the review:

1. Neonatal vitamin A supplementation compared to no supplementation or placebo: we considered only oral synthetic vitamin A

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supplementation. There was no restriction on the dosage and frequency of the medicine. The comparison group could include a placebo or standard of care.

- 2. Oral dextrose gel supplementation during the neonatal period compared to no supplementation: we placed no limits on the dose or frequency of the dextrose supplementation. We only considered dextrose gel as the intervention and excluded dextrose given in other forms such as intravenous, nasogastric tube, or mixed with infant formula. The reason to exclude forms other than dextrose gel was that administration of dextrose in those forms may require special circumstances (like trained staff to place an IV) or special delivery vehicles, such as formula, that may not be available in LMIC. The comparison group included placebo or standard of care.
- 3. Neonatal oral probiotics/synbiotics compared to no probiotic supplementation: probiotics are live microbial organisms that are given to promote the growth of commensal gut bacteria and prevent the growth of pathogenic bacteria. Prebiotics are dietary supplements that promote the growth of commensal bacteria. Synbiotics are a combination of prebiotics and probiotics (Millar et al., 2003; Patel & Denning, 2015). We placed no limits on the dose or frequency of probiotics. We included studies that used probiotics and synbiotics supplementation and excluded studies that used only prebiotics. Comparison groups included placebo or standard of care.

Each of the above interventions (i.e., vitamin A, dextrose, or probiotics) was summarized separately, and the interventions were not compared to each other directly or indirectly.

4.1.4 | Types of outcome measures

Primary outcomes The primary outcomes were:

- 1. All-cause neonatal mortality (death between 0 and 28 days of life)
- All cause infant mortality at 6 months (death between 0 days and 6 months of life)
- 3. All-cause infant mortality at 12 months (death between 0 days and 12 months life).

We anticipated that studies might not report the outcomes in the follow-up period mentioned above for the primary outcomes. If a study did not report mortality outcomes at day 28, 6 months, or 12 months, we contacted authors for data for the same. If segregated data were not available from authors, we included mortality data as follows: mortality in the first 6 weeks of life was included as neonatal mortality at day 28; between 3 and 6 months was included as 6 months, and between 9 and 12 months was included as 12 months. If the follow up was not clear, we included the mortality data at the longest follow-up.

Secondary outcomes

The secondary outcomes included:

- 1. Sepsis-specific mortality measured between 0 and 28 days, 0 days and 6 months, and 0 days and 12 months of life
- 2. Neonatal sepsis (as defined by authors) in the first 6 weeks of life
- 3. NEC as defined by authors
- 4. VAD
- 5. Prevention of Hypoglycemia (as defined by authors) during the neonatal period
- Treatment of Hypoglycemia (recurrence of hypoglycaemia after the episode treated)
- 7. Any adverse reactions during the intervention period
- 8. Serious adverse events
- 9. Neurodevelopmental outcomes at 12 and 24 months and at the longest follow-up.

The term neurodevelopment is a composite term that refers to cognitive, neurologic, and/or sensory outcomes. The term neurodevelopment may include intellectual disability as measured on the Mental Developmental Index of the Bayley Scales of Infant Development; gross motor delay measured on Gross Motor Function Classification System, and hearing and vision loss requiring amplification devices.

In order to be eligible for inclusion in the review, a study should have reported at least one of the primary or secondary outcomes. This was assessed at the full-text review stage.

Duration of follow-up. We included all participants in eligible studies that had outcomes of interest measured. There were no restrictions based on the duration of exposure, duration of followup, or timing of the outcome measurement. If the duration of treatment exceeded the neonatal period (i.e., 28 days), we considered another 2 weeks maximum but did not include studies in which the treatment went beyond 6 weeks of supplementation. We included mortality outcomes measured at 28 days, 6 months, 12 months of life, and at the longest follow-up as reported by authors.

Type of settings. We included studies conducted in LMIC. Low-income countries were defined as those with a gross national income (GNI) per capita of USD 1005 or less in 2016, and middle-income economies were those with a GNI per capita between USD 1006 and 3955 in 2016 (World Bank, 2017).

4.2 | Search methods for identification of studies

The identification of studies included various methods, such as electronic and other sources. We did not exclude any based on the outcome at the screening stages.

4.2.1 | Electronic searches

The electronic search for relevant studies was done in the following databases: PubMed, CINAHL, LILACS, SCOPUS, and CENTRAL (Cochrane Central Register for Controlled Trials).

Appendix 1 gives the search strategy for PubMed, CINAHL, LI-LACS, SCOPUS, and CENTRAL. It includes keywords and MeSH terms as appropriate. This approach includes a search strategy for the population (neonates) and interventions of interest. We planned to run searches for each intervention separately. We first ran the search for the population, which is the same for each intervention. Then we ran the search for each intervention. We then combined both searches by using "AND" and kept the searches in a separate EndNote file.

An example of a search strategy for vitamin A for PubMed was as follows:

(((("Vitamin A"[Mesh]) OR (Vitamin A[tiab] OR Aquasol A[tiab] OR Retinol[tiab] OR All Trans Retinol[tiab] OR All-Trans-Retinol[tiab] OR Vitamin A1[tiab] OR Vitamin A 1[tiab] OR 11-cis-Retinol[tiab] OR 11 cis Retinol[tiab] OR Tretinoin[tiab])AND Supplement*[tiab]))AND (("Infant"[Mesh] OR "Premature Birth"[Mesh]) OR (Neonat*[tiab] OR neo nat*[tiab]) OR (newborn* OR new Born*[tiab] OR newly born* [tiab]) OR (preterm[tiab] OR preterms[tiab] OR pre term[tiab] OR pre terms[tiab]) OR (premature*[tiab] AND (birth*[tiab] OR born[tiab] OR deliver*[tiab])) OR (low[tiab] AND (birthweight*[tiab]) OR birth weight*[tiab])) OR (lbw[tiab] OR vlbw[tiab] OR elbw[tiab]) OR infant* [tiab] OR (baby[tiab] OR babies[tiab])))) NOT ("Animals"[Mesh] NOT ("Animals"[Mesh] AND "Humans"[Mesh]))

We applied restriction of "humans" to searches. We did not apply any restrictions on searches based on outcomes, study design, or language. There was no restriction on date of publication.

The searches were conducted for vitamin A on December 10, 2018 (updated on November 13, 2019); probiotics on February 8, 2019 (updated on November 27, 2019); and dextrose on April 25, 2019 (updated on November 26, 2019).

4.2.2 | Searching other resources

Other resources included the search for ongoing trials at www. clinicaltrials.gov and WHO's ICTRP trials database. We also searched websites of international agencies such as WHO (including WHO's Reproductive Health Library), UNICEF, Global Alliance for Improved Nutrition (GAIN), International Food Policy Research Institute, International Initiative for Impact Evaluation (3ie), Nutrition International (NI), World Bank, USAID and USAID affiliates (e.g., FANTA, SPRING), and the World Food Programme.

Grey literature search sources included NI, GAIN, International Food Policy and Research Institute (IFPRI), and the WHO library database (WHOLIS).

We searched the reference lists of all included studies. We did citation searches of included studies in Google Scholar and Web of Science. We also searched the reference sections of previously published systematic reviews and the latest published studies. We contacted the experts and authors of the newest published studies to ask about any additional studies. Duplicates were removed.

4.3 | Data collection and analysis

4.3.1 | Selection of studies

Two authors independently screened titles/abstracts using prespecified inclusion/exclusion criteria. A full text was reviewed for the studies selected in the initial screening, and the same inclusion/exclusion criteria were applied. If there was a conflict about the inclusion of a study between the two reviewers, a third reviewer (ZAB) was consulted. We used a web-based software "Covidence" (Covidence, 2019) to do both title/abstract and full-text screening. This software allows simultaneous independent screening of studies, and inter-reviewer reliability can be assessed by checking the number of conflicts in the resolved conflict page following each stage of screening.

Description of methods used in primary research

We expected that the majority of the included studies would be randomized or cluster-randomized. We extracted the information on study design explicitly and made a careful differentiation between experimental and observational studies. We aimed to analyze randomized and nonrandomized studies separately.

Criteria for determination of independent findings

We anticipated that authors might report the results of a study in multiple publications. We coded such trials as a single study to avoid double counting of the data and included all the relevant outcomes decided a priori for this review. If a pilot study was done before the larger study, we included the two studies separately unless the data from the pilot study was included in the main trial. When a clinical trial registration number was available for a study, we searched that number on PubMed to locate all the published studies linked to that trial number.

4.3.2 | Data extraction and management

Details of study coding categories

The data from included studies were abstracted into a standardized data abstraction form by two authors. We extracted data in duplicates, and any discrepancies were resolved by discussion firs. A third reviewer (ZAB) was consulted if the conflict existed after the initial discussion.

The data extraction sheet had the following information.

- General study information: authors, publication year, study design
- Study setting: World Bank region, country, World Bank income level, city/town, urban/urban slum/rural/mixed setting, duration of data collection, date of data collection

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- Study population: sample size recruited, sample size analysed, female (%), description of participants (i.e., inclusion/exclusion criteria applied to recruitment)
- Intervention characteristics: type of intervention, duration of intervention, unit of randomization (where applicable), dose, frequency of provision, duration of follow up, attrition rate
- Quality assessment

Each quantitative outcome sheet contained the following:

- Subgroup (if applicable)
- Subgroup sample size
- Outcome type
- Outcome units
- Outcomes
- a. Outcome measure treatment group
- b. Outcome measure comparison group
- c. Standard deviation
- Effect size:
- a. Effect measure
- b. 95% CI

4.3.3 | Assessment of ROB in included studies

We used the Cochrane ROB tool (Higgins & Green, 2011) for randomized studies. The Cochrane ROB tool includes the following items:

- Selection bias: random sequence generation and allocation concealment
- Performance bias: blinding of participants and personnel
- Detection bias: blinding of outcome assessment
- Attrition bias: incomplete outcome data
- Reporting bias: selective reporting
- Other sources of bias

Two authors independently performed the ROB assessments for each study. A third reviewer was involved to resolve any disagreements (ZAB). An overall score was not provided.

4.3.4 | Measures of treatment effect

We performed a meta-analysis for the synthesis of quantitative data when the included studies had comparable participants, interventions, and outcomes. We did not assess the effect on outcome across the interventions, such as is done in network meta-analysis. Each intervention was analysed separately. We analysed continuous and dichotomous data separately. For dichotomous outcomes, results were presented as summary risk ratios with 95% CI. We combined risk ratios (events per child) and rate ratios (events per child year) for incidence data because of their similar interpretation and scale. For continuous outcomes, we presented the summary results as the mean difference with 95% CI when data were available on the same scale across the studies. We used the standardized mean difference with 95% CI when data were presented in different scales across the studies.

To avoid reviewer bias, we planned to predetermine the preference for specific data for certain outcomes. For example, for mortality outcomes, we gave preference to denominators in the following order: number with the definite outcome known, number randomized, and child-years. For morbidity data such as neonatal sepsis where both survivors and nonsurvivors might have contributed data, we gave preference to child years, number with the definite outcome known, and number randomized. For randomized trials, we gave preference to data that required the least manipulation by authors or inference by reviewers. We extracted the raw values (e.g., means and *SD*s) and built the intention-to-treat (ITT) analysis where applicable.

We anticipated that cause-specific morbidity or mortality data might not be readily available, as febrile illness due to respiratory, urinary, or central nervous system infection during the neonatal period are often categorized under a broader term of neonatal sepsis (WHO, 2017b).

4.3.5 | Unit of analysis issues

As we planned to include multiple interventions, all interventions and outcomes within those interventions, were meta-analyzed separately.

For randomized trials, we meta-analyzed individual and clusterrandomized trials in the same analysis. We assessed analyses in the cluster-randomized trials to ensure that clustering was appropriately accounted for within the analysis of the primary study, such that study precision was not over or under-estimated within our analysis. If the authors adjusted for cluster randomization, no further adjustment was made. In case a cluster-randomized study was not adjusted by primary authors, we adjusted effect estimates by using the mean cluster size (M) and the intra-cluster correlation coefficient (ICC) to calculate the design effect as follows: design effect = 1 + (M - 1) ICC. We then used the design effect to adjust the study data such that a trial was reduced to its effective sample size or standard error of the summary estimate was inflated. We used the ICC given in the published studies. If the ICC was not available from the published study, we contacted the authors for the same. If the ICC was not available from the authors, we used ICC from the similar studies done in the similar region and on a similar population or took it from the previously published reviews (Haider et al., 2017).

Multiple-arm trials

We included studies with multiple intervention arms, but we only included the arms that were eligible for the review. We selected one pair (with appropriate intervention and control group) that satisfied the inclusion criteria of the review and excluded the rest. In case there were more than two groups eligible for inclusion, we combined these groups into a single pair-wise comparison. In multiple-arm trials using two different doses of the same intervention, we combined the two groups to avoid double counting the participants in the control group.

4.3.6 | Dealing with missing data

Any missing data were noted including loss to follow-up and dropouts. The reasons for the missing data were taken from the studies, and if it was not mentioned in the studies, the authors were contacted for the same. If the authors reported the adjusted values for missing data, we used the adjusted values.

4.3.7 | Assessment of heterogeneity

Statistical heterogeneity was assessed using τ^2 , l^2 , and significance of the χ^2 test. We also assessed statistical heterogeneity by visually inspecting the forest plots.

4.3.8 | Assessment of reporting biases

A funnel plot and its symmetry were used to assess publication bias if the number of included studies for intervention was more than 10. If the funnel plot was suggestive of publication bias, we further investigated the publication bias with the use of Egger's test (Higgins & Green, 2011).

4.3.9 | Data synthesis

Synthesis procedures and statistical analysis

We used the software Review Manager 5.3 (Review Manager, 2019) to conduct the statistical analysis. For randomized trials, we followed the ITT analysis. If ITT was not available, and the author reported the analyses as specified in the protocol, we reconstructed the data to create an ITT analysis.

We used a random-effect model to account for expected heterogeneity in the intervention, comparisons, or setting within studies included in a given synthesis. We used the generic inverse variance method of meta-analysis for fixed effect models and random effect models. This method of meta-analysis gives weight to studies based on their variance in a way that a study with low variance gets a high weight and vice versa.

We interpreted the results of the meta-analysis based on p value at the 95% confidence level (a value <0.05 was considered statistically significant) and reported both significant and non-significant results. For subgroup analysis, we used an interaction

test to determine if there was a relevant difference in effect across subgroups.

We assessed the quality of overall evidence using the GRADE approach. This method of quality assessment considers study type, within-study ROB (methodological quality), directness of evidence, heterogeneity, precision of effect estimates, and risk of publication bias (Guyatt et al., 2011). We rated the quality of the body of evidence for each key outcome as "high," "moderate," "low," or "very low."

4.3.10 | Subgroup analysis and investigation of heterogeneity

Neonatal vitamin A supplementation

Although we had planned a number of subgroup analyses for neonatal vitamin A supplementation; however, a recent IPD analysis (West et al., 2019) covered both individual and study level subgroup analyses, so we did not perform any subgroup analysis for vitamin A supplementation at this stage

Neonatal probiotic supplementation

- 1. Gestational age: term and preterm
- 2. Strains used in probiotics: single strain versus multiple strain and of type of strain used in each probiotic
- 3. Strains used in probiotics: contains *Lactobacillus* versus *Bifidobacterium* versus both
- 4. Settings: community-based versus hospital setting
- 5. Type of feedings: breastmilk versus formula milk versus mixed.

Oral dextrose gel supplementation

- Gestational age: term and postterm versus late preterm (35–36 weeks) versus moderately preterm (30–34 weeks) versus extremely preterm (<30 weeks)
- 2. Dose: equal or <200 mg/kg versus >200 mg/kg
- 3. Frequency: one versus more than one dose
- Time of administration: ≤1 h of age versus after 1 h of age versus after 2 h of age.

4.3.11 | Sensitivity analysis

- High quality studies versus low quality studies. The quality of study was subjectively based on the ROB assessment. Even though we considered all the domains included in the Cochrane ROB tool, we gave higher importance to sequence generation and allocation concealment, as most of the outcomes for this review were objective, and it was less likely that the results of the included studies would have been biased by a lack of blinding.
- 2. Random versus fixed effect models. We chose this sensitivity analysis to assess if the summary estimates will change

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significantly based on use of random versus fixed effect model. There is no exact criterion to choose between the two models, and we wanted to make sure that estimates were not significantly different between the two models.

Treatment of qualitative research We did not plan to include qualitative research.

4.3.12 | Summary of findings and assessment of the certainty of the evidence

Summary of findings' tables

We constructed "Summary of findings" tables for all of the primary outcomes using the Grading of Recommendations Assessment, Development and Evaluation (GRADE) criteria (GRADEpro GDT 2015). These covered consideration of within-study ROB (methodological quality), directness of evidence, heterogeneity, precision of effect estimates and risk of publication bias. We rated the certainty of evidence for each key outcome as "high," "moderate," "low," or "very low." The GRADE evidence is described in Table 1. Nonrandomised studies were initially rated as "low" quality. If there were no serious methodological flaws, we upgraded the evidence for studies with a large magnitude of effect; presence of a dose response relationship; and effect of plausible residual confounding.

We used GRADE and prepared the summary of findings tables for the following primary outcomes:

- Stillbirth defined as baby born with no signs of life at or after 28 weeks' gestation
- Perinatal mortality (stillbirth and deaths ≤7 days)
- Neonatal mortality (death < 28 days)
- Infant mortality (deaths between 0 and 12 months)
- Under-five mortality (deaths between 0 and 59 months)
- Miscarriage
- Mean maternal body mass index

5 | RESULTS

5.1 | Description of studies

5.1.1 | Results of the search

Figure 1 shows the PRISMA flow diagram for our literature search.

Vitamin A supplementation during neonatal period

Sixteen studies reported in 45 publications assessed the effect of neonatal vitamin A supplementation (Ahmad et al., 2019; Basu et al., 2019; Benn et al., 2008, 2010, 2014; Edmond et al., 2015; Giridhar et al., 2019; Humphrey et al., 1996; Klemm et al., 2008; Malaba et al., 2005; Masanja et al., 2015; Mazumder et al., 2015; Rahmathullah et al., 2003; Soofi et al., 2017; Sun et al., 2019; West et al., 1995). These studies included a total of about 16,366 participants. All the studies were RCTs and published in a peer-reviewed journal.

Dextrose gel supplementation during neonatal period

We did not identify any studies that assessed the use of dextrose gel supplementation during neonatal period for prevention or treatment of hypoglycaemia in LMIC.

Probiotic supplementation during neonatal period

Thirty-three studies reported in 37 publications evaluated the effect of probiotic supplementation during the neonatal period and included a total of 11,595 participants (probiotics, 5854 and controls, 5741; Amini et al., 2017; Braga et al., 2011; Chowdhury et al., 2016; Cooper et al., 2017; Cui et al., 2019; Dashti et al., 2014; Demirel et al., 2013; Dilli et al., 2015; Dongol Singh et al., 2017; Dutta et al., 2015; Fernández-Carrocera et al., 2013; Hariharan et al., 2016; Kaban et al., 2019, Mazumder et al., 2015; Nandhini et al., 2016; Guney-Varal et al., 2017; Oncel et al., 2014; Panigrahi et al., 2017; Rojas et al., 2012; Roy et al., 2014; Samanta et al., 2009; Shashidhar et al., 2017; Sinha et al., 2015; Sari et al., 2011; Serce et al., 2013; Shadkam et al., 2015; Saengtawesin et al., 2014; Tewari et al., 2015; Xu et al., 2016).

Three of the studies were available in the form of abstracts (Hariharan et al., 2016; Huaxian, 2013; Rehman et al., 2018). The rest of the studies were published in a peer-reviewed journal.

5.1.2 | Included studies

The characteristics of included studies are available in the table Characteristics of included studies.

Vitamin A supplementation during neonatal period

Type of studies. All the included studies were RCTs. Thirteen studies were individually randomized (Ahmad et al., 2019; Basu et al., 2019; Benn et al., 2008, 2010, 2014; Edmond et al., 2015; Giridhar et al., 2019; Humphrey et al., 1996; Malaba et al., 2005; Masanja et al., 2015; Mazumder et al., 2015; Rahmathullah et al., 2003), and three studies were cluster-randomized (Klemm et al., 2008; Soofi et al., 2017; Sun et al., 2019; West et al., 1995). Three trials had multiple arms of interventions (Benn et al., 2010, 2014; Malaba et al., 2005).

Country. Studies were conducted in 10 different countries with four studies conducted in India (Basu et al., 2019; Giridhar et al., 2019; Mazumder et al., 2015; Rahmathullah et al., 2003), three studies in Guinea-Bissau (Benn et al., 2008, 2010, 2014), two studies in Bangladesh (Ahmad et al., 2019; Klemm et al., 2008), and one each in China (Sun et al., 2019), Ghana (Edmond et al., 2015), Indonesia (Humphrey et al., 1996), Nepal (West et al., 1995), Pakistan (Soofi et al., 2017), Tanzania (Masanja et al., 2015), and Zimbabwe (Malaba et al., 2005).

TABLE 1 Effect of probiotic supplementation during neonatal period: Subgroup analysis



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Alt-cause mortality: subgroup analysis: type of feedings Baby received breastmilk only 14 0.41 (0.62, 1.05) $p = .44$ Baby received formula milk only 1 1.38 (0.31, 6.06) $l^2 = 0\%$ Baby received both both breastmilk and formula milk 8 0.69 (0.48, 0.99) $l^2 = 0\%$ Type of feeding was unclear 3 1.33 (0.63, 2.81) $l^2 = 0\%$ Preparation contained Bil/dobacterium 1 0.43 (0.17, 1.09) $l^2 = 0\%$ Preparation contained Bil/dobacterium 1 0.43 (0.17, 1.09) $l^2 = 0\%$ Bil/dobacterium 1 0.43 (0.17, 1.09) $l^2 = 0\%$ Preparation contained bil/dobacterium 2 1.12 (0.46, 2.71) $l^2 = 0\%$ Necrotizing enterocolitis: subgroup analysis: probiotic preparation 1 0.29 (0.25, 0.61) $p = .05$ Preparation contained bil/dobacterium 1 0.20 (0.09, 0.47) $l^2 = 60.5\%$ Bil/dobacterium 1 0.20 (0.09, 0.47) $l^2 = 60.5\%$ Preparation contained bil/dobacterium 1 0.20 (0.09, 0.47) $l^2 = 60.5\%$ Bid/dobacterium 1 0.21 (0.03, 1.06) $p = .74$ Baby received formula milk	Study include preterm/low birth weight babies	24	0.79 [0.65, 0.95]	p = .47
Baby received breastmilk only 14 0.81 [0.62, 1.05] p 44 Baby received formula milk only 1 3.133 [0.31, 0.63] p Type of feeding was unclear 3 1.33 [0.63, 2.81] p Preparation contained Lactobacillus 10 0.82 [0.63, 1.05] p = .47 Preparation contained both Lactobacillus and 13 0.71 [0.47, 1.08] l^2 = 0% Bifdobacterium 1 0.43 [0.17, 1.09] l^2 = 0% Preparation contained both Lactobacillus and 13 0.71 [0.47, 1.08] l^2 = 0% Bifdobacterium 1 0.20 [0.09, 0.47] p = .05 Preparation contained Lactobacillus and 14 0.49 [0.36, 0.68] l^2 = 60.5% Bifdobacterium 1 0.20 [0.09, 0.47] p = .05 Preparation contained Lactobacillus and 14 0.49 [0.31, 0.59] p = .74 Baby received breastmilk only 1 0.20 [0.09, 0.47] p Preparation contained Subulardifi only 2 0.41 [0.11, 1.00] p Baby received breastmilk only complanalysis: type of feeding 2	Study included term infants only	1	1.38 [0.31, 6.08]	$I^2 = 0\%$
Baby received formula milk only 1 1.38 [0.31, 6.08] //* = 0% Baby received both both breastmilk and formula milk 8 0.69 [0.48, 0.99] /* = 0% Baby received formula milk 3 1.33 [0.63, 2.81] /* 0 All-cause mortality: subgroup analysis: probiotics preparation 1 0.43 [0.17, 1.09] p= .47 Preparation contained bath Lockbacillus and 13 0.43 [0.17, 1.09] /* = 0% Bithdobacterium 1 0.43 [0.17, 1.09] /* = 0% Bithdobacterium 1 0.43 [0.27, 1.047, 1.08] /* = 0% Bithdobacterium 1 0.42 [0.63, 0.68] /* = 0% Preparation contained bath Lockbacillus and 14 0.49 [0.36, 0.68] /* = 60.5% Bithdobacterium 1 0.42 [0.03, 1.05] p = .74 Preparation contained Schulardii only 1 0.43 [0.31, 0.59] p = .74 Bithdobacterium 1 0.43 [0.31, 0.59] p = .74 Bithdobacterium 1 0.43 [0.31, 0.59] p = .74 Bithdobacterium 1 0.43 [0.31, 0.59] p = .74 </td <td>All-cause mortality: subgroup analysis: type of feedings</td> <td></td> <td></td> <td></td>	All-cause mortality: subgroup analysis: type of feedings			
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Type of feeding was unclear 3 1.33 [0.63, 2.81] All-cause mortality: subgroup analysis: probiotics preparation Preparation contained Lactobacillus 10 0.82 [0.63, 1.05] p = 47 Preparation contained Lactobacillus and 13 0.71 [0.47, 1.08] l² = 0% Bildobacterium Preparation contained Saccharomyces boulardii only 2 1.12 [0.46, 2.71] Preparation contained Lactobacillus and 13 0.39 [0.25, 0.61] p = .05 Preparation contained Saccharomyces boulardii only 2 0.42 [0.09, 0.47] Preparation contained Lactobacillus and 14 0.20 [0.09, 0.47] Preparation contained bath Lactobacillus and 14 0.49 [0.36, 0.68] P= 605 Preparation contained Lactobacillus and 14 0.49 [0.31, 0.59] p = .05 Preparation contained Saccharomyces brule freeding 13 0.43 [0.31, 0.59] p = .74 Baby received borts brule diffuebacterium 13 0.43 [0.31, 0.59] p = .74 Baby received borts brule diffuebacterium 10 0.43 [0.31, 0.59] p = .74 Baby received borts brule diffuebacterium 10 0.43 [0.31, 0.57] p = .76 <t< td=""><td>Baby received formula milk only</td><td>1</td><td>1.38 [0.31, 6.08]</td><td></td></t<>	Baby received formula milk only	1	1.38 [0.31, 6.08]	
All-cause mortality: subgroup analysis: probiotics preparation100.82 [0.63, 1.05] $p = .47$ Preparation contained Lactobacillus and130.71 [0.47, 1.08] $l^2 = 0\%$ Bifidobacterium130.71 [0.47, 1.08] $l^2 = 0\%$ Preparation contained back backbackliks and130.71 [0.47, 1.08] $l^2 = 0\%$ Bifidobacterium10.39 [0.25, 0.61] $p = .05$ Preparation contained backbackliks and140.49 [0.36, 0.68] $l^2 = 60.5\%$ Bifidobacterium10.20 [0.09, 0.47] $p = .05$ Preparation contained bifidobacterium140.49 [0.36, 0.68] $l^2 = 60.5\%$ Bifidobacterium10.20 [0.09, 0.47] $p = .74$ Preparation contained bifidobacterium130.43 [0.31, 0.59] $p = .74$ Preparation contained S. boulardii only20.94 [0.45, 1.95] $p = .74$ Raby received formula only10.21 [0.03, 1.76] $p = .74$ Baby received formula only10.21 [0.03, 1.76] $p = .50$ Preparation contained a single strain of probiotics150.48 [0.30, 0.76] $p = .50$ Preparation contained a single strain of probiotics150.48 [0.35, 0.67] $l^2 = 0\%$ Preparation contained a single strain of probiotics150.48 [0.35, 0.67] $l^2 = 0\%$ Preparation contained a single strain of probiotics150.48 [0.35, 0.67] $l^2 = 0\%$ Preparation contained a single strain of probiotics160.74 [0.62, 0.87] $p = .79$ Preparation contained synbiotics10 </td <td>Baby received both both breastmilk and formula milk</td> <td>8</td> <td>0.69 [0.48, 0.99]</td> <td>$l^2 = 0\%$</td>	Baby received both both breastmilk and formula milk	8	0.69 [0.48, 0.99]	$l^2 = 0\%$
Preparation contained Lactobacillus100.82 [0.63, 1.05] $p = .47$ Preparation contained Bifdobacterium10.43 [0.17, 1.09] $l^2 = 0\%$ Preparation contained bif Lactobacillus and130.71 [0.47, 1.08] $l^2 = 0\%$ Bifdobacterium21.12 [0.46, 2.71] $l^2 = 0\%$ Preparation contained Saccharomyces boulardii only21.21 [0.46, 2.71]Necrotizing enterocolitis: subgroup analysis: probiotic preparation $0.39 [0.25, 0.61]$ $p = .05$ Preparation contained Bifdobacterium10.20 [0.09, 0.47] $l^2 = 60.5\%$ Preparation contained bif Lactobacillus and140.49 [0.36, 0.68] $l^2 = 60.5\%$ Bifdobacterium10.21 [0.03, 1.76] $l^2 = 0\%$ Preparation contained S. boulardii only20.94 [0.45, 1.95] $l^2 = 0\%$ Necrotizing enterocolitis: subgroup analysis: type of feeding $l^2 = 0.5 [0.33, 0.57]$ $p = .74$ Baby received formula only10.21 [0.03, 1.76] $l^2 = 0\%$ Baby received formula only10.21 [0.03, 0.76] $p = .50$ Type of feeding was unclear70.41 [0.17, 1.00] $l^2 = 0\%$ Necrotizing enterocolitis: subgroup analysis: type of probiotics120.48 [0.30, 0.76] $p = .50$ Preparation contained a single strain of probiotics120.48 [0.30, 0.76] $p = .50$ Preparation contained a single strain of probiotics120.48 [0.30, 0.76] $p = .50$ Preparation contained single strain of probiotics120.48 [0.30, 0.68] $l^2 = 0\%$ Prepara	Type of feeding was unclear	3	1.33 [0.63, 2.81]	
Preparation contained bif/dobacterium10.43 [0.17, 1.09] 0.7 [0.47, 1.08]1² = 0% 2² = 0%Preparation contained both Lactobacillus and Bifidobacterium21.12 [0.46, 2.71]Preparation contained Saccharomyces boulardii only20.39 [0.25, 0.61] 0.29 [0.25, 0.61]p = 0.5Preparation contained Lactobacillus10.20 [0.09, 0.47] 1² = 60.5%p = 0.5Preparation contained bifidobacterium10.49 [0.36, 0.68]l² = 60.5%Preparation contained bifidobacterium20.49 [0.36, 0.68]p = .74Preparation contained S. boulardii only20.43 [0.31, 0.59]p = .74Baby received breastmilk only130.43 [0.31, 0.59]p = .74Baby received formula only10.21 [0.03, 1.76]Baby received formula only10.21 [0.03, 1.76]Baby received formula only10.48 [0.30, 0.76]p = .50Preparation contained single strain of probioticsP = .50Preparation contained single strain of probiotics150.48 [0.30, 0.76]p = .50Preparation contained single strain of probiotics120.48 [0.30, 0.76]p = .50Preparation contained single strain of probiotics120.48 [0.30, 0.66]l² = 0%Preparation contained bifidobacterium130.43 [0.31, 0.59]p = .50Preparation contained single strain of probiotics150.48 [0.30, 0.67]p = .50Preparation contained single strain of probiotics150.48 [0.30, 0.67]p = .50Preparation contained bifidobacterium14 <t< td=""><td>All-cause mortality: subgroup analysis: probiotics preparation</td><td></td><td></td><td></td></t<>	All-cause mortality: subgroup analysis: probiotics preparation			
Preparation contained both Lactobacillus and Bildibacterium 13 $0.71 [0.47, 1.06]$ $l^2 = 0\%$ Bildibacterium Preparation contained Saccharomyces boulardii onjy 1 $1.12 [0.46, 2.71]$ Necrotizing enterocolitis: subgroup analysis: probiotic preparation 13 $0.39 [0.25, 0.61]$ $p = .05$ Preparation contained Lactobacillus and Bildibbacterium 14 $0.20 [0.09, 0.47]$ $p = .05$ Preparation contained both Lactobacillus and Bildibbacterium 14 $0.20 [0.09, 0.47]$ $p = .05$ Preparation contained Suburdii only 2 $0.94 [0.45, 1.95]$ $p = .05$ Necrotizing enterocolitis: subgroup analysis: type of feeding $p = .74$ $p = .74$ Baby received formula only 13 $0.21 [0.03, 1.76]$ $p = .74$ Baby received formula only 12 $0.48 [0.30, 0.76]$ $p = .50$ Preparation contained a single strain of probiotics 12 $0.48 [0.30, 0.67]$ $p = .50$ Preparation contained Milpib strains of probiotics 13 $0.28 [0.12, 0.67]$ $p = .79$ Preparation contained Milpib strain of probiotics 12 $0.48 [0.30, 0.67]$ $p = .50$ Preparation contained Milpib strain of probiotics 12 $0.48 [0.3$	Preparation contained Lactobacillus	10	0.82 [0.63, 1.05]	p = .47
Bifdobacterium21.12 [0.46, 2.71]Preparation contained Saccharomyces boulardii only21.12 [0.46, 2.71]Necrotizing enterocolitis: subgroup analysis: probiotic preparation130.39 [0.25, 0.61] $p = .05$ Preparation contained Bifdobacterium10.20 [0.09, 0.47] $p = .05$ Preparation contained Bifdobacterium140.49 [0.36, 0.68] $l^2 = 60.5\%$ Bifdidbacterium20.94 [0.45, 1.95] $p = .74$ Preparation contained S. boulardii only130.43 [0.31, 0.59] $p = .74$ Baby received breastmilk only130.43 [0.31, 0.59] $p = .74$ Baby received breastmilk and formula milk90.55 [0.33, 0.92] $l^2 = 0\%$ Type of feeding was unclear70.410.11 $p = .50$ Preparation contained single strain of probiotics120.48 [0.30, 0.76] $p = .50$ Preparation contained single strain of probiotics120.48 [0.30, 0.76] $p = .50$ Preparation contained symbiotics120.48 [0.30, 0.67] $l^2 = 0\%$ Preparation contained symbiotics110.74 [0.62, 0.87] $p = .79$ Preparation contained symbiotics110.74 [0.62, 0.87] $p = .79$ Preparation contained bif Bifdobacterium10.81 [0.60, 1.09] $l^2 = 0\%$ Preparation contained bith Bifdobacterium30.73 [0.57, 0.44]Preparation contained bith Bifdobacterium30.73 [0.57, 0.94]Preparation contained S. boulardii only20.59 [0.22, 1.56]Preparation contained S. b	Preparation contained Bifidobacterium	1	0.43 [0.17, 1.09]	
Necrotizing enterocolitis: subgroup analysis: probiotic preparation Preparation contained Lactobacillus and 13 0.39 [0.25, 0.61] $p = .05$ Preparation contained both Lactobacillus and 14 0.49 [0.36, 0.68] $l^2 = 60.5\%$ Biffdbbacterium Preparation contained both Lactobacillus and 14 0.49 [0.36, 0.68] $l^2 = 60.5\%$ Biffdbbacterium Preparation contained S. boulardii only 2 0.94 [0.45, 1.95] Necrotizing enterocolitis: subgroup analysis: type of feeding Baby received breastmilk only 1 0.21 [0.03, 1.76] Baby received formula only 1 0.21 [0.03, 0.76] $p = .74$ Baby received formula only 1 0.21 [0.03, 0.76] $l^2 = 0\%$ Nype of feeding was unclear 7 0.41 [0.17, 1.00] Necrotizing enterocolitis: subgroup analysis: type of probiotics Preparation contained a single strain of probiotics 15 0.48 [0.30, 0.76] $p = .50$ Preparation contained a single strain of probiotics 15 0.48 [0.30, 0.76] $p = .50$ Preparation contained synbiotics 0 12 0.48 [0.30, 0.76] $p = .50$ Preparation contained synbiotics 15 0.48 [0.35, 0.67] $l^2 = 0\%$ Necrotizing enterocolitis: subgroup analysis: type of probiotics 15 0.48 [0.30, 0.76] $p = .79$ Preparation contained Bifidbbacterium 1 0.74 [0.62, 0.87] $p = .79$ Preparation contained Elifdbbacterium 1 0.81 [0.60, 1.09] Preparation contained Bifidbbacterium and 6 0.38 [0.68, 1.02] $l^2 = 0\%$ Preparation contained S. boulardii only 3 0.73 [0.57, 0.94] Neonatal sepsis: subgroup analysis: type of feeding Baby received both formula and breastmilk only 8 0.71 [0.61, 0.83] $p = .04$ Baby received breastmilk only 6 0.77 [0.65, 0.90] $l^2 = 65\%$ Type of feeding was unclear 4 0.95 [0.82, 1.09]	Preparation contained both <i>Lactobacillus</i> and <i>Bifidobacterium</i>	13	0.71 [0.47, 1.08]	$I^2 = 0\%$
Preparation contained Lactobacillus 13 0.39 [0.25, 0.61] $p = .05$ Preparation contained Bifidobacterium 1 0.20 [0.09, 0.47] Preparation contained both Lactobacillus and 14 0.49 [0.36, 0.68] $l^2 = 60.5\%$ Bifidobacterium Preparation contained S. boulardii only 2 0.94 [0.45, 1.95] Pereparation contained S. boulardii only 13 0.43 [0.31, 0.59] $p = .74$ Baby received breastmilk only 13 0.43 [0.31, 0.59] $p = .74$ Baby received formula only 1 0.21 [0.03, 1.76] Peparation contained S. boulardii only Necrotizing enterocolitis: subgroup analysis: type of probiotics Preparation contained a single strain of probiotics Preparation contained a single strain of probiotics P= 50 Preparation contained symbiotics 12 0.48 [0.30, 0.76] $p = .50$ Preparation contained symbiotics 12 0.48 [0.30, 0.76] $p = .50$ Preparation contained symbiotics 12 0.48 [0.30, 0.76] $p = .50$ Preparation contained symbiotics 12 0.48 [0.30, 0.76] $p = .79$ Preparation contained symbiotics 10 0.74 [0.62, 0.87] $p = .79$ Preparation contained bifidobacterium	Preparation contained Saccharomyces boulardii only	2	1.12 [0.46, 2.71]	
Preparation contained Bifidobacterium 1 0.20 [0.09, 0.47] Preparation contained both Lactobacillus and Bifidobacterium 14 0.49 [0.36, 0.68] l^2 = 60.5% Preparation contained S. boulardii only 2 0.94 [0.45, 1.95] 1 Necrotizing enterocolitis: subgroup analysis: type of feeding Baby received breastmilk only 13 0.43 [0.31, 0.59] $p = .74$ Baby received breastmilk and formula milk 9 0.55 [0.33, 0.92] $l^2 = 0\%$ Type of feeding was unclear 7 0.41 [0.17, 1.00] 12 Necrotizing enterocolitis: subgroup analysis: type of probiotics Preparation contained a single strain of probiotics 12 0.48 [0.30, 0.76] $p = .50$ Preparation contained a single strain of probiotics 12 0.48 [0.30, 0.76] $p^2 = 0\%$ Preparation contained symbiotics (prebiotics + probiotics) 3 0.28 [0.12, 0.67] $p^2 = .9\%$ Preparation contained both Bifdobacterium and Lactobacillus 11 0.74 [0.62, 0.87] $p = .79$ Preparation contained both Bifdobacterium and Lactobacillus 6 0.33 [0.68, 1.02] $l^2 = 0\%$ Preparation contained both Bifdobacterium and Lactobacillus 0.33 [0.68, 1.02] $l^2 = 0\%$ Preparation contained S.	Necrotizing enterocolitis: subgroup analysis: probiotic preparation			
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BifidobacteriumPreparation contained S. boulardii only20.94 [0.45, 1.95]Necrotizing enterocolitis: subgroup analysis: type of feeding 13 0.43 [0.31, 0.59] $p = .74$ Baby received breastmilk only10.21 [0.03, 1.76] $p = .74$ Baby received formula only10.21 [0.03, 1.76] $p = .74$ Baby received both breastmilk and formula milk90.55 [0.33, 0.92] $l^2 = 0\%$ Necrotizing enterocolitis: subgroup analysis: type of probiotics70.41 [0.17, 1.00]Necrotizing enterocolitis: subgroup analysis: type of probiotics120.48 [0.30, 0.76] $p = .50$ Preparation contained a single strain of probiotics150.48 [0.30, 0.67] $l^2 = 0\%$ Preparation contained synbiotics150.48 [0.30, 0.67] $l^2 = 0\%$ Preparation contained synbiotics100.74 [0.62, 0.87] $p = .50$ Preparation contained synbiotics110.74 [0.62, 0.87] $p = .79$ Preparation contained lactobacillus10.74 [0.62, 0.87] $p = .79$ Preparation contained bifidobacterium and Lactobacillus60.83 [0.68, 1.02] $l^2 = 0\%$ Preparation contained S. boulardii only30.75 [0.94] $l^2 = 0\%$ Neonatal sepsis: subgroup analysis: type of feeding Baby received formula milk only80.71 [0.61, 0.83] $p = .04$ Baby received formula milk only60.77 [0.65, 0.90] $l^2 = 6\%$ Baby received formula milk only60.77 [0.65, 0.90] $l^2 = 6\%$ Baby received formula milk on	Preparation contained Bifidobacterium	1		
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Baby received breastmilk only 8 0.71 [0.61, 0.83] p = .04 Baby received formula milk only 2 0.59 [0.22, 1.56] Baby received both formula and breastmilk only 6 0.77 [0.65, 0.90] l ² = 65% Type of feeding was unclear 4 0.95 [0.82, 1.09] l	Preparation contained S. boulardii only	3	0.73 [0.57, 0.94]	
Baby received formula milk only 2 0.59 [0.22, 1.56] Baby received both formula and breastmilk only 6 0.77 [0.65, 0.90] $l^2 = 65\%$ Type of feeding was unclear 4 0.95 [0.82, 1.09] $l^2 = 65\%$	Neonatal sepsis: subgroup analysis: type of feeding			
Baby received both formula and breastmilk only6 $0.77 [0.65, 0.90]$ $l^2 = 65\%$ Type of feeding was unclear4 $0.95 [0.82, 1.09]$	Baby received breastmilk only	8	0.71 [0.61, 0.83]	<i>p</i> = .04
Type of feeding was unclear 4 0.95 [0.82, 1.09]	Baby received formula milk only	2	0.59 [0.22, 1.56]	
	Baby received both formula and breastmilk only			$l^2 = 65\%$
Neonatal sepsis: type of probiotics	Type of feeding was unclear	4	0.95 [0.82, 1.09]	
	Neonatal sepsis: type of probiotics			

TABLE 1 (Continued)

Outcome or subgroup	No. of studies	Effect estimate: relative risk	Test for subgroup difference
Preparation contained single strain of probiotics	8	0.84 [0.74, 0.96]	p = .21
Preparation contained multiple strains of probiotics	9	0.81 [0.68, 0.97]	$l^2 = 35\%$
Preparation contained synbiotics (prebiotics + probiotics)	4	0.67 [0.54, 0.83]	
Neonatal sepsis: subgroup analysis: settings			
Hospital based	19	0.83 [0.76, 0.91]	p = .19
Community based	2	0.67 [0.49, 0.91]	$I^2 = 42\%$

Settings. Thirteen studies were conducted in the community setting, while three studies were conducted in the hospital setting (Basu et al., 2019; Giridhar et al., 2019; Sun et al., 2019).

Participants. Most of the studies included live born infants who were otherwise healthy. One study included only low birth weight babies (Benn et al., 2010), two studies included newborns with very low birth weight (Basu et al., 2019; Giridhar et al., 2019), and one study included extremely premature babies (Sun et al., 2019). The sample size of each study ranged from 120 (Giridhar et al., 2019) to 44,948 (Mazumder et al., 2015).

Dose. Most of the included studies for the use of neonatal vitamin A supplementation used a dose of 50,000 IU. Rahmathullah et al. (2003) gave 24,000 IU daily, and Benn et al. (2010) used 25,000 IU. Benn et al. (2014) compared doses of 50,000 IU versus 25,000 IU. Basu et al. (2019) used a daily dose of 1,500 IU.

Comparison. In all the included studies for neonatal vitamin A supplementation a placebo was given to the control group.

Probiotic supplementation during neonatal period

Type of studies. All the studies that evaluated the effect of probiotic supplementation during the neonatal period were individual RCTs. Two studies had multiple intervention groups (Dilli et al., 2015; Dutta et al., 2015). One of these studies compared different combinations of probiotics with prebiotics (Dilli et al., 2015), and the other study compared different doses of probiotics (Dutta et al., 2015). For the study by Dilli et al. (2015), we included the data in a way that the only difference between the two groups was probiotics. For the study by Dutta et al. (2015), we combined all the groups that compared different doses and compared them with the placebo to avoid double-counting of the placebo group data.

One study included neonates with and without exposure to human immunodeficiency virus (HIV; based on maternal history of HIV). We included the data for these groups separately in the metaanalysis (Niekerk et al., 2015 (HIV exposed); Niekerk et al., 2015 (HIV nonexposed).

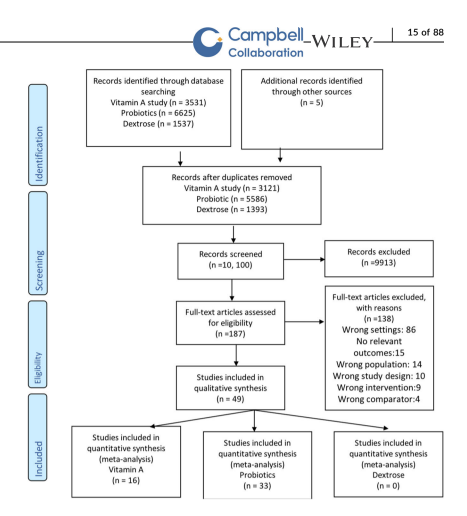
Country. Studies were conducted in 13 different countries with 10 studies conducted in India (Dutta et al., 2015; Hariharan et al., 2016; Mazumder et al., 2015; Nandhini et al., 2016; Panigrahi et al., 2017; Roy et al., 2014; Samanta et al., 2009; Shashidhar et al., 2017; Sinha et al., 2015; Tewari et al., 2015), six studies in Turkey (Demirel et al., 2013; Dilli et al., 2015; Guney-Varal et al., 2017; Oncel et al., 2014; Sari et al., 2011; Serce et al., 2013), three studies each in China (Cui et al., 2019; Huaxian, 2013; Xu et al., 2016) and Iran (Amini et al., 2017; Dashti et al., 2014; Shadkam et al., 2015), two each in Mexico (Fernández-Carrocera et al., 2013; Hernández-Enríquez et al., 2016) and South Africa (Cooper et al., 2017; Niekerk et al., 2015 (HIV exposed)), and one each in Bangladesh (Chowdhury et al., 2016), Brazil (Braga et al., 2011), Colombia (Rojas et al., 2012), Indonesia (Kaban et al., 2019), Nepal (Dongol Singh et al., 2017), Pakistan (Hussain et al., 2016), and Thailand (Saengtawesin et al., 2014).

Settings. All the studies were conducted in the hospital setting except for three studies where participants were followed in the community setting (Cooper et al., 2017; Dongol Singh et al., 2017; Panigrahi et al., 2017).

Participants. Only one study (Cooper et al., 2017) included neonates that were full term. The rest of the studies included participants that were either low birth weight, preterm, or both. The participants were recruited from neonatal intensive care units except in three studies (Cooper et al., 2017; Dongol Singh et al., 2017; Panigrahi et al., 2017), where participants were recruited from the community.

The intervention. Thirteen studies used a single strain of probiotics (Cui et al., 2019; Demirel et al., 2013; Dongol Singh et al., 2017; Hernández-Enríquez et al., 2016; Hussain et al., 2016; Kaban et al., 2019; Oncel et al., 2014; Rojas et al., 2012; Roy et al., 2014; Serce et al., 2013; Shadkam et al., 2015; Tewari et al., 2015; Xu et al., 2016), and 13 studies used a preparation that contained multiple strains of probiotics (Amini et al., 2017; Braga et al., 2011; Chowdhury et al., 2016; Dashti et al., 2014; Dutta et al., 2015; Fernández-Carrocera et al., 2013;

FIGURE 1 PRISMA flow diagram



Hariharan et al., 2016; Niekerk et al., 2015 (HIV exposed); Roy et al., 2014; Saengtawesin et al., 2014; Samanta et al., 2009; Shadkam et al., 2015; Sinha et al., 2015). Five studies used a preparation that had a probiotic + prebiotic (synbiotic) (Cooper et al., 2017; Dilli et al., 2015; Guney-Varal et al., 2017; Nandhini et al., 2016; Panigrahi et al., 2017; Dilli et al., 2015; Panigrahi et al., 2017) used a probiotic preparation that had a single strain of bacteria, and the other two studies used a preparation that had multiple strains of bacteria (Guney-Varal et al., 2017; Nandhini et al., 2016). One study did not report the strain of probiotic supplementation (Huaxian, 2013).

Ten studies used a probiotic preparation that contained *Lactobacillus* (Cooper et al., 2017; Cui et al., 2019; Dongol Singh et al., 2017; Hernández-Enríquez et al., 2016; Kaban et al., 2019; Oncel et al., 2014; Panigrahi et al., 2017; Rojas et al., 2012; Roy et al., 2014; Shadkam et al., 2015), and two studies used a preparation that contained *Bifidobacterium* (Dilli et al., 2015; Hussain et al., 2016). Fourteen studies used a preparation that had both *Lactobacillus* and *Bifidobacterium* (Amini et al., 2017; Braga et al., 2011; Chowdhury et al., 2016; Dashti et al., 2014; Fernández-Carrocera et al., 2013; Guney-Varal et al., 2017; Hariharan et al., 2016; Nandhini et al., 2016; Niekerk et al., 2015 (HIV exposed); Roy et al., 2014; Saengtawesin et al., 2014; Samanta et al., 2009; Shashidhar et al., 2017; Sinha et al., 2015. Three studies used *Saccharomyces boulardii* (Demirel et al., 2013; Serce et al., 2013; Xu et al., 2016), and one study used *Bacillus clausii* (Tewari et al., 2015). The probiotics were mostly given with breastmilk or formula feedings and started when the baby was able to tolerate minimal enteral feeds. The duration and dose of probiotic supplementation varied among the studies.

Comparison. Sixteen studies used a placebo (Cui et al., 2019; Dashti et al., 2014; Demirel et al., 2013; Dilli et al., 2015; Dongol Singh et al., 2017; Dutta et al., 2015; Kaban et al., 2019; Niekerk et al., 2015 (HIV exposed); Oncel et al., 2014; Panigrahi et al., 2017; Rojas et al., 2012; Roy et al., 2014; Serce et al., 2013; Shadkam et al., 2015; Sinha et al., 2015; Tewari et al., 2015); the rest of the studies used a control group receiving standard of care only.

Outcomes. All the studies reported data for at least one outcome that could be included in the meta-analysis. Twenty five studies reported data for all-cause mortality, 29 studies reported data for NEC, and 21 studies reported data for the incidence of neonatal sepsis. See Section 5.3 for more details.

5.1.3 | Excluded studies

Overall, 138 studies were excluded. See the Characteristics of excluded studies for reasons for exclusion of studies.

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Among excluded studies, 86 studies were excluded because of the wrong settings, and most of these studies were conducted in high-income countries. Fifteen studies were excluded because no relevant clinical outcomes were available from the abstract or full text of the studies. Fourteen studies were excluded for the wrong population, and 10 studies were excluded because of the wrong study design. Nine studies had a wrong intervention, and four studies had a wrong comparator.

5.2 | ROB in included studies

5.2.1 | Vitamin A supplementation during the neonatal period

Figure 2 shows the ROB in the 16 included studies that addressed vitamin A supplementation during the neonatal period.

5.2.2 | Probiotic supplementation during the neonatal period

Figure 3 show the ROB in the 33 included studies that addressed probiotic supplementation during the neonatal period.

5.2.3 | Allocation (selection bias)

Vitamin A supplementation during the neonatal period All the studies for vitamin A supplementation were at low ROB for sequence generation and allocation concealment.

Probiotic supplementation during the neonatal period

Two studies were judged to be at high ROB due to inadequate randomization (Guney-Varal et al., 2017; Kaban et al., 2019), and four studies did not provide enough information to allow a judgment about methods of randomization; these were labelled as having an unclear ROB (Hariharan et al., 2016; Hernández-Enríquez et al., 2016; Huaxian, 2013; Rehman et al., 2018). The rest of the studies had a low ROB for sequence generation.

Two studies were considered at high ROB due to inability to conceal the allocation (Dongol Singh et al., 2017; Guney-Varal et al., 2017). Ten studies had an unclear ROB, as these studies did not provide enough information to assess methods of allocation concealment (Amini et al., 2017; Cui et al., 2019; Hariharan et al., 2016; Hernández-Enríquez et al., 2016; Huaxian, 2013; Hussain et al., 2016; Kaban et al., 2019, Rehman et al., 2018; Samanta et al., 2009; Shadkam et al., 2015). The rest of the studies had a low ROB for allocation concealment.

5.2.4 | Blinding (performance bias and detection bias)

Vitamin A supplementation during neonatal period

None of the included studies for neonatal vitamin A supplementation was at increased ROB for blinding.

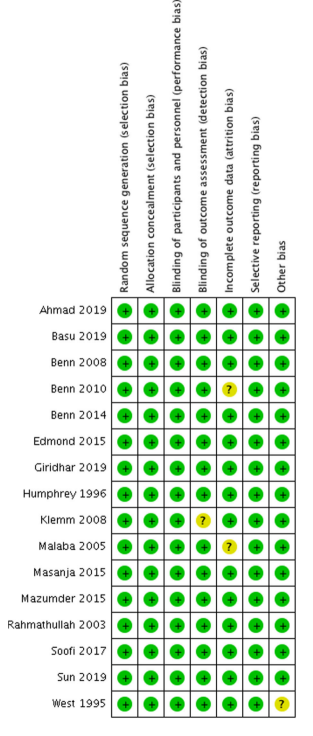


FIGURE 2 Risk of bias: neonatal vitamin A supplementation

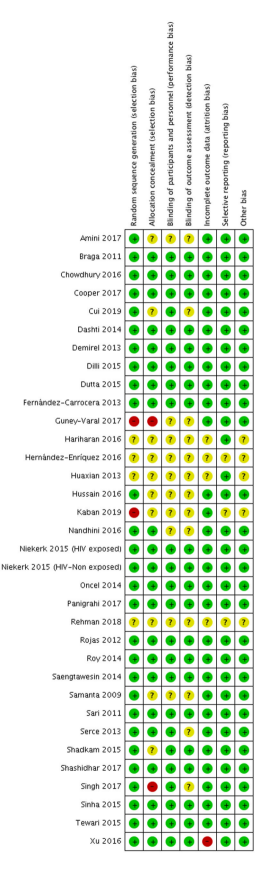


FIGURE 3 Risk of bias: probiotic supplementation during neonatal period



Probiotic supplementation during neonatal period

Ten studies had an unclear ROB due to blinding of the participants (Amini et al., 2017; Guney-Varal et al., 2017; Hariharan et al., 2016; Hernández-Enríquez et al., 2016; Huaxian, 2013; Hussain et al., 2016; Kaban et al., 2019; Nandhini et al., 2016; Rehman et al., 2018; Samanta et al., 2009). The rest of the studies had a low ROB due to the inability to do blinding of the participants.

Thirteen studies had an unclear ROB for blinding of the outcome assessors (Amini et al., 2017; Cui et al., 2019; Dongol Singh et al., 2017; Guney-Varal et al., 2017; Hariharan et al., 2016; Hernández-Enríquez et al., 2016; Huaxian, 2013; Hussain et al., 2016; Kaban et al., 2019; Nandhini et al., 2016; Rehman et al., 2018; Serce et al., 2013; Samanta et al., 2009). The rest of the studies had a low ROB.

5.2.5 | Incomplete outcome data (attrition bias)

Vitamin A supplementation during the neonatal period

All studies were at low risk for attrition bias except two studies that had an unclear ROB (Benn et al., 2010; Malaba et al., 2005).

Probiotic supplementation during the neonatal period

Most of the studies had a minimal loss to follow-up. One study was at high ROB where more than 20% of the participants were lost to follow-up (Xu et al., 2016). Four studies had an unclear ROB, as there was not enough information to make an assessment in these studies (Hariharan et al., 2016; Hernández-Enríquez et al., 2016; Huaxian, 2013; Rehman et al., 2018).

5.2.6 | Selective reporting (reporting bias)

Vitamin A supplementation during neonatal period All studies were considered to have low ROB for selective outcome reporting.

Probiotic supplementation during neonatal period

Most of the studies reported all relevant outcomes, and we did not consider any particular study at high ROB. Three studies had unclear ROB for selective outcome reporting (Hernández-Enríquez et al., 2016; Kaban et al., 2019; Rehman et al., 2018).

5.2.7 Other potential sources of bias

Vitamin A supplementation during neonatal period No study was considered at high ROB due to other reasons.

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Probiotic supplementation during neonatal period No other major source of ROB was noted. Five studies had unclear ROB due to limited available information (Hariharan et al., 2016; Hernández-Enríquez et al., 2016; Huaxian, 2013; Kaban et al., 2019; Rehman et al., 2018).

5.3 | Effects of interventions

5.3.1 | VItamin A supplementation during neonatal period

All-cause mortality during the neonatal period

Five studies from community settings reported the effect of vitamin A supplementation on all-cause neonatal mortality. These combined results showed no significant difference between the intervention and the control group (RR, 0.99; 95% CI, 0.90–1.08; six studies, 126,548 participants, heterogeneity: $\tau^2 = 0.00$; $\chi^2 = 3.64$, (p = 0.46); $l^2 = 0$ %). The grade rating for this outcome was "high." Summary of findings Table 2.

Sensitivity analysis: Fixed effect model. Use of a fixed effect model did not change the summary estimate for neonatal mortality (RR, 0.99; 95% CI, 0.90–1.08).

All-cause mortality at 6 months

Twelve studies from community settings reported the data for the effect of neonatal vitamin A supplementation on all-cause mortality at 6 months. The combined results showed no difference between the intervention and control group (RR, 0.98; 95% CI, 0.89–1.07; 12 studies, 154,940 participants; heterogeneity: $\tau^2 = 0.01$; $\chi^2 = 19.14$, (p = 0.06); $l^2 = 43\%$). The Grade rating for this outcome was "high." Summary of findings Table 2. A funnel plot for publication bias was symmetrical.

Sensitivity analysis: Fixed effect model. Use of a fixed effect model led to minimal change in the summary estimate (RR, 0.97; 95% CI, 0.91–1.03).

All-cause mortality at 12 months

Eight studies from community settings reporting on the impact of neonatal vitamin A supplementation reported data for all-cause mortality at 12 months (RR, 1.04; 95% CI, 0.94–1.14; eight studies, 118,376 participants; Heterogeneity: $\tau^2 = 0.01$; $\chi^2 = 12.99$, df = 7 (p = 0.07); $l^2 = 46\%$]. We rated this evidence as high certainty Summary of findings Table 2.

Sensitivity analysis: Fixed effect model. Use of a fixed effect model did not change the summary estimate significantly (RR, 1.02; 95% Cl; 0.96–1.08).

Three studies from hospital settings also reported data on mortality. The time to event for mortality was not clear in these studies. We did not pool data from the hospital-based studies with other studies, as the community-based studies had participants that were very different from hospital-based studies. We briefly describe the results of these studies below.

A study by Basu et al. (2019) reported the primary outcome; this was a composite incidence of all-cause mortality and oxygen requirement for 28 days. The results showed a reduction in mortality in the vitamin A group compared to the control group (RR, 0.44; 95% CI, 0.23–0.84).

No difference in mortality was reported by Sun et al. (2019) in the Vitamin A group versus placebo (RR, 0.49; 95% CI, 0.45–5.32).

Similarly, no difference was noted in all-cause mortality by Giridhar et al. (2019) (RR, 2; 95% CI, 0.63–6.30).

Adverse outcomes: Bulging fontanelle

Six studies reported on the effect of neonatal vitamin A supplementation on incidence of bulging fontanelle. The combined results showed a 53% increased risk of bulging fontanelle in the intervention group compared to control (RR, 1.53; 95% CI, 1.12–2.09; heterogeneity: $\tau^2 = 0.08$; $\chi^2 = 14.20$, (*p* = 0.01); $l^2 = 65\%$). We have high certainty in this evidence. Summary of findings Table 2.

Adverse outcomes: Vomiting

The combined results from six studies showed that neonatal vitamin A supplementation did not increase the risk of vomiting (RR, 1.00; 95% CI, 0.93–1.07; heterogeneity: $\tau^2 = 0.00$; $\chi^2 = 3.90$, (*p* = 0.42); $l^2 = 0$ %).

Vitamin A deficiency

One study from the community setting (Benn et al., 2008) reported VAD at 6 weeks and 4 months post neonatal supplementation. No significant difference was noted between the two groups at 6 weeks (RR, 0.94; 95% CI, 0.75–1.19) or 4 months (RR, 1.02; 95% CI, 0.64–1.62).

Another study from the hospital setting (Giridhar et al., 2019) showed a significant decrease in VAD in the intervention group compared to control (RR, 0.09; 95% CI, 0.024–0.38).

Neurodevelopment outcomes

Two studies reported long-term neurodevelopmental outcomes after use of vitamin A supplementation during the neonatal period. As the outcomes measured and duration of follow up were different, we did not pool the studies.

Humphrey et al. (1996) reported on neurodevelopmental outcomes at 3 year follow-up after neonatal vitamin A supplementation by using Bayley Scales of Infant Development. The study authors analysed the data for children with (n = 91) and without (n = 432)bulging fontanelle who received vitamin A versus placebo. The results showed that neonatal vitamin A supplementation did not have any adverse effect on development in the presence or absence of bulging fontanelle. Neonatal vitamin A supplementation had a posi**TABLE 2** Vitamin A compared to placebo for neonatal health

Vitamin A compared to placebo for neonatal health

Patient or population: neonates (0-28 days)

Setting: low and middle income countries

Intervention: vitamin A

Comparison: placebo

Comparison. placebo			
Outcomes	Relative effect (95% CI)	No. of participants (studies)	Certainty of the evidence (GRADE)
All-cause neonatal mortality	RR, 0.99 (0.90-1.08)	126,548 (6 RCTs)	⊕⊕⊕⊕ HIGH
All-cause mortality at 6 months of age	RR, 0.98 (0.89-1.07)	154,940 (12 RCTs)	⊕⊕⊕⊕ HIGH
All-cause mortality at 12 months of age	RR, 1.04 (0.94-1.14)	118,376 (8 RCTs)	⊕⊕⊕⊕ HIGH
Adverse Events: Bulging Fontanelle 48–72 h	RR, 1.53 (1.12-2.09)	100,562 (6 RCTs)	⊕⊕⊕⊕ HIGH

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*The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% Cl).

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

- **Moderate certainty:** We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different
- Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Abbreviations: CI, confidence interval; OR, odds ratio; RR, risk ratio.

tive effect on all developmental scores. The reported developmental scores addressed developmental areas such as orientationengagement, emotional regulation, and motor quality.

Klemm et al. (2008) reported data on neurodevelopmental outcomes 8 years after vitamin A supplementation. The authors followed a cohort of participants (n = 1613) who either directly received neonatal vitamin A or whose mother received vitamin A during pregnancy. The results showed no significant difference in intelligence, memory, and motor function; however, when the neonates and their mothers were supplemented with vitamin A versus placebo, it increased their performance in reading, spelling, and math computation.

5.3.2 | Probiotic supplementation during the neonatal period

Data were available for the effect of probiotic supplementation for all-cause mortality, NEC, sepsis, and sepsis-specific mortality.

All-cause mortality at longest follow-up

Twenty-five studies that included 10,998 subjects (probiotics 5548, control 5450) reported data for all-cause mortality for the effect of

probiotic supplementation on all-cause mortality. Our meta-analysis found a reduction of 20% in all-cause mortality in the probiotic group compared to control (RR, 0.80; 95% CI, 0.66–0.96; heterogeneity: $\tau^2 = 0.00$; (p = 0.55); $l^2 = 0$ %) (Figure 4). The number needed to treat was 100. The GRADE rating for this outcome was "high." Summary of findings Table 3.

Publication bias. A funnel plot for publication bias looked symmetrical (Figure 5).

Subgroup analyses. Table 1 gives the summary of data for subgroup analyses. For the outcome of all-cause mortality, data were available to perform subgroup analyses based on settings (hospital vs. community-based studies), type of probiotics (single strains vs. multiple strain vs. synbiotic), type of participants (term vs. preterm/low birth weight), type of feeding (breastfeeding vs. formula feeding vs. mixed feeding), and probiotic preparation (preparation containing *Lactobacillus* vs. *Bifidobacterium* vs. both *Lactobacillus* and *Bifidobacterium* vs. *S. boulardii*). No significant difference was noted among the subgroups; however, the number of studies varied for each group within the subgroup analysis.



		E	Experimental	Control		Risk Ratio	Risk Ratio
Study or Subgroup	log[Risk Ratio]	SE	Total	Total	Weight	IV, Random, 95% Cl	IV, Random, 95% Cl
Braga 2011	-0.046	0.243	122	121	15.2%	0.96 [0.59, 1.54]	
Chowdhury 2016	-0.3365	0.5563	60	60	2.9%	0.71 [0.24, 2.13]	
Cooper 2017	0.3209	0.7575	207	204	1.6%	1.38 [0.31, 6.08]	
Dashti 2014	0.6637	0.5879	69	67	2.6%	1.94 [0.61, 6.15]	
Demirel 2013	0.0144	0.621	138	140	2.3%	1.01 [0.30, 3.43]	
Dilli 2015	-0.8473	0.4776	200	200	3.9%	0.43 [0.17, 1.09]	
Dutta 2015	0.2054	0.7666	114	35	1.5%	1.23 [0.27, 5.52]	
Fernández-Carrocera 2013	-1.9459	1.0565	75	75	0.8%	0.14 [0.02, 1.13]	
Guney-Varal 2017	-2.7668	1.0367	76	43	0.8%	0.06 [0.01, 0.48]	·
Hariharan 2016	-0.121	0.6554	93	103	2.1%	0.89 [0.25, 3.20]	
Kaban 2019	-1.3863	1.0988	47	47	0.7%	0.25 [0.03, 2.15]	
Nandhini 2016	0.1054	0.4392	110	110	4.6%	1.11 [0.47, 2.63]	
Niekerk 2015 (HIV exposed)	1.0986	1.1311	37	37	0.7%	3.00 [0.33, 27.54]	
Niekerk 2015 (HIV non-exposed)	-0.8799	0.8146	54	56	1.3%	0.41 [0.08, 2.05]	
Oncel 2014	-0.2971	0.2353	213	211	16.2%	0.74 [0.47, 1.18]	
Panigrahi 2017	0.4055	0.6448	2278	2278	2.2%	1.50 [0.42, 5.31]	
Rojas 2012	-0.2252	0.2754	372	378	11.8%	0.80 [0.47, 1.37]	
Roy 2014	-0.1335	0.4818	56	56	3.9%	0.88 [0.34, 2.25]	
Samanta 2009	-1.2097	0.5476	91	95	3.0%	0.30 [0.10, 0.87]	
Sari 2011	-0.2113	0.2999	121	121	10.0%	0.81 [0.45, 1.46]	
Serce 2013	0.2231	0.6563	104	104	2.1%	1.25 [0.35, 4.52]	
Shadkam 2015	-0.6931	1.1972	30	30	0.6%	0.50 [0.05, 5.22]	
Shashidhar 2017	-1.0986	1.1379	52	52	0.7%	0.33 [0.04, 3.10]	
Singh 2017	-0.0556	0.7817	37	35	1.5%	0.95 [0.20, 4.38]	
Sinha 2015	-0.6872	1.2235	668	672	0.6%	0.50 [0.05, 5.53]	
Tewari 2015	-0.1869	0.372	124	120	6.5%	0.83 [0.40, 1.72]	
Total (95% CI)			5548	5450	100.0%	0.80 [0.66, 0.96]	•
Heterogeneity: Tau ² = 0.00; Chi ² = 3	23.67, df = 25 (P =	0.54); l ² =	= 0%				
Test for overall effect: Z = 2.41 (P =	0.02)						0.01 0.1 1 10 100 Favours Probiotics Favours Control
							Favours Frobiolics Favours Control

FIGURE 4 (Analysis 2.1) Forest plot of comparison: 2 probiotics versus control, outcome: 2.1 all-cause mortality

Sensitivity analysis.

Random versus fixed effect models. Use of a fixed-effect model did not change the summary estimate for the effect of probiotics on all-cause mortality (RR, 0.80; 95% Cl, 0.66–0.96; heterogeneity: χ^2 = 22.55, (*p* = 0.55); *I*² = 0%) Analysis 2.7.

Risk of bias. Exclusion of three studies (Dongol Singh et al., 2017; Fernández-Carrocera et al., 2013; Kaban et al., 2019) that were at high ROB for randomizations/allocation concealment did not change the summary estimate to a great extent, and results remained statistically significant Analysis 2.8.

Incidence of NEC

This outcome was reported by twenty-nine studies that included a total of 5574 (probiotics, 2843; control, 2731) participants. The combined results showed that the probiotics group had a relative reduction in NEC prevalence of 54% compared to control group (RR, 0.46; 95% CI, 0.35–0.59; heterogeneity: $\tau^2 = 0.11$; (p = 0.12); $I^2 = 24\%$) (Figure 6). The number needed to treat was 17. We have high certainty in this evidence. Summary of findings Table 3.

Publication bias. A funnel plot for publication bias looked symmetrical.

Subgroup analyses. Table 1 shows the results of subgroup analyses. For NEC, data were available to perform the following 3 subgroup

analyses: type of probiotics (single strain vs. multiple strain vs. synbiotic), type of feeding (breastfeeding vs. formula feeding vs. mixed feeding), and probiotic preparation (preparation containing Lactobacillus vs. Bifidobacterium vs. both Lactobacillus and Bifidobacterium vs. S. boulardii). No significant difference was noted among the subgroup analyses except the one based on probiotics preparation (p value for subgroup difference 0.05). The probiotics preparation that has Lactobacillus in it, either as a single strain (RR, 0.39; 95% CI, 0.25-0.61; 13 studies) or in combination with Bifidobacterium (RR, 0.44; 95% CI, 0.36-0.68; 13 studies), had a significant effect compared to Bifidobacterim alone (RR, 0.20; 95% CI, 0.09, -0.47) or S. boulardii alone (RR, 0.94; 95% CI, 0.45-1.95; two studies). This subgroup analysis should be interpreted carefully, as the number of studies was not uniformly distributed among all the subgroups, and the statistical difference might be due to the small number of studies in two of the subgroups.

Sensitivity analyses.

Random versus fixed effect models. The use of a fixed effect model did not change the summary estimate for the effect of probiotics on NEC (RR, 0.45; 95% Cl, 0.37–0.56; heterogeneity: χ^2 = 37.92, df = 29 (*p* = 0.12); *l*² = 24%).

Risk of bias. The exclusion of three studies (Dongol Singh et al., 2017; Fernández-Carrocera et al., 2013; Kaban et al., 2019) with a high



TABLE 3 Probiotics supplementation during neonatal period

Probiotics supplementation compared to control during neonatal period

Patient or population: neonates (Mmost of the included studies had preterm and low birth weight neonates)

Setting: low and middle income countries

Intervention: probiotics/synbiotics

Comparison: control

				Anticipated absolute effects* (95% CI)
Outcomes	No. of participants (studies) Follow up	Certainty of the evidence (GRADE)	Relative effect (95% Cl)	Risk with Risk difference with control probiotics (intervention)
All-cause mortality	10904 (25 RCTs)	$\oplus \oplus \oplus \oplus HIGH^{a,b,c,d}$	RR, 0.80 (0.66-0.96)	Study population 47 per 1000 9 fewer per 1000 (15 fewer to 1 fewer)
Neonatal sepsis	8918 (21 RCTs)	⊕⊕⊕⊕ HIGH ^{a,d,e}	RR, 0.78 (0.70-0.86)	Study population 205 per 1000 45 fewer per 1000 (62 fewer to 29 fewer)
Necrotizing enterocolitis	55574 (29 RCTs)	⊕⊕⊕⊕ HIGH ^{a,d,f}	RR, 0.46 (0.35–0.61)	Study population 101 per 1000 55 fewer per 1000 (66 fewer to 41 fewer)

*The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: Confidence interval; RR: Risk ratio; OR: Odds ratio;

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

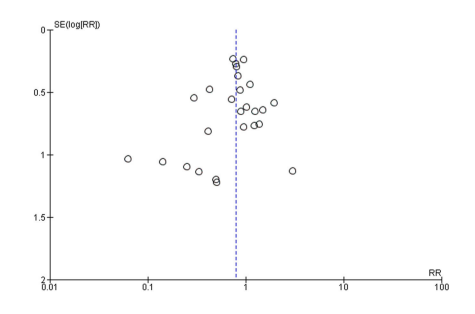
^aEven though three (Dongol Singh et al., 2017; Fernández-Carrocera et al., 2013; Kaban et al., 2019) of the included studies in the analysis had high ROB related to randomizations, the exclusion of these studies did not have much effect on the magnitude of the summary estimate or its statistical significance. ^b l^2 was 0%.

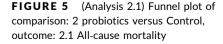
^cAll-cause mortality is an objective outcome and there were no concerns about the indirect measurement of the outcome.

^dThe confidence interval of the summary estimate did not include 1.

 $^{\rm e}l^2$ was 23% and the p. value for heterogeneity was 0.16.

^fThe *I*² was 24%.





		Ex	perimental	Control		Risk Ratio	Risk Ratio
Study or Subgroup	log[Risk Ratio]	SE	Total	Total	Weight	IV, Random, 95% CI	IV, Random, 95% Cl
Amini 2017	-1.0296	0.32	60	60	8.3%	0.36 [0.19, 0.67]	
Braga 2011	-2.2054	1.4852	122	121	0.8%	0.11 [0.01, 2.02]	
Chowdhury 2016	-1.7918	1.0646	60	60	1.4%	0.17 [0.02, 1.34]	
Cui 2019	-1.5449	1.0756	45	48	1.4%	0.21 [0.03, 1.76]	
Dashti 2014	0.1937	0.4422	69	67	5.8%	1.21 [0.51, 2.89]	
Demirel 2013	-0.1398	0.5433	138	140	4.4%	0.87 [0.30, 2.52]	
Dilli 2015	-1.6094	0.4359	200	200	5.9%	0.20 [0.09, 0.47]	_ -
Dutta 2015	1.4035	1.4551	114	35	0.8%	4.07 [0.23, 70.49]	
Fernández-Carrocera 2013	-0.6931	0.4726	75	75	5.3%	0.50 [0.20, 1.26]	
Guney-Varal 2017	-2.7568	1.4787	76	43	0.8%	0.06 [0.00, 1.15]	
Hariharan 2016	0.1021	0.8039	93	103	2.4%	1.11 [0.23, 5.35]	
Hernández-Enríquez 2016	-2.4849	1.0042	24	20	1.6%	0.08 [0.01, 0.60]	
Huaxian 2013	-1.5533	0.769	65	55	2.5%	0.21 [0.05, 0.95]	
Hussain 2016	-1.665	0.3957	150	150	6.6%	0.19 [0.09, 0.41]	_ - -
Kaban 2019	-1.9459	1.498	47	47	0.8%	0.14 [0.01, 2.69]	
Nandhini 2016	-0.6931	0.4115	110	110	6.3%	0.50 [0.22, 1.12]	
Niekerk 2015 (HIV exposed)	0	1.395	37	37	0.9%	1.00 [0.06, 15.40]	
Niekerk 2015 (HIV non-exposed)	0.0364	1.4013	54	56	0.9%	1.04 [0.07, 16.17]	
Oncel 2014	-0.2326	0.4643	213	211	5.4%	0.79 [0.32, 1.97]	
Rehman 2018	-1.3863	0.773	73	73	2.5%	0.25 [0.05, 1.14]	
Rojas 2012	-0.4948	0.4153	372	378	6.3%	0.61 [0.27, 1.38]	
Roy 2014	0.0001	0.982	56	56	1.7%	1.00 [0.15, 6.85]	
Saengtawesin 2014	-0.0667	1.3904	31	29	0.9%	0.94 [0.06, 14.27]	
Samanta 2009	-1.0556	0.4951	91	95	5.0%	0.35 [0.13, 0.92]	
Sari 2011	-0.5108	0.5001	121	121	4.9%	0.60 [0.23, 1.60]	
Serce 2013	-0.0001	0.5162	104	104	4.7%	1.00 [0.36, 2.75]	
Shadkam 2015	-1.7047	0.724	30	30	2.8%	0.18 [0.04, 0.75]	
Shashidhar 2017	-1.0986	0.7926	52	52	2.4%	0.33 [0.07, 1.58]	
Singh 2017	-0.5664	0.4594	37	35	5.5%	0.57 [0.23, 1.40]	
Tewari 2015	0.6604	1.218	124	120	1.1%	1.94 [0.18, 21.07]	
Total (95% CI)			2843	2731	100.0%	0.46 [0.35, 0.59]	◆
Heterogeneity: Tau ² = 0.11; Chi ² =	Heterogeneity: Tau ² = 0.11; Chi ² = 37.92, df = 29 (P = 0.12); l ² = 24%					0.005 0.1 1 10 200	
Test for overall effect: Z = 5.88 (P <							
•	,						Favours [Probiotics] Favours Control

FIGURE 6 (Analysis 2.9) Forest plot of comparison: 2 probiotics versus control, outcome: 2.9 necrotizing enterocolitis (any type)

ROB due to randomization/allocation concealment did not change the summary estimate significantly.

Incidence of neonatal sepsis

The effect of probiotics on the incidence of neonatal sepsis was reported by 21 studies that included 9105 (probiotics, 4606; control, 4499) participants. The combined results showed a statistically significant reduction in incidence of sepsis of 22% in the intervention group compared to control (RR, 0.78; 95% CI, 0.70–0.86; heterogeneity: $\tau^2 = 0.01$; (p = 0.16); $l^2 = 23\%$). The number needed to treat was 14. The grade rating for this outcome was "high." Summary of findings Table 3.

Publication bias. A funnel plot for publication bias looked symmetrical.

Subgroup analysis. Table 1 shows the results of subgroup analyses. Subgroup analysis could be done for the incidence of neonatal sepsis according to study setting, type of probiotic (single strain vs. multiple strain vs. synbiotic), type of feeding (breastfeeding vs. formula feeding vs. mixed feeding), and probiotic preparation (preparation containing *Lactobacillus* vs. *Bifidobacterium* vs. both *Lactobacillus* and *Bifidobacterium* vs. *S. boulardii*). Only the type of feeding differed among subgroups, probiotic supplementation seemed to have a significant effect for the prevention of neonatal

sepsis when the supplementation was given to babies who received breastmilk only (RR, 0.76; 95% CI, 0.61–0.83; 8 studies) or breastmilk in combination with formula milk (RR, 0.76; 95% CI, 0.64–0.90; seven studies), p value for subgroups difference was 0.04.

Sensitivity analysis.

Random versus fixed effect models. Use of a fixed effect model had a minimal effect on summary estimates for the effect of probiotics on the incidence of neonatal sepsis (RR, 0.79; 95% Cl, 0.73–0.85; heterogeneity: χ^2 = 26.06, df = 20 (*p* = 0.16); *l*² = 23%).

Risk of bias. No significant difference was noted when two studies (Fernández-Carrocera et al., 2013; Kaban et al., 2019) with a high ROB were excluded from the analysis Analysis 2.21.

Sepsis specific mortality

Two studies reported the data for sepsis specific mortality. The combined results showed a reduction of sepsis specific mortality of 89% in the intervention group compared to control (RR, 0.21; 95% Cl, 0.04–1.01; heterogeneity: $\tau^2 = 0.00$; $\chi^2 = 0.91$, df = 1 (p = 0.34); $l^2 = 0$ %). This is limited by a wide CI of the estimate including 1.

Adverse events

No adverse event was reported in any of the included studies.

Neurodevelopmental outcomes

Two studies assessed neurodevelopmental outcomes after neonatal probiotic supplementation. We did not perform a meta-analysis for these outcomes, as these studies used different scales.

Sari et al. (2011) reported the data for neurodevelopmental outcomes at 18–22 months post neonatal probiotic supplementation for extremely low birth weight infants. Their results did not show any difference in growth and neurodevelopmental outcomes between the two groups. The growth outcomes were reported as weight (probiotic group 10.5 ± 1.7 kg vs. control 10.5 ± 1.7 kg, *p* value .92), length (probiotic group 79.4 ± 7.8 cm vs. control 81.0 ± 5.3 cm, *p* value .32), and head circumference (probiotic group 47.5 ± 6.5 cm vs. control 46.7 ± 1.8 cm, *p* value .53). The neurodevelopmental outcomes were reported as mental development index (probiotic group 90.7 ± 15.5 vs. control 90.4 ± 14.5 , *p* value .88) and Psychomotor Development Index (probiotic group 95.4 ± 17.2 vs. control 93.2 ± 16.4 , *p* value .39).

Oncel et al. (2014) also followed a cohort of extremely low birth weight infants after neonatal supplementation with probiotics or placebo. There was no significant difference in the neurodevelopmental outcomes at 18–24 months of age post supplementation, mental development index (probiotic 81 (median), 49 (min) to 124 (max) vs. placebo 82, 53–128; *p* value .48) or Psychomotor Development Index (probiotic 80, 49–112 vs. placebo 79, 49–107; *p* value .67).

6 | DISCUSSION

6.1 | Summary of main results

This review evaluated three neonatal nutritional interventions. Vitamin A supplementation during the neonatal period in the community setting did not have any significant effect on all-cause mortality at 1, 6, or 12 months. We did not identify any studies from LMICs that assessed the use of dextrose gel for the treatment or prevention of neonatal hypoglycaemia. Probiotic supplementation during the neonatal period mainly given to low birth weight and preterm babies was shown to reduce all-cause mortality, NEC, and neonatal sepsis.

6.2 | Overall completeness and applicability of evidence

The evidence of neonatal vitamin A supplementation included 16 studies, and the number of participants in these studies exceeded



150,000. Overall, most of the included studies were at low ROB across many of the ROB items assessed using the Cochrane ROB scale. The statistical heterogeneity in the pooled data for mortality outcomes at 6 and 12 months was noted for neonatal vitamin A supplementation; however, the value of l^2 was <50%. Subgroup analyses done in a recent IPD analysis could potentially explain the reasons for heterogeneity and are discussed in Section 6.5.

The results for the use of probiotics during the neonatal period seem very promising. Almost all the studies in this metaanalysis included preterm and/or low birth weight neonates. The effect of probiotics on all-cause mortality was reported in 25 studies, and the analysis included more than 10,000 participants. The forest plot showed a homogenous effect in favor of the intervention with I^2 of 0%. We think that this effect is biologically plausible, and the most likely pathway of reduction in neonatal mortality from probiotic supplementation is via a reduction in sepsis and NEC, as shown in Analysis 2.15 and Analysis 2.9, respectively. We also think that these results are less likely due to bias. The studies by Guney-Varal et al. (2017), Kaban et al. (2019), and Dongol Singh et al. (2017) were at high ROB due to inadequate randomization methods. The exclusion of these three studies from meta-analysis for the effect of probiotics on allcause mortality did not change the summary estimate significantly (RR, 0.82; 95% CI, 0.68-0.99). A funnel plot for publication bias was symmetrical. The use of a fixed versus random effect model also did not change the results significantly. In addition to these observations, the fact that the effect of probiotics on the reduction of neonatal sepsis and NEC was mostly homogenous (I^2 of 23% and 24%, respectively) indicates that the use of probiotics could be beneficial for babies with low birth weight and preterm birth. We notice, however, that there was significant clinical heterogeneity in the dose, duration, and strains of probiotics used; this indicates that more research is needed to determine the appropriate dose and duration of probiotic supplementation in neonates.

The use of probiotics for the prevention of NEC, sepsis, and mortality in preterm and/or low birth weight babies has been debated in the past. The use of probiotics was advocated after the publication of a Cochrane review in 2011 (Alfaleh et al., 2011, later updated in 2014, AlFaleh & Anabrees, 2014) that showed that probiotics reduced NEC (stage II or more) (RR. 0.35: 95% CI. 0.24-0.52) and mortality (RR, 0.40; 95% CI, 0.27-0.60) in preterm/ low birth weight neonates (Ofek Shlomai et al., 2014; Robinson, 2014). Others challenged the appropriateness of the meta-analysis in the setting of clinical heterogeneity (Mihatsch et al., 2012; Mihatsch, 2011) and called for large trials before this intervention could be recommended in clinical practice (Mihatsch, 2011). This led to two large clinical trials, the ProPrems trial conducted in Australia and New Zealand (Jacob et al., 2013) and the PiPS trial conducted in the United Kingdom (Costeloe et al., 2016). The ProPrem trial used a mixture of probiotics (Bifidobacterium infantis,

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Streptococcus thermophilus, and Bifidobacterium lactis) and included 1099 preterm (<32 weeks) and very low birth weight (<1500 g) neonates. The results of the ProPrems trial showed that the use of probiotics did not reduce the incidence of sepsis and mortality, but did reduce the incidence of NEC (RR. 0.46: 95% CI. 0.23-0.93) (Jacob et al., 2013). The PiPS trial included 1315 neonates between the gestational age of 23-30 weeks who were randomised within 24-48 h to a single probiotic (Bifidobacterium breve BBG-001) or placebo. The results of the PiPS trial showed no difference between the intervention and the control group for the outcomes of NEC (RR, 0.93; 95% CI, 0.68-1.27), sepsis (RR, 0.97; 95% CI. 0.73-1.29), or death (RR. 0.93; 95% CI. 0.67-1.30) (Costeloe et al., 2016). The results of these trials were surprising in the setting of known evidence from meta-analyses of the available studies. A closer look at the results of the PiPS trial showed that there was significant contamination of the control group, as about 49% of the neonates from the control group had the same type of probiotic bacteria in their stool as those who were in the intervention group. This decreased the power of the study, and it was argued that a potential lack of effect might be explained by the cross-contamination of the control group (Deshpande et al., 2016; McKinlay et al., 2016). The debate of appropriateness of probiotic supplementation in the neonatal period has continued, and a recent network meta-analysis concluded that there is not enough evidence in favor of either a single or a mixture of strains of probiotics that could be suggested for routine clinical use for the prevention of NEC, sepsis, and/or mortality in preterm/low birth weight babies (van den Akker et al., 2018).

Our review focused on studies from LMIC. The effect of probiotic supplementation on mortality, sepsis, and NEC was significant when we pooled studies from these countries only. We noticed clinical heterogeneity in the use of probiotics in terms of type of probiotics, single versus multiple strains used, and baseline interventions such as the use of breastmilk. We think that the use of a meta-analysis is appropriate to pool these studies, as the primary target of all the studies was the same, that is, correction of dysbiosis. We further demonstrated that subgroup analysis based on certain clinical factors reveals no significant difference in results. We notice, however, the relative lack of studies from the community setting. This is likely due to the increased rate of community over hospitalbased births in LMIC. For this reason, we read the results of a community-based study by Panigrahi et al. (2017) with great interest. This study was the largest study conducted on the use of probiotics (synbiotics) and had a sample size of 4556; this is three times larger than the sample size of the PiPS trial. Panigrahi et al. (2017) used a synbiotic which was a mixture of the probiotic Lactobacillus plantarum ATCC-202195 and the prebiotic fructooligosaccharide. This study recruited neonates who were at least 35 weeks of gestation and weighed at least 2000 g from rural settings from India. The results of the study showed a significant reduction in the primary outcome (combination of sepsis and death) in the intervention group compared to control (RR, 0.60; 95% CI, 0.48–0.74), culture-positive and culture-negative sepsis, and lower respiratory tract infections (Panigrahi et al., 2017). More such community-based studies are needed from other countries examining term, preterm, and low birth weight infants.

Are probiotics safe for use during the neonatal period? We did not find any substantial evidence of adverse events with the use of probiotics in neonatal age group. It is important, however, to consider the safety considerations of probiotic supplementation in this vulnerable population. Probiotics are not regulated as a medication and are thus susceptible to variations in quality within and between countries. If not carefully produced and handled, probiotics may also contain pathogenic contaminants that may then lead to neonatal sepsis. Probiotic supplementation has been linked to both bacterial (Dani et al., 2016) as well as fungal sepsis (Vallabhaneni et al., 2014). Despite their rare occurrence, neonates receiving probiotics should be followed in a registry to ensure any reporting and observation of the potential risk of sepsis in a large sample size in the real world setting.

6.3 | Quality of the evidence

The GRADE quality of evidence was considered high for most of the outcomes for neonatal vitamin A and probiotic supplementation. The GRADE method of assessment of overall evidence considers the type of study, ROB, statistical heterogeneity, indirectness, and imprecision of the summary estimates as well as the risk of publication bias (Guyatt et al., 2011).

All the included studies for neonatal vitamin A supplementation were randomized and had minimal ROB. The statistical heterogeneity for pooled studies for neonatal vitamin A supplementation was noticeable but was not significant enough to decrease our confidence in the summary estimate. Similarly, the CIs around the summary estimate were narrow, and there was no increased risk of publication bias for studies that assessed neonatal vitamin A supplementation.

All the included studies for neonatal probiotic supplementation were also randomized. High ROB was noted for three of the included studies (Dongol Singh et al., 2017; Fernández-Carrocera et al., 2013; Kaban et al., 2019), but exclusion of these studies did not change the results of any of the outcomes, including all-cause mortality, NEC, and neonatal sepsis. The pooled results were mostly homogenous, and the summary estimates were precise with narrow Cls. The funnel plots for publication bias were symmetrical for the outcomes of all-cause mortality, NEC, and neonatal sepsis.

6.4 | Potential biases in the review process

We used standard methods of Campbell and Cochrane collaborations to conduct the review. Two review authors screened the titles and abstracted the data from the included studies. Our inclusion/ exclusion criteria were decided a priori, and a peer reviewed protocol was published giving details of methods of conduct of this review.

We performed two posthoc subgroup analyses for the effect of probiotic supplementation during the neonatal period. This analysis was based on the type of feeding, as our team thought it was essential to establish any differential effect of probiotics when the intervention was delivered with breastmilk or formula or both. The results for this analysis were similar among the subgroups for outcomes of all-cause mortality and NEC; however, there was significant heterogeneity among subgroups for the outcome of neonatal sepsis Analysis 2.18. A close examination of the data showed that the difference among subgroups was due to the group where the status of the feeding was "unclear." Exclusion of this subgroup showed a homogenous protective effect in the case of breastmilk or formula milk-fed babies against neonatal sepsis (data not shown). So, we think that probiotics may have a significant protective effect against neonatal sepsis, NEC, and mortality irrespective of the type of food offered. The second posthoc subgroup analysis was based on study setting. We were interested in knowing if the probiotics had a similar effect on infants born in the hospital setting compared to those in the community setting. We were interested in this analysis because a significant number of births happen at home in LMIC. There were a limited number of studies conducted in the community setting that addressed the effect of probiotics; therefore, no solid conclusion could be drawn at this time for any of the outcomes.

6.5 | Agreements and disagreements with other studies or reviews

The effects of neonatal vitamin A supplementation have been reviewed in two Cochrane reviews (Darlow et al., 2016; Haider et al., 2017). The Cochrane review by Haider et al. (2017) focused on randomized studies from the community setting only. We considered studies from both community and hospital settings. We updated the literature search and found one additional study from the community setting (Ahmad et al., 2019) and added three studies from the hospital setting (Basu et al., 2019; Giridhar et al., 2019; Sun et al., 2019). The studies from hospital settings were done in the neonatal intensive care setting for very low birth weight babies. We did not pool the results of these studies with those of the community-based studies. The Cochrane review on neonatal vitamin A supplementation for very low birth weight infants included 11 trials and reported a reduction in risk of death or oxygen requirement at 1 month of age (RR, 0.93; 95% CI, 0.88-0.99). Of the three studies we included in our review from LMIC, only one study showed a reduction in composite outcome of incidence of all-cause mortality and oxygen requirement



for 28 days (RR, 0.44; 95% CI, 0.229–0.844)] from vitamin A supplementation in very low birth weight infants (Basu et al., 2019). The new study for neonatal vitamin A supplementation from the community was small and included 306 participants. The addition of this study did not change the results significantly compared to those published in the 2017 Cochrane review (Haider et al., 2017).

An individual participant meta-analysis of neonatal vitamin A studies conducted in the community setting was published during the preparation of this review (West et al., 2019). This review addressed multiple subgroup analyses both at the study- and individual-level characteristics and used the original data from individual trials to pool the studies. The overall results were similar to our results for all-cause mortality at 6 and 12 months (West et al., 2019). The subgroup analysis based on study-level characteristics showed that neonatal vitamin A supplementation significantly reduced 6-month mortality among the trials conducted in South Asia (RR, 0.87; 95% CI, 0.77-0.98) but not in Africa; they also showed a potential for increased risk of mortality in African countries (RR, 1.07; 95% CI, 1.00-1.15). Further subgroup analyses showed that neonatal vitamin A supplementation reduced all-cause mortality in the context of moderate or severe maternal VAD (defined as 10% or higher proportion of women with serum retinol <0.7 μ mol/L or 5% or more women with night blindness) (RR, 0.87; 95% CI, 0.80-0.94), in settings where baseline (control group) early infant mortality was 30 or more per 1000 live births (RR, 0.91; 95% CI, 0.85-0.98), and in the context of lack of maternal education (>32% mothers had no schooling) (RR, 0.88; 95% CI, 0.80-0.96). The subgroup analyses conducted based on individual-level characteristics such as sex, birth weight, gestational age and size, age at dosing, parity, time of breastfeeding initiation, maternal education, and maternal vitamin A supplementation did not show any significant differential effect of neonatal vitamin A supplementation compared to placebo for these groups. As most of the subgroup analyses that we prespecified in our review were addressed in this study (West et al., 2019), we did not repeat these analysis in our current study.

Oral dextrose as a treatment of hypoglycaemia and prevention of hypoglycaemia in high-risk neonates has been evaluated in two Cochrane reviews (Hegarty et al., 2017; Weston et al., 2016). The review by Weston et al. (2016) addressed treatment of neonatal hypoglycaemia and included two studies, one from New Zealand and another from Ireland. They did not show any major difference in episodes of hypoglycaemia between the two study groups. The review by Hegarty et al. (2017) addressed the prevention of hypoglycaemia in high-risk neonates and included one study from New Zealand. The included study showed a significant reduction in hypoglycaemia episodes in the intervention group compared to control (RR, 0.76; 95% CI, 0.62–0.94). No randomized study was available from LMIC in either of the two reviews mentioned above. We updated the searches and did not find any study from LMIC.

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Other reviews have been published that assess the effect of probiotic supplementation during the neonatal period. A Cochrane review was published in 2011 (Alfaleh et al., 2011) with an update in 2014 (AlFaleh & Anabrees, 2014). This has not since been updated. The Cochrane review included studies from LMIC and high-income countries and concluded that probiotic supplementation reduced NEC (stage II or more) (RR, 0.43; 95% CI, 0.33-0.56) and mortality (RR. 0.65: 95% CI. 0.52-0.81) but showed no effect for nosocomial sepsis (RR, 0.91; 95% CI, 0.80-1.03). More studies have been published since the publication of the 2014 Cochrane review update. Deshpande et al. (2017) reviewed studies from LMIC that addressed probiotic supplementation during the neonatal period. This review included twenty-three studies and concluded that probiotic supplementation reduced allcause mortality (RR, 0.73; 95% CI, 0.59-0.90), NEC (RR, 0.46, 95% CI, 0.34-0.61), and neonatal sepsis (RR, 0.80; 95% CI, 0.71-0.91). We included thirty-three studies and updated the meta-analyses. With this new data, the magnitude and the statistical significance remained the same, but the summary estimates became more precise for the outcomes of all-cause mortality, NEC, and sepsis. We also conducted additional subgroup analyses that were not previously performed.

7 | AUTHORS' CONCLUSIONS

7.1 | Implications for practice

Neonatal vitamin A supplementation in the community setting does not appear to reduce infant mortality at 1, 6, or 12 months of age. Vitamin A supplementation during neonatal period increases the risk of bulging fontanelle. No data were available for dextrose gel supplementation for the prevention or treatment of neonatal hypoglycaemia in LMIC.

Probiotic supplementation is a promising intervention and can reduce all-cause mortality, neonatal sepsis, and NEC in low birth weight and/or preterm babies in LMIC in the hospital setting. Though we observed no adverse effects, infants receiving probiotics should be entered into a registry in order to observe any concerns for safety in large sample sizes in the real world setting.

7.2 | Implications for research

There was significant clinical heterogeneity in terms of strains and dose of probiotics used in the included studies. More studies are needed to decide upon the right strain and optimal dose and duration of probiotics supplementation. Most of the included studies for probiotics supplementation were conducted in preterm/low birth weight babies in the neonatal intensive care unit. It is not clear if the similar protective effects would be seen when probiotics are given in the community setting or to term babies. It is also unclear if the supplementation of prebiotics and probiotics together (synbiotics) is more effective than probiotics alone. More studies are needed to answer these questions and to assess the effect of dextrose supplementation for the prevention and treatment of neonatal hypoglycaemia in LMIC.

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CONTRIBUTIONS OF AUTHORS

Aamer Imdad, Deepika Ranjit wrote the first draft of the protocol. Gamael S. S. Surin participated in the design of the search strategy and writing of the protocol. Abigail Smith and Sarah Lawler designed the search strategy. Aamer Imdad wrote the manuscript for the final review. Aamer Imdad, Faseeha Rehman, Evans Davis, Gamael S. S. Surin, Deepika Ranjit, and Suzanna L. Attia helped with data extraction and analysis. Suzanna L. Attia and Aamer Imdad edited the manuscript. Zulfiqar A. Bhutta supervised and gave feedback for the design of the protocol and the main review.

DECLARATIONS OF INTEREST

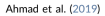
Zulfiqar A. Bhutta was principle investigator of study (Soofi et al., 2017). He was not involved in the selection of the study for this review and also did not participate in the data extraction from this study. All other authors declare that they do not have any conflict of interest.

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

- The planned subgroup analyses for vitamin A were not conducted, as the same analyses were available from a recent IPD analysis (West et al., 2019).
- We did not use EPOC methodology for the ROB assessment, as all the included studies were RCTs and the Cochrane ROB assessment tool was used for the same.
- We did two posthoc subgroup analyses for probiotic supplementation during the neonatal period. These included type of feeding and study setting.
- We did the sensitivity analysis for ROB based on sequence generation and allocation concealment.

PUBLISHED NOTES

CHARACTERISTICS OF STUDIES CHARACTERISTICS OF INCLUDED STUDIES





Methods	A block-randomized, double-masked, placebo-controlled intervention trial conducted in Bangladesh
Participants	 Inclusion criteria: "consent of the mother and willingness to have their infant participate; singleton birth at MCHTI clinic and eligible for vaccination according to the national and MCHTI clinic policy". Exclusion criteria: "planned at home delivery because of the low likelihood of vaccination at MCHTI within 48 h of birth, (2) congenital disease or a serious infection showing that the infant was not healthy; infant with birth weight <1500 g and inability to enrol within 48 h of birth due to lack of timely notification or other exceptional circumstances".
Interventions	Intervention group: 50,000 IU vitamin A (retinyl palmitate) Comparison: Placebo (unfortified soya based oil) The intervention was delivered within 48 h of birth
Outcomes	Neonatal mortality, adverse events, microbiome changes, thymus size
Notes	Data on mortality and bulging fontanelle were taken from publication (J Nutr 2019;00:1–8). Data on mortality was reported at 15 weeks. We included the data with "all-cause mortality at 6 months".

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The randomization lists of vitamin A and placebo within each group were generated by WHO, using Stata, v11" $$
Allocation concealment (selection bias)	Low risk	Quote: "Preplanned statistical analyses using arbitrary group identifiers" Comment: Most likely done
Blinding of participants and personnel (performance bias)	Low risk	Quote: "dose of VA in oil or an identical placebo (PL) within 48" Comment: Most likely done
Blinding of outcome assessment (detection bias)	Low risk	Quote: "Preplanned statistical analyses using arbitrary group identifiers (group 1, group 2) were completed on 4 April 2014 before unblinding"
Incomplete outcome data (attrition bias)	Low risk	Attrition rate: 5.2%
Selective reporting (reporting bias)	Low risk	Author prespecified the outcomes. Trial was registered as ClinicalTrials.gov: NCT01583972.
Other bias	Low risk	No other risk of bias was noted

Amini et al. (2017)

Methods	Prospective randomized control trial conducted in Iran
Participants Interventions	 Inclusion Criteria: "All premature newborns (n = 115) weighting 750–1500 g or <32 weeks' gestation who received antibiotics and total parenteral nutrition in NICU of Vali Asr Hospital were included" Exclusion Criteria: "Premature babies <750 and more than 1500 g and neonates with congenital heart disease, congenital malformations, and immune system deficiency, even in their family members, were excluded from the study."
	Intervention: Multistrain powder probiotic infant formula containing Streptococcus thermophilus, Lactobacillus rhamnosus, Lactobacillus acidophilus, Lactobacillus bulgaricus, Bifidobacterium infantis, Lactobacillus casei. The dose was 0.8–1 g per day in 8–10 doses given for 13 days Comparison: Enteral feed without probiotic
Outcomes	NEC
Notes	Data were taken from table 3

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Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "In this double blind randomized clinical trial (RCT), block randomization was used and 60 cases were randomly divided into 2 groups."
Allocation concealment (selection bias)	Unclear risk	No clear information was available about allocation concealment
Blinding of participants and personnel (performance bias)	Unclear risk	No clear information available
Blinding of outcome assessment (detection bias)	Unclear risk	No clear information available
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the outcomes irrespective of their statistical significance
Other bias	Low risk	No other risk of bias was noted

Basu et al. (2019)

Methods	A randomized double-blind placebo-controlled trial India
Participants	 Inclusion Criteria: Inborn, VLBW (birth weight (BW) < 1500 g) neonates admitted in NICU and requiring respiratory support in the form of oxygen inhalation through nasal prongs or head box, continuous positive airway pressure (CPAP), high flow nasal cannula (HFNC), or mechanical ventilation (MV) at the age of 24 h, were included. Exclusion Criteria: Neonates with major congenital malformation, any life-threatening condition such as reversal of umbilical artery end-diastolic blood flow on antenatal Doppler, perinatal asphyxia with moderate to severe hypoxic ischemic encephalopathy, shock with escalating doses of vasopressors, recurrent seizures, and suspected inborn errors of metabolism
Interventions	Intervention: 10,000 IU of retinol/dose, alternate days, 28 days or until discharge Compairson: Placebo
Outcomes	All-cause mortality, sepsis, NEC
Notes	Study conducted in very low birth weight babies

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Randomization into vitamin A or placebo group was done using random permuted blocks of 4, 6, and 8, prepared by an independent statistician not involved in the study."
Allocation concealment (selection bias)	Low risk	Quote: "Allocation into vitamin A or placebo group was done using serially numbered opaque and sealed envelopes by on-duty residents who were appropriately trained for the process beforehand. Allocation concealment was maintained throughout the study."
Blinding of participants and personnel (performance bias)	Low risk	Quote: "Vitamin A and placebo oral solutions were supplied in identical bottles of 20 mL with dropper marked at 1 mL"
Blinding of outcome assessment (detection bias)	Low risk	Quote: "Treating physicians, nursing staffs, and the parents were unaware about the composition of the bottles."
Incomplete outcome data (attrition bias)	Low risk	Three patients from the intervention and two patients from the placebo group left against medical advice
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Benn et al. (2008)



Methods	Randomized placebo controlled trial conducted in Guinea-Bissau
Participants	Inclusion Criteria: Weight at least 2500 g at presentation and no signs of overt illness or malformations Exclusion Criteria: Weight <2500 g at presentation and/or signs of overt illness and/or malformations. Also, infants who died in the maternity ward before the vaccination team could arrive Total number randomized to the intervention group: 2145 Total number randomized to the control group: 2200
Interventions	Intervention: 50,000 IU vitamin A intradermally Control: 0.5 ml vegetable oil intradermally Common intervention given to all groups: 10 IU vitamin E intradermally
Outcomes	Primary outcomes: Overall mortality Other outcomes: Bulging fontanelles, vomiting, irritability, infections, fever, skin problems, and healthcare contacts
Notes	Vitamin A supplementation appeared to benefit boys but was harmful to girls

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The mother drew a lot from an envelope prepared by the study supervisor. Each envelope contained 100 lots—50 marked "1" and 50 marked "2"—indicating from which of two numbered bottles, "1" or "2," the child should receive the supplement" Comment: Most likely done
Allocation concealment (selection bias)	Low risk	Quote: "The lots were folded, making it impossible to tell what was written on them before they were opened" Comment: Most likely done
Blinding of participants and personnel (performance bias)	Low risk	Quote: "When asked, none of the three assistants who were responsible for the randomisation procedures at the hospital and at the heath centres had any idea which bottles contained vitamin A and which placebo. We concluded that the blinding of mothers and assistants was successful" Comment: Most likely done
Blinding of outcome assessment (detection bias)	Unclear risk	Quote: "Accumulating evidence for sex differential effects of vitamin A supplementation during the trial made us hypothesise before we started the analyses that supplementation would be particularly beneficial for boys" Comment: Most likely done
Incomplete outcome data (attrition bias)	Low risk	Total number of loss to follow up: 70 (1.6%) The loss to follow up was not balanced
Selective reporting (reporting bias)	Low risk	Authors seem to report all the outcomes irrespective of their statistical significance
Other bias	Low risk	No other risk of bias was noted

Benn et al. (2010)

Methods	Randomized placebo controlled two by two factorial trial conducted in Guinea-Bissau
Participants	Inclusion criteria: Weight <2500g at presentation Exclusion criteria: Weight >2500g at presentation Total number randomized to the intervention group: 864 Total number randomized to the control group: 872
Interventions	Intervention: 25,000 IU vitamin A intradermally Control: 0.5 ml vegetable oil intradermally Common intervention given to all groups: 10 IU vitamin E intradermally
Outcomes	Primary outcomes: Infant mortality Other outcomes: Fever, septicaemia, malaria, malnutrition, and respiratory infections
Notes	

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Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Once consent was provided, the mother drew an envelope from a bag. Each bag was prepared by the study supervisor and contained 48 envelopes; each envelope contained a lot name. Within each bag were 12 envelopes with lots marked "BCG 6," 12 marked "BCG 7," 12 marked "no BCG 6," and 12 marked "no BCG 7." The numbers "6" and "7" indicated from which of two numbered bottles, "6" or "7," the child should receive treatment (that is, either 25,000 IU vitamin A or placebo)" Comment: Most likely done
Allocation concealment (selection bias)	Low risk	Quote: "The envelopes were closed and non-transparent, making it impossible to identify the allocation before the envelopes were opened" Comment: Most likely done
Blinding of participants and personnel (performance bias)	Unclear risk	Quote: "Once consent was provided, the mother drew an envelope from a bag" "Each bag was prepared by the study supervisor and contained 48 envelopes; each envelope contained a lot name" Comment: Most likely done
Blinding of outcome assessment (detection bias)	Low risk	Quote: "Follow-up was performed by assistants who were unaware of the allocated treatment" Comment: Most likely done
Incomplete outcome data (attrition bias)	Unclear risk	Total number of loss to follow up: 145 (8.4%) The loss to follow up was not balanced
Selective reporting (reporting bias)	Low risk	Authors seem to report all the outcomes irrespective of their statistical significance
Other bias	Low risk	No other risk of bias was noted

Benn et al. (2014)

Methods	Double-blind, placebo-controlled randomized trial conducted in Guinea-Bissau	
Participants	Inclusion criteria: Normal birth-weight neonates who were healthy and due for BCG vaccination Exclusion criteria: Birth weight <2500 g at presentation or overt illness and/or malformations Total number randomized to the intervention group 1 (50,000 IU vitamin A): 2015 Total number randomized to the intervention group 25 (25,000 IU vitamin A): 2011 Total number randomized to the control group: 2022	
Interventions	Intervention: 50,000 IU vitamin A intradermally or 25,000 IU vitamin A intradermally Control: 0.5 ml Vegetable oil intradermally Common intervention given to all groups: 10 IU vitamin E intradermally	
Outcomes	Primary outcome: Infant mortality Other outcomes: None measured	
Notes		

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	 Quote. "Each envelope was prepared by the data manager, who did not take part in the enrolment procedures, and contained 48 folded lots indicating from which of 3 numbered bottles—"3," "4," or "5"—the child should receive his or her supplement" Comment: Most likely done
Allocation concealment (selection bias)	Low risk	Quote. "48 folded lots indicating from which of 3 numbered bottles—"3," "4," or "5"—the child should receive his or her supplement" Comment: Most likely done
Blinding of participants and personnel (performance bias)	Low risk	Quote. "At each inclusion site, the randomization procedure was carried out by 1 carefully trained assistant every day except during short vacations. After providing consent, the mother drew a lot from an envelope. Each envelope was



		prepared by the data manager, who did not take part in the enrolment procedures" Comment: Most likely done
Blinding of outcome assessment (detection bias)	Low risk	 Quote. "The registration system assistants and the special team were unaware of the allocated treatment, because they were not present during enrolment, and the information was not transferred to the children's vaccination card or follow-up forms" Comment: Most likely done
Incomplete outcome data (attrition bias)	Unclear risk	Quote: "Of 6053 children invited to participate, 6048 were randomly allocated to each of the 3 groups (50,000 IU vitamin A, 25,000 IU vitamin A, or placebo) (Figure 1). The 3 randomly assigned groups were similar in terms of their background characteristics (Table 1). A total of 176 deaths occurred; 2 of these were due to accidents and were censored. Fourteen deaths occurred after the child had been eligible for a national vitamin A campaign. Hence, censoring for accidents and subsequent VAS, the cohort had 160 deaths during 4125 person- years of risk, corresponding to an MR of 39 per 1000 person-years"
Selective reporting (reporting bias)	Low risk	Authors seem to report all the outcomes irrespective of their statistical significance
Other bias	Low risk	No other risk of bias was noted

Braga et al. (2011)

Methods	Randomized, double blind control study conducted in Brazil
Participants	Inclusion criteria: All infants included in this study were born locally and admitted to the Neonatal Intensive Care Unit (NICU) with a birth weight from 750 to 1499 g, and had no major congenital malformations, life threatening chromosomal alterations, or congenital infections
Interventions	 Intervention: Probiotic supplementation: <i>Bifidobacterium breve</i> and <i>Lactobacillus casei</i>: The intervention was started on the second day of life and was maintained until 30 d of life, a diagnosis of NEC, discharge from the hospital, or death, whichever occurred first. The dose was 3 ml human milk from the bank milk to which L. casei and B. breve had been added providing 3.5 × 10⁷ to 3.5 × 10⁹ CFU Comparison: The control group received the same volume of human milk without probiotics
Outcomes	Mortality, NEC and sepsis
Notes	Authors did not do intention to treat analysis. We created the intention to treat analysis by taking the number randomized as denominators. The data on outcomes was taken from table 2 of the main manuscript

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomization was carried out in blocks of 10, and the list of random numbers was generated by the subprogram Epitable from Epi-Info 6.04
Allocation concealment (selection bias)	Low risk	A sealed envelope with the identification number in ascending order, containing information about which group they belonged to, was provided for each infant and sent to the hospital's nutritional centre
Blinding of participants and personnel (performance bias)	Low risk	Neither the medical and nursing staff responsible for monitoring the infants nor the researchers were aware of which group the infants were allocated to
Blinding of outcome assessment (detection bias)	Low risk	Neither the medical and nursing staff responsible for monitoring the infants nor the researchers were aware of which group the infants were allocated to
Incomplete outcome data (attrition bias)	Low risk	Attrition in intervention group was 2% and 7% in the control group
Selective reporting (reporting bias)	Low risk	Most of the outcomes were reported
Other bias	Low risk	No other bias was noted

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Chowdhury et al. (2016)

Methods	A randomized controlled trial conducted in Bangladesh
Participants	 Inclusion crietria: Preterm (<33 woks), VLBW (<1500 g) infants who are able to tolerate oral feeds and survive beyond 48 h Exclusion criteria: Babies with suspicion of clinical sepsis, presence of prenatal asphyxia, major congenital anomaly and babies who expired due to other neonatal illness were excluded
Interventions	 Intervention: Probiotic supplementation: Bifidobacterium breve and Lactobacillus casei: The dose was 3 ml once daily of solution containing Bifidobacterium breve and Lactobacillus casei 106 CFU. The intervention was continued for at least 10 days Comparison: No probiotics
Outcomes	NEC, all-cause mortality
Notes	Authors did not perform intention to treat analysis, however, we created the intention to treat analysis from Figure 1. The data for mortality was taken from Figure 1 and the data for NEC was taken from table 2

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Coding to group 1 and 2 was done by a faculty of another department not related to this study. First case was selected to 1 group by lottery method and subsequent group was continued accordingly
Allocation concealment (selection bias)	Low risk	Participants and investigators did not know group allocation
Blinding of participants and personnel (performance bias)	Low risk	Probiotics were added to breast milk by registrar or assistant registrar of the corresponding unit before feeding
Blinding of outcome assessment (detection bias)	Low risk	Participants and investigators did not know group allocation
Incomplete outcome data (attrition bias)	Low risk	15% attrition. Reasons for loss to follow up reported.
Selective reporting (reporting bias)	Low risk	All expected outcomes were reported
Other bias	Low risk	No concerns for other risk of bias

Cooper et al. (2017)

Methods	A randomized double-blind controlled trial conducted in South Africa in community settings
Participants	 Inclusion criteria: "healthy", full term (37-42 weeks), born to HIV + formula feeding mothers, ≤3 days old, 2500-4500 g, singleton birth Exclusion criteria: Congenital illness or malformation affecting growth; significant perinatal disease, antibiotics in 1st 3 days of life, caregivers could not comply, or in another trial
Interventions	Intervention: Probiotics: Formula containing prebiotic (bovine milk-derived oligosaccharides) and probiotic ((B. lactis strain CNCM-I-3446 with 1 × 107 cfu/g. of powder formula). The duration of intervention was 6 months Comparison: Formula without prebiotic and probiotic
Outcomes	All-cause mortality
Notes	The data was taken from the last paragraph of the result section. We note that the intervention group received both prebiotics and probiotics and the control group did not receive any prebiotics or probiotics. Some of the study participants were HIV positive



Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The randomization was performed using the in-house TrialSys software".
Allocation concealment (selection bias)	Low risk	Formulas labelled similarly
Blinding of participants and personnel (performance bias)	Low risk	Parents (caregivers), investigators, study support staff, and the clinical project managers were blinded to the identity of the products.
Blinding of outcome assessment (detection bias)	Low risk	Likely same care-team
Incomplete outcome data (attrition bias)	Low risk	Total loss to follow up was 1%
Selective reporting (reporting bias)	Low risk	Authors do not seem to selectively report outcomes
Other bias	Low risk	No other risk of bias was noted.

Cui et al. (2019)

Methods	A prospective, double-blinded randomized study conducted in China
Participants	 Inclusion criteria: Formula-fed preterm infants, gestational age ≥30 and <37 weeks; birthweight ≥1500 g and ≤ 2000 g with vital sign and hemodynamic parameters stable Exclusion criteria: Congenital diseases, expected hospitalisations <2 weeks and maternal or neonatal antibiotics or other probiotics before admission
Interventions	Intervention: Lactobacillus reuteri DSM 17938, five drops daily for minim of 7 days. Each drop had 1×10 ⁸ colony-forming units Comparison: Placebo
Outcomes	Sepsis, NEC, growth
Notes	Data were taken from table 2

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Randomization was conducted according to a random computer- determined allocation order considering gestational age" Comment: Most likely done
Allocation concealment (selection bias) Blinding of participants and personnel (performance bias)	Unclear risk Low risk	No clear information available for allocation concealment Quote: "Blinding was possible because the nurses who administered L. reuteri to the infants were not involved in the daily care and the attending neonatal team was unaware of the randomization assignments"
Blinding of outcome assessment (detection bias)	Unclear risk	It was not clear if the families of the participating neonates were aware of the treatment assignments
Incomplete outcome data (attrition bias)	Low risk	About 18% attrition reported that was balanced in two groups
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

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Dashti et al. (2014)

Methods	Prospective triple-blinded, interventional, randomized clinical trial
Participants	 Inclusion criteria: Birth weight of 700-1800 g, stable hemodynamic, be able to have enteral feeding, and written parental consent. Exclusion criteria: Evidence or suspicion of congenital intestinal obstruction or perforation, prenatal or postnatal diagnosis of gastroschisis, large omphalocele, or congenital diaphragmatic hernia, and major congenital anomalies."
Interventions	 Intervention: Protexin (probiotics). Protexin (Restore): 1 × 109 CFU (colony forming unit), 1 g (one sachet) contains: Lactobacillus acidophilus, Lactobacillus rhamnosus, Bifidobacterium longum, Lactobacillus bulgaricus, Lactobacillus casei, Streptococcus thermophilus, Bifidobacterium breve, and Bifidobacterium The dose was as follows Neonates weighing 1000 g were fed with a half of sachet once daily (5×10⁸ CFU of probiotics), Neonates weighing 1001-1500 g were fed with 3/4 of a sachet once daily (7.5×10⁸ CFU of probiotics) Neonates weighing more than 1500 g were fed with a full sachet once daily (1×10⁹ CFU of probiotics). Control: "placebo that was physically indistinguishable from the probiotic powder"
Outcomes	NEC, mortality and sepsis
Notes	The data for NEC, mortality and sepsis was taken from table 2. We assumed that group A was the intervention group and group B was the control. Authors did not mention clearly in the paper which group is the intervention group and which one is the control group.The duration of intervention was not clearly stated

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	No clear data is available to support the assessment however the two groups were comparable after randomizations and allocation seems to be concealed. So less likely that randomizations was not done properly
Allocation concealment (selection bias)	Low risk	Quote: "To blind the trial the probiotic and placebo sachets were set in similar indistinguishable packages"
Blinding of participants and personnel (performance bias)	Low risk	Quote: "The control group was fed with milk and a placebo that was physically indistinguishable from the probiotic powder"
Blinding of outcome assessment (detection bias)	Low risk	Quote: "After starting the feeding, infants were observed continuously by a chart containing basic information like daily weight, feeding volume, abdominal girth, appearance of erythema of abdominal wall, loose stools with blood, vomiting, and orogastric tube suction volume. The amount of feeding was advanced slowly, if tolerated, with no more than a 20 ml/kg/d"
Incomplete outcome data (attrition bias)	Low risk	Quote: "Feeding was discontinued if there was any sign of feeding intolerance (defined as the presence of gastric aspirate in the amount that was more than a half of the previous feeding or abdominal distension)"
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Demirel et al. (2013)

Methods	Prospective, blinded, randomized control trial conducted in Turkey
Participants	Inclusion criteria: Neonates born ≤32 weeks and birthweight ≤1500 g who survived to start enteral feedings. Exclusion criteria: Major congenital anomalies, lack of parental consent, death in first seven days after study start
Interventions	Intervention: Probiotic supplementation: <i>S. boulardii</i> : Dose was 250 mg (5 billion cfu), added to breastmilk or formula, frequency was once daily and supplementation continued till discharge Comparison : Placebo
Outcomes	NEC, sepsis, mortality
Notes	Data was not analysed as intention to treat analysis. We created the intention to treat analysis by using the number randomized and the outcome numbers given in table 3

Risk of bias table



Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote, "Randomisation was simple and unadjusted and was performed using sequential numbers generated at the computer centre of the NICU"
Allocation concealment (selection bias)	Low risk	Quote, "The allocations were sealed in opaque, sequentially numbered envelopes"
Blinding of participants and personnel (performance bias)	Low risk	Quote, "The supplements were prepared by personnel on the breast milk team following the instructions in the sealed envelope. These individuals were the only personnel who were aware of the group assignments, and they were not involved in the care of the infants"
Blinding of outcome assessment (detection bias)	Low risk	Quote, "The supplements were prepared by personnel on the breast milk team following the instructions in the sealed envelope. These individuals were the only personnel who were aware of the group assignments, and they were not involved in the care of the infants"
Incomplete outcome data (attrition bias)	Low risk	7/278 dropped out. The intervention and the control group were fairly similar
Selective reporting (reporting bias)	Low risk	Authors seems to report all the outcomes mentioned in the analysis plan
Other bias	Low risk	No other bias was noted

Dilli et al. (2015)

Methods	Prospective, randomized, controlled trial conducted in TurkeyQuote
Participants	 Inclsuin criteria:: "VLBW infants with a gestational age of <32 weeks and a birth weight of <1500 g, born at or transferred to the NICU within the 1st week of life and fed enterally before inclusion, were eligible Exclusion criteria: Infants with any disease other than those linked to prematurity or congenital anomalies of the intestinal tract, not fed enterally or who died before the seventh day after birth, whose mothers had taken nondietary probiotic supplements, and whose parents refused to participate were excluded"
Interventions	 Intervention: Probitics/Prebiotics Multiple Arm trial 1) Probiotic (<i>Bifiidobacterium lactis</i>, 5×10⁹ colony-forming units) 2) Prebiotic (inulin, 900 mg) 3) Synbiotic (<i>Bifidobacterium lactis</i>, 5×10⁹ colony-forming units, 30 mg plus inulin, 900 mg 4) The control group: maltodextrin powder as a placebo The dose was 1 sachet per day of pre/probiotics with breast milk or formula until discharge or death, for a maximum of 8 weeks, whichever comes first
Outcomes	NEC, sepsis and mortality
Notes	The data were taken from table 3. We included data as probiotics+synbiotics vs. prebiotics+placebo

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Infants were randomized by balanced blocks using sealed envelopes"
Allocation concealment (selection bias)	Low risk	Quote: "Infants were randomized by balanced blocks using sealed envelopes"
Blinding of participants and personnel (performance bias)	Low risk	Quote: " In feeding units, sachets were opened and mixed with 1 ml of sterile water or breastmilk immediately before administration to infants who were receiving enteral feeding on the day of the supplementation. The feeding team was not involved in the care of the infant and followed directions on the sealed envelopes"

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Blinding of outcome assessment (detection bias)	Low risk	Quote: "The only personnel who knew of the infants group assignments were the investigators". As the outcomes were mostly objective, it is less likely that study had significant detection bias
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up reasons reported
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other sources of bias were noted

Dutta et al. (2015)

Methods	A randomized, placebo-controlled trial conducted in India
Participants	 Inclusion criteria: Neonates born at 27–33 weeks gestation in our hospital aged <96 h of life who were likely to either remain admitted in hospital or reside within 30 km of the hospital for the next 28 days who were tolerating at least 15 ml/kg/day of milk feeds Exclusion criteria: a gastro-intestinal malformation prior NEC or sepsis any life-threatening malformation that limited estimated life expectancy to less than a month
Interventions	 Intervention group: Probiotic Lactobacillus acidophilus (662.5 million), Lactobacillus rhamnosus (362.5 million), Bifidobacterium longum (87.5 million), and Saccharomyces boulardii (137.5 million) Four groups A. High-dose long course 10¹⁰ cells 12 hourly for 21 days B. High-dose short course 10¹⁰ cells 12 hourly for days 1-14; followed by placebo from days 15-21 C. Low-dose long course 10⁹ cells 12 hourly for 21 days D. Control group: Placebo for 21 days
Outcomes	NEC, mortality and sepsis
Notes	We combined all the probiotics groups (group $A + B + C$) to avoid the double counting of the placebo.

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "A block randomized sequence was generated online by an investigator who was not involved in the recruitment of subjects"
Allocation concealment (selection bias)	Low risk	Quote: "A block randomized sequence was generated online by an investigator who was not involved in the recruitment of subjects"
Blinding of participants and personnel (performance bias)	Low risk	Quote: "The external appearance and the contents of the sachets of high dose, low dose, and placebo were identical looking.
Blinding of outcome assessment (detection bias)	Low risk	As the intervention was concealed properly, less likely that outcomes assessors were aware of the allocation
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seems to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Edmond et al. (2015)

Methods	Randomized, double-blind, placebo-controlled trial conducted in Ghana
Participants	Recruited from the community, at least 2 h old, able to tolerate oral feeds and the family was likely to stay in the area. Parental consent was needed for inclusion to study.
Interventions	Intervention: Vitamin A: single dose: each dose was 50,000 IU Control: Placebo Common intervention given to all groups: Vitamin E

Outcomes	Primary outcome: All-cause mortality Other outcomes: Diarrhea, vomiting, bulging fontanelle, irritability, fever
Notes	Some of the participants were HIV positive

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The computerised block randomisation scheme was done with a block size of 20, so that in each block ten infants received vitamin A and ten received placebo" Comment: Most likely done
Allocation concealment (selection bias)	Low risk	Quote: "An independent statistician who was not part of the trial prepared the randomisation code at the WHO offices in Geneva, Switzerland. The code was available only to the Data Safety and Monitoring Board (DSMB) and their statistician" Comment: Most likely done
Blinding of participants and personnel (performance bias)	Low risk	Quote: "The research team and parents were fully unaware of the content of the capsules, which were only labelled with the infant number. Amanufacturer (StridesArcolab Limited, Bangalore, India) supplied the capsules. Separate staff, who were not part of the trial, labelled all capsules" Comment: Most likely done
Blinding of outcome assessment (detection bias)	Unclear risk	Quote: "" Comment: Most likely done
Incomplete outcome data (attrition bias)	Low risk	Quote: "Our loss to follow-up was only 1.1% at the time of ascertainment of our primary outcome at 6 months and only 2.9% at 12 months"
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Fernández-Carrocera et al. (2013)

Methods	Propsective, double-blind, randomized clinical trial conducted in Mexico
Participants	Inclusion criteria: Preterm newborns who weighed <1500 g admitted at the intensive and intermediate care units Exclusion criteria: Preterm newborns weighing <1500 g with a low Apgar score (<6 at 5 min), gastrointestinal malformations, genetic syndromes, asphyxia and IA–IB NEC stages according to Bell's
Interventions	 Intervention: Probiotic supplementation: "Lacidophilus 1.0×10⁹ colony forming units (CFU) CFU/g, Lactobacillus rhamnosus 4.4×108 CFU/g, Lactobacillus casei 1.0×10⁹ CFU/g, Lactobacillus plantarum 1.76×108 CFU/g, Bifidobacteruim infantis 2.76×107 CFU/g, Streptococcus theremophillus 6.6×105 CFU/g, each pack (Laboratorio Italmex SA)" Control: "The control group received their regular feeds from their mother's own milk when available with nothing added, or a premature infant formula"
Outcomes	All-cause mortality, NEC
Notes	Data were taken from table 3

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Infants were prospectively and randomly assigned to one of two groups using a random digit table
Allocation concealment (selection bias)	Low risk	"which was handled by the Human Milk Bank staff that was not involved in the care of the patients and adhered to proper trial procedures."
Blinding of participants and personnel (performance bias)	Low risk	"As allocation concealment measure, the study group received a suspension that matched the physical appearance of milk and the bottles were labelled only with the patient's name and identification number as usual."
Blinding of outcome assessment (detection bias)	Low risk	Attending physicians and nurses caring for the infants were blinded to the group assignments.

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Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up	
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes	
Other bias	Low risk	No other risk of bias was noted	

Giridhar et al. (2019)

Methods	Randomized, parallel group, placebo controlled trial
Participants	 Inclusion criteria: All infants admitted to the neonatal intensive care unit with birth weight between 750 and 1250 g and between 24 to 96 h of life Exclusion criteria: Lethal congenital malformations, terminal illness characterized by shock or bradycardia for more than 2 h, refusal of consent
Interventions	 Intervention 1: Vitamin A: 5000 IU (0.125 ml) IM on alternate days till establishment of adequate enteral feeds followed by oral vitamin A 10,000 IU (1 ml) once daily for a total duration of 28 days Control intervention 1: Placebo: 0.125 ml 0.9% normal saline IM alternate days till establishment of adequate enteral feeds followed by 1 ml oral dose of inert pharmacy made substance once daily for a total duration of 28 days
Outcomes	Proportion of infants with vitamin A deficiency (plasma retinol <200 mcg/L), mortality, sepsis
Notes	

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The random sequence was generated online from the web site www. randomizer. org"
Allocation concealment (selection bias)	Low risk	Quote: "Each stratum had permuted, even-numbered, randomly varying block sizes"
Blinding of participants and personnel (performance bias)	Low risk	Quote: "The investigators, supervisors, caregivers, laboratory personnel, and statistician were blinded to the intervention"
Blinding of outcome assessment (detection bias)	Low risk	Quote: "The investigators, supervisors, caregivers, laboratory personnel, and statistician were blinded to the intervention"
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Guney-Varal et al. (2017)

Methods	Prospective, randomized controlled trial conducted in Turkey
Participants	Inclusion Criteria: preterm infants with a gestational age ≤32 week and a birth weight ≤ 1500 g Exclusion Crietria: "detected chromosomal abnormalities, previous gastrointestinal system surgery, a diagnosis of metabolic disease, babies lost in the first postnatal week and babies with severe sepsis episode were excluded from the study"
Interventions	Intervention: Probiotic supplementation: Lactobacillus rhamnosus (4.1×10 ⁸ cfu) + Lactobacillus casei (8.2×108 cfu) + Lactobacillus plantorum (4.1×10 ⁸ cfu) + Bifidobacterium animalis (4.1×10 ⁸ cfu) Comparison: No probiotics
Outcomes	Mortality, NEC, sepsis
Notes	Data were taken from table 2





Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"Alternate randomization was used to enrol the infants to the study arms" This method of randomization is not adequate
Allocation concealment (selection bias)	High risk	Less likely to be done as the sequence generation was done on alternate basis
Blinding of participants and personnel (performance bias)	Unclear risk	No details are provided about blinding
Blinding of outcome assessment (detection bias)	Unclear risk	No details are provided about blinding
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Hariharan et al. (2016)

Methods	Prospective, randomized control trial conducted in India
Participants	Inclusion criteria: "Infants with birth weight <1250 g, gestation <32 weeks"
Interventions	Intervention: Lactobacillus acidophilus, Bifidobacterium bifidum, Saccharomyces boulardii 2.5 ×10 ⁹ UFC of each twice a day, from the 3rd day of life, for 6 week courses Comparison: No probiotic
Outcomes	Mortality, NEC, sepsis
Notes	Only abstract available

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	No details were available as only the abstract was available
Allocation concealment (selection bias)	Unclear risk	No details were available as only the abstract was available
Blinding of participants and personnel (performance bias)	Unclear risk	No details were available as only the abstract was available
Blinding of outcome assessment (detection bias)	Unclear risk	No details were available as only the abstract was available
Incomplete outcome data (attrition bias)	Unclear risk	No details were available as only the abstract was available
Selective reporting (reporting bias)	Low risk	Minimal loss to follow up
Other bias	Unclear risk	No details were available as only the abstract was available

Hernández-Enríquez et al. (2016)

Methods	Prospective randomized controlled trial conducted in Mexico
Participants	Inclusion criteria: Infants with very low birth weight
Interventions	 Intervention: Lactobacillus reuteri 5 drops, equivalent to 100 million colony forming units (1 × 10⁸ CFU) daily, whether they were newborns with weight >1000 to 1500 g. In the case of newborns with weight <1000 g them 3 drops were administered Lactobacillus reuteri (60 million CFU) daily The duration of supplementation was 20 days Compasrion: Group B (control group) received no probiotic.
Outcomes	NEC, sepsis
Notes	Study published in Spanish. Data extracted from the abstract

Risk of bias table

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Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Study published in Spanish. Details not available
Allocation concealment (selection bias)	Unclear risk	Study published in Spanish. Details not available
Blinding of participants and personnel (performance bias)	Unclear risk	Study published in Spanish. Details not available
Blinding of outcome assessment (detection bias)	Unclear risk	Study published in Spanish. Details not available
Incomplete outcome data (attrition bias)	Unclear risk	Study published in Spanish. Details not available
Selective reporting (reporting bias)	Unclear risk	Study published in Spanish. Details not available
Other bias	Unclear risk	Study published in Spanish. Details not available

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Huaxian, 2013

Methods	Randomized controlled trial conducted in China
Participants	Inclusuion criteria: Preterm babies admitted to NICU
Interventions	Intervention: Probiotics and early minimal feeding Control: Early minimal feedings
Outcomes	NEC
Notes	Only abstract was available

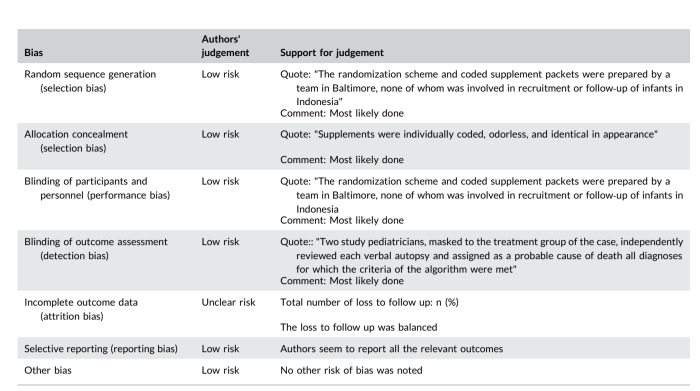
Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	No details were available as only the abstract was available
Allocation concealment (selection bias)	Unclear risk	No details were available as only the abstract was available
Blinding of participants and personnel (performance bias)	Unclear risk	No details were available as only the abstract was available
Blinding of outcome assessment (detection bias)	Unclear risk	No details were available as only the abstract was available
Incomplete outcome data (attrition bias)	Unclear risk	No details were available as only the abstract was available
Selective reporting (reporting bias)	Low risk	Minimal loss to follow up
Other bias	Unclear risk	No details were available as only the abstract was available

Humphrey et al. (1996)

Methods	A placebo-controlled trial conducted in Indonesia
Participants	Inclusion criteria: All infants born at Hasan Sadikin Hospital in Bandung, Indonesia from June 18, 1992, to June 3, 1993 Exclusion criteria: Infants that were considered very low birthweight (<1500 g) and infants with life-threatening conditions Total number randomized in the intervention group: 1034 Total number randomized in the control group: 1033
Interventions	Intervention: 1 dose of 52 μmol of vitamin A (as retinyl palmitate) orally Control: Placebo (<0.10 μmol of vitamin A) orally Common intervention given to all groups: 23 μmol vitamin E (as <i>dl-α-</i> tocopherol) orally
Outcomes	Primary outcome : Infant Morbidity & Mortality Other outcomes : Diarrhea, fever, cough, rapid breathing, wheezing, otitis media, pneumonia, sepsis
Notes	

Risk of bias table



Hussain et al. (2016)

Methods	Propsective, randomized controlled trial conducted in Pakistan
Participants	 Inclusion criteria: Pre-term neonates <36 weeks gestation Low birth weight neonates <2.5 Kg Both genders Both NG feed and bottle feed neonates All neonates that were admitted at day 1 of life Exclusion criteria: Neonates <30 weeks low birth weight neonates, <1.5 Kg Neonate on mechanical ventilatory support IUGR (gestational age>36 weeks and weight <2.5 kg Patients with congenital cyanotic heart diseases or has birth asphyxia and persistent cyanosis and need of oxygen inhalation.
Interventions	Intervention: Probiotic supplementation: Bifidobacteria. Comparison: No probiotics
Outcomes	NEC
Notes	The strain of the probiotic was not clearly stated. The dose and duration of the intervention was not given

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "neonates were divided in two groups by using random number tables"
Allocation concealment (selection bias)	Unclear risk	Seems unlikely, but no clear statement is made regarding this matter
Blinding of participants and personnel (performance bias)	Unclear risk	No supporting statement is age in this regard
Blinding of outcome assessment (detection bias)	Unclear risk	No clear statement is made in this regard



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Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up	
Selective reporting (reporting bias)	Low risk	Authors seem to report the relevant outcomes	
Other bias	Low risk	No other source of bias was noted	

Kaban et al. (2019)

Methods	A double-blind randomized controlled clinical trial conducted in Indonesia
Participants	Inclusion criteria: Gestational age of 28–34 weeks, birth weight of 1000–1800 g in a stable condition Exclusion criteria: Lower gastrointestinal tract obstruction, massive gastrointestinal tract bleeding, NEC, sepsis and shock, and refusal of the infants' parents to participate in the study
Interventions	 Intervention: Lactobacillus reuteri DSM 17938, duration of at least 7 days or until the subject was discharged, experienced NEC, or died, five drops per day, 10⁸ colony-forming units/day Comparison: Placebo: The placebo contains a mixture of pharmaceutical-grade medium-chain triglycerides and sunflower oil together with pharmaceutical-grade silicon
Outcomes	Mortality, sepsis and death
Notes	Data were taken from table 4

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Quote: "Subjects were allocated to the groups by a third party using a simple alternating randomization technique" Comment: Alternation allocation of patients is not random
Allocation concealment (selection bias)	Unclear risk	Quote: "Subjects were allocated to the groups by a third party using a simple alternating randomization technique" Comment: It is not clear if the allocation was revealed before the patient were allocated to intervention or placebo
Blinding of participants and personnel (performance bias)	Unclear risk	Authors mentioned that it was a double blind trial but no details were provided on how the blinding was done
Blinding of outcome assessment (detection bias)	Unclear risk	Authors mentioned that it was a double blind trial but no details were provided on how the blinding was done
Incomplete outcome data (attrition bias)	Low risk	No attrition was reported
Selective reporting (reporting bias)	Unclear risk	Authors seem to report all the relevant outcomes
Other bias	Unclear risk	No other risk of bias was noted

Klemm et al. (2008)

Methods	Community-based, double-masked, cluster-randomized, and placebo-controlled trial conducted in Bangladesh
Participants	 Inclusion criteria: Infants born to consenting mothers who were participating in the parent trial Exclusion criteria: Infants of consenting mothers who had died before they could be supplemented by staff, infants born outside of the study area, and infants who could not be reached to receive a supplement during the first 30 days after birth Total number randomized in the intervention group: 8525 Total number randomized in the control group: 8591
Interventions	Intervention: 50,000 IU vitamin A Control: Placebo Common intervention given to all groups:
Outcomes	Primary outcome: All-Cause infant mortality Other outcomes: Bulging fontanel
Notes	 "Follow-up of the trial cohort at 3 years of age revealed no evidence of adverse effects associated with having had a perinatal bulging fontanel in terms of cognitive, motor, and behavioral test outcomes" This study was concluded early by direction of the Data Safety and Monitoring Board

Risk of bias table



Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Sectors were listed in geographically contiguous order and were randomized in blocks of 4 within each of 3 previously randomized maternal supplementation trial treatment arms" Comment: Most likely done
Allocation concealment (selection bias)	Unclear risk	Quote: "Community maps of the area were developed, homes were issued numeric addresses, and married women of reproductive age were enumerated and issued unique study identification numbers" Comment: Most likely done
Blinding of participants and personnel (performance bias)	Low risk	Quote: "The supplements for both groups were opaque gelatinous capsules identical in shape, size, and color containing edible oil" Comment: Most likely done
Blinding of outcome assessment (detection bias)	Unclear risk	Quote: "Infant vital status was assessed weekly at home for the first 12 weeks of life by field staff and then again at 24 weeks of age" Comment: Most likely done
Incomplete outcome data (attrition bias)	Low risk	Total number of loss to follow up: 11 (0.07%) The loss to follow up was not balanced
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Malaba et al. (2005)

Methods	Randomized, placebo-controlled, 2-by-2 factorial design trial was conducted in Zimbabwe
Participants	 Inclusion criteria: Neither the mother nor the infant had an acutely life-threatening condition, the infant was a singleton with a birth weight of >1500 g, and the mother planned to stay in the region after delivery Exclusion criteria: Either the mother and/or the infant had an acutely life-threatening condition, the infant was not a singleton or had a birth weight of <1500 g, and the mother did not plan to stay in the region after delivery Total number randomized in the intervention group: Aa: 3529 Ap: 3529 Pa: 3530 Total number randomized in the control group: Pp: 3522
Interventions	 Intervention: Mothers received 400,000 IU vitamin A (as retinyl palmitate) and infants received 50,000 IU vitamin A (Aa group) Mothers received 400,000 IU vitamin A and infants received placebo (Ap group) Mothers received placebo and infants received 50,000 IU vitamin A (Pa group) Control: Both mothers and infants received placebo (Pp group) Common intervention given to all groups: Soy oil base with vitamin E as a preservative (50 IU per maternal capsule; 10 IU per infant capsule)
Outcomes	Primary outcome: Infant mortality Other outcomes: None
Notes	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "A separate team at Johns Hopkins University prepared the study capsule packets. Study identification numbers were randomly allocated to the treatment groups by computer in blocks of 12" Comment: Most likely done
Allocation concealment (selection bias)	Low risk	Quote: "Lists linking the study number to the treatment were kept in sealed envelopes and encrypted computer files" "Treatment and placebo capsules appeared identical" Comment: Most likely done

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Blinding of participants and personnel (performance bias)	Unclear risk	Quote: "Treatment and placebo capsules appeared identical"
	TISK	Comment: Most likely done
Blinding of outcome assessment (detection bias)	Low risk	Quote: "Cause of death was determined from medical records for infants who died in a hospital or from a review of verbal autopsy information by a study pediatrician, who was masked to treatment group, for infants dying at home" Comment: Most likely done
Incomplete outcome data (attrition bias)	Unclear	Total number of loss to follow up: n (%)
	risk	The loss to follow up was not balanced
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Masanja et al. (2015)

Methods	A randomized, double-blind, placebo-controlled trial conducted in Tanzania
Participants	 Inclusion criteria: Able to feed orally, parents planned to stay in the study area for at least 6 months, and informed written consent was provided Exclusion criteria: Infants enrolled in another trial Total number randomized in the intervention group: 15,995 Total number randomized in the control group: 16,004
Interventions	Intervention: 50,000 IU of vitamin A (as retinol palmitate) orally Control: Placebo (minute amounts of vitamin E (9.5–12.6 IU) in soybean oil) orally Common intervention given to all groups: Minute amounts of vitamin E (9.5–12.6 IU) in soybean oil
Outcomes	Primary outcome: Mortality between supplementation and 6 months of age Other outcomes: Mortality between supplementation and 28 days of age, Mortality between supplementation and 365 days of age, and hospital admission in the first 6 months of life
Notes	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "We randomly assigned infants to receive either vitamin A or a placebo. The unit of randomisation was the individual infant. Block randomisation was done at WHO (Geneva, Switzerland) in block sizes of 20 (ten infants received vitamin A and ten received placebo)" Comment: Most likely done
Allocation concealment (selection bias)	Low risk	Quote: "The vitamin A and placebo capsules were identical in taste and appearance. Capsules were individually packed in blister packs of two capsules each; one for the dose and the second for the backup dose. Labels for the capsules were printed at WHO with country and infant study number in sequential order" Comment: Most likely done
Blinding of participants and personnel (performance bias)	Low risk	Quote: "Codes for the experimental regimens were kept with the data and safety monitoring board and broken during the analysis after a cleaned and locked database for the study was submitted to WHO" Comment: Most likely done
Blinding of outcome assessment (detection bias)	Low risk	 Quote: "All reported deaths of children were investigated and trained field staff visited the family at least 6 weeks after the date of death to do a verbal autopsy interview" "Trained field interviewers visited enrolled infants at home (or in health facilities for cases in which the mother and child were not discharged after delivery) 1 day and 3 days after dosing to monitor possible adverse events after supplementation" Comment: Most likely done
Incomplete outcome data (attrition bias)	Unclear risk	Total number of loss to follow up: n (%) The loss to follow up was not balanced
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Mazumder et al. (2015)



Methods	Randomized, double-blind, placebo-controlled trial conducted in India
Participants	Inclusion criteria : Livebirths born in the study area Exclusion criteria : Died before screening, serious illness, and/or were admitted into the intensive care unit Total number randomized in the intervention group: 22,493 Total number randomized in the control group: 22,491
Interventions	Intervention: 50,000 IU vitamin A plus vitamin E 9.5–12.6 IU Control: Placebo (vitamin E 9.5–12.6 IU) Common intervention given to all groups: Vitamin E 9.5–12.6 IU
Outcomes	Primary outcome : Infant mortality from supplementation to 6 months Other outcomes : Neonatal mortality, mortality between supplementation and 12 months of age, infant hospital admission one or more times due to any illness between supplementation and 6 months of age, potential adverse events in the 3-day period following supplementation, and vitamin A status in a sub-sample of infants at 2 weeks and 3 months of age
Notes	

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The unit of randomisation was the individual infant. We randomly assigned infants using a block randomisation scheme with a block size of 20, so that in each block ten infants received vitamin A and ten received placebo. The randomisation list was prepared offsite at WHO (Geneva, Switzerland) by a statistician not otherwise involved with the trial" Comment: Most likely done
Allocation concealment (selection bias)	Low risk	Quote: "The vitamin A and placebo capsules were identical in colour, shape, and size. Capsules were individually packaged in identical blister packs with two capsules, one for the dose and the other as a backup" Comment: Most likely done
Blinding of participants and personnel (performance bias)	Low risk	Quote: "Investigators, participants' families, and the data analysis team were masked to treatment allocation" Comment: Most likely done
Blinding of outcome assessment (detection bias)	Low risk	Quote: "Research staff were trained to do surveillance, interview families, obtain informed consent, give capsules, collect baseline and follow-up information, and data capture" Comment: Most likely done
Incomplete outcome data (attrition bias)	Low risk	Total number of loss to follow up: 40 (0.09%) The loss to follow up was not balanced
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Nandhini et al. (2016)

Methods	A prospective, double blind. controlled trail conducted in India
Participants	Inclusion criteria: Preterm infants, enterally fed, 28–34 weeks, birthweight >1000 g, admitted to the NICU Exclusion crietria: major congenital anomalies, surgical problems of the GI tract, severe birth asphyxia, early onset sepsis
Interventions	 Intervention: Synbiotic supplementation: probiotics (Lactobacillus acidophilus (700 million CFU), Bifidobacterium longum (400 million CFU), Lactobacillus rhamnosus (400 million CFU), Lactobacillus plantaris (300 million CFU), Lactobacillus casei (300 million CFU), Lactobacillus bulgaricus (300 million CFU), Bifidobacterium infantis (300 million CFU) and Bifidobacterium breve (300 million CFU) + Prebiotic (fructo-oligosaccharide) The dose was given two times daily for 7 days. The Probitiotics were mixed with breastmilk Comparison: Standard of care without synbiotics
Outcomes	Mortality, neonatal sepsis, NEC
Notes	Data were included from table 2. We created the intervention to treat analysis



Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	After obtaining informed consent from the parents, neonates satisfying the inclusion criteria were randomized prior to starting enteral feeds into two groups using computer generated random numbers kept in opaque sealed envelopes
Allocation concealment (selection bias)	Low risk	After obtaining informed consent from the parents, neonates satisfying the inclusion criteria were randomized prior to starting enteral feeds into two groups using computer generated random numbers kept in opaque sealed envelopes
Blinding of participants and personnel (performance bias)	Unclear risk	No details are provided in the study
Blinding of outcome assessment (detection bias)	Unclear risk	No details are provided in the study
Incomplete outcome data (attrition bias)	Low risk	Miminal loss to follow up
Selective reporting (reporting bias)	Low risk	Author seem to report all the outcomes
Other bias	Low risk	No other bias was noted

Niekerk et al. (2015) (HIV exposed)

Methods	A randomized, double blind, placebo-controlled clinical trial conducted in South Africa
Participants	 Inclsuion criteria: "(i) HIV-positive or HIV-negative mothers who gave birth to a premature and VLBW baby at TBCH and consented to participate in the study; (ii) only breastfeeding mothers, regardless of their HIV status; and (iii) HIV-positive mothers that were on the prevention of mother to child transmission treatment schedule. Babies were included if they (i) had a birth weight of 500 g and 1250 g; (ii) were either HIV-exposed or HIV-unexposed; and (iii) received breast milk (either from their mothers or donor breast milk)" Exclusion criteria: abnormalities such as gastroschisis, a large omphalocele or congenital diaphragmatic hernia
Interventions	Intervention: Probiotics: L. rhamnosus GG [0.35 ×10^9 colony-forming units (CFU)] and B. infantis (0.35 ×10 ⁹ CFU), 5 drops daily for 28 days Control: Placebo
Outcomes	Mortality and NEC
Notes	Data were taken from table 3. This was a multiple arm trial and we included the data for HIV exposed and HIV nonexposed separately

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Participants were randomized into either the study or control groups (probiotic vs. placebo supplementation) with a random-number sequence allocated to each participant number"
Allocation concealment (selection bias)	Low risk	Quote: "No differences in the colour and appearance of the probiotic and placebo were noted. The probiotic and placebo were blinded with the use of a colour- coded label (orange or purple)"
Blinding of participants and personnel (performance bias)	Low risk	Quote: "The attending physician, nurses, researcher, research assistant and study participants were blinded to the group assignment"
Blinding of outcome assessment (detection bias)	Low risk	Quote: "The attending physician, nurses, researcher, research assistant and study participants were blinded to the group assignment"
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other bias was noted

Niekerk et al. (2015) (HIV nonexposed)



Methods	Same as above study
Participants	
Interventions	
Outcomes	
Notes	

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Participants were randomized into either the study or control groups (probiotic vs. placebo supplementation) with a random-number sequence allocated to each participant number"
Allocation concealment (selection bias)	Low risk	Quote: "No differences in the colour and appearance of the probiotic and placebo were noted. The probiotic and placebo were blinded with the use of a colour- coded label (orange or purple)"
Blinding of participants and personnel (performance bias)	Low risk	Quote: "The attending physician, nurses, researcher, research assistant and study participants were blinded to the group assignment"
Blinding of outcome assessment (detection bias)	Low risk	Quote: "The attending physician, nurses, researcher, research assistant and study participants were blinded to the group assignment"
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other bias was noted

Oncel et al. (2014)

Methods	A prospective, double-blinded, randomized, placebo controlled trial conducted in Turkey			
Participants	Inclusion criteria: Preterm infants with a gestational age ≤32 weeks and birth weight ≤1500 g, which survived to feed enterally, were eligible for the study. Exclusion crietria: Major congenital malformations and lack of parental consent.			
Interventions	Intervention group: Probiotic supplementation:Infants in the probiotic group received 5 drops of oil-based suspension containing 1×10 ⁸ colony-forming units of <i>Lactobacillus reuteri</i> DSM 17938". The probiotic was given once a day, until death or discharge from the hospital Comparison: Placebo			
Outcomes	Mortality, NEC			
Notes	We created the intention to treat analysis by using the numbers from fig. 1 and the table 2. We included all the patients who were randomized to intervention or control group			

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Study infants were randomly assigned to probiotic or placebo by using sequential numbers generated at the computer centre of the NICU by 1:1 allocation ratio
Allocation concealment (selection bias)	Low risk	The allocations were contained in opaque, sequentially numbered sealed envelopes
Blinding of participants and personnel (performance bias)	Low risk	Identical vial containing only oil base were administered following the same protocol as the probiotic group.
Blinding of outcome assessment (detection bias)	Low risk	Seems less likely

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Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other bias was noted

Panigrahi et al. (2017)

Methods	A community-based, double-blind, placebo controlled randomized trial conducted in India
Participants	 Inclusion criteria: Neonate >24 h and <96 h old, ≥2000 g at birth, breastfeeding begun by 24 h of life, ability to tolerate oral feeds, informed consent by parent or guardian Exclusion criteria: Evidence or suspicion of clinical sepsis before the infant was randomized, gestational age reported voluntarily by the mother to be <35 weeks, infant >96 h old, infant did not cry immediately after birth, mother had fever (>38°C) within 2 days of delivery, mother had foul-smelling amniotic discharge within 2 days of delivery, mother had foul-smelling amniotic discharge within 2 days of delivery, mother unlikely to stay in the village for 60 days, difficulty in carrying out study (maternal sickness etc.), or presence of major congenital anomalies (defined as any malformation that was felt to be life-threatening or that required surgical intervention)"
Interventions	Intervention: Synbiotic supplementation: The synbiotic preparation consisted of a capsule containing ~109 <i>Lactobacillus</i> <i>plantarum</i> ATCC strain 202195 and 150 mg of fructooligosaccharide with 100 mg maltodextrin as excipient The synbiotic was administered orally to the newborns for 7 days beginning on day 2-4 of life Comparison : Placebo capsules contained only 250 mg of maltodextrin
Outcomes	Mortality, sepsis
Notes	This was a community based study. Data were taken from table 2

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Each assignment was the product of a random permutation scheme that assigned 2 intervention and 2 placebo slots to each of 38 consecutive blocks of 4 assignments for each village. This numbered list and corresponding bar codes were created by the GCRC (General Clinical Research Center) at the University of Maryland with assistance from the Department of Bioinformatics"
Allocation concealment (selection bias)	Low risk	Quote: "This numbered list and corresponding bar codes were created by the GCRC (General Clinical Research Center) at the University of Maryland with assistance from the Department of Bioinformatics and given to the clinical trial supplier (Laxai USA, South Plainfield, New Jersey, USA) for labelling of the synbiotics and to prepare packages for each village to be assigned consecutively to enrolled subjects"
Blinding of participants and personnel (performance bias)	Low risk	Quote: "physicians had no access to randomization, distribution, or administration of the intervention making them completely blinded to the intervention"
Blinding of outcome assessment (detection bias)	Low risk	Quote: "physicians had no access to randomization, distribution, or administration of the intervention making them completely blinded to the intervention"
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Rahmathullah et al. (2003)

 Methods
 Community based, randomized, double-blind, placebo controlled trial conducted in southern India

 Participants
 Inclusion criteria: Liveborn infants from all pregnancies in the participating villages

 Exclusion criteria: Stillbirths, Miscarriages, Any delivery more than 20 km outside the study area, and infants who died before the study team arrived

 Total number randomized in the intervention group: 6624

 Total number randomized in the control group: 6570



Interventions	Intervention: 24,000 IU vitamin A Control: Placebo Common intervention given to all groups: Edible oil solution	
Outcomes	Primary outcome: Infant mortality at 6 months Other outcomes: None	

Notes

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Randomisation was at the individual level, stratified by geographical area in blocks of four. Because births were likely in a variety of locations, randomisation was conducted at the time of recruitment" Comment: Most likely done
Allocation concealment (selection bias)	Low risk	Quote: "Treatment codes were kept in a sealed envelope in a locked filing cabinet in Baltimore" Comment: Most likely done
Blinding of participants and personnel (performance bias)	Low risk	Quote: "Investigators, study staff, and mothers were masked to the assigned treatment" Comment: Most likely done
Blinding of outcome assessment (detection bias)	Unclear risk	 Quote: "Project staff visited the household every 2 weeks to assess the vital status of the child and any morbidity. "Investigators, study staff, and mothers were masked to the assigned treatment" Comment: Most likely done
Incomplete outcome data (attrition bias)	Unclear risk	Total number of loss to follow up: 143 (1.1%) The loss to follow up was not balanced
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Rehman et al. (2018)

Methods	A randomized control conducted in Pakistan
Participants	 Inclsuion criteria: "preterm infants having gestation of 27 to 36 + 7 weeks; they were VLBW (<1500 g) and they survived to feed enterally" Exclusion criteria: None available as only abstract was available for data extraction
Interventions	Authors mentioned that they supplemented the neonates with Probiotic mixtures. Exact strain was not clear
Outcomes	NEC
Notes	Study only available as abstract

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Study available as abstract
Allocation concealment (selection bias)	Unclear risk	Study available as abstract
Blinding of participants and personnel (performance bias)	Unclear risk	Study available as abstract
Blinding of outcome assessment (detection bias)	Unclear risk	Study available as abstract
Incomplete outcome data (attrition bias)	Unclear risk	Study available as abstract
Selective reporting (reporting bias)	Unclear risk	Study available as abstract
Other bias	Unclear risk	Study available as abstract

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Rojas et al. (2012)

Methods	A multicenter, double-blinded, randomized, placebo-controlled trial conducted in Colombia
Participants	 Inclusion Cciteria: "Preterm infants admitted to NICU, birth weight ≤ 2000 g, haemodynamically stable (blood pressure not requiring boluses or pressors), and ≤ 48 h of age" Exclusion criteria: "Infants with evidence or suspicion of congenital intestinal obstruction or perforation, gastroschisis, large omphalocele, congenital diaphragmatic hernia, major congenital heart defects, or anticipated transfer to a NICU not participating in the study were excluded"
Interventions	Intervention: Probiotic administration:"Infants in the probiotic group received 5 drops of an oil-based suspension containing 10 ⁸ colony-forming units of L reuteri DSM 17938" Comparison: Oil based placebo
Outcomes	Mortality, NEC
Notes	Data were taken from table 2. The duration of the intervention was not clearly stated

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: " Study participants were randomly assigned to probiotic or placebo by the use of a computer-generated balanced block randomization scheme"
Allocation concealment (selection bias)	Low risk	Quote: "Treatment assignment was performed by using sealed, sequentially numbered, opaque envelopes, color-coded for strata, available in each NICU pharmacy"
Blinding of participants and personnel (performance bias)	Low risk	Quote: "Infants were administered probiotic or placebo regardless of whether enteric feeds were started"
Blinding of outcome assessment (detection bias)	Low risk	The authors do not report clearly if the outcome assessment was blinded, however in the setting of adequate allocation concealment and blinding of interventions it is less likely the outcome assessors knew the intervention vs placebo group
Incomplete outcome data (attrition bias)	Low risk	Quote: "The study was terminated before the completion of the targeted sample because of a substantial drop in patient recruitment among participating institutions as well as funding restrictions that limited our ability to recruit the required additional subjects"
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Roy et al. (2014)

Methods	A prospective, randomized, double blind, placebo-controlled trial in India
Participants	 Inclusion criteria: "Admission to the NICU, a stable oral feeding within 72 h of birth and an informed parental consent; gestational age (GA) < 37 weeks; birth weight <2500 g; adequate renal and liver function; a postnatal age <2 week; did not have baseline fungal colonization at enrolment (with colonization defined by isolation of fungi from a culture specimen obtained from any site during the first 3 days of life); did not receive any form of antifungal prophylaxis other than the probiotic used" Exclusion criteria: "presence of major congenital malformation; antenatal and perinatal risk factors for sepsis, major congenital malformation; stigma of congenital infection; severe lesions diagnosed by cranial ultrasound (e.g. intraventricular haemorrhage (IVH) grade 3 and 4 and major ischemic lesions); altered liver and renal function; likely to die within 72 h of birth; and babies of mothers taking supplemental probiotics by capsule/powder"
Interventions	Intervention: "6×109 CFU Lactobacillus: half a sachet of Lactobacillus acidophilus 1.25 billion, B. longum 0.125billion, B. bifidum 0.125billion, and B. lactis 1 billion/1g sachet, daily for 6 weeks or NICU discharge Control:sterile water in breastmilk
Outcomes	NEC, mortality and sepsis
Notes	Data were taken from table 2





Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The newborns were randomized into two groups by a random-generated (computer-generated), predetermined number table"
Allocation concealment (selection bias)	Low risk	Quote: "All doctors, nurses, laboratory staff, and parents are blind to the randomized allocation"
		Less likely that allocation was revealed
Blinding of participants and personnel (performance bias)	Low risk	Quote: "All doctors, nurses, laboratory staff, and parents are blind to the randomized allocation"
Blinding of outcome assessment (detection bias)	Low risk	Quote: "All doctors, nurses, laboratory staff, and parents are blind to the randomized allocation"
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Saengtawesin et al. (2014)

Methods	A prospective, randomized control trial conducted in Thailand
Participants	Inclusion criteria: "all preterm infants with gestational age ≤34 weeks and birth weight ≤1500 g" Exclusion criteria: "Very low birth weight preterm infants who had severe birth asphyxia, chromosome anomalies, cyanotic congenital heart disease, congenital intestinal obstruction, gastroschisis, omphalocele, nil per oral >3 weeks and parents who declined consent"
Interventions	Intervention: Probiotics: "Infloran(1×10^9 Lactobacillus acidophilus and 1×109 Bifidobacterium bifida): 125 mg/kg/dose two times daily for 6 weeks Control: "unsupplemented breast milk or preterm formula"
Outcomes	Neonatal Sepsis, NEC
Notes	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "randomized by blocks of four into two groups, study and control group" Comment: Most likely done
Allocation concealment (selection bias)	Low risk	Quote: "Infants in the control group were fed either breast milk or premature formula alone. Infloran® was stored in a refrigerator at 4°C to 8°C, at the hospital pharmacy and sent then to the neonatal unit according to prescription" Comment: The sequence was kept at a central location
Blinding of participants and personnel (performance bias)	Low risk	Seems less likely that participants were aware of the intervention
Blinding of outcome assessment (detection bias)	Low risk	Quote: "Whenever an infant was suspected to have NEC, he/she was evaluated by two attending neonatologist" Seems less likely that outcome assessment was biased
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Samanta et al. (2009)

Methods	A prospective randomized double-blind control trial conducted in India
Participants	 Inclusion criteria: "preterm (<32 weeks) VLBW (<1500 g) born between October 2007 and March 2008, started feed enterally, survived beyond 48 h of life" Exclusion criteria: "babies with major congenital and GI anomalies and babies who expired due to other neonatal illnesses"
Interventions	Intervention: Probitotics containing Bifidobacteria infantis, Bifidobacteria bifidum, Bifidobacteria longum, Lactobacillus acidophilus: Dose 2.5 billion cfu each of 4 strains twice daily until NICU discharge Control: Breastmilk without probiotics
Outcomes	Mortality, sepsis
Notes	

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "the infants were randomly assigned to two groups by random number table sequence"
Allocation concealment (selection bias)	Unclear risk	No details are provided
Blinding of participants and personnel (performance bias)	Unclear risk	No details are provided
Blinding of outcome assessment (detection bias)	Unclear risk	No details are provided
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Sari et al. (2011)

Methods	Prospective, randomized, controlled trial Turkey
Participants	Inclusion criteria: Infants with gestational age of <33 weeks or birth weight of 1500 g Exclusion criteria: Major congenital malformations and lack of parental consent
Interventions	Intervention: "Lactobacillus sporogenes with a dose of 350 000 000 c.f.u. once a day until discharge" Control: "The control group was fed with breast milk or formula without the probiotics
Outcomes	NEC, mortality, sepsis
Notes	Data were taken from table 2 of the study. We created the intention to treat analysis

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "using sequential numbers generated at the computer center of"
Allocation concealment (selection bias)	Low risk	Quote: "The allocations were contained in opaque, sequentially numbered sealed envelopes"
Blinding of participants and personnel (performance bias)	Low risk	Quote: "Thus, the only personnel who knew of the infants' group assignments were the investigators and those in the breast-milk team who were not involved in the care of the study infants"
Blinding of outcome assessment (detection bias)	Low risk	

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Quote: "Whenever an infant was suspected to have NEC, the infant was
evaluated by two senior-attending neonatologists who did not know the
group assignment of the infant"

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		group assignment of the man
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the outcomes irrespective of their statistical significance
Other bias	Low risk	No other risk of bias was noted

Serce et al. (2013)

Methods	A prospective, double blind, placebo controlled trial conducted in Turkey
Participants	 Inclusion criteria: "VLBW infants (gestational age ≤32 weeks; birth weight ≤1500 g) who survived to feed enterally were eligible for the trial" Exclusion criteria: "Infants who had severe asphyxia (stage III), major congenital anomalies, those who had been fasted for more than 3 weeks, died in the first postnatal 14 days and infants who used antifungal therapy were excluded"
Interventions	 Intervention: "The study group received Saccharomyces boulardii (50 mg/kg equal to 0.5×10⁹ cell/kg per dose twice daily until discharge Comparison: placebo (distilled water; 1 ml per dose twice daily)
Outcomes	Mortality, NEC, sepsis
Notes	Data were taken from table 3

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Randomization was performed by using sequential numbers generated at the computer"
Allocation concealment (selection bias)	Low risk	Quote: "The allocations were contained in opaque, sequentially numbered sealed envelopes"
Blinding of participants and personnel (performance bias)	Low risk	Quote: "The allocations were contained in opaque, sequentially numbered sealed envelopes" Comment less likely that participants were aware
Blinding of outcome assessment (detection bias)	Unclear risk	No information was provided on who made the assessment
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the outcomes irrespective of their statistical significance
Other bias	Low risk	No other risk of bias was noted

Shadkam et al. (2015)

Methods	A prospective, triple blind, placebo controlled trial conducted in Iran
Participants	 Inclusion criteria: "premature infants admitted at the neonatal intensive care unit (NICU) during October 2012-March 2013. Gestational age of infants was estimated at 28–34 weeks using the Dubowitz method, and birth weight of infants was calculated to be 1000–1800 g" Exclusion criteria: "presence of disorders such as digestive obstruction, GI bleeding, gastroschisis, omphalocele, withdrawal syndrome, neonatal proven or clinical sepsis, congenital heart defect and asphyxia (degree II or III)"
Interventions	 Intervention: "5 ml of a mixture containing Lactobacillus reuteri DSM 17938. One drop of this product holds a minimum of 20 million live Lactobacillus reuteri protectis." The intervention was given two times daily. The intervention was given until the child achieved full enteral feedings Control: "placebo group received 0.5 ml of distilled water every 12 hours"
Outcomes	Mortality, sepsis, NEC
Notes	Data were taken from table 2



Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "using the random allocation software"
Allocation concealment (selection bias)	Unclear risk	No details were provided on where the allocation was placed
Blinding of participants and personnel (performance bias)	Low risk	Quote: "the intervention was implemented by nurses, and the physician was not aware of the condition of neonates in detail"
Blinding of outcome assessment (detection bias)	Low risk	Quote: "the intervention was implemented by nurses, and the physician was not aware of the condition of neonates in detail"
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the outcomes irrespective of their statistical significance
Other bias	Low risk	No other risk of bias was noted

Shashidhar et al. (2017)

Methods	A double blind randomized controlled trial in India	
Participants	Inclusion criteria: "All neonates with a birth weight between 750 and 1499g admitted to the NICU" Exclusion criteria: "Neonates with gastrointestinal anomalies, severe congenital malformation, and those not started on enteral feeds by day 14 of life were excluded"	
Interventions	 Intervention: "Multicomponent probiotic formulation of <i>Lactobacillus acidophilus</i>, <i>Lactobacillus rhamnosus</i>, <i>Bifidobacterium longum</i>, and <i>Saccharomyces boulardii</i> in the form of powdered sachets of 1 g each. The intervention was administered once a day at a dose of 1.25×109 CFU starting within 24 h of initiation of feeds. The probiotic supplementation was continued till discharge given once a day if the volume of feeds was 2 ml or more, and in two divided doses if the baby received <2 ml/feed Control: "The no probiotic group received only breast milk and served as the control" 	
Outcomes	Mortality, NEC	
Notes	Data were taken from table 2	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The subjects were randomly allocated into two groups using computer generated random numbers by an investigator not directly involved in the study"
Allocation concealment (selection bias)	Low risk	Quote: "Sequentially numbered opaque sealed envelopes were used for allocation concealment"
Blinding of participants and personnel (performance bias)	Low risk	Quote: "The two groups were coded as A and B and the group code was kept off site in an opaque sealed envelope and opened only after the final analysis was done"
Blinding of outcome assessment (detection bias)	Low risk	Quote: "The two groups were coded as A and B and the group code was kept off site in an opaque sealed envelope and opened only after the final analysis was done"
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the outcomes irrespective of their statistical significance
Other bias	Low risk	No other risk of bias was noted

Dongol Singh et al. (2017)



Methods	A randomized, double blind, placebo controlled conducted in Nepal
Participants	Inclusion criteria: Preterm babies admitted to the NICU Exclusion criteria: "Sick infants (neonates with clinical or proven sepsis), those with congenital malformation especially (central nervous system) malformation and other such as gastrointestinal obstruction, gastrointestinal bleeding, gastroschisis, omphalocoele, congenital heart defect and birth asphyxia (grade III). Out born babies were also excluded in this study"
Interventions	 Intervention: "probiotics Lactobacillus casei var. rhamnosis (LCR 35) 0.8 mg (half packet) dissolved in 2 ml of EBM in infant more than 1500 g and 0.4 mg probiotics (1/4th packet) dissolved in 1 ml of EBM in infants <1500 g was given twice a day until they reached full feeding" Control: "placebo as expressed breast milk only"
Outcomes	NEC and mortality
Notes	Data were taken from the last two paragraphs of the result section

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "using random selection by lottery"
Allocation concealment (selection bias)	High risk	Quote: "Intervention was instructed by the researcher and conducted by nursing staff of NICU"
Blinding of participants and personnel (performance bias)	Low risk	Less likely that neonates were aware of the intervention
Blinding of outcome assessment (detection bias)	Unclear risk	No clear information was provided on who made the assessment for clinical outcomes
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the outcomes irrespective of their statistical significance
Other bias	Low risk	No other risk of bias was noted

Sinha et al. (2015)

Methods	Randomized, double-blind, placebo-controlled trial conducted in India in community settings
Participants	Inclusion criteria: infants aged 3 days, born in the hospitals weighing 1500–2500 g" Exclusion criteria: "extremely premature infants (<34 weeks), sick infants, those with congenital malformations incompatible with life, and those with guardians not giving consent and belonging to out of study areas"
Interventions	 Intervention: "VSL#3 (a mix of eight strains: Streptococcus thermophilus, Bifidobacterium breve, Bifidobacterium longum, Bifidobacterium infantis, Lactobacillus acidophilus, Lactobacillus plantarum, Lactobacillus paracasei, and Lactobacillus delbrueckii spp bulgaricus, at a dose of 10 billion cfu for 30 days, starting on the third day of life and continued for 30 days" Control: "A similar-looking maltodextrin preparation in the same outer packing was administered to the control group"
Outcomes	Mortality, sepsis
Notes	Data were taken from table 2 and table 4. For sepsis, we included the numbers for suspected sepsi

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "A team of scientists at INCLEN Trust, New Delhi, used a computer- generated table for subject allocation"
Allocation concealment (selection bias)	Low risk	Quote: "Allocation concealment was ensured by sequentially numbering the sachet packets containing VSL#3 or placebo after block

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		randomisation. Identical packaging of VSL#3 and a placebo with similar consistency and colour was provided"
Blinding of participants and personnel (performance bias)	Low risk	Quote: "Parents of enrolled infants, investigators and field workers were masked to treatment allocation. Data analysis was performed in a blinded manner"
Blinding of outcome assessment (detection bias)	Low risk	Quote: "Parents of enrolled infants, investigators and field workers were masked to treatment allocation. Data analysis was performed in a blinded manner"
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the outcomes irrespective of their statistical significance
Other bias	Low risk	No other risk of bias was noted

Soofi et al. (2017)

Methods	A cluster randomized, placebo controlled trial conducted in Pakistan
Participants	Inclusion criteria: All infants born in the study village were eligible for inclusion. Exclusion criteria: Infants with congenital anomalies were excluded
Interventions	Intervention: The intervention group received a single dose of vitamin A 50,000 IU Comparison: the comparison group received placebo. Both the groups received vitamin E.
Outcomes	All-cause mortality, febrile illness, diarrhoea or pneumonia
Notes	

Bias	Authors'	Current for Judgement
Random sequence generation (selection bias)	judgement Low risk	Support for judgement Quote: "This was a cluster randomized, placebo-controlled trial"; and "an external consultant generated the computerized allocation sequence of clusters to each study intervention to either group using Epi Info 3.5.3 with restricted randomization based on population size, expected births and LHW presence"
Allocation concealment (selection bias)	Low risk	Quote: "An external consultant generated the computerized allocation sequence of clusters to each study intervention to either group using Epi Info 3.5.3 with restricted randomization based on population size, expected births and LHW presence"
Blinding of participants and personnel (performance bias)	Low risk	Quote: "The capsules were identical in appearance (Banner Pharmacaps, Canada) and supplied through the courtesy of the Micronutrient Initiative (Ottawa, Canada). The capsules were packaged in containers labelled as A & B. The content of the capsules were masked from field staff and supervisors, and the codes were only known to the external consultant responsible for cluster randomizations and the chair of the DSMB. The masking was maintained until the completion of the study"
Blinding of outcome assessment (detection bias)	Low risk	Quote: "The capsules were identical in appearance (Banner Pharmacaps, Canada) and supplied through the courtesy of the Micronutrient Initiative (Ottawa, Canada). The capsules were packaged in containers labelled as A & B. The content of the capsules were masked from field staff and supervisors, and the codes were only known to the external consultant responsible for cluster randomization and the chair of the DSMB. The masking was maintained until the completion of the study"
Incomplete outcome data (attrition bias)	Low risk	Quote: "We were able to follow 10,286 (93%) infants until death or 6 months of age"
Selective reporting (reporting bias)	Low risk	Author seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted





Methods	A prospective, randomized study conducted in China
Participants	Inclusion criteria: Admitted to the neonatal intensive care unit at a gestational age of 28 weeks, 96 h of age Exclusion criteria: Genetic metabolic diseases; congenital major abnormalities; congenital TORCH infections with overt signs at birth; terminal stage of illness (pH 7.0 or hypoxia with bradycardia 2 h); or the lack of parental consent
Interventions	Intervention: Vitamin A, 1500 IU/day and continued if the infant tolerated the milk, for 28 days or until discharge Comparison: Placebo
Outcomes	Mortality, NEC
Notes	

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "A blocked randomization method stratified by the neonatal intensive care unit size was used to assign infants to either the control or oral VA group"
Allocation concealment (selection bias)	Low risk	Quote: "solutions, which were only labelled with the study site and infant number"
Blinding of participants and personnel (performance bias)	Low risk	Quote: "The medical and nursing teams caring for the infants were thus completely unaware of the content of the solutions"
Blinding of outcome assessment (detection bias)	Low risk	Quote: "The medical and nursing teams caring for the infants were thus completely unaware of the content of the solutions, which were only labelled with the study site and infant number"
Incomplete outcome data (attrition bias)	Low risk	No loss to follow up was noted
Selective reporting (reporting bias)	Low risk	Authors seem to report all the relevant outcomes
Other bias	Low risk	No other risk of bias was noted

Tewari et al. (2015)

Methods	A double-blinded, placebo-controlled, randomized trial conducted in India
Participants	Inclusion criteria: "Preterm neonates <34 weeks admitted to the NICU" Exclusion criteria: "i. Extramural preterm neonates >10 day age with clinical or lab marker of sepsis ii. Preterm babies with necrotizing enterocolitis (NEC) or an intestinal surgical anomaly iii. Preterm babies with a lethal congenital anomaly, dysmorphism or aneuploidy"
Interventions	 Intervention: "Bacillus clausii containing 2×10⁹ spores in 5 ml minibottle in a dose of 2 ml per-oral every 8 h mixed with the enteral feeds through orogastric tube or oral feeds, giving them 2.4×10⁹ spores per day. Probiotic supplementation was continued till postnatal age of 6 weeks, or till discharge or death or occurrence of LOS, whichever was earlier for babies in both" Control: "Babies in the placebo group received sterile water, 2 ml per-oral every 8 h mixed with feeds"
Outcomes	Mortality, sepsis, NEC
Notes	Data were taken from table 3 and 4. We combined the numbers for very preterm and extreme preterm babies

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Randomization was done using an online service (www. randomization.com)"
Allocation concealment (selection bias)	Low risk	Quote: "Serially numbered opaque sealed envelopes with the allocation were available with the in-charge nurse of the NICU, who dispensed the intervention in a syringe for oral administration"

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Blinding of participants and personnel (performance bias)	Low risk	Quote: "All probiotic and sterile water mini bottles were coded and labels concealed"
Blinding of outcome assessment (detection bias)	Low risk	Quote: "All the investigators were blinded to the intervention"
Incomplete outcome data (attrition bias)	Low risk	Minimal loss to follow up
Selective reporting (reporting bias)	Low risk	Authors seem to report all the outcomes irrespective of their statistical significance
Other bias	Low risk	No other risk of bias was noted

West et al. (1995)

Methods	A randomized, double-masked trial conducted in Nepal
Participants	Inclusion criteria: Infants 5 months of age or younger Exclusion criteria: Infants >5 months of age Total number randomized in the intervention group: 5832 Total number randomized in the control group: 6086
Interventions	Intervention: - 15,000 RE (50,000 IU) vitamin A administered orally in approximately 3 drops of oil for neonates (<1 month old)
Outcomes	Primary outcome: Infant mortality, all-cause mortality Other outcomes: Malnutrition, ALRI, diarrhea or dysentery, whooping cough, meningitis, sudden death
Notes	All analyses were completed on an intent-to-treat basis

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "After a random start, wards were systematically assigned, blocked on VDAs, for infants to receive" Comment: Most likely done
Allocation concealment (selection bias)	Low risk	Quote: "from gelatinous capsules of identical appearance" Comment: Most likely done
Blinding of participants and personnel (performance bias)	Low risk	Quote: "The protocol and procedures for the trial were reviewed and approved by the Nepal Medical Research Council, Kathmandu, and the Joint Committee on Clinical Investigation at the Johns Hopkins University School of Medicine, Baltimore" Comment: Most likely done.
Blinding of outcome assessment (detection bias)	Unclear risk	Quote: "Verbal autopsy reports were independently reviewed by two physicians (SKK and RA) who standardized their reviews for 'S0 pre- study death reports" Comment: Most likely done
Incomplete outcome data (attrition bias)	Unclear risk	Total number of loss to follow up: n (%) The loss to follow up was not balanced
Selective reporting (reporting bias)	Low risk	Authors seem to report all the outcomes irrespective of their statistical significance
Other bias	Unclear risk	No other risk of bias was noted

Xu et al. (2016)



Methods	A prospective, randomized, case-controlled trial conducted in China
Participants	 Inclusion criteria: "hospital-born formula-fed infants with a gestational age of 30–37 weeks and a birth weight between 1500 and 2500 g" Exclusion criteria: "severe neonatal pathologies, such as severe birth complications, GI malformations, chromosomal abnormalities, known immunodeficiency, hydrops fetalis, central venous catheter, antifungal drugs, and probiotics"
Interventions	Intervention: "The intervention group received Saccharomyces boulardii CNCM I-745, administered two times per day as separate medication, not mixed with formula, at a dosage of 50 mg/kg. The study period ended at the 28th day after birth or when the infant was discharged from the hospital. Minimum duration of intervention was 7 days" Control: "Nothing was administered to the control group"
Outcomes	Neonatal sepsis
Notes	Data were taken from table 2

Risk of bias table

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Randomization was conducted according to a random computer- determined allocation order considering birth weight"
Allocation concealment (selection bias)	Low risk	 Quote: "Blinding was possible because the nursing staff who administered S. boulardii to the infants was not involved in the daily care and the attending neonatal team was unaware of the randomization assignments" Comment: Less likely that randomization sequence was revealed
Blinding of participants and personnel (performance bias)	Low risk	Quote: "Blinding was possible because the nursing staff who administered S. boulardii to the infants was not involved in the daily care and the attending neonatal team was unaware of the randomization assignments"
Blinding of outcome assessment (detection bias)	Low risk	Quote: "Blinding was possible because the nursing staff who administered S. boulardii to the infants was not involved in the daily care and the attending neonatal team was unaware of the randomization assignments"
Incomplete outcome data (attrition bias)	High risk	Quote: "25 (20%) patients were considered dropouts"
Selective reporting (reporting bias)	Low risk	Authors seem to report all the outcomes irrespective of their statistical significance
Other bias	Low risk	No other risk of bias was noted

Characteristics of excluded studies

Abdulkadir et al. (2016)	
Reason for exclusion	Study conducted in a high income country (UK)
Abrahamse-Berkeveld et al. (2016)	
Reason for exclusion	Study conducted in a high income country (Germany and Italy)
Abrahamsson et al. (2005)	
Reason for exclusion	No relevant outcomes were found
ADAPTS trial (2019)	
Reason for exclusion	Wrong settings: Ongoing study in Australia
Agarwal et al. (2003)	
Reason for exclusion	No relevant clinical outcomes were reported
Agarwal (2018)	



Reason for exclusion
Ahmadipour et al. (2019)
Reason for exclusion
Ahmadpour Kacho et al. (2005)
Reason for exclusion
Al-Hosni et al. (2012)
Reason for exclusion
Ala-Houhala et al.(1988)
Reason for exclusion
Allen et al. (2010)
Reason for exclusion
Armanian et al. (2014)
Reason for exclusion
Arthur et al. (1992)
Reason for exclusion
Aryayev et al. (2018)
Reason for exclusion
Athalye-Jape et al. (2018)
Reason for exclusion
Awad et al. (2010)
Reason for exclusion
Ayah et al. (2007)
Reason for exclusion
Aydin et al. (2012)
Reason for exclusion
Baglatzi et al. (2016)
Reason for exclusion
Bakker (2005)
Reason for exclusion
Bakker Zierikzee (2005)
Reason for exclusion
Bin-Nun et al. (2005)
Reason for exclusion
Bocquet et al. (2013)
Reason for exclusion
Bonati (1993a)
Reason for exclusion
Bonati (1996b)
Reason for exclusion
Bora & Deori (2019)
Reason for exclusion
Cekola et al. (2015)
Reason for exclusion
Chabra et al. (2013)
Reason for exclusion
Chandel et al. (2017)
Reason for exclusion
Chi (2019)
Reason for exclusion

Chouraqui et al. (2008)

Study conducted in a high income country (Australia)

Study conducted to treat neonatal Jaundice.

No relevant outcomes were reported

Study conducted in a high income country (USA)

Study conducted in a high income country (Finland)

Study conducted in high income country (UK)

Study participants were given only Prebiotics and no probiotics

Study population did not include Neonates

Study conducted in high income country

Study conducted in high income country

Study was retracted

Vitamin A was given at 14 weeks

Population included children with Congential Heart disease only

Study conducted in a high income country

Study continued supplementation of probiotics for 4 months

Study conducted in a high income country (Netherland)

Study conducted in Israel

Study conducted in a high income country (France)

Study conducted in a high income country (Italy)

Study conducted in a high income country (Italy)

Study compared two forms of the same intervention

Study conducted in a high income country (USA)

Study conducted in a high income country (USA)

No relevant clinical outcomes were available

No clinical outcomes were available

IMDAD ET AL. Reason for exclusion Chrzanowska-Liszewska (2011) Reason for exclusion Chua et al. (2017) Reason for exclusion Corkins & Kovacevich (2001) Reason for exclusion Costalos et al. (2003) Reason for exclusion Costeloe (2016) Reason for exclusion Coutsoudis et al. (1996) Reason for exclusion Dani et al. (2002) Reason for exclusion Darboe et al. (2007) Reason for exclusion Delimont et al. (2019) Reason for exclusion Delvin et al. (2000) Reason for exclusion Deng & Chen (2010) Reason for exclusion Denkel et al. (2017) Reason for exclusion Deshpande 2016 Reason for exclusion Diaby et al. (2018) Reason for exclusion Dilli et al. (2013) Reason for exclusion Elom et al. (2019) Reason for exclusion Escribano et al. (2018) Reason for exclusion Galderisi et al. (2016)

Reason for exclusion Garg et al. (2017) Reason for exclusion Garland et al. (2011)

Reason for exclusion Garofoli et al. (2018)

Reason for exclusion

Gomber (1996)

Reason for exclusion

Gomez-Rodriguez et al. (2019)

Reason for exclusion

Campbell__{W/II E}v Collaboration

Study conducted in a high income country (France)

Study conducted in a high income country (Poland)

Study conducted in a high income country (Netherland)

Study conducted UK

Study conducted in a high income country (Greece)

Study conducted in a high income country (UK)

No clinical outcomes were available

Study conducted in high income country (

Study conducted on wrong study population (infants)

Study conducted on older children and used Sorghum-Based and Corn-Based Fortified Blended Foods

Study conducted in a high income country (US)

No abstract or full text available and no relevant outcomes were available.

Study conducted in a high income country (Germany)

Study conducted in high income country

Observational study assessing the coverage of vitamin A supplementation

Study conducted on wrong patient population (infants with congenital heart diseases)

Wrong study population

Study conducted in a high income country (Spain)

The study investigated glucose monitoring and not the dextrose gel

Wrong study design (retrospective cohort study)

Study conducted in a high income country (Australia)

Wrong setting

Wrong study design



Gonchar et al. (2016)

Reason for exclusion

Guo-Qiang et al. (2016) Reason for exclusion Hammerman & Bin-nun (2007) Reason for exclusion Harris et al. (2016) Reason for exclusion Hays (2015) Reason for exclusion Hoy-Schulz et al. (2016) Reason for exclusion Hoyos (1999) Reason for exclusion

Hua et al. (2014)

Reason for exclusion

Huang et al. (2016) Reason for exclusion Hunter et al. (2012) Reason for exclusion Härtel (2019) Reason for exclusion Idindili et al. (2007) Reason for exclusion Indrio et al. (2008) Reason for exclusion IRCT (2015) Reason for exclusion Jacobs (2017) Reason for exclusion Janvier et al. (2014) Reason for exclusion Kahbazi et al. (2019) Reason for exclusion Kanic et al. (2015) Reason for exclusion Karthikeyan & Bhat (2017) Reason for exclusion Kiatchoosakun (2014) Reason for exclusion Kirkwood et al. (2010) Reason for exclusion Kliegman (2005) Reason for exclusion Koksal et al. (2015)

Reason for exclusion

Compared two different regimens of probiotics. No placebo group was included.

No relevant outcomes were available. Study only available in the form of abstract. Authors were contacted for full text but no response

Wrong study design

Wrong intervention

Study conducted in New Zealand

Study conducted in a high income country

Study conducted on wrong patient population

Wrong study design

Only abstract available and no relevant clinical outcomes were available

No relevant outcomes were available

Wrong study design

The study is being conducted in a high income country (Germany)

Study conducted on wrong patient population

Study conducted in a high income country

Wrong comparison

Study conducted in a high income country

Study conducted in a high income country

Wrong study population

Study conducted in a high income country

Study conducted in a high income country

No relevant outcomes were available

Wrong study design

Wrong study design

Only abstract was available and no analyzable data were reported

Kukkonen et al. (2008) Reason for exclusion Leele et al. (2015) Reason for exclusion Li (2019) Reason for exclusion Lin (2009) Reason for exclusion Long & Dempsey (2018) Reason for exclusion Lozano (2008) Reason for exclusion Lund et al. (2014)

Reason for exclusion

Lundelin et al. (2017) Reason for exclusion Mactier et al. (2012) Reason for exclusion Maldonado-Lobon (2015) Reason for exclusion Manzano et al. (2017) Reason for exclusion Manzoni et al. (2006) Reason for exclusion Manzoni et al. (2009) Reason for exclusion Marissen et al. (2019) Reason for exclusion Martins et al. (2009) Reason for exclusion Materna (2010) Reason for exclusion McCulloch et al. (2012) Reason for exclusion McKinlay (2016) Reason for exclusion Meyer & Gortner (2014) Reason for exclusion Mg (2011) Reason for exclusion Mihatsch et al. (2010) Reason for exclusion Millar (2017) Reason for exclusion Moles et al. (2015) Reason for exclusion Nadella et al. (2019)

Reason for exclusion



Collaboration
Wrong study design
Study conducted in a high income country (Singapore)
The intervention continued for 4 months
Study conducted in a high income country (Taiwan)
Study conducted in a developed country
Study conducted in a high income country
The control group did not receive the placebo but polio vaccine. It is difficult to tease out the effect of vitamin A supplementation vs. No vitamin A supplementation.
Study conducted in a high income country
Study conducted in a high income country
Study conducted in a high income country
Study conducted in a high income country (Spain)
Study conducted in a high income country (Italy)
Study conducted in a high income country (italy)
This is an ongoing study in Germany which is a high income country.
Wrong study population
Study conducted in a high income country
Study conducted in a high income country (UK)
Study conducted in a high income country
Study conducted in a high income country (Germany)
Study conducted in a high income country (Italy)
Study conducted in a high income country (Germany)
Study conducted in a high income country
Study conducted in high income country
Wrong intervention

-WILEY-C Campbell Collaboration

Collaboration	
Nct (2006)	
Reason for exclusion	Study conducted in high income country
Nct (2016)	
Reason for exclusion Papagaroufalis et al. (1991)	Wrong settings
Reason for exclusion	Study conducted in high income country (Greece)
Papagaroufalis et al. (2014)	
Reason for exclusion	Study conducted in high income country (Greece)
Patole et al. (2016)	
Reason for exclusion	Study conducted in a high income country (Australia)
Pearson et al. (1992)	
Reason for exclusion	Study conducted in a high income country (USA)
Plummer et al. (2018)	
Reason for exclusion	Study conducted in a high income country (Australia)
Puccio et al. (2007)	
Reason for exclusion	Study conducted in a high income country (Italy)
Qiao et al. (2017)	
Reason for exclusion	No relevant outcomes were available
Radke et al. (2017)	
Reason for exclusion	Study conducted in a high income country (Germany)
Raguž et al. (2016)	
Reason for exclusion	Wrong study design
Rakshasbhuvankar et al. (2017)	
Reason for exclusion	Study conducted in a high income country (Australia)
Rawat et al. (2016)	
Reason for exclusion	
	Study conducted in the USA
Repa et al. (2015)	
Repa et al. (2015) Reason for exclusion	Study conducted in a high income country (Austria)
Repa et al. (2015) Reason for exclusion Robbins & Fletcher (1993)	Study conducted in a high income country (Austria)
Repa et al. (2015) Reason for exclusion Robbins & Fletcher (1993) Reason for exclusion	
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Repa et al. (2015) Reason for exclusion Robbins & Fletcher (1993) Reason for exclusion Rodriguez-Herrera et al. (2019) Reason for exclusion	Study conducted in a high income country (Austria)
Repa et al. (2015)Reason for exclusionRobbins & Fletcher (1993)Reason for exclusionRodriguez-Herrera et al. (2019)	Study conducted in a high income country (Austria) wrong comparator Study conducted in a high income country Study compared two forms of Probiotics and no comparison with
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Repa et al. (2015)Reason for exclusionRobbins & Fletcher (1993)Reason for exclusionRodriguez-Herrera et al. (2019)Reason for exclusionRodríguez (2015)Reason for exclusion	Study conducted in a high income country (Austria) wrong comparator Study conducted in a high income country Study compared two forms of Probiotics and no comparison with
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Repa et al. (2015)Reason for exclusionRobbins & Fletcher (1993)Reason for exclusionRodriguez-Herrera et al. (2019)Reason for exclusionRodríguez (2015)Reason for exclusionRohan (2016)Reason for exclusionRoss et al. (1993)	Study conducted in a high income country (Austria) wrong comparator Study conducted in a high income country Study compared two forms of Probiotics and no comparison with placebo was available Study conducted in a high income country (UK)
Repa et al. (2015)Reason for exclusionRobbins & Fletcher (1993)Reason for exclusionRodriguez-Herrera et al. (2019)Reason for exclusionRodríguez (2015)Reason for exclusionRohan (2016)Reason for exclusionRoss et al. (1993)Reason for exclusion	Study conducted in a high income country (Austria) wrong comparator Study conducted in a high income country Study compared two forms of Probiotics and no comparison with placebo was available Study conducted in a high income country (UK)
Repa et al. (2015)Reason for exclusionRobbins & Fletcher (1993)Reason for exclusionRodriguez-Herrera et al. (2019)Reason for exclusionRodríguez (2015)Reason for exclusionRohan (2016)Reason for exclusionRoss et al. (1993)Reason for exclusionRouge et al. (2009)	Study conducted in a high income country (Austria) wrong comparator Study conducted in a high income country Study compared two forms of Probiotics and no comparison with placebo was available Study conducted in a high income country (UK) Study included children 6 months and older
Repa et al. (2015)Reason for exclusionRobbins & Fletcher (1993)Reason for exclusionRodriguez-Herrera et al. (2019)Reason for exclusionRodríguez (2015)Reason for exclusionRohan (2016)Reason for exclusionRoss et al. (1993)Reason for exclusionRouge et al. (2009)Reason for exclusion	Study conducted in a high income country (Austria) wrong comparator Study conducted in a high income country Study compared two forms of Probiotics and no comparison with placebo was available Study conducted in a high income country (UK) Study included children 6 months and older
Repa et al. (2015)Reason for exclusionRobbins & Fletcher (1993)Reason for exclusionRodriguez-Herrera et al. (2019)Reason for exclusionRodríguez (2015)Reason for exclusionRohan (2016)Reason for exclusionRoss et al. (1993)Reason for exclusionRouge et al. (2009)Reason for exclusionRubaltelli et al. (2000)	Study conducted in a high income country (Austria) wrong comparator Study conducted in a high income country Study compared two forms of Probiotics and no comparison with placebo was available Study conducted in a high income country (UK) Study included children 6 months and older Study conducted in a high income country (France)
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Reason for exclusion	Study conducted in a high income country (USA)
Storm et al. (2019)	
Reason for exclusion	Study conducted in a high income country USA
Stratiki et al. (2007)	
Reason for exclusion	Study conducted in a high income country (Greece)
Strus et al. (2018)	
Reason for exclusion	Study conducted in a high income country (Poland)
Ter (2017)	
Reason for exclusion	Study conducted in Australia
Thanhaeuser et al. (2014)	
Reason for exclusion	Study conducted in a high income country (Austria)
Totsu et al. (2014)	
Reason for exclusion	Study conducted in a high income country (Japan)
Tyson et al. (1999)	
Reason for exclusion	Study conducted in a high income country (USA)
Venkatarao et al. (1996)	
Reason for exclusion	Infant received vitamin A at 6 months
Vlieger et al. (2009)	
Reason for exclusion	Study conducted in high income country (Netherland)
Wardle et al. (2001)	
Reason for exclusion	Study conducted in high income country (UK)
West et al. (1991)	
Reason for exclusion	Study included children 6-59 months of age
Yang et al. (2011)	
Reason for exclusion	Full text not available and no abstract was available either so no relevant outcomes were available.

Characteristics of studies awaiting classification

Barclay et al. (2003)

Methods	Randomized controlled trial
Participants	Term infants
Interventions	Formula with pre-pro and synbiotics
Outcomes	Growth
Notes	Study available only in the abstract form

Chubarova and Sharyafetdinova (2017)

Methods	Randomized Controlled trial
Participants	Preterm neonates
Interventions	A combination of freeze-dried strains of lactobacilli and bifidobacteria: Lactobacillus rhamnosus, Bifidobacterium infantis, Bifidobacterium longum, Lactobacillus acidophilus (at time of manufacture 2 billion CFUs per 3 g of the preparation), also containing the additional component maltodextrin. The comparison group received placebo
Outcomes	NEC and stay in NICU and others
Notes	Only abstract available. It is not clear on where the study was done



Characteristics of ongoing studies

DelPiano (2016)

Study name	Our open-label, randomized controlled study has the primary endpoint of reducing diarrhea and infectious diseases (number of episodes/severity) and the secondary endpoint of decreasing infant mortality
Methods	Randomized control trial
Participants	The trial is currently conducted in Luzira, a suburb of Kampala, the capital of Uganda, and in Gulu and Lira, in the north of Uganda. The study is projected to enrol 4000 babies (control = 2000 and treatment=2000) who will be followed till 1 year of life. As controls, 2000 babies of the same community are planned to be considered
Interventions	The probiotic product selected for the trial is composed of 3 designated microorganisms, namely <i>Bifidobacterium breve</i> BR03 (DSM 16604), <i>B. breve</i> B632 (DSM 24706), and <i>Lactobacillus delbrueckii</i> subsp. delbrueckii LDD01 (DSM 22106)
Outcomes	Incidence of diarrhea and mortality
Starting date	
Contact information	Gastroenterology Department, Santa Rita Hospital-Policlinico di Monza †Department of Gynecology and Obstetrics, Santa Rita Hospital-Policlinico di Monza, Vercelli ‡Biolab Research Ltd, Novara, Italy
Notes	

Goodman (2015)

Participants

Study name	Prevention of vitamin A deficiency by supplementation alongside routine vaccinations: a randomised controlled trial in Ghana infants
Methods Participants	 Randomised controlled trial Participant inclusion criteria 1. Mothers normally resident in the study area 2. Informed consent obtained from the mother Participant exclusion criteria 1. Mothers unable to give informed consent 2. Mothers considered to be at high risk of adverse outcome in puerperal period 3. Multiple deliveries 4. Severe adverse reaction to vitamin A supplementation
Interventions	 1st Group: Mothers 200,000 IU vitamin A shortly after delivery Infants: 25,000 IU vitamin A with each Diphtheria, Pertussis, Tetanus (DPT) vaccine 1, 2 and 3 2nd Group: Mothers 200,000 IU vitamin A at infant's Bacillus Calmette-Guerin (BCG) vaccine and another 200,000 IU vitamin A at infant's 1st DPT Infants: 50,000 IU vitamin A with each DPT 1, 2 and 3
Outcomes	 Primary outcome measure Serum retinol levels, assessed by carrying out mRDR testing of infants at 6 weeks, 6 and 9 months Modified Retinol Dose Response (mRDR) tests Incidence of side effects such as bulging of the anterior fontanel and vomiting Incidence of severe morbidity Secondary outcome measures Breast milk retinol concentrations, assessed at 6 weeks, 6 and 9 months for an assessment of the impact of the different supplementation regimes mRDR testing of infants at 9 months of age
Starting date	01/01/2004
Contact information	World Health Organization Geneva-27 CH-1211 Switzerland
Notes	No results are available
Heydarabad (2018)	
Study name	Evaluation of the effect of probiotics on late-onset sepsis in very preterm newborns
Methods	A randomized, triple blinded, placebo controlled clinical trial with two parallel groups

Enrolled within 48 h of birth

Inclusion criteria include: 1. Preterm infants weighting 1000-1500 g and <32 weeks' gestational age at birth 2.

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	 Exclusion criteria: (a) Major congenital anomalies (Esophageal atresia,omphalocele, imperforate anus). (2) Major congenital heart malformations. (3) Genetic anomalies(e.g. Trisomy 21 or other trisomies). (4) Considered likely to die within 72 h of birth. (5) Death before minimal entral feeding (10-20cc/kg/day). (6) Parents from whom informed consent cannot be obtained. (7) Sepsis in admission (CRP > 10 mg/dl in 1th day of admission). (8) Asphyxia (grade II, III). (9) Maternal chorioamnionitis
Interventions	Very low-birth-weight preterm infants with a gestational age of <32 weeks and a weight of 1000-1500 gr who are admitted to the NICU of Shahid Motahari Hospital in Urmia during the first 48 h of their birth. Patients are randomly divided into two groups; the intervention group receive probiotic and the control group receive Dish water as placebo
Outcomes	Late onset sepsis
Starting date	2018-03-05
Contact information	Kamran Dehghan Iran (Islamic Republic of) dehghan.k@umsu.ac.ir
Notes	

Kaur (2018)

Study name	Effect of probiotic supplementation on feed tolerance and weight gain in low birth weight infants on tube feeds
Methods	Randomized, parallel group trial
Participants	Inclusion criteria: All neonates with a birth weight between 1000 and 1800 gm admitted to the NICU in whom enteral feeds can be started. Exclusion criteria: Neonates with weight <1000 gm Neonates with gastrointestinal anomalies Neonates with major congenital malformations Neonates in whom the feed could not be started by day 14 of life
Interventions	Intervention 1: Probiotic: Probiotic containing Lactibacillus acidophilus, Lactibacillus rhamnosus, Bifidobacterium longum, Saccharomyces boulardi in form of powdered sachet of 1g each Control Intervention 1: Nil: Nil
Outcomes	To compare the time taken (in days) to reach full enteral feeds (150 ml/kg/day) in low birth weight infants on orogastric feeds between the probiotic and no probiotic group To compare episodes of feed intolerance and weight gain in both the groups.Timepoint: Time taken to reach full enteral feeds
Starting date	31-07-2018
Contact information	Neonatal Intensive Care Unit, Department of pediatrics,GGS medical college and hospital faridkot Faridkot PUNJAB 151203 India dr.amarpreet12@gmail.com
Notes	

Londhe (2019)

Study name	Use of zinc and pre-probiotics as a therapeutic adjunct in neonatal sepsis in preterms- An Open label randomized controlled trial
Methods	Randomized, parallel group, multiple arm trial
Participants	 All intramural preterm neonates from 28 week 1day to 36 week 6 days admitted to NICU at GMCH Aurangabad with proven sepsis during the study period, whose parents consented to be part of the study were included Diagnostic criteria for sepsis was (a) Positive "sepsis screen," that is, presence of at least two of the following three parameters, namely, Total leucocyte count <5000/mm3, Low absolute neutrophil count (as per standard charts), C-reactive protein>1 mg/dl, (b) Radiological evidence of pneumonia (c) Culture positive sepsis (d) Meningitis

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Interventions	Zinc and prebiotics: zinc and pre-probiotic group was given both zinc 10 mg per day and pre-probiotics as syrup 5 ml per day containing <i>L. acidophilus</i> (1.25 billion), <i>B. longum</i> (0.125 billion), B. bifidum (0.125 billion), <i>B. lactis</i> (1 billion) and Inulin (25 mg) till discharge Zinc: Zinc group was given oral zinc 10 mg once a day irrespective of age of newborn till discharge. Pre-biotic group: Pre-probiotics group was given as Syrup 5 ml per day containing <i>L. acidophilus</i> (1.25 billion), <i>B. longum</i> (0.125 billion), <i>B. bifidum</i> (0.125 billion), <i>B. lactis</i> (1 billion) and Inulin (25 mg) till discharge Control: Not receiving any of the above
Outcomes	Reduction in mortality
Starting date	Date Completed: 31/08/2016
Contact information	Division of Neonatology Department of Pediatrics Govt Medical College Aurangabad 431001 Aurangabad, MAHARASHTRA India atul.londhe1982@gmail.com
Notes	Description results are available at the web site where the clinical trial was registered. We wrote to authors to obtain the results

Mirmohammdi (2018)

Study name	Determination of the effect of probiotics on prevention of necrotizing enterocolitis in preterm infants
Methods	This randomized clinical trial will be conducted on preterm infants with low birth weight and very low birth weight.
Participants	 Inclusion criteria: VLBW infants (gestational age ≤34 weeks and birth weight ≤1500 g) survive until the onset of oral nutrition Exclusion criteria: (1) Severe asphyxia (stage III) (2) Congenital major anomalies (3) Babies who have not started oral feeding for 3 weeks after birth. (4) Infants receiving antifungal treatment
Interventions	The intervention group consisted of newborn infants with probiotic podilakat made by Iran Fertilizer Company in a quantity of 1 drops per kg of weight every 12 h with breast milk or milk powder The control group is a neonate who will receive 0.5 cc normal saline every 12 h with breastfeeding or breastfeeding
Outcomes	Baby weight at the end of the 3rd month
Starting date	2019-07-29
Contact information	mir farhad mirmohammdi Ibn sina ave- imam reza hospital- mashhad- Iran Emailmirmohammadif951@mums.ac.ir
Notes	

Mukhtar, 2019

Study name	Role of prophylactic microbial supplements in prevention of blood stream infection and intestinal tract injury in premature neonates
Methods Participants	Randomized, parallel group trial All infants admitted to our NICU with birth weight <2 kg or gestational age <35 weeks as assessed by EDD and further confirmed by Ballards score who survive the first 24 hrs of life will be enrolled in the study
Interventions	Mixture of Lactobacillus acidophilus, Lactobacillus rhamnosus, Bifidobacterium longum, and Streptomyces boulardii in a dose of 1.25 billion CFU twice daily started with the first feed which can be in the form of expressed breast milk or preterm formula feed and continued till discharge The control group will receive expressed breast milk or preterm formula feed with no supplements added
Outcomes	 Necrotising colitis Nosocomial sepsis In preterm low birth weight babies
Starting date	01/12/2012
Contact information	Seniour resident, Department of Pediatrics and Neonatology, Sher-i-Kashmir Institute of Medical Sciences, Soura Srinagar JAMMU & KASHMIR 190005 India gousiamukhtar@gmail.com,
Notes	Study started in 2012 but no results have been published yet



Nandhini 2	012
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Study name	A clinical study to analyse the effectiveness of administration of harmless bacteria in preventing infection of intestine in preterm babies
Methods	Randomized, parallel group trial
Participants	Inclusion criteria: birth weight >1000 g gestational age-28-34 weeks enterally fed Exclusion criteria: surgical conditions of gastrointestinal tract severe birth asphyxia major congenital malformations chromosomal anomalies
Interventions	Intervention 1: probiotics: Bifidobacterium and Lactobacilli species- capsule(available as pre pro HS capsules marketed by fourrts limited) one capsule, administered twice a day orally mixed with breast milk for 7 days Control Intervention 1: none
Outcomes	Incidence and severity of NEC. Timepoint: during period of hospital stay
Starting date	15-02-2011
Contact information	Department of pediatrics JIPMER, Dhanvanthri Nagar 605006 Pondicherry, PONDICHERRY India drnbiswal@yahoo.com
Notes	The web site said that the study is completed. We wrote to authors to ask for the results
Study name	Effect of oral probiotic supplementation on the rate of hospital acquired infection and necrotizing enterocolitis in preterm very low birth weight infants
Methods	Randomized controlled study
Participants	 Participant inclusion criteria 1. VLBW preterm infants (gestational age (GA) < 35 weeks, body weight (BW) < 1.5 kg) 2. Admitted to the neonatal intensive care unit (NICU) who survived the first 3 days of life Participant exclusion criteria 1. Infants with chromosome abnormality 2. Infants with severe congenital defects 3. Infants with gastrointestinal anomalies (e.g., omphalocele, gastroschisis, intestinal obstruction) 4. Infants with unstable hemodynamic status
Interventions	Daily enteral probiotic supplementation of live Lactobacillus acidophilus and Bifidobacterium infantis at a dose of 2.5 ×108 CFU of each strain once a day for at least 28 days versus placebo
Outcomes	Primary outcome measure Nosocomial Infections Secondary outcome measures 1. Necrotizing enterocolitis (NEC) 2. Feeding tolerance 3. Time to reach full enteral feeding
Starting date	28/04/2011
Contact information	Rama IV Road Pathumwan Bangkok 10330 Thailand grad@chula.ac.th
Notes	
Rathod, 2019	
e. 1	

 Study name
 Probiotics for prevention of necrotising enterocoilitis in preterm neonates

 Methods
 Randomized, parallel group, placebo controlled trial

 Participants
 1. gestational age 28-34 weeks

 2. Birth weight of <2 kg</th>

-WILEY-C Campbell_ Collaboration

	Collaboration
	Exclusion criteria: Preterm newborn
	1. Birth weight >2 kg
	2. Lethal Congenital malformation
	3. Newborn on ventilator
Interventions	Intervention 1: giving probiotics: 50% of preterm newborn are given probiotics while rest 50% are not given
	probiotics Intervention 2: giving probiotics: 50% of preterm newborn are given probiotics while rest 50% are not given
	probiotics
Outcomes	Control Intervention 1: not applicable Occurrence of necrotising enterocolitis
Starting date	28/09/2017
Contact information	Mahendra Rathod
	Department of Pediatrics,sir Takhtsinhji hospital, Bhavnagar 364001
	Bhavnagar, GUJARAT
	India jayendragohil@gmail.com
Notes	It is not clear if the study is completed.
5	
Razavi, 2014	
Study name	Effect of probiotic in prevention of necroziting entrocolitis in preterm infants in Hafez hospital
Methods	Randomized, blinding: double blinded, placebo
Participants	Inclusion criteria: premature neonates with 1500 gr and below that are stable and tolerate (10cc/kg/day) formula or breast milk.Exclusion criteria: decline to participate; severe congenital anomaly; death
Interventions	Intervention group: add probiotics (Probiotic drops Pedilact Manufacturing zist takhmir) milk at the rate of 0/
	1cc/kg/day until hospital Control group: The control group was used as control and did not receive drug
Outcomes	Sign and symptom of NEC
Starting date	2014-05-22
Contact information	Dr.Seyed Mostajab Razavi Neonatal Research Center, Neonatal Department, Namazi Hospital, Zand Street, Shiraz Shiraz Iran (Islamic Republic of)
	porarish@sums.ac.ir
Notes	
Shashidhar, 2019	
Study name	A study of the effect of probiotic organism administration on feeding tolerance in very low birth weight newborn babies
Methods	Randomized, parallel group, active controlled trial
Participants	Inclusion criteria: All neonates (infants in the first 28 days of life) with a birth weight <1.5 kg, admitted to NICU
	SJMCH Bangalore. Postnatal age <2 wks and started enteral feeds. Exclusion criteria: Neonates GI tract anomalies, severe congenital malformations.
Interventions	Intervention 1: probiotic sachets: <i>Lactobacillus</i> Spp, <i>Bifidobacter</i> Spp, <i>Saccharomyces boulardi</i> oral 1 sachet once a day orally mixed in breast milk till discharge Control Intervention 1: breast milk: breast milk only
Outcomes	Incidence of feed intolerance Incidence of NEC stage 2 or more Duration of hospital stay Days on TPN Weight gain Mortality
Starting date	31-08-2012
Contact information	Senior resident Dept.of Neonatology St.Johns Medical College Koramangala Bangalore Bangalore KARNATAKA 560034 India shashiishere@gmail.com

Sinha, 2019



Study name	Phase III, multicentre, randomized, double-blind, placebo-controlled study to evaluate efficacy of probiotic supplementation for prevention of neonatal sepsis in 0–2 months old low birth weight infants in India
Methods	Randomized, parallel group, placebo controlled trial
Participants	 Inclusion criteria: Birth weight: 1500 g to 500 g Age of the new-born Day 3-7 on recruitment, that is, not later than 7th day Stable clinical condition as assessed by physician and accepting feeds orally (where stable is defined as, does not require intravenous fluids and vasopressor medication to maintain circulation and accepts oral feeding or breastfeeding). The mother (with the new-born) is planning to stay in study area for a period of at least 2 months Exclusion criteria: New born with extreme prematurity, that is, <34 weeks New born with illness requiring prolonged hospitalisation and interference with oral feeding Presence of a gross congenital malformation incompatible with life Parent or Legally authorized representative (LAR) not providing written consent
Interventions	 Intervention 1: Vivomixx Drops, that is, Lactic acid bacteria and Bifidobacteria drop with medium chain triglyderide oil. 1. Each bottle cap contains at least 50 Billion Lactic acid bacteria and Bifidobacteria a. Streptococcus thermophilus DSM 24731 b. Bifidobacterium longum DSM 24736, B. breve DSM 24732 and B. infantis DSM 24737) c. Lactobacillus acidophilus DSM 24735, L. plantarum DSM 24730, L. paracasei DSM 24733, and L. delbruecki subs. bulgaricus DSM 24734 2. Each bottle contains MCT, that is, Medium chain triglyceride oil 5 ml. The contents of the cap should be mixed with MCT oil and shaken well prior to administration. Store in refrigerator at 2-8°C: 1 ml per day for 30 days (corresponding to NLT 10 billion CFU per day) Control Intervention 1: Placebo Drops 1. Each Bottle contains MCT, that is, Medium chain triglyceride oil 5 ml The contents of the cap should be mixed with MCT oil and shaken well prior to administration. Store in refrigerator at 2-8°C: 1 ml per day for 30 days (corresponding to NLT 10 billion CFU per day) Control Intervention 1: Placebo Drops 1. Each Bottle contains MCT, that is, Medium chain triglyceride oil 5 ml The contents of the cap should be mixed with MCT oil and shaken well prior to administration. Store in refrigerator at 2-8 degrees centigrade.: 1 ml per day for 30 days
Outcomes	 Sepsis Possible serious bacterial infections (PSBIs). Timepoint: 60 days
Starting date	Date of first enrolment: 01-12-2019
Contact information	Division of Reproductive, Maternal and Child Health Ansari Nagar, New Delhi South DELHI 110029 India
Notes	Study started in Jan 2019

Additional tables

DATA AND ANALYSES

1. Vitamin A versus control

Outcome or subgroup	Studies	Participants	Statistical method	Effect estimate
11. All-cause neonatal mortality	6	126548	Risk ratio (IV, Random, 95% CI)	0.99 [0.90, 1.08]
12. All-cause infant mortality at 6 months	12	154940	Risk ratio (IV, Random, 95% CI)	0.98 [0.89, 1.07]
13. All-cause infant mortality at 12 months	8	118376	Risk ratio (IV, Random, 95% CI)	1.04 [0.94, 1.14]
14. Adverse events: bulging fontanelle	6	100256	Risk ratio (IV, Random, 95% CI)	1.53 [1.12, 2.09]
15. Adverse events: vomiting	5	99582	Risk ratio (IV, Random, 95% CI)	1.00 [0.93, 1.07]
16. All-cause neonatal mortality: sensitivity analysis: fixed effect model	5	126242	Risk ratio (IV, Fixed, 95% CI)	0.99 [0.90, 1.08]
model	5	120272		0.77 [0.70, 1.00]

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months: sensitivity analysis: fixed effect model	12	154940	Risk ratio (IV, Fixed, 95% CI)	0.97 [0.91, 1.03]
 All-cause infant mortality at 12 months: sensitivity analysis: fixed effect model 	8	118376	Risk ratio (IV, Fixed, 95% CI)	1.02 [0.96, 1.08]

2. Probiotics versus control

Outcome or subgroup	Studies	Participants	Statistical method	Effect estimate
21. All-cause mortality	26	10998	Risk ratio (IV, Random, 95% CI)	0.80 [0.66, 0.96]
22. All-cause mortality: subgroup analysis: settings	26	10998	Risk ratio (IV, Random, 95% CI)	0.80 [0.66, 0.96]
221. Hospital based	23	4691	Risk ratio (IV, Random, 95% CI)	0.78 [0.65, 0.94]
222. Community based	3	6307	Risk ratio (IV, Random, 95% CI)	1.25 [0.51, 3.05]
23. All-cause mortality: subgroup analysis: type of probiotics	26	10998	Risk ratio (IV, Random, 95% CI)	0.80 [0.66, 0.96]
231. Preparation contain a single strain of probiotics	9	2242	Risk ratio (IV, Random, 95% CI)	0.80 [0.61, 1.05]
232. Preparation contained multiple strains of probiotics	12	3050	Risk ratio (IV, Random, 95% CI)	0.80 [0.58, 1.09]
233. Preparation contained synbiotics (prebiotics + probiotics)	5	5706	Risk ratio (IV, Random, 95% CI)	0.69 [0.29, 1.61]
24. All-cause mortality: subgroup analysis: type of participants	26	10998	Risk ratio (IV, Random, 95% CI)	0.80 [0.66, 0.96]
241. Study include preterm/low birth weight babies	25	10587	Risk ratio (IV, Random, 95% CI)	0.79 [0.65, 0.95]
242. Study included term infants only	1	411	Risk ratio (IV, Random, 95% CI)	1.38 [0.31, 6.08]
25. All-cause mortality: subgroup analysis: type of feedings	26	10998	Risk ratio (IV, Random, 95% CI)	0.80 [0.66, 0.96]
251. Baby received breastmilk only	14	7721	Risk ratio (IV, Random, 95% CI)	0.81 [0.62, 1.05]
252. Baby received formula milk only	1	411	Risk ratio (IV, Random, 95% CI)	1.38 [0.31, 6.08]
253. Baby recieved both both breastmilk and formula milk	8	2385	Risk ratio (IV, Random, 95% CI)	0.69 [0.48, 0.99]
254. Type of feeding was unclear	3	481	Risk ratio (IV, Random, 95% CI)	1.33 [0.63, 2.81]
26. All-cause mortality: subgroup analysis: probiotics preparation	26	10998	Risk ratio (IV, Random, 95% CI)	0.80 [0.66, 0.96]
261. Preparation contained Lactobacillus	10	7002	Risk ratio (IV, Random, 95% CI)	0.82 [0.63, 1.05]
262. Preparation contained <i>Bifidobacterium</i>	1	400	Risk ratio (IV, Random, 95% CI)	0.43 [0.17, 1.09]
263. Preparation contained both Lactobacillus and Bifidobacterium	13	3110	Risk ratio (IV, Random, 95% CI)	0.71 [0.47, 1.08]
264. Preparation contained Saccharomyces boulardii only	2	486	Risk ratio (IV, Random, 95% CI)	1.12 [0.46, 2.71]
27. All-cause mortality: sensitivity analysis: fixed effect models	26	10998	Risk ratio (IV, Fixed, 95% CI)	0.80 [0.66, 0.96]
28. All-cause mortality: sensitivity analysis: risk of bias	26	10998	Risk ratio (IV, Random, 95% CI)	0.80 [0.66, 0.96]
281. High Risk of bias for randomisation/allocation concealment	3	285	Risk ratio (IV, Random, 95% CI)	0.27 [0.05, 1.40]
282. Low or Unclear Risk of bias for randomisation/allocation concealment	23	10713	Risk ratio (IV, Random, 95% CI)	0.82 [0.68, 0.99]

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29. Necrotizing enterocolitis (any type)	30	5574	Risk ratio (IV, Random, 95% CI)	0.46 [0.35, 0.59]
210. Necrotizing enterocolitis subgroup analysis: probi preparation		5574	Risk ratio (IV, Random, 95% CI)	0.46 [0.35, 0.59]
2101. Preparation contained <i>Lactobacillus</i>	13	2738	Risk ratio (IV, Random, 95% CI)	0.39 [0.25, 0.61]
2102. Preparation contained <i>Bifidobacterium</i>	1	400	Risk ratio (IV, Random, 95% CI)	0.20 [0.09, 0.47]
2103. Preparation contained <i>Lactobacillus</i> and <i>Bifidobacterium</i>	both 14	1950	Risk ratio (IV, Random, 95% CI)	0.49 [0.36, 0.68]
2104. Preparation contained Saccharomyces boulardii	only 2	486	Risk ratio (IV, Random, 95% CI)	0.94 [0.45, 1.95]
211. Necrotizing enterocolitis subgroup analysis: type	of			
feeding	30	5574	Risk ratio (IV, Random, 95% CI)	0.46 [0.35, 0.59]
2111. Baby received breastr		1945	Risk ratio (IV, Random, 95% CI)	0.43 [0.31, 0.59]
2112. Baby receieved formula milk only	1	93	Risk ratio (IV, Random, 95% CI)	0.21 [0.03, 1.76]
2113. Baby received both bre and formula milk	eastmilk 9	2445	Risk ratio (IV, Random, 95% CI)	0.55 [0.33, 0.92]
2114. Type of feeding was ur		1091	Risk ratio (IV, Random, 95% CI)	0.41 [0.17, 1.00]
212. Necrotizing enterocolitis		2072		0.12 [0.27, 2.00]
subgroup analysis: type probiotics		5574	Risk ratio (IV, Random, 95% CI)	0.46 [0.35, 0.59]
2121. Preparation contained strain of probiotics	a single 12	2679	Risk ratio (IV, Random, 95% CI)	0.48 [0.30, 0.76]
2122. Preparation contained strains of probiotics	multiple 15	2156	Risk ratio (IV, Random, 95% CI)	0.48 [0.35, 0.67]
2123. Preparation contained synbiotics				
(prebiotics + probiotics)		739	Risk ratio (IV, Random, 95% CI)	0.28 [0.12, 0.67]
213. Necrotizing enterocolitis sensitivity analysis: fixed models		5574	Risk ratio (IV, Fixed, 95% CI)	0.45 [0.37, 0.56]
214. Necrotizing enterocolitis sensitivity analysis: risk	:	5574	Risk ratio (IV, Random, 95% CI)	0.46 [0.35, 0.59]
2141. Low or unclear risk of randomisation/allocatic	bias for			
concealment	27	5289	Risk ratio (IV, Random, 95% CI)	0.46 [0.35, 0.61]
2142. High risk of bias for randomisation/allocatio				
concealment	3	285	Risk ratio (IV, Random, 95% CI)	0.33 [0.09, 1.13]
215. Neonatal sepsis216. Neonatal sepsis: subgroup	21 Ip	9105	Risk ratio (IV, Random, 95% CI)	0.78 [0.70, 0.86]
analysis: settings	21	9105	Risk ratio (IV, Random, 95% CI)	0.78 [0.70, 0.86]
2161. Hospital based	19	3209	Risk ratio (IV, Random, 95% CI)	0.83 [0.76, 0.91]
2162. Community based	2	5896	Risk ratio (IV, Random, 95% CI)	0.67 [0.49, 0.91]
217. Neonatal sepsis: type of probiotics	21	9105	Risk ratio (IV, Random, 95% CI)	0.78 [0.70, 0.86]
2171. Preparation contained strain of probiotics	8	1328	Risk ratio (IV, Random, 95% CI)	0.84 [0.74, 0.96]
2172. Preparation contained strains of probiotics	multiple 9	2482	Risk ratio (IV, Random, 95% CI)	0.81 [0.68, 0.97]
2173. Preparation contained synbiotics (prebiotics + probiotics)	4	5295	Risk ratio (IV, Random, 95% CI)	0.67 [0.54, 0.83]
	Ŧ	5275		0.07 [0.04, 0.00]

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218. Neonatal sepsis: subgroup analysis: type of feeding	21	9105	Risk ratio (IV, Random, 95% CI)	0.78 [0.70, 0.86]
2181. Baby received breastmilk only	8	6961	Risk ratio (IV, Random, 95% CI)	0.71 [0.61, 0.83]
2182. Baby received formula milk o nly	2	218	Risk ratio (IV, Random, 95% CI)	0.59 [0.22, 1.56]
2183. Baby received both formula and breastmilk only	7	1401	Risk ratio (IV, Random, 95% CI)	0.76 [0.64, 0.90]
2184. Type of feeding was unclear	4	525	Risk ratio (IV, Random, 95% CI)	0.95 [0.82, 1.09]
219. Neonatal sepsis: subgroup analysis: probiotic preparation	21	9105	Risk ratio (IV, Random, 95% CI)	0.78 [0.70, 0.86]
2191. Preparation contained <i>Lactobacillus</i>	11	7068	Risk ratio (IV, Random, 95% CI)	0.74 [0.62, 0.87]
2192. Preparation contained <i>Bifidobacterium</i>	1	400	Risk ratio (IV, Random, 95% CI)	0.81 [0.60, 1.09]
2193. Preparation contained both <i>Bifidobacterium</i> and <i>Lactobacillus</i>	6	1026	Risk ratio (IV, Random, 95% CI)	0.83 [0.68, 1.02]
2194. Preparation contained Saccharomyces boulardii only	3	611	Risk ratio (IV, Random, 95% CI)	0.73 [0.57, 0.94]
220. Neonatal Sepsis: Sensitivity analysis: Fixed Effect Model	21	9105	Risk ratio (IV, Fixed, 95% CI)	0.79 [0.73, 0.85]
221. Neonatal sepsis: sensitivity analysis: risk of bias	21	9105	Risk ratio (IV, Random, 95% CI)	0.78 [0.70, 0.86]
2211. Low or unclear risk of bias for randomization/allocation concealment	19	8892	Risk ratio (IV, Random, 95% CI)	0.79 [0.71, 0.87]
2212. High risk of bias due to randomization/allocation	2	212	Diele ratio (IV) Dandam (05% CI)	0.47 [0.25, 0.00]
concealment 222. Sepsis specifc mortality	2 2	213 4672	Risk ratio (IV, Random, 95% CI) Risk ratio (IV, Random, 95% CI)	0.47 [0.25, 0.90] 0.21 [0.04, 1.01]

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Feedback

REFERENCES TO STUDIES

INCLUDED STUDIES

Aage, S., Kiraly, N., Da Costa, K., Byberg, S., BjerregaardAndersen, M., Fisker, A. B., Aaby, P., & Benn, C. S. (2015). Neonatal vitamin A supplementation associated with increased atopy in girls. *Allergy*, *70*(8), 985–994.

- Agoestina, T., Humphrey, J. H., Taylor, G. A., Usman, A., Subardja, D., Hidayat, S., Nurachim, M., Wu, L., Friedman, D. S., & West, K. P., Jr (1994). Safety of one 52-mumol (50,000 IU) oral dose of vitamin A administered to neonates. *Bulletin of the World Health Organization*, 72(6), 859–868.
- Ahmad, S. M., Raqib, R., Huda, M. N., Alam, M. J., Monirujjaman, M., Akhter, T., Wagatsuma, Y., Qadri, F., Zerofsky, M. S., & Stephensen, C. B. (2019). High-dose neonatal vitamin A supplementation transiently decreases thymic function in early infancy. *Journal of Nutrition*, 150(1), 176–183.
- Ahmad, S. M., Raqib, R., Qadri, F., & Stephensen, C. B. (2014). The effect of newborn vitamin A supplementation on infant immune functions: trial design, interventions, and baseline data. *Contemporary Clinical Trials*, 39(2), 269–279.
- Ali, H., Hamadani, J., Mehra, S., Tofail, F., Hasan, M. I., Shaikh, S., Shamim, A. A., Wu, L. S., West, K. P., Jr, & Christian, P. (2017). Effect of maternal antenatal and newborn supplementation with vitamin A on cognitive development of school-aged children in rural Bangladesh: A follow-up of a placebo-controlled, randomized trial. American Journal of Clinical Nutrition, 106(1), 77–87.
- Amini, E., Dalili, H., Niknafs, N., Shariat, M., Nakhostin, M., & Jedari-Attari, S. (2017). The effect of probiotics in prevention of necrotising enterocolitis in preterm neonates in comparison with control group. *Iranian Journal of Pediatrics*, 27(6), 1–4.
- Bahl, R., Bhandari, N., Dube, B., Edmond, K., Fawzi, W., Fontaine, O., Kaur, J., Kirkwood, B. R., Martines, J., Masanja, H., Mazumder, S., Msham, S., Newton, S., Oleary, M., Ruben, J., Shannon, C., Smith, E., Taneja, S., & Yoshida, S., NEOVITA Study Author Group. (2013). Efficacy of early neonatal vitamin A supplementation in reducing

mortality during infancy in Ghana, India and Tanzania: Study protocol for a randomized controlled trial. *Trials*, 13, 22.

- Basu, S., Khanna, P., Srivastava, R., & Kumar, A. (2019). Oral vitamin A supplementation in very low birth weight neonates: A randomized controlled trial. *European Journal of Pediatrics*, 178(8), 1255–1265.
- Benn, C. S., Diness, B. R., Roth, A., Nante, E., Fisker, A. B., Lisse, I. M., Yazdanbakhsh, M., Whittle, H., Rodrigues, A., & Aaby, P. (2008). Effect of 50,000 IU vitamin A given with BCG vaccine on mortality in infants in Guinea-Bissau: Randomised placebo controlled trial. *BMJ*, 336(7658), 1416–1420.
- Benn, C. S., Fisker, A. B., Napirna, B. M., Roth, A., Diness, B. R., Lausch, K. R., Ravn, H., Yazdanbakhsh, M., Rodrigues, A., Whittle, H., & Aaby, P. (2010). Vitamin A supplementation and BCG vaccination at birth in low birthweight neonates: two by two factorial randomised controlled trial. *BMJ*, 340, c1101.
- Benn, C. S., Diness, B. R., Balde, I., Rodrigues, A., Lausch, K. R., Martins, C. L., Fisker, A. B., & Aaby, P. (2014). Two different doses of supplemental vitamin A did not affect mortality of normal-birth-weight neonates in Guinea-Bissau in a randomized controlled trial. *The Journal of Nutrition*, 144(9), 1474–1479.
- Biering-Sørensen, S., Fisker, A. B., Ravn, H., Camala, L., Monteiro, I., Aaby, P., & Benn, C. S. (2013). The effect of neonatal vitamin A supplementation on growth in the first year of life among low-birthweight infants in Guinea-Bissau: two by two factorial randomised controlled trial. *BMC Pediatrics*, 13, 87.
- Biering-Sorensen, S., Andersen, A., Ravn, H., Monterio, I., Aaby, P., & Benn, C. (2015). Early BCG vaccine to low-birth-weight infants and the effects on growth in the first year of life: A randomised controlled trial. *BMC Pediatrics*, 15, 137.
- Braga, T. D., da Silva, G. A., de Lira, P. I., & de Carvalho Lima, M. (2011). Efficacy of Bifidobacterium breve and Lactobacillus casei oral supplementation on necrotizing enterocolitis in very-low-birthweight preterm infants: A double-blind, randomized, controlled trial. American Journal of Clinical Nutrition, 93(1), 81–86.
- Chowdhury, T., Ali, M. M., Hossain, M. M., Singh, J., Yousuf, A. N., Yasmin, F., & Chowdhury, F. R. (2016). Efficacy of probiotics versus placebo in the prevention of necrotizing enterocolitis in preterm very low birth weight infants: A double-blind randomized controlled trial. *Journal of the College of Physicians and Surgeons*, 26(9), 770–774.
- Coles, C. L., Labrique, A., Saha, S. K., Ali, H., Al-Emran, H., Rashid, M., Christian, P., West, K. P., Jr, & Klemm, R. (2011). Newborn vitamin A supplementation does not affect nasopharyngeal carriage of *Streptococcus pneumoniae* in Bangladeshi infants at age 3 months. *Journal of Nutlrition*, 141(10), 1907–1911.
- Cooper, P., Bolton, K. D., Velaphi, S., de Groot, N., Emady-Azar, S., Pecquet, S., & Steenhout, P. (2017). Early benefits of a starter formula enriched in prebiotics and probiotics on the gut microbiota of healthy infants born to HIV + mothers: A randomized double-blind controlled trial. *Clinical Medicine Insights*, 8(10), 119–130.
- Cui, X., Shi, Y., Gao, S., Xue, X., & Fu, J. (2019). Effects of Lactobacillus reuteri DSM 17938 in preterm infants: A double-blinded randomized controlled study. *Italian Journal of Pediatrics*, 45(1), 140.
- Dashti, A. S., Afjeh, S. A., Basiry, A., Shirvani, F., Seifi, K., & Taheri, Z. M. (2014). Prophylactic probiotics for prevention of necrotizing enterocolitis (NEC) in low birth weight neonates. Archives of Pediatric Infectious Diseases, 2(1), 174–179.
- Demirel, G., Erdeve, O., Celik, I. H., & Dilmen, U. (2013). Saccharomyces boulardii for prevention of necrotizing enterocolitis in preterm infants: A randomized, controlled study. Acta Paediatrica, 102(12), e560–e565.
- Diness, B. R., Christoffersen, D., Pedersen, U. B., Rodrigues, A., Fischer, T. K., Andersen, A., Whittle, H., Yazdanbakhsh, M., Aaby, B., & Benn, C. S. (2010). The effect of high-dose vitamin A supplementation given with bacille CalmetteGuerin vaccine at birth on infant rotavirus infection and diarrhea: A randomized prospective study from GuineaBissau. *The Journal of infectious diseases*, 202, S243–S251.



- Diness, B. R., Fisker, A. B., Roth, A., Yazdanbakhsh, M., Sartono, E., Whittle, H., Nante, J. E., Lisse, I. M., Ravn, H., Rodrigues, A., Aaby, P., & Benn, C. S. (2007). Effect of high-dose vitamin A supplementation on the immune response to Bacille Calmette-Guerin vaccine. *The American Journal of Clinical Nutrition*, 86(4), 1152–1159.
- Diness, B. R., Martins, C. L., Bale, C., Garly, M. L., Ravn, H., Rodrigues, A., Whittle, H., Aaby, P., & Benn, C. S. (2011). The effect of high-dose vitamin A supplementation at birth on measles incidence during the first 12 months of life in boys and girls: An unplanned study within a randomised trial. *British Journal of Nutrition*, 105(12), 1819–1822.
- Dilli, D., Aydin, B., Fettah, N. D., Özyazıcı, E., Beken, S., Zenciroğlu, A., Okumuş, N., Özyurt, B. M., İpek, M. Ş., Akdağ, A., Turan, Ö., & Bozdağ, Ş. (2015). The propre-save study: Effects of probiotics and prebiotics alone or combined on necrotizing enterocolitis in very low birth weight infants. *Journal of Pediatrics*, 166(3), 545–551.
- Dutta, S., Ray, P., & Narang, A. (2015). Comparison of stool colonization in premature infants by three dose regimes of a probiotic combination: A randomized controlled trial. *American Journal of Perinatology*, 32(8), 733–740.
- Edmond, K. M., Newton, S., Shannon, C., O'Leary, M., Hurt, L., Thomas, G., Amenga-Etego, S., Tawiah-Agyemang, C., Gram, L., Hurt, C. N., Bahl, R., Owusu-Agyei, S., & Kirkwood, B. R. (2015). Effect of early neonatal vitamin A supplementation on mortality during infancy in Ghana (Neovita): A randomised, double-blind, placebo-controlled trial. *Lancet*, 385(9975), 1315–1323.
- Fernández-Carrocera, L. A., Solis-Herrera, A., Cabanillas-Ayón, M., Gallardo-Sarmiento, R. B., García-Pérez, C. S., Montaño-Rodríguez, R., & Echániz-Aviles, M. O. (2013). Double-blind, randomised clinical assay to evaluate the efficacy of probiotics in preterm newborns weighing less than 1500 g in the prevention of necrotising enterocolitis. Archives of Disease in Childhood. Fetal and Neonatal Edition, 98(1), F5-F9.
- Fisker, A. B., Aaby, P., Rodrigues, A., Frydenberg, M., Bibby, B. M., & Benn, C. S. (2011). Vitamin A supplementation at birth might prime the response to subsequent vitamin A supplements in girls. Three year follow-up of a randomized trial. *PLOS One*, *6*(8), e23265.
- Fisker, A. B., Benn, C. S., Diness, B. R., Martins, C., Rodrigues, A., Aaby, P., & Bibby, B. M. (2011). The effect of 50000IU vitamin A with BCG vaccine at birth on growth in the first year of life. *Journal of Tropical Medicine*, 2011, 1–9.
- Fisker, A. B., Lisse, I. M., Aaby, P., Erhardt, J. G., Rodrigues, A., Bibby, B. M., & Benn, C. S. (2007). Effect of vitamin A supplementation with BCG vaccine at birth on vitamin A status at 6 wk and 4 mo of age. *The American Journal of Clinical Nutrition*, 86(4), 1032–1039.
- Giridhar, S., Kumar Savita, J., Verma, A., Dutta, S., & Kumar, P. (2019). Intramuscular followed by oral vitamin A supplementation in neonates with birth weight from 750 to 1250g: A randomized controlled trial. *Indian Journal of Clinical Biochemistry*, 1–16.
- Giridhar, S. A clinical trial to study the effect of vitamin A supplementation in babies with birth weight from 750 grams to 1250 grams. http://www.ctri.nic.in/ Clinicaltrials/pmaindet2.php?trialid=294CTRI/2009/091/000010
- Guney-Varal, I., Koksal, N., Ozkan, H., Bagci, O., & Dogan, P. (2017). The effect of early administration of combined multi-strain and multispecies probiotics on gastrointestinal morbidities and mortality in preterm infants: A randomized controlled trial in a tertiary care unit. *Turkish Journal Of Pediatrics*, 59(1), 13–19.
- Hariharan, D., Balasubramanian, L., Kannappan, V., & Veluswami, G. (2016). Probiotic supplementation in VLBW preterm infants improves feeding tolerance and reduces risk of gram negative sepsis. *Journal of Pediatric Gastroenterology and Nutrition*, 62, 655.
- Hernández-Enríquez, N. P., Rosas-Sumano, A. B., Monzoy-Ventre, M. A., & Galicia-Flores, L. (2016). *Lactobacillus reuteri* DSM 17938 in preventing necrotizing enterocolitis in preterm newborns. Pilot study of efficacy and safety. *Revista Mexicana de Pediatria*, 83(2), 37–43.

WILEY-Campbell Collaboration

- Huaxian, Lin (2013). Analysis of the effect of probiotics combined with early minimal enteral feeding on prevention of neonatal necrotizing enterocolitis. *Journal of Pediatric Pharmacy*, 6, 18–20.
- Huda, M. N., Ahmad, S. M., Kalanetra, K. M., Taft, D. H., Alam, M. J., Khanam, A., Raqib, R., Underwood, M. A., Mills, D. A., & Stephensen, C.
 B. (2019). Neonatal vitamin A supplementation and vitamin A status are associated with gut microbiome composition in Bangladeshi infants in early infancy and at 2 years of age. *Journal of Nutrition*, 149(6), 1075–1088.
- Humphrey, J. H., Agoestina, T., Juliana, A., Septiana, S., Widjaja, H., Cerreto, M. C., Wu, L. S., Ichord, R. N., Katz, J., & West, K. P., Jr (1998). Neonatal vitamin A supplementation: Effect on development and growth at 3 y of age. *The American Journal of Clinical Nutrition*, 68(1), 109–117.
- Humphrey, J. H., Agoestina, T., Wu, L., Usman, A., Nurachim, M., Subardja, D., Hidayat, S., Tielsch, J., West, K. P., Jr, & Sommer, A. (1996). Impact of neonatal vitamin A supplementation on infant morbidity and mortality. *The Journal of Pediatrics*, 128(4), 489-496.
- Hussain, M., Jabeen, S., & Subhani, R. U. H. (2016). Role of probiotics in prevention of nectrotizing enterocolitis in preterm low birth weight neonates. *Pakistan Journal of Medical and Health Sciences*, 10(2), 455-459.
- Jorgensen, M. J., Fisker, A. B., Sartono, E., Andersen, A., Erikstrup, C., Lisse, I. M., Yazdanbakhsh, M., Aaby, P., & Benn, C. S. (2013). The effect of at-birth vitamin A supplementation on differential leucocyte counts and in vitro cytokine production: An immunological study nested within a randomised trial in Guinea-Bissau. *The British Journal* of Nutrition, 109(3), 467–477.
- Kaban, R. K., Wardhana, Hegar, B., Rohsiswatmo, R., Handryastuti, S., Amelia, N., Muktiarti, D., Indrio, F., & Vandenplas, Y. (2019). *Lactobacillus reuteri* DSM 17938 improves feeding intolerance in preterm infants. *Pediatric Gastroenterology, Hepatology & Nutrition*, 22(6), 545–553.
- Kiraly, N., Benn, C. S., Biering-Sorensen, S., Rodrigues, A., Jensen, K. J., Ravn, H., Allen, K. J., & Aaby, P. (2013). Vitamin A supplementation and BCG vaccination at birth may affect atopy in childhood: Longterm follow-up of a randomized controlled trial. *Allergy*, 68(9), 1168–1176.
- Klemm, R. D., Labrique, A. B., Christian, P., Rashid, M., Shamim, A. A., Katz, J., Sommer, A., & West, K. P., Jr. (2008). Newborn vitamin A supplementation reduced infant mortality in rural Bangladesh. *Pediatrics*, 122(1), 242–250.
- Nante, J. E., Diness, B. R., Ravn, H., Roth, A., Aaby, P., & Benn, C. S. (2008). No adverse events after simultaneous administration of 50 000 IU vitamin A and Bacille Calmette-Guerin vaccination to normal-birthweight newborns in GuineaBissau. *European Journal of Clinical Nutrition*, 62(7), 842–848.

PUBLISHED AND UNPUBLISHED DATA

- Akar, M., Eras, Z., Oncel, M. Y., Arayici, S., Guzoglu, N., Canpolat, F. E., Uras, N., & Oguz, S. S. (2017). Impact of oral probiotics on neurodevelopmental outcomes in preterm infants. *The Journal of Maternal-Fetal & Neonatal Medicine*, 30(4), 411–415.
- Bahl, R., Bhandari, N., Dube, B., Edmond, K., Fawzi, W., Fontaine, O., Kaur, J., Kirkwood, B. R., Martines, J., Masanja, H., Mazumder, S., Msham, S., Newton, S., Oleary, M., Ruben, J., Shannon, C., Smith, E., Taneja, S., & Yoshida, S. (2012). Efficacy of early neonatal vitamin A supplementation in reducing mortality during infancy in Ghana, India and Tanzania: Study protocol for a randomized controlled trial. *Trials*, 13(22), 22.
- Bahl, R., Bhandari, N., Dube, B., Edmond, K., Fawzi, W., Fontaine, O., Kaur, J., Kirkwood, B. R., Martines, J., Masanja, H., Mazumder, S., Msham, S., Newton, S., Oleary, M., Ruben, J., Shannon, C., Smith, E., Taneja, S., & Yoshida, S. (2012). Efficacy of early neonatal vitamin A supplementation in

reducing mortality during infancy in Ghana, India and Tanzania: Study protocol for a randomized controlled trial. *Trials*, 13(22), 22.

- Coles, C. L., Rahmathullah, L., Kanungo, R., Thulasiraj, R. D., Katz, J., Santhosham, M., & Tielsch, J. M. (2001). Vitamin A supplementation at birth delays pneumococcal colonization in South Indian infants. *Journal of Nutrition*, 131(2), 255–261.
- Dongol Singh, S. S., Klobassa, D. S., Resch, B., Urlesberger, B., & Shrestha, R. P. (2017). Placebo controlled introduction of prophylactic supplementation of probiotics to decrease the incidence of necrotizing enterocolitis at Dhulikhel Hospital in Nepal. *Kathmandu* University Medical Journal, 15(60), 319–323.
- Humphrey, J. H., Iliff, P. J., Marinda, E. T., Mutasa, K., Moulton, L. H., Chidawanyika, H., Ward, B. J., Nathoo, K. J., Malaba, L. C., Zijenah, L. S., Zvandasara, P., Ntozini, R., Mzengeza, F., Mahomva, A. I., Ruff, A. J., Mbizvo, M. T., & Zunguza, C. D. (2006). Effects of a single large dose of vitamin A, given during the postpartum period to HIVpositive women and their infants, on child HIV infection, HIV-free survival, and mortality. *The Journal of infectious diseases*, 193(6), 860–871.
- Iliff, P. J., Humphrey, J. H., Mahomva, A. I., Zvandasara, P., Bonduelle, M., Malaba, L., & Nathoo, K. J. (1999). Tolerance of large doses of vitamin A given to mothers and their babies shortly after delivery. *Nutrition Research*, 19(10), 1437–1446.
- Katz, J., West, K. P., Khatry, S. K., Thapa, M. D., LeClerq, S. C., Pradhan, E. K., Pokhrel, R. P., & Sommer, A. (1995). Impact of vitamin A supplementation on prevalence and incidence of xerophthalmia in Nepal. *Investigative Ophthalmology and Visual Science*, 36(13), 2577–2583.
- Malaba, L. C., Iliff, P. J., Nathoo, K. J., Marinda, E., Moulton, L. H., Zijenah, L. S., Zvandasara, P., Ward, B. J., ZVITAMBO Study Group, & Humphrey, J. H. (2005). Effect of postpartum maternal or neonatal vitamin A supplementation on infant mortality among infants born to HIV-negative mothers in Zimbabwe. *The American Journal of Clinical Nutrition*, 81(2), 454–460.
- Masanja, H., Smith, E. R., Muhihi, A., Briegleb, C., Mshamu, S., Ruben, J., Noor, R. A., Khudyakov, P., Yoshida, S., Martines, J., Bahl, R., & Fawzi, W. W. (2015). Effect of neonatal vitamin A supplementation on mortality in infants in Tanzania: A randomised, double-blind, placebocontrolled trial. *Lancet*, 385, 1324–1332.
- Mazumder, S., Taneja, S., Bhatia, K., Yoshida, S., Kaur, J., Dube, B., Toteja, G. S., Bahl, R., Fontaine, O., Martines, J., & Bhandari, N. Neovita India Study Group (2015). Efficacy of early neonatal supplementation with vitamin A to reduce mortality in infancy in Haryana, India (Neovita): A randomised, double-blind, placebo-controlled trial. *Lancet*, 385(9975), 1333–1342.
- Miller, M. F., Stoltzfus, R. J., Iliff, P. J., Malaba, L. C., Mbuya, N. V., & Humphrey, J. H. (2006). Zimbabwe vitamin A for mothers and babies project (ZVITAMBO) study group. Effect of maternal and neonatal vitamin A supplementation and other postnatal factors on anemia in Zimbabwean infants: a prospective, randomized study. *The American Journal of Clinical Nutrition*, 84(1), 212–222.
- Nandhini, L. P., Biswal, N., Adhisivam, B., Mandal, J., Bhat, B. V., & Mathai, B. (2016). Synbiotics for decreasing incidence of necrotizing enterocolitis among preterm neonates - a randomized controlled trial. *The Journal of Maternal-Fetal & Neonatal Medicine*, 29(5), 821–825.
- Niekerk, E., Nel, D. G., Blaauw, R., & Kirsten, G. F. (2015). Probiotics reduce necrotizing enterocolitis severity in HIV-exposed premature infants. *Journal of Tropical Pediatrics*, 61(3), 155–164. (HIV exposed).
- Niekerk, E., Nel, D. G., Blaauw, R., & Kirsten, G. F. (2015). Probiotics Reduce Necrotizing Enterocolitis Severity in HIV-exposed Premature Infants. *Journal of Tropical Pediatrics*, 61(3), 155–164. (HIV nonexposed).
- Oncel, M. Y., Sari, F. N., Arayici, S., Guzoglu, N., Erdeve, O., Uras, N., Oguz, S. S., & Dilmen, U. (2014). *Lactobacillus reuteri* for the prevention of necrotising enterocolitis in very low birthweight infants: A randomised controlled trial. *Archives of Disease in Childhood. Fetal and Neonatal Edition*, 99(2), F110–F115.

- Panigrahi, P., Parida, S., Nanda, N. C., Satpathy, R., Pradhan, L., Chandel, D. S., Baccaglini, L., Mohapatra, A., Mohapatra, S. S., Misra, P. R., Chaudhry, R., Chen, H. H., Johnson, J. A., Morris, J. G., Paneth, N., & Gewolb, I. H. (2017). A randomized synbiotic trial to prevent sepsis among infants in rural India. *Nature*, 548(7668), 407–412.
- Rahmathullah, L., Tielsch, J. M., Thulasiraj, R. D., Katz, J., Coles, C., Devi, S., John, R., Prakash, K., Sadanand, A. V., Edwin, N., & Kamaraj, C. (2003). Impact of supplementing newborn infants with vitamin A on early infant mortality: community based randomised trial in southern India. *BMJ*, 327(7409), 254.
- Rehman, S., Iqbal, A., & Ali, W. (2018). Role of probiotics in reducing frequency of necrotizing enterocolitis in preterm neonates. *Pakistan Paediatric Journal*, 42(3), 172–177.
- Rojas, M. A., Lozano, J. M., Rojas, M. X., Rodriguez, V. A., Rondon, M. A., Bastidas, J. A., Perez, L. A., Rojas, C., Ovalle, O., Garcia-Harker, J. E., Tamayo, M. E., Ruiz, G. C., Ballesteros, A., Archila, M. M., & Arevalo, M. (2012). Prophylactic probiotics to prevent death and nosocomial infection in preterm infants. *Pediatrics*, 130(5), e1113–e1120.
- Rojas, M., & Lozano, J. Prophylactic probiotics to prevent death or nosocomial infection in preterm infants in Colombia. Pediatric Academic Societies Annual Meeting 2012.
- Roy, A., Chaudhuri, J., Sarkar, D., Ghosh, P., & Chakraborty, S. (2014). Role of enteric supplementation of probiotics on late-onset sepsis by Candida species in preterm low birth weight neonates: A randomized, double blind, placebo-controlled trial. *North American Journal of Medical Sciences*, 6(1), 50–57.
- Saengtawesin, V., Tangpolkaiwalsak, R., & Kanjanapattankul, W. (2014). Effect of oral probiotics supplementation in the prevention of necrotizing enterocolitis among very low birth weight preterm infants. *Journal of the Medical Association of Thailand*, 97(6), S20–S25.
- Samanta, M., Sarkar, M., Ghosh, P., Ghosh, Jk, Sinha, Mk, & Chatterjee, S. (2009). Prophylactic probiotics for prevention of necrotizing enterocolitis in very low birth weight newborns. *Journal of Tropical Pediatrics*, 55(2), 128–131.
- Sari, F. N., Dizdar, E. A., Oguz, S., Erdeve, O., Uras, N., & Dilmen, U. (2010). Oral probiotics: *Lactobacillus sporogenes* in prevention of necrotizing enterocolitis in very low birth weight infants: A randomized, controlled trial. *Early Human Development*, *86*, S872010.
- Sari, F. N., Dizdar, E. A., Oguz, S., Erdeve, O., Uras, N., & Dilmen, U. (2011). Oral probiotics: *Lactobacillus sporogenes* for prevention of necrotizing enterocolitis in very low-birth weight infants: a randomized, controlled trial. *European Journal of Clinical Nutrition*, 65(4), 434–439.
- Sari, F. N., Eras, Z., Dizdar, E. A., Erdeve, O., Oguz, S. S., Uras, N., & Dilmen, U. (2012). Do oral probiotics affect growth and neurodevelopmental outcomes in very low-birth-weight preterm infants? *American Journal of Perinatology*, 29(8), 579–586.
- Serce, O., Benzer, D., Gursoy, T., Karatekin, G., & Ovali, F. (2013). Efficacy of Saccharomyces boulardii on necrotizing enterocolitis or sepsis in very low birth weight infants: a randomised controlled trial. Early Human Development, 89(12), 1033–1036.
- Serce, O., Gursoy, T., Karatekin, G., & Ovali, F. (2013). Effects of prebiotic and probiotic combination on necrotizing enterocolitis and sepsis prophylaxis in very low birth weight infants. *Journal of Perinatal Medicine*, 41, 219–289.
- Shadkam, M. N., Jalalizadeh, F., & Nasiriani, K. (2015). Effects of probiotic Lactobacillus reuteri (DSM 17938) on the incidence of necrotizing enterocolitis in very low birth weight premature infants. Iranian Journal of Neonatology, 6(4), 15–20.
- Shashidhar, A., Suman Rao, P. N., Nesargi, S., Bhat, S., & Chandrakala, B. S. (2017). Probiotics for promoting feed tolerance in very low birth weight neonates—A randomized controlled trial. *Indian Pediatrics*, 54(5), 363–367.
- Sinha, A., Gupta, S. S., Chellani, H., Maliye, C., Kumari, V., Arya, S., Garg, B. S., Gaur, S. D., Gaind, R., Deotale, V., Taywade, M., Prasad, M. S., Thavraj, V., Mukherjee, A., & Roy, M. (2015). Role of probiotics VSL#3

Collaboration

in prevention of suspected sepsis in low birthweight infants in India: A randomised controlled trial. *BMJ Open*, *5*(7), e006564.

- Soofi, S., Ariff, S., Sadiq, K., Habib, A., Bhatti, Z., Ahmad, I., Hussain, M., Ali, N., Cousens, S., & Bhutta, Z. A. (2017). Evaluation of the uptake and impact of neonatal vitamin A supplementation delivered through the Lady Health Worker programme on neonatal and infant morbidity and mortality in rural Pakistan: an effectiveness trial. Archives of Disease in Childhood, 102(3), 216–223.
- Smith, E. R., Muhihi, A., Mshamu, S., Sudfeld, C. R., Noor, R. A., Spiegelman, D., Shapiro, R. L., Masanja, H., & Fawzi, W. (2016). The effect of neonatal vitamin A supplementation on morbidity and mortality at 12 months: A randomized trial. *International Journal of Epidemiology*, 45(6), 2112–2121.
- Sun, H., Cheng, R., & Wang, Z. (2019). Early vitamin A supplementation improves the outcome of retinopathy of prematurity in extremely preterm infants. *Retina*, 40, 1176–1184.
- Tewari, V. V., Dubey, S. K., & Gupta, G. (2015). Bacillus clausii for prevention of late-onset sepsis in preterm infants: A randomized controlled trial. Journal of Tropical Pediatrics, 61(5), 377–385.
- Tielsch, J. M., Rahmathullah, L., Thulasiraj, R. D., Katz, J., Coles, C., Sheeladevi, S., John, R., & Prakash, K. (2007). Newborn vitamin A dosing reduces the case fatality but not incidence of common childhood morbidities in South India. *Journal of Nutrition*, 137(11), 2470–2474.
- West, K. P., Jr, Katz, J., Shrestha, S. R., LeClerq, S. C., Khatry, S. K., Pradhan, E. K., Adhikari, R., Wu, L. S., Pokhrel, R. P., & Sommer, A. (1995). Mortality of infants <6 mo of age supplemented with vitamin A: A randomized, double-masked trial in Nepal. *American Journal of Clinical Nutrition*, 62(1), 143–148.
- West, K. P., Jr, Khatry, S. K., LeClerq, S. C., Adhikari, R., See, L., Katz, J., Shrestha, S. R., Pradhan, E. K., Pokhrel, R. P., & Sommer, A. (1992). Tolerance of young infants to a single, large dose of vitamin A: A randomized community trial in Nepal. Bulletin of the World Health Organization, 70(6), 733–739.
- Xu, L., Wang, Y., Wang, Y., Fu, J., Sun, M., Mao, Z., & Vandenplas, Y. (2016). A double-blinded randomized trial on growth and feeding tolerance with *Saccharomyces boulardii* CNCM I-745 in formula-fed preterm infants. *Jornal De Pediatria*, 92(3), 296–301.

EXCLUDED STUDIES

- Abdulkadir, B., Nelson, A., Skeath, T., Marrs, E. C., Perry, J. D., Cummings, S. P., Embleton, N. D., Berrington, J. E., & Stewart, C. J. (2016). Routine use of probiotics in preterm infants: Longitudinal impact on the microbiome and metabolome. *Neonatology*, 109(4), 293-247.
- Abrahamse-Berkeveld, M., Alles, M., Franke-Beckmann, E., Helm, K., Knecht, R., Köllges, R., Sandner, B., Knol, J., Ben Amor, K., & Bufe, A. (2016). Infant formula containing galacto-and fructo-oligosaccharides and *Bifidobacterium breve* M-16V supports adequate growth and tolerance in healthy infants in a randomised, controlled, double-blind, prospective, multicentre study. *Journal of Nutritional Science*, 28(5), e42.
- Abrahamsson, T., Jakobsson, T., Sinkiewicz, G., Fredriksson, M., & Bjorksten, B. (2005). Intestinal microbiota in infants supplemented with the probiotic bacterium *Lactobacillus reuteri*. *Journal of Pediatric Gastroenterology and Nutrition*, 40(5), 692.
- ADAPTS trial. (2019). A randomised controlled trial: Effect of probiotics on gut microbiome and vaccine responses in newborns with antibiotic-induced dysbiosis (ADAPTS: Antibiotic Dysbiosis and Probiotics Trial in infantS). ACTRN12619000369123p.
- Agarwal, R., Sharma, N., Chaudhry, R., Deorari, A., Paul, V. K., Gewolb, I. H., & Panigrahi, P. (2003). Effects of oral *Lactobacillus* GG on enteric

microflora in low-birth-weight neonates. Journal of Pediatrics Gastroenterology and Nutrition, 36(3), 397–402.

- Agrawal, S., Rao, S., Nathan, E. A., & Patole, S. (2018). Effect of probiotics on C-reactive protein levels in preterm infants: Secondary analysis of a randomized controlled trial. *Journal of Neonatal-Perinatal Medicine*, 11(2), 165–171.
- Ahmadipour, S., Baharvand, P., Rahmani, P., Hasanvand, A., & Mohsenzadeh, A. (2019). Effect of synbiotic on the treatment of jaundice in full term neonates: A randomized clinical trial. *Pediatric Gastroenterology*, *Hepatology* & *Nutrition*, 22(5), 453–459.
- Ahmadpour Kacho, M., Zahedpasha, Y., Ahmadzadeh Amiri, A., Hajiahmadi, M., & Firoozi, M. (2005). Effect of vitamin A on prevention of retinopathy of prematurity (ROP). *Pediatric Research*, 58, 355.
- Ala-Houhala, M., Koskinen, T., Mäki, R., & Rinkari, S. (1988). Serum vitamin A levels in mothers and their breast-fed term infants with or without supplemental vitamin A. Acta Paediatrica Scandinavica, 77(2), 198–201.
- Al-Hosni, M., Duenas, M., Hawk, M., Stewart, L. A., Borghese, R. A., Cahoon, M., Atwood, L., Howard, D., Ferrelli, K., & Soll, R. (2012). Probiotics-supplemented feeding in extremely low-birth-weight infants. *Journal of Perinatology*, 32(4), 253–259.
- Allen, S. J., Jordan, S., Storey, M., Thornton, C. A., Gravenor, M., Garaiova, I., Plummer, S. F., Wang, D., & Morgan, G. (2010). Dietary supplementation with *Lactobacilli* and *Bifidobacteria* is well tolerated and not associated with adverse events during late pregnancy and early infancy. *Journal of Nutrition*, 140(3), 483–488.
- Armanian, A. M., Sadeghnia, A., Hoseinzadeh, M., Mirlohi, M., Feizi, A., Salehimehr, N., Saee, N., & Nazari, J. (2014). The effect of neutral oligosaccharides on reducing the incidence of necrotizing enterocolitis in preterm infants: A randomized clinical trial. *International Journal of Preventive Medicine*, 5(11), 1387–1395.
- Arthur, P., Kirkwood, B., Ross, D., Morris, S., Gyapong, J., Tomkins, A., & Addy, H. (1992). Impact of vitamin A supplementation on childhood morbidity in northern Ghana. *Lancet*, 8(339), 361–362.
- Aryayev, M. L., Senkivska, L. I., Bredeleva, N. K., & Talashova, I. V. (2018). Prophylaxis of acute respiratory infections via improving the immune system in late preterm newborns with E. coli strain Nissle 1917: A controlled pilot trial. *Pilot and Feasibility Studies*, 4, 79.
- Athalye-Jape, G., Minaee, N., Nathan, E., Simmer, K., & Patole, S. (2018). Outcomes in preterm small versus appropriate for gestation infants after *Bifidobacterium breve* M-16 V supplementation. *The Journal of Maternal-Fetal & Neonatal Medicine*, 33, 1–7.
- Awad, H., Mokhtar, H., Imam, S. S., Gad, G. I., Hafez, H., & Aboushady, N. (2010). Comparison between killed and living probiotic usage versus placebo for the prevention of necrotizing enterocolitis and sepsis in neonates. *Pakistan Journal of Biological Sciences*, 13(6), 253–262.
- Ayah, R. A., Mwaniki, D. L., Magnussen, P., Tedstone, A. E., Marshall, T., Alusala, D., Luoba, A., Kaestel, P., Michaelsen, K. F., & Friis, H. (2007). The effects of maternal and infant vitamin A supplementation on vitamin A status: A randomised trial in Kenya. *British Journal of Nutrition*, 98(2), 422-430.
- Aydin, B., Dilli, D., Erol, S., Ozyazici, E., Beken, S., Cullas Ilarslan, N. E., Zenciroglu, A., & Okumus, N. (2012). The effects of synbiotics on morbidity and mortality in newborns with cyanotic congenital heart disease: A prospective randomized controlled trial. Archives of Disease in Childhood, 97, A462.
- Baglatzi, L., Gavrili, S., Stamouli, K., Zachaki, S., Favre, L., Pecquet, S., Benyacoub, J., & Costalos, C. (2016). Effect of infant formula containing a low dose of the probiotic *Bifidobacterium lactis* CNCM I-3446 on immune and gut functions in C-section delivered babies: A pilot study. *Clinical Medicine Insights. Pediatrics*, 13(10), 9–11.
- Bakker-Zierikzee, A. M., Alles, M. S., Knol, J., Kok, F. J., Tolboom, J. J., & Bindels, J. G. (2005). Effects of infant formula containing a mixture of galacto- and fructo-oligosaccharides or viable *Bifidobacterium animalis* on the intestinal microflora during the first 4 months of life. *British Journal of Nutrition*, 94(5), 902–905.

- Bakker-Zierikzee, A. M., Alles, M. S., Knol, J., Kok, F. J., Tolboom, J. J., & Bindels, J. G. (2005). Effects of infant formula containing a mixture of galacto- and fructo-oligosaccharides or viable *Bifidobacterium animalis* on the intestinal microflora during the first 4 months of life. *British Journal of Nutrition*, 94(5), 783–790.
- Bin-Nun, A., Bromiker, R., Wilschanski, M., Kaplan, M., Rudensky, B., Caplan, M., & Hammerman, C. (2005). Oral probiotics prevent necrotizing enterocolitis in very low birth weight neonates. *Journal* of *Pediatrics*, 147(2), 192–196.
- Bocquet, A., Lachambre, E., Kempf, C., & Beck, L. (2013). Effect of infant and follow-on formulas containing *B lactis* and galacto- and fructooligosaccharides on infection in healthy term infants. *Journal of Pediatrics Gastroenterology and Nutrition*, 57(2), 180-187.
- Bonati, M., Barzago, M. M., Stellari, F. F., Bortolotti, A., Introvini, P., Pagani, G., & Siliprandi, N. (1996). Vitamin A supplementation in premature neonates with postnatal lung injury. Italian Collaborative Group on Preterm Delivery (ICGPD). International Journal of Clinical Pharmacology and Therapeutics, 34(8), 362–365.
- Bora, R., & Deori, S. (2019). Transitional hypoglycaemia management in small for gestational age neonates with sucrose enriched expressed breastmilk in resource poor setting. *Journal of Tropical Pediatrics*, 66(3), 267–274.
- Cekola, P. L., Czerkies, L. A., Storm, H. M., Wang, M. H., Roberts, J., & Saavedra, J. M. (2015). Growth and tolerance of term infants fed formula with probiotic *Lactobacillus reuteri*. *Clinical Pediatrics* (*Cleveland*, *OH*), 54(12), 1175–1184.
- Chabra, S., Mayock, D. E., Zerzan, J., Bittner, R., Neufeld, M. D., & Gleason, C. A. (2013). Vitamin A status after prophylactic intramuscular vitamin A supplementation in extremely low birth weight infants. *Nutrition in Clinical Practice*, 28(3), 381–386.
- Chandel, D. S., Perez-Munoz, M. E., Yu, F., Boissy, R., Satpathy, R., Misra, P. R., Sharma, N., Chaudhry, R., Parida, S., Peterson, D. A., Gewolb, I. H., & Panigrahi, P. (2017). Changes in the gut microbiota after early administration of oral synbiotics to young infants in India. *Journal of Pediatrics Gastroenterology and Nutrition*, 65(2), 218–224.
- Chi, C., Xue, Y., Liu, R., Wang, Y., Lv, N., Zeng, H., Buys, N., Zhu, B., Sun, J., & Yin, C. (2019). Effects of a formula with a probiotic *Bifidobacterium lactis* supplement on the gut microbiota of low birth weight infants. *European Journal Nutrition*, 59, 1493–1503.
- Chouraqui, J. P., Grathwohl, D., Labaune, J. M., Hascoet, J. M., de Montgolfier, I., Leclaire, M., Giarre, M., & Steenhout, P. (2008). Assessment of the safety, tolerance, and protective effect against diarrhea of infant formulas containing mixtures of probiotics or probiotics and prebiotics in a randomized controlled trial. American Journal of Clinical Nutrition, 87(5), 1365–1372.
- Chrzanowska-Liszewska, D., Seliga-Siwecka, J., & Kornacka, M. K. (2012). The effect of *Lactobacillus rhamnosus* GG supplemented enteral feeding on the microbiotic flora of preterm infants-double blinded randomized control trial. *Early Human Development*, *88*(1), 57–60.
- Chua, M. C., Ben-Amor, K., Lay, C., Neo, A. G. E., Chiang, W. C., Rao, R., Chew, C., Chaithongwongwatthana, S., Khemapech, N., Knol, J., & Chongsrisawat, V. (2017). Effect of synbiotic on the gut microbiota of cesarean delivered infants: A randomized, double-blind, multicenter study. *Journal of Pediatrics Gastroenterology and Nutrition*, 65(1), 102–106.
- Chou, I. C., Kuo, H. T., Chang, J. S., Wu, S. F., Chiu, H. Y., Su, B. H., & Lin, H. C. (2010). Lack of effects of oral probiotics on growth and neurodevelopmental outcomes in preterm very low birth weight infants. *Journal of Pediatrics*, 156(3), 393–396.
- Corkins, M. R., & Kovacevich, D. S. (2001). Randomised controlled trial of oral vitamin A supplementation in preterm infants to prevent chronic lung disease. Nutrition in Clinical Practice, 16(4), 265.
- Costalos, C., Skouteri, V., Gounaris, A., Sevastiadou, S., Triandafilidou, A., Ekonomidou, C., Kontaxaki, F., & Petrochilou, V. (2003). Enteral feeding of premature infants with *Saccharomyces boulardii*. *Early Human Development*, 74(2), 89–96.

- Costeloe, K., Bowler, U., Brocklehurst, P., Hardy, P., Heal, P., Juszczak, E., King, A., Panton, N., Stacey, F., Whiley, A., Wilks, M., & Millar, M. R. (2016). A randomised controlled trial of the probiotic *Bifidobacterium breve* BBG-001 in preterm babies to prevent sepsis, necrotising enterocolitis and death: the Probiotics in Preterm infantS (PiPS) trial. *Health Technology Assessment*, 20(66), 1–194.
- Costeloe, K., Hardy, P., Juszczak, E., Wilks, M., & Millar, M. R., Probiotics in Preterm Infants Study Collaborative Group. (2016). *Bifidobacterium breve* BBG-001 in very preterm infants: A randomised controlled phase 3 trial. *Lancet*, 387(10019), 649–660.
- Coutsoudis, A., Adhikari, M., Pillay, K., & Coovadia, H. M. (1996). Absorption of high-dose enteral vitamin A in low-birth-weight neonates. South African Medical Journal, 86(10), 1337–1339.
- Dani, C., Biadaioli, R., Bertini, G., Martelli, E., & Rubaltelli, F. F. (2002). Probiotics feeding in prevention of urinary tract infection, bacterial sepsis and necrotizing enterocolitis in preterm infants. A prospective double-blind study. *Biology of the Neonate*, 82(2), 103–108.
- Darboe, M. K., Thurnham, D. I., Morgan, G., Adegbola, R. A., Secka, O., Solon, J. A., Jackson, S. J., Northrop-Clewes, C., Fulford, T. J., Doherty, C. P., & Prentice, A. M. (2007). Effectiveness of an early supplementation scheme of high-dose vitamin A versus standard WHO protocol in Gambian mothers and infants: A randomised controlled trial. *Lancet*, 369(9579), 2088–2096.
- Delimont, N. M., Vahl, C. I., Kayanda, R., Msuya, W., Mulford, M., Alberghine, P., Praygod, G., Mngara, J., Alavi, S., & Lindshield, B. L. (2019). Complementary feeding of sorghum-based and corn-based fortified blended foods results in similar iron, vitamin A, and anthropometric outcomes in the MFFAPP Tanzania efficacy study. *Current Developments in Nutrition*, 3(6), nzz027.
- Delvin, E. E., Salle, B. L., Reygrobellet, B., Mellier, G., & Claris, O. (2000). Vitamin A and E supplementation in breast-fed newborns. *Journal of Pediatrics Gastroenterology and Nutrition*, 31(5), 562–565.
- Deng, J., & Chen, K. (2010). Early minimal feeding combined with probiotics toprevent necrotizing enterocolitis in preterm infant. *Chinese Journal of Modern Drug Application*, 4, 13–14.
- Denkel, L. A., Schwab, F., Garten, L., Geffers, C., Gastmeier, P., & Piening, B. (2017). Dual-strain probiotics reduce NEC, mortality and neonatal bloodstream infections among extremely low birthweight infants. Archives of Disease in Childhood. Fetal and Neonatal Edition, 102(6), f559–f560.
- Deshpande, G., Rao, S., Athalye-Jape, G., Conway, P., & Patole, S. (2016). Probiotics in very preterm infants: The PiPS trial. *Lancet*, *388*(10045), 655.
- Diaby, A., Mohamed, A. S., Camara, B., Sall, G., & Youssouf, M. (2018). Coverage of vitamin A supplementation in children 6 to 59 months after two successive campaigns in Senegal. *Journal de Pediatrie et de La Puericulture*, 31(6), 277–281.
- Dilli, D., Aydin, B., Zenciroglu, A., Ozyazici, E., Beken, S., & Okumus, N. (2013). Treatment outcomes of infants with cyanotic congenital heart disease treated with synbiotics. *Pediatrics*, 132(4), e932–e938.
- Elom, M. O., Okafor, F. C., Eyo, J. E., Usanga, V. U., & Umoh, N. O. (2019). Maternal vitamin a supplementation delays time to first episode of parasitaemia and reduces malaria parasite densities among infants in rural communities in ebonyi state, Nigeria. *Tropical Biomedicine*, 36(1), 194–200.
- Escribano, E., Zozaya, C., Madero, R., Sanchez, L., van Goudoever, J., Rodriguez, J. M., & de Pipaon, M. S. (2018). Increased incidence of necrotizing enterocolitis associated with routine administration of Infloran in extremely preterm infants. *Benef Microbes*, 9(5), 683–690.
- Galderisi, A., Facchinetti, A., Steil, G. M., Ortiz-Rubio, P., Cobelli, C., & Trevisanuto, D. (2016). Neonatal hypoglycemia continuous glucose monitoring: A randomized controlled trial in preterm infants. *Diabetes Technology and Therapeutics*, 18, A57.
- Garg, B. D., Kabra, N. S., Balasubramanian, H., Avasthi, B. S., Sharma, S. R., Ahmed, J., & Dash, S. K. (2017). Prophylactic probiotics for prevention



of necrotizing enterocolitis in preterm neonates: A cohort study. *Perinatology*, 18(2), 41-49.

- Garland, S. M., Tobin, J. M., Pirotta, M., Tabrizi, S. N., Opie, G., Donath, S., Tang, M. L., Morley, C. J., Hickey, L., Ung, L., & Jacobs, S. E. (2011). The ProPrems trial: Investigating the effects of probiotics on late onset sepsis in very preterm infants. *BMC Infectious Diseases*, 11, 210.
- Garofoli, F., Mazzucchelli, I., Decembrino, L., Bartoli, A., Angelini, M., Broglia, M., Tinelli, C., Banderali, G., & Stronati, M. (2018). Levels and effectiveness of oral retinol supplementation in VLBW preterm infants. *International Journal of Immunopathology & Pharmacology*, 32, 2058738418820484.
- Gomber, S. (1996). Potential toxicity of vitamin A supplementation in infancy. *Indian Pediatrics*, 33(12), 1065–1067.
- Gomez-Rodriguez, G., Amador-Licona, N., Daza-Benitez, L., Barbosa-Sabanero, G., Carballo-Magdaleno, D., Aguilar-Padilla, R., & Gonzalez-Ramirez, E. (2019). Single strain versus multispecies probiotic on necrotizing enterocolitis and faecal IgA levels in very low birth weight preterm neonates: A randomized clinical trial. *Pediatrics and Neonatology*, 60(5), 564–569.
- Gonchar, N. V., Lo Skiavo, L. A., & Suvorov, A. N. (2016). Effects of using a probiotic strain of enterococcus on the dynamics of body weight, reduction of the incidence of complications and change of gut microbiota in deeply premature newborn infants. *Voprosy Detskoi Dietologii*, 14(1), 5–14.
- Guo-Qiang, Z., Hua-Jian, H., Chuan-Yang, L., Shristi, S., & Zhong-Yue, L. (2016). Probiotics for preventing late-onset sepsis in preterm neonates. *Medicine (Baltimore)*, 95(8), 1–11.
- Hammerman, C., & Bin-nun, A. (2007). Prebiotics vs. placebo in the prevention of necrotizing enterocolitis in premature neonates. Clinicaltrials.gov identifier: NCT00437567.
- Harris, D. L., Weston, P. J., & Harding, J. E. (2016). A good breast-feed does not always result in an increased in blood glucose concentration, in hypoglycaemic babies. *Journal of Paediatrics and Child Health*, 52, 105.
- Härtel, C. (2019). PRIMAL clinical study: Efficacy of probiotics to prevent gut dysbiosis in very preterm infants (VPIs) and moderate preterm infants of 28 + 0-32 + 6 weeks of gestation: A randomized, placebo-controlled doubleblind study. http://www.drks.de/DRKS00013197DRKS00013197
- Hays, S., Jacquot, A., Gauthier, H., Kempf, C., Beissel, A., & Pidoux, O. (2015). Probiotics and growth in preterm infants: a randomized controlled trial. *Clinical Nutrition*, 35(4), 802–811.
- Hays, S., Jacquot, A., Gauthier, H., Kempf, C., Beissel, A., Pidoux, O., Jumas-Bilak, E., Decullier, E., Lachambre, E., Beck, L., Cambonie, G., Putet, G., Claris, O., & Picaud, J.-C. (2015). Probiotics and growth in preterm infants: a randomized controlled trial, PREMAPRO study. *Clinical Nutrition*, 35(4), 802–811.
- Hays, S., Jacquot, A., Gauthier, H., Kempf, C., Beissel, A., Pidoux, O., Jumas-Bilak, E., Decullier, E., Lachambre, E., Beck, L., Cambonie, G., Putet, G., Claris, O., & Picaud, J.-C. (2016). Probiotics and growth in preterm infants: A randomized controlled trial, PREMAPRO study. *Clinical Nutrition*, 35(4), 802–811.
- Hoyos, A. B. (1999). Reduced incidence of necrotizing enterocolitis associated with enteral administration of *Lactobacillus acidophilus* and *Bifidobacterium infantis* to neonates in an intensive care unit. *International Journal of Infectious Diseases, Summer, 3*(4), 197–202.
- Hoy-Schulz, Y. E., Jannat, K., Roberts, T., Zaidi, S. H., Unicomb, L., Luby, S., & Parsonnet, J. (2016). Safety and acceptability of *Lactobacillus reuteri* DSM 17938 and *Bifidobacterium longum* subspecies infantis 35624 in Bangladeshi infants: a phase I randomized clinical trial. *BMC Complementary and Alternative Medicine*, 16, 44.
- Hua, X. T., Tang, J., Mu, D. Z., Dang, D., Er, K., & Za, Z. (2014). Effect of oral administration of probiotics on intestinal colonization with drugresistant bacteria in preterm infants. *Zhongguo Dang Dai Er Ke Za Zhi*, 16, 606–609.
- Huang, N. N., Wang, G. Z., Wang, J. F., & Yuan, Y. X. (2016). Risk factors for neonatal nosocomial enteric infection and the effect of intervention

with BIFICO. European Review for Medical and Pharmacological Sciences, 20(17), 3713–3719.

- Hunter, C., Dimaguila, M. A., Gal, P., Wimmer, J. E., Jr., Ransom, J. L., Carlos, R. Q., Smith, M., & Davanzo, C. C. (2012). Effect of routine probiotic, *Lactobacillus reuteri* DSM 17938, use on rates of necrotizing enterocolitis in neonates with birthweight <1000 grams: A sequential analysis. *BMC Pediatrics*, 12, 142.
- Idindili, B., Masanja, H., Urassa, H., Bunini, W., van Jaarsveld, P., Aponte, J. J., Kahigwa, E., Mshinda, H., Ross, D., & Schellenberg, D. M. (2007). Randomized controlled safety and efficacy trial of 2 vitamin A supplementation schedules in Tanzanian infants. *American Journal of Clinical Nutrition*, 85(5), 1312–1319.
- Indrio, F., Riezzo, G., Raimondi, F., Bisceglia, M., Cavallo, L., & Francavilla, R. (2008). The effects of probiotics on feeding tolerance, bowel habits, and gastrointestinal motility in preterm newborns. *Journal of Pediatrics*, 152(6), 801–806.
- Indrio, F., Riezzo, G., Tafuri, S., Ficarella, M., Carlucci, B., Bisceglia, M., Polimeno, L., & Francavilla, R. (2017). Probiotic supplementation in preterm: Feeding intolerance and hospital cost. *Nutrients*, 9(9), 965.
- Irct201505253915N (2015). Comparing the efficacy of two doses of vitamin A in preterm infants (Other: IRCT201505253915N). http://www.who. int/trialsearch/Trial2.aspx
- Jacobs, S. E., Hickey, L., Donath, S., Opie, G. F., Anderson, P. J., Garland, S. M., & Cheong, J. L. Y. (2017). Probiotics, prematurity and neurodevelopment: Follow-up of a randomised trial. *BMJ Paediatr Open*, 1(1), 176.
- Jacobs, S. E., Tobin, J. M., Opie, G. F., Donath, S., Tabrizi, S. N., Pirotta, M., Morley, C. J., & Garland, S. M. (2013). Probiotic effects on late-onset sepsis in very preterm infants: a randomized controlled trial. *Pediatrics*, 132(6), 1055–1062.
- Jacobs, S. E., Tobin, J. M., Opie, G., Donath, S., Pirotta, M., Tabrizi, S. N., Morley, C. J., & Garland, S. M. (2012). The proprems randomised trial investigating the effects of probiotics on late onset sepsis in very preterm infants. Archives of Disease in Childhood, 97, A50.
- Jacobs, S., Hickey, L., Donath, S., Opie, G., Anderson, P., Garland, S., & Cheong, J. (2016). Probiotics and neurodevelopment in very preterm infants: Follow-up of a randomized trial. *European Journal of Pediatrics*, 175(11), 1440–1441.
- Jacobs, S., Tobin, J., Opie, G., Donath, S., Tabrizi, S., & Pirotta, M. The ProPrems Randomised Trial investigating the effects of probiotics on late onset sepsis in very preterm infants. Pediatric Academic Societies Annual Meeting 2013.
- Janvier, A., Malo, J., & Barrington, K. J. (2014). Cohort study of probiotics in a North American neonatal intensive care unit. *Journal of Pediatrics*, 164(5), 980–985.
- Kahbazi, M., Sharafkhah, M., Yousefichaijan, P., Taherahmadi, H., Rafiei, M., Kaviani, P., Abaszadeh, S., Massoudifar, A., & Mohammadbeigi, A. (2019). Vitamin A supplementation is effective for improving the clinical symptoms of urinary tract infections and reducing renal scarring in girls with acute pyelonephritis: a randomized, double-blind placebo-controlled, clinical trial study. *Complementary Therapies in Medicine*, 42, 429–437.
- Kanic, Z., Micetic Turk, D., Burja, S., Kanic, V., & Dinevski, D. (2015). Influence of a combination of probiotics on bacterial infections in very low birthweight newborns. *Wiener Klinische Wochenschrift*, 127(5), S210–S215.
- Karthikeyan, G., & Bhat, B. V. (2017). The PiPS (Probiotics in Preterm Infants Study) Trial—Controlling the confounding factor of crosscontamination unveils significant benefits. *Indian Pediatrics*, 54(2), 162.
- Kiatchoosakun, P., Jirapradittha, J., Panthongviriyakul, C., Khampitak, T., Yongvanit, P., & Boonsiri, P. (2014). Vitamin a supplementation for prevention of bronchopulmonary dysplasia in very-low-birth-weight premature thai infants: A randomized trial. *Chotmaihet Thangphaet* (Journal of the Medical Association of Thailand), 97, S82–S88.

- Kiatchoosakun, P., Jirapradittha, J., Panthongviriyakul, M. C., Khampitak, T., Yongvanit, P., & Boonsiri, P. (2014). Vitamin A supplementation for prevention of bronchopulmonary dysplasia in very-low-birth-weight premature Thai infants: A randomized trial. Journal of the Medical Association of Thailand, 97, S82–S88.
- Kirkwood, B., Humphrey, J., Moulton, L., & Martines, J. (2010). Neonatal vitamin A supplementation and infant survival. *Lancet*, 376(9753), 1643–1644.
- Kliegman, R. M. (2005). Oral probiotics reduce the incidence and severity of necrotizing enterocolitis in very low birth weight infants. *Journal of Pediatrics*, 146(5), 710.
- Koksal, N., Varal, I., Ozkan, H., Bagci, O., & Dotan, P. (2015). Effect of probiotic support on feeding intolerance and mortality at preterm infants. *Journal of Perinatal Medicine*, 43.
- Kukkonen, K., Savilahti, E., Haahtela, T., Juntunen-Backman, K., Korpela, R., Poussa, T., Tuure, T., & Kuitunen, M. (2008). Long-term safety and impact on infection rates of postnatal probiotic and prebiotic (synbiotic) treatment: Randomized, double-blind, placebocontrolled trial. *Pediatrics*, 122(1), 8–12.
- Leele, Y., Bharani, R., Biswas, A., Lee, J., Tran, L. A., Pecquet, S., & Steenhout, P. (2015). Normal growth of infants receiving an infant formula containing *Lactobacillus reuteri*, galacto-oligosaccharides, and fructo-oligosaccharide: A randomized controlled trial. *Maternal Health*, *Neonatology and Perinatology*, 1, 9.
- Li, X., Peng, Y., Li, Z., Christensen, B., Heckmann, A. B., Stenlund, H., Lonnerdal, B., & Hernell, O. (2019). Feeding infants formula with probiotics or milk fat globule membrane: A double-blind, randomized controlled trial. *Frontiers in Pediatrics*, *7*, 347.
- Lin, H. C., Hsu, C. H., Chen, H. L., Chung, M. Y., Hsu, J. F., Lien, R. I., Tsao, L. Y., Chen, C. H., & Su, B. H. (2009). Oral probiotics prevent necrotizing enterocolitis in very low birth weight preterm infants: A multicenter, randomized, controlled trial. *Obstetrical and Gynecological Survey*, 64(2), 84–85.
- Lin, H. C., Hsu, C. H., Chen, H. L., Chung, M. Y., Hsu, J. F., Lien, R. I., Tsao, L. Y., Chen, C. H., & Su, B. H. (2008). Oral probiotics prevent necrotizing enterocolitis in very low birth weight preterm infants: A multicenter, randomized, controlled trial. *Pediatrics*, 122(4), 693–700.
- Lin, H. C., Su, B. H., Chen, A. C., Lin, T. W., Tsai, C. H., Yeh, T. F., & Oh, W. (2005). Oral probiotics reduce the incidence and severity of necrotizing enterocolitis in very low birth weight infants. *Pediatrics*, 115(1), 1–4.
- Lin, H. C., Su, B. H., & Oh, W. (2006). Oral probiotics prevent necrotizing enterocolitis. Journal of Pediatrics, 148(6), 849.
- Long, A., & Dempsey, E. (2018). Oral probiotic supplementation in the prevention of necrotising enterocolitis among very preterm infants. *Irish Journal of Medical Science*, 187(3), S75.
- Lozano, J. M. (2008). Prophylactic probiotics for the prevention of sepsis and NEC in premature infants in Colombia. A randomized double-blind, multicenter trial (Clinicaltrials.gov identifier: NCT00727363).
- Lund, N., Biering-Sorensen, S., Andersen, A., Monteiro, I., Camala, L., Jorgensen, M. J., Aaby, P., & Benn, C. S. (2014). Neonatal vitamin A supplementation associated with a cluster of deaths and poor early growth in a randomised trial among low-birth-weight boys of vitamin A versus oral polio vaccine at birth. *Boston Medical Center Pediatrics August, 28*, 14.
- Lund, N., Biering-Sørensen, S., Andersen, A., Monteiro, I., Camala, L., Jul Jørgensen, M., Aaby, P., & Stabell Benn, C. (2016). Neonatal vitamin A supplementation associated with a cluster of deaths and poor early growth in a randomised trial among low-birth-weight boys of vitamin A versus oral polio vaccine at birth. *Neonatal Intensive Care*, 29(4), 44–52.
- Lundelin, K., Poussa, T., Salminen, S., & Isolauri, E. (2017). Long-term safety and efficacy of perinatal probiotic intervention: Evidence from a follow-up study of four randomized, double-blind, placebocontrolled trials. *Pediatric Allergy and Immunology*, 28(2), 170–175.

- Mactier, H., McCulloch, D. L., Hamilton, R., Galloway, P., Bradnam, M. S., Young, D., Lavy, T., Farrell, L., & Weaver, L. T. (2012). Vitamin A supplementation improves retinal function in infants at risk of retinopathy of prematurity. *Journal of Pediatrics*, 160(6), 954–959.
- Maldonado-Lobon, J. A., Gil-Campos, M., Maldonado, J., Lopez-Huertas, E., Flores-Rojas, K., Valero, A. D., Rodriguez-Benitez, M. V., Banuelos, O., Lara-Villoslada, F., Fonolla, J., & Olivares, M. (2015). Long-term safety of early consumption of *Lactobacillus fermentum* CECT5716: A 3-year follow-up of a randomized controlled trial. *Pharmacological Research*, 95-96, 12–19.
- Maldonado-Lobón, J. A., Gil-Campos, M., Maldonado, J., López-Huertas, E., Flores-Rojas, K., Valero, A. D., Rodríguez-Benítez, M. V., Bañuelos, O., Lara-Villoslada, F., Fonollá, J., & Olivares, M. (2015). Long-term safety of early consumption of *Lactobacillus fermentum* CECT5716: A 3-year follow-up of a randomized controlled trial. *Pharmacological Research*, 95-96, 12–19.
- Manzano, S., De Andres, J., Castro, I., Rodriguez, J. M., Jimenez, E., & Espinosa-Martos, I. (2017). Safety and tolerance of three probiotic strains in healthy infants: A multi-centre randomized, double-blind, placebo-controlled trial. *Benef Microbes*, 8(4), 569–578.
- Manzoni, P., Mostert, M., Leonessa, M. L., Priolo, C., Farina, D., Monetti, C., Latino, M. A., & Gomirato, G. (2006). Oral supplementation with *Lactobacillus casei* subspecies rhamnosus prevents enteric colonization by Candida species in preterm neonates: A randomized study. *Clinical Infectious Diseases*, 42(12), 1735–1742.
- Manzoni, P., Rinaldi, M., Cattani, S., Pugni, L., Romeo, M. G., Messner, H., Stolfi, I., Decembrino, L., Laforgia, N., Vagnarelli, F., Memo, L., Bordignon, L., Saia, O. S., Maule, M., Gallo, E., Mostert, M., Magnani, C., Quercia, M., Bollani, L., ... Farina, D. (2009). Bovine lactoferrin supplementation for prevention of late-onset sepsis in very low-birthweight neonates: A randomized trial. *Journal of the American Medical Association*, 302(13), 1421–1428.
- Marissen, J., Haiss, A., Meyer, C., Van Rossum, T., Bunte, L. M., Frommhold, D., Gille, C., Goedicke-Fritz, S., Gopel, W., Hudalla, H., Pagel, J., Pirr, S., Siller, B., Viemann, D., Vens, M., Konig, I., Herting, E., Zemlin, M., Gehring, S., ... Hartel, C. (2019). Efficacy of *Bifidobacterium longum*, *B. infantis* and *Lactobacillus acidophilus* probiotics to prevent gut dysbiosis in preterm infants of 28 + 0-32 + 6 weeks of gestation: a randomised, placebo-controlled, double-blind, multicentre trial: The PRIMAL Clinical Study protocol. *BMJ Open*, 9(11), e032617.
- Martins, T. M., Martinelli, C. E., Daneluzzi, J. C., Del Ciampo, L. A., Ricco, R. G., Vannucchi, H., Jordão, A. A., Portari, G. V., De Souza Meirelles, M. S., Ovidio, P. P., & Ferraz, I. S. (2009). Impact of maternal vitamin a supplementation on mother-infant pair: A randomized double-blind placebo-controlled trial. *Pediatric Research*, *65*(4), 478–481.
- Materna Laboratories. (2010). Evaluation of the effect of milk based infant formula supplemented either with probiotic microorganisms and/or with prebiotic on the intestinal microflora during the first 4 months of life of healthy, full term infants and it's long term effect on morbidity up to the age of 9 months (Clinicaltrials.gov identifier: NCT00836771).
- McCulloch, D. L., Mactier, H., Farrell, L., & Hamilton, R. (2012). Lightadapted ERGs in the VitAL study: a randomised controlled trial of early high-dose vitamin A in infants at risk of retinopathy of prematurity (ROP). *Documenta Ophthalmologica*, 124(1), 22–23.
- McKinlay, C. J., Rebello, C., & Tarnow-Mordi, W. (2016). Probiotics in very preterm infants: The PiPS trial. *Lancet*, *388*(10045), 655.
- Meyer, S., & Gortner, L. (2014). Early postnatal additional high-dose oral vitamin A supplementation versus placebo for 28 days for preventing bronchopulmonary dysplasia or death in extremely low birth weight infants. *Neonatology*, 105(3), 182–188.
- Mihatsch, W. A., Vossbeck, S., Eikmanns, B., Hoegel, J., & Pohlandt, F. (2010). Effect of *Bifidobacterium lactis* on the incidence of nosocomial infections in very-low-birth-weight infants: A randomized controlled trial. *Neonatology*, 98(2), 156–163.



- Mihatsch, W. A., Vossbeck, S., Franz, A. R., Kron, M., & Pohlandt, F. (2004). Effect of enteral administration of a probiotic strain of bifidobacteria on the incidence of nosocomial infections in preterm infants. *Pediatric Research*, 55, 46.
- Millar, M., Seale, J., Greenland, M., Hardy, P., Juszczak, E., Wilks, M., Panton, N., Costeloe, K., & Wade, W. G. (2017). The microbiome of infants recruited to a randomised placebo-controlled probiotic trial (PiPS Trial). *EBioMedicine*, 20, 255–262.
- Millar, M., Seale, J., Greenland, M., Hardy, P., Juszczak, E., Wilks, M., Panton, N., Costeloe, K., & Wade, W. G. (2017). The microbiome of infants recruited to a randomised placebo-controlled probiotic trial (PiPS Trial). *EBioMedicine*, 20, 255–262.
- Moles, L., Escribano, E., de Andres, J., Montes, M. T., Rodriguez, J. M., Jimenez, E., Saenz de Pipaon, M., & Espinosa-Martos, I. (2015). Administration of *Bifidobacterium breve* PS12929 and *Lactobacillus salivarius* PS12934, two strains isolated from human milk, to very low and extremely low birth weight preterm infants: A pilot study. *Journal of Immunology Research*, 2015, 538171.
- Nadella, P., Smith, E. R., Muhihi, A., Noor, R. A., Masanja, H., Fawzi, W. W., & Sudfeld, C. R. (2019). Determinants of delayed or incomplete diphtheriatetanus-pertussis vaccination in parallel urban and rural birth cohorts of 30,956 infants in Tanzania. *BMC Infectious Disease*, 19(1), 188.
- NCT. (2006). Vitamin A and very low birthweight babies (VitAL). https:// clinicaltrials.gov/show/nct00417404
- NCT. (2011). Bifidobacterium supplementation for very low birth weight infants. https://clinicaltrials.gov/show/nct01375309
- NCT (2016). Functional evaluation of two infant formula supplemented with probiotics isolated from breast milk. https://clinicaltrials.gov/show/ nct03204630
- Papagaroufalis, C., Megreli, C., Hagjigeorgi, C., & Xanthou, M. (1991). A trial of vitamin A supplementation for the prevention of intraventricular haemorrhage in very low birth weight neonates. *Journal of Perinatal Medicine*, 19(1), 382–387.
- Papagaroufalis, C., Spyropoulos, G., Stamocosta, E., Megreli, C., & Xanthou, M. (1992). A trial of vitamin A (VA) supplementation for the prevention of retinopathy of prematurity (ROP) in very low birth weight (VLBW) infants. *Journal of Perinatal Medicine*, 20(1), 35.
- Papagaroufalis, K., Fotiou, A., Egli, D., Tran, L. A., & Steenhout, P. (2014). A randomized double blind controlled safety trial evaluating d-lactic acid production in healthy infants fed a *Lactobacillus reuteri*-containing formula. *Nutrition and Metabolic Insights*, 7, 19–27.
- Patole, S. K., Keil, A. D., Nathan, E., Doherty, D., Esvaran, M., Simmer, K. N., & Conway, P. (2016). Effect of *Bifidobacterium breve* M-16V supplementation on faecal bifidobacteria in growth restricted very preterm infants - analysis from a randomised trial. *The Journal of Maternal-Fetal & Neonatal Medicine*, 29(23), 3751–3755.
- Patole, S., Keil, A. D., Chang, A., Nathan, E., Doherty, D., Simmer, K., Esvaran, M., & Conway, P. (2014). Effect of *Bifidobacterium breve* M-16V supplementation on fecal bifidobacteria in preterm neonates—A randomised double blind placebo controlled trial. *PLOS One*, *9*(3), e89511.
- Pearson, E., Bose, C., Snidow, T., Ransom, L., Young, T., Bose, G., & Stiles, A. (1992). Trial of vitamin A supplementation in very low birth weight infants at risk for bronchopulmonary dysplasia. *Journal of Pediatrics*, 121(3), 420–427.
- Plummer, E. L., Bulach, D. M., Murray, G. L., Jacobs, S. E., Tabrizi, S. N., & Garland, S. M. (2018). Gut microbiota of preterm infants supplemented with probiotics: Sub-study of the ProPrems trial. *BMC Microbiology*, 18(1), 184.
- Puccio, G., Cajozzo, C., Meli, F., Rochat, F., Grathwohl, D., & Steenhout, P. (2007). Clinical evaluation of a new starter formula for infants containing live *Bifidobacterium longum* BL999 and prebiotics. *Nutrition*, 23(1), 1–8.
- Qiao, L. X., Zhu, W. Y., Zhang, H. Y., & Wang, H. (2017). Effect of early administration of probiotics on gut microflora and feeding in pre-term

-WILEY-Campbell Collaboration

infants: a randomized controlled trial. The Journal of Maternal-Fetal & Neonatal Medicine, 30(1), 13–16.

- Radke, M., Picaud, J. C., Loui, A., Cambonie, G., Faas, D., Lafeber, H. N., de Groot, N., Pecquet, S. S., Steenhout, P. G., & Hascoet, J. M. (2017).
 Starter formula enriched in prebiotics and probiotics ensures normal growth of infants and promotes gut health: A randomized clinical trial. *Pediatric Research*, 81(4), 622–631.
- Raguž, M. J., Brzica, J., Rozić, S., Glamuzina, D. S., Mustapić, A., Bošnjak, M. N., & Božić, T. (2016). The impact of probiotics (*Lactobacillus reuteri* Protectis) on the treatment, course and outcome of premature infants in the intensive care unit in Mostar. *Journal of Pediatric and Neonatal Individualized Medicine*, 5(2), e050228.
- Rakshasbhuvankar, A., Patole, S., Simmer, K., & Pillow, J. J. (2017). Enteral vitamin A for reducing severity of bronchopulmonary dysplasia in extremely preterm infants: a randomised controlled trial. BMC Pediatrics, 17(1), 204.
- Rawat, M., Chandrasekharan, P., Turkovich, S., Barclay, N., Perry, K., Schroeder, E., Testa, L., & Lakshminrusimha, S. (2016). Oral dextrose gel reduces the need for intravenous dextrose therapy in neonatal hypoglycemia. *Biomed Hub*, 1(3), 1–9.
- Repa, A., Thanhaeuser, M., Endress, D., Weber, M., Kreissl, A., Binder, C., Berger, A., & Haiden, N. (2015). Probiotics (*Lactobacillus acidophilus* and *Bifidobacterium infantis*) prevent NEC in VLBW infants fed breast milk but not formula [corrected]. *Pediatric Research*, 77(2), 381–388.
- Robbins, S. T., & Fletcher, A. B. (1993). Early vs delayed vitamin A supplementation in very-low-birth-weight infants. *Journal of Parenteral* and Enteral Nutrition, 17(3), 220–225.
- Rodríguez (2015). Prevention of necrotizing enterocolitis of premature newborns under less than 1500 g using probiotics. 2014.https:// clinicaltrials.gov/show/nct02245815
- Rodriguez-Herrera, A., Mulder, K., Bouritius, H., Rubio, R., Munoz, A., Agosti, M., Lista, G., Corvaglia, L., Ludwig, T., Abrahamse-Berkeveld, M., & Perez-Navero, J. L. (2019). Gastrointestinal tolerance, growth and safety of a partly fermented formula with specific prebiotics in healthy infants: A double-blind, randomized, controlled trial. *Nutrients*, 11(7), 1530.
- Rohan, Annie J. (2016). Bifidobacterium breve BBG-001 in very preterm infants: A randomised controlled phase 3 trial. The American Journal of Maternal Child Nursing, 41(4), 258.
- Romeo, M. G., Romeo, D. M., Trovato, L., Oliveri, S., Palermo, F., Cota, F., & Betta, P. (2011). Role of probiotics in the prevention of the enteric colonization by Candida in preterm newborns: incidence of late-onset sepsis and neurological outcome. *Journal of Perinatology*, 31(1), 63–69.
- Ross, D. A., Dollimore, N., Smith, P. G., Kirkwood, B. R., Arthur, P., Morris, S. S., Binka, F., Arthur, P., Gyapong, J. O., & Tomkins, A. M., Ghana VAST Study Team. (1993). Vitamin A supplementation in northern Ghana: Effects on clinic attendances, hospital admissions, and child mortality. *Lancet*, 342(8862), 7–12.
- Rouge, C., Piloquet, H., Butel, M. J., Berger, B., Rochat, F., Ferraris, L., Des Robert, C., Legrand, A., de la Cochetiere, M. F., N'Guyen, J. M., Vodovar, M., Voyer, M., Darmaun, D., & Roze, J. C. (2009). Oral supplementation with probiotics in very-low-birth-weight preterm infants: a randomized, double-blind, placebo-controlled trial. *American Journal of Clinical Nutrition*, 89(6), 1828–1835.
- Rubaltelli, F., Biadaioli, R., & Dani, C. (2000). Probiotics feeding prevents necrotizing enterocolitis in preterm infants: a prospective doubleblind study. *Pediatric Research*, 47(4), 346A.
- Sadowska-Krawczenko, I., Korbal, P., Polak, A., Wietlicka-Piszcz, M., & Szajewska, H. (2012). Lactobacilllus rhamnosus ATC A07FA for preventing necrotizing enterocolitis in very-low-birth-weight preterm infants: A randomized controlled trial (preliminary results). *Pediatria Polska*, 87(2), 139–145.
- Samuels, N., van de Graaf, R., Been, J. V., de Jonge, R. C., Hanff, L. M., Wijnen, R. M., Kornelisse, R. F., Reiss, I. K., & Vermeulen, M. J. (2016). Necrotising enterocolitis and mortality in preterm infants after

introduction of probiotics: a quasi-experimental study. *Scientific Reports*, *6*, 31643.

- Shenai, J. P., Kennedy, K. A., Chytil, F., & Stahlman, M. T. (1987). Clinical trial of vitamin A supplementation in infants susceptible to bronchopulmonary dysplasia. *Journal of Pediatrics*, 111(2), 269–277.
- Shujie, Y., Haiying, Y., Bin, G., Shu, X., Xianglan, D., & Jiang, W. (2011). The clinical application value of endangered preterm infants given earlier amounts of micro feedings and adding probiotics. *The Journal of Pediatric Pharmacology*, 17, 21–24.
- Smilowitz, J. T., Moya, J., Breck, M. A., Cook, C., Fineberg, A., Angkustsiri, K., & Underwood, M. A. (2017). Safety and tolerability of *Bifidobacterium longum* subspecies infantis EVC001 supplementation in healthy term breastfed infants: A phase I clinical trial. *BMC Pediatrics*, 17(1), 133.
- Storm, H. M., Shepard, J., Czerkies, L. M., Kineman, B., Cohen, S. S., Reichert, H., & Carvalho, R. (2019). 2'-Fucosyllactose is well tolerated in a 100% whey, partially hydrolyzed infant formula with *Bifidobacterium lactis*: A randomized controlled trial. *Global Pediatric Health*, *6*, 6.
- Stratiki, Z., Costalos, C., Sevastiadou, S., Kastanidou, O., Skouroliakou, M., Giakoumatou, A., & Petrohilou, V. (2007). The effect of a bifidobacter supplemented bovine milk on intestinal permeability of preterm infants. *Early Human Development*, 83(9), 575–579.
- Strus, M., Helwich, E., Lauterbach, R., Rzepecka-Weglarz, B., Nowicka, K., Wilinska, M., Szczapa, J., Rudnicka, M., Slawska, H., Szczepanski, M., Wasko, A., Mikolajczyk-Cichonska, A., Tomusiak-Plebanek, A., & Heczko, P. B. (2018). Effects of oral probiotic supplementation on gut *Lactobacillus* and *Bifidobacterium* populations and the clinical status of low-birth-weight preterm neonates: a multicenter randomized, double-blind, placebo-controlled trial. *Infection and Drug Resistance*, 11, 1557–1571.
- Ter, M., Halibullah, I., Leung, L., & Jacobs, S. (2017). Implementation of dextrose gel in the management of neonatal hypoglycaemia. *Journal of Paediatrics and Child Health*, 53(4), 408–411.
- Thanhaeuser, M., Repa, A., Weber, M., Endress, D., Kreissl, A., Binder, C., Berger, A., & Haiden, N. (2014). Probiotics (infloran) for NEC prevention: Influence of enteral nutrition. Archives of Disease in Childhood, 99, A176–A177.
- Totsu, S., Yamasaki, C., Terahara, M., Uchiyama, A., & Kusuda, S. (2014). *Bifidobacterium* and enteral feeding in preterm infants: clusterrandomized trial. *Pediatria Internazionale*, 56(5), 714–719.
- Tyson, J. E., Wright, L. L., Oh, W., Kennedy, K. A., Mele, L., Ehrenkranz, R. A., Stoll, B. J., Lemons, J. A., Stevenson, D. K., Bauer, C. R., Korones, S. B., Donovan, E. F., Carlo, W. A., Shankaran, S., Stark, A. R., Papile, L. A., Jobe, A., Stacewicz-Sapuntzakis, M., Verter, J., & Fanaroff, A. A. (1999). Vitamin A supplementation for extremely-low-birth-weight infants. New England Journal of Medicine, 340(25), 1962–1968.
- Venkatarao, T., Ramakrishnan, R., Nair, N. G., Radhakrishnan, S., Sundaramoorthy, L., Koya, P. K., & Kumar, S. K. (1996). Effect of vitamin A supplementation to mother and infant on morbidity in infancy. *Indian Pediatrics*, 33(4), 279–286.
- Vlieger, A. M., Robroch, A., van Buuren, S., Kiers, J., Rijkers, G., Benninga, M. A., & te Biesebeke, R. (2009). Tolerance and safety of *Lactobacillus paracasei* ssp. paracasei in combination with *Bifidobacterium animalis* ssp. lactis in a prebiotic-containing infant formula: a randomised controlled trial. *British Journal of Nutrition*, 102(6), 869-875.
- Wardle, S. P., Hughes, A., Chen, S., & Shaw, N. J. (2001). Randomised controlled trial of oral vitamin A supplementation in preterm infants to prevent chronic lung disease. Archives of Disease in Childhood. Fetal and Neonatal Edition, 84(1), F9-F13.
- West, K. P., Jr, Pokhrel, R. P., Katz, J., LeClerq, S. C., Khatry, S. K., Shrestha, S. R., Pradhan, E. K., Tielsch, J. M., Pandey, M. R., & Sommer, A. (1991). Efficacy of vitamin A in reducing preschool child mortality in Nepal. *Lancet*, 338(8759), 67–71. 13.

STUDIES AWAITING CLASSIFICATION

- Barclay, D., Puccio, G., Fazzolari-Nesci, A., Giammanco, A., Raiha, N., Carrie Fassler, A. L., Brown, C., Chauffard, F., Grathwohl, D., Hager, C., & Endres, W. (2003). Growth and tolerance of a whey-based starter infant formula with enhanced protein efficiency and containing pro-, pre- or synbiotics. A randomized controlled trial in term infants. *Journal of Pediatrics Gastroenterology and Nutrition*, *37*, 388.
- Chubarova, A. I., & Sharyafetdinova, G. R. (2017). An experience of using a preparation containing combined probiotic strains of bifidobacteria and lactobacilli in premature newborns in neonatal resuscitation and intensive care units. *Voprosy Detskoi Dietologii*, 15(4), 5–13.

ONGOING STUDIES

- Del Piano, M., Coggiola, F., Pane, M., Amoruso, A., Nicola, S., & Mogna, L. (2015). Can probiotics reduce diarrhea and infant mortality in africa?: The project of a pilot study. *Journal of Clinical Gastroenterology*, S120–S123.
- Goodman, T. (2015). Prevention of vitamin A deficiency by supplementation alongside routine vaccinations: A randomised controlled trial in Ghana infants. http://isrctn.com/ISRCTN97670178ISRCTN97670178
- Kaur, A. (2018). Effect of probiotic supplementation on feed tolerance and weight gain in low birth weight infants on tube feeds. http://www.ctri.nic. in/Clinicaltrials/pmaindet2.php?trialid=26719
- Londhe, A. (2019). Use of zinc and pre-probiotics as a therapeutic adjunct in neonatal sepsis in preterms—An Open label randomized controlled trial. http://www.ctri.nic.in/Clinicaltrials/pmaindet2.php?trialid=16167
- Mirmohammdi, M. F. (2018). The effect of probiotics on prevention of necrotizing enterocolitis in neonates. http://en.irct.ir/trial/ 25078IRCT20170121032075N2
- Mukhtar, G. (2019). Role of prophylactic microbial supplements in prevention of blood stream infection and intestinal tract injury in premature neonates (CTRI/2018/04/013401)http://www.ctri.nic.in/Clinicaltrials/ pmaindet2.php?trialid=23706
- Nandhini, I. P. (2012). A clinical study to analyse the effectiveness of administration of harmless bacteria in preventing infection of intestine in preterm babies. http://www.ctri.nic.in/Clinicaltrials/pmaindet2.php? trialid=3864CTRI/2012/09/003008
- Punnahitananda, S. (2011). Effect of oral probiotic supplementation on the rate of hospital acquired infection and necrotizing enterocolitis in preterm very low birth weight infants. http://isrctn.com/ISRCTN39142169ISRCTN39142169
- Rathod, M. (2019). Probiotics for prevention of necrotising enterocoilitis in preterm neonates. http://www.ctri.nic.in/Clinicaltrials/pmaindet2.php? trialid=1924
- Razavi, S. M. (2014). Effect of probiotic in prevention of necroziting entrocolitis in preterm infants in Hafez hospital. http://en.irct.ir/trial/ 16814IRCT2014072618591N1
- Shashidhar, A. (2019). A study of the effect of probiotic organism administration on feeding tolerance in very low birth weight newborn babies. http://www.ctri.nic.in/Clinicaltrials/pmaindet2.php?trialid= 4955CTRI/2012/08/002853
- Siamack Karimi Heydarabad (2018). Evaluation of the effect of probiotics on late-onset sepsis in very preterm newborns. http://en.irct.ir/trial/ 29386IRCT20171218037936N1
- Sinha, A. (2019). Phase III, multicentre, randomized, double-blind, placebocontrolled study to evaluate efficacy of probiotic supplementation for prevention of neonatal sepsis in 0-2 months old low birth weight infants in India. http://www.ctri.nic.in/Clinicaltrials/pmaindet2.php?trialid=33881

OTHER REFERENCES-ADDITIONAL REFERENCES

van den Akker, C. H. P., van Goudoever, J. B., Szajewska, H., Embleton, N. D., Hojsak, I., Reid, D., & Shamir, R. (2018). Probiotics for Preterm Infants: A strain specific systematic review and network meta-



83 of 88

analysis. Journal of Pediatric Gastroenterology and Nutrition, 67, 1–122. https://doi.org/10.1097/MPG.00000000001897

- Alfaleh, K., Anabrees, J., Bassler, D., & Al-Kharfi, T. (2011). Probiotics for prevention of necrotizing enterocolitis in preterm infants. *Cochrane Database of Systematic Reviews*, 16(3), CD005496.
- AlFaleh, K., & Anabrees, J. (2014). Probiotics for prevention of necrotizing entercolitis in preterm infants. *Cochrane Database of Systematic Reviews*, 4, CD005496. https://doi.org/10.1002/14651858. CD005496.pub4
- Alkema, L., Chao, F., You, D., Pedersen, J., & Sawyer, C. (2014). National, regional, and global sex ratios of infant, child and under-5 mortality and identification of countries with outlying ratios: a systematic assessment. *The Lancet Global Health*, 2(9), e521–e530. https://doi. org/10.1016/52214-109X(14)70280-3
- Arrieta, M. C., Stiemsma, L. T., Amenyogbe, N., Brown, E. M., & Finlay, B. (2014). The intestinal microbiome in early life: Health and disease. *Frontiers in Immunology*, 427(5), 427.
- Bates, C. J. (1995). Vitamin A. Lancet, 345(8941), 31-35.
- Bhutta, Z. A., Das, J. K., Rizvi, A., Gaffey, M. F., Walker, N., & Horton, S. (2013). Evidence-based interventions for improvement of maternal and child nutrition: what can be done and at what cost? *The Lancet*, 382(9890), 452–477. https://doi.org/10.1016/S0140-6736(13)60996-4
- Cochrane Effective Practice and Organisation of Care (EPOC). Suggested risk of bias criteria for EPOC reviews. EPOC Resources for Review Authors. 2017. https://epoc.cochrane.org/resources/epoc-resourcesreview-authors
- Conde-Agudelo, A., & Díaz-Rossello, J. L. (2016). Kangaroo mother care to reduce morbidity and mortality in low birthweight infants. *Cochrane Database of Systematic Reviews*, 8, CD002771. https://doi.org/10. 1002/14651858.CD002771.pub4
- Coors, S. M., Cousin, J. J., Hagan, J. L., & Kaiser, J. R. (2018). Prophylactic dextrose gel does not prevent neonatal hypoglycemia: A quasiexperimental pilot study. *The Journal of Pediatrics*, 198, 156–161. https://doi.org/10.1016/j.jpeds.2018.02.025
- Costeloe, K., Bowler, U., Brocklehurst, P., Hardy, P., Heal, P., Juszczak, E., King, A., Panton, N., Stacey, F., Whiley, A., Wilks, M., & Millar, M. R. (2016). A randomised controlled trial of the probiotic *Bifidobacterium breve* BBG-001 in preterm babies to prevent sepsis, necrotising enterocolitis and death: the Probiotics in Preterm infantS (PiPS) trial. *Health Technology Assessment*, 20(66), 1–194. https://doi.org/10.3310/ hta20660.
- Covidence systematic review software. Veritas Health Innovation, Melbourne, Australia. www.covidence.org 2019.
- Dani, C., Coviello, C. C., Corsini, I. I., Arena, F., Antonelli, A., & Rossolini, G. M. (2016). *Lactobacillus sepsis* and probiotic therapy in newborns: Two new cases and literature review. *AJP Reports*, 6(1), e25–e29. https:// doi.org/10.1055/s-0035-1566312
- Darlow, B. A., Graham, P. J., & Rojas-Reyes, M. X. (2016). Vitamin A supplementation to prevent mortality and short- and long-term morbidity in very low birth weight infants. *Cochrane Database of Systematic Reviews*, 8, CD000501. https://doi.org/10.1002/14651858. CD000501.pub4
- Del Piano, M., Coggiola, F., Pane, M., Amoruso, A., Nicola, S., & Mogna, L. (2015). Can probiotics reduce diarrhea and infant mortality in africa? The project of a pilot study. *Journal of Clinical Gastroenterology*, S120–S123.
- Deshmukh, H. S., Liu, Y., Menkiti, O. R., Mei, J., Dai, N., O'Leary, C. E., Oliver, P. M., Kolls, J. K., Weiser, J. N., & Worthen, G. S. (2014). The microbiota regulates neutrophil homeostasis and host resistance to *Escherichia coli* K1 sepsis in neonatal mice. *Nature Medicine (New York,* NY, United States), 20, 524–530.
- Deshpande, G., Rao, S., Athalye-Jape, G., Conway, P., & Patole, S. (2016). Probiotics in very preterm infants: The PiPS trial. *Lancet*, 388(10045), 655.
- Deshpande, G., Jape, G., Rao, S., & Patole, S. (2017). Benefits of probiotics in preterm neonates in low-income and medium-income countries: A

WILEY-Campbell Collaboration

systematic review of randomised controlled trials. *BMJ Open*, 7(12), e017638. https://doi.org/10.1136/bmjopen-2017-017638

- Faverge, B., Palix, C., & Coignet, J. (1985). Serum levels of vitamins A and E following the oral administration of Uvesterol. Studies in premature and term neonates. *Revue. de Pediatrie*, 21(5), 225–229.
- Gewolb, I. H., Schwalbe, R. S., Taciak, V. L., Harrison, T. S., & Panigrahi, P. (1999). Stool microflora in extremely low birthweight infants. Archives of Disease in Childhood. Fetal and Neonatal Edition, 80, F167–F173.
- Gogia, S., & Sachdev, H. S. (2009). Neonatal vitamin A supplementation for prevention of mortality and morbidity in infancy: systematic review of randomised controlled trials. *BMJ (Clinical Research Ed.)*, 338, b919. https://doi.org/10.1136/BMJ.B919
- Guyatt, G., Oxman, A. D., Akl, E. A., Kunz, R., Vist, G., Brozek, J., Norris, S., Falck-Ytter, Y., Glasziou, P., & deBeer, H. (2011). GRADE guidelines:1. Introduction–GRADE evidence profiles and summary of findings tables. *Journal of Clinical Epidemiology*, 64(4), 383–394.
- Güney-Varal, İ., Köksal, N., Özkan, H., Bağcı, O., & Doğan, P. (2017). The effect of early administration of combined multi-strain and multispecies probiotics on gastrointestinal morbidities and mortality in preterm infants: A randomized controlled trial in a tertiary care unit. *The Turkish Journal of Pediatrics*, 59(1), 13–19. https://doi.org/10. 24953/turkjped.2017.01.003
- Haider, B. A., & Bhutta, Z. A. (2011). Neonatal vitamin A supplementation for the prevention of mortality and morbidity in term neonates in developing countries. *Cochrane Database of Systematic Reviews*, 10, CD006980. https://doi.org/10.1002/14651858.CD006980.pub2
- Haider, B. A., Sharma, R., & Bhutta, Z. A. (2017). Neonatal vitamin A supplementation for the prevention of mortality and morbidity in term neonates in low and middle income countries. *Cochrane Database* of Systematic Reviews, 2, CD006980. https://doi.org/10.1002/ 14651858.CD006980.pub3
- Hegarty, J. E., Harding, J. E., Gamble, G. D., Crowther, C. A., Edlin, R., & Alsweiler, J. M. (2016). Prophylactic oral dextrose gel for newborn babies at risk of neonatal hypoglycaemia: A randomised controlled dose-finding trial (the Pre-hPOD Study). *PLOS Med*, 13(10), e1002155. https://doi.org/10.1371/journal.pmed.1002155
- Hegarty, J. E., Harding, J. E., Crowther, C. A., Brown, J., & Alsweiler, J. (2017). Oral dextrose gel to prevent hypoglycaemia in at-risk neonates. *Cochrane Database of Systematic Reviews*, 7, CD012152.
- Higgins, J. P. T. & Green, S. (Eds.). Cochrane handbook for systematic reviews of interventions version 5.1.0 (updated September 2011). The Cochrane Collaboration 2011. www.cochrane-handbook.org
- Imdad, A., Ahmed, Z., & Bhutta, Z. A. (2016). Vitamin A supplementation for the prevention of morbidity and mortality in infants one to six months of age. *Cochrane Database of Systematic Reviews*, 9, CD007480. https://doi.org/10.1002/14651858.CD007480.pub3
- Imdad, A., Mayo-Wilson, E., Herzer, K., & Bhutta, Z. A. (2017). Vitamin A supplementation for preventing morbidity and mortality in children from six months to five years of age. *Cochrane Database of Systematic Reviews*, 3, CD008524. https://doi.org/10.1002/14651858.CD008524.pub3
- Imdad, A., Ranjit, D., Surin, G. S., Lawler, S., Smith, A. A., & Bhutta, Z. A. (2019). Effects of neonatal nutrition interventions on neonatal mortality and child health and development outcomes: A systematic review. *Campbell Systematic Reviews*, 15(e1021).
- Jacobs, S. E., Tobin, J. M., Opie, G. F., Donath, S., Tabrizi, S. N., Pirotta, M., Morley, C. J., & Garland, S. M. ProPrems Study Group (2013). Probiotic effects on late-onset sepsis in very preterm infants: a randomized controlled trial. *Pediatrics*, 132(6), 1055–1062.
- Johnson-Henry, K. C., Abrahamsson, T. R., You Wu, R., & Sherman, P. M. (2016). Probiotics, prebiotics, and synbiotics for the prevention of necrotizing enterocolitis. Advances in Nutrition, 7(5), 928–937. https:// doi.org/10.3945/an.116.012237
- Kaiser, J. R., Bai, S., Gibson, N., Holland, G., Lin, T. M., Swearingen, C. J., Mehl, J. K., & ElHassan, N. O. (2015). Association between transient newborn hypoglycemia and fourth-grade achievement test

proficiency: A population-based study. JAMA Pediatr, 169(10), 913–921.

- Lee, AC, Kozuki, N, Cousens, S, Stevens, GA, Blencowe, H, & Silveira CHERG Small-for-Gestational-Age-Preterm Birth Working Group (2017). Estimates of burden and consequences of infants born small for gestational age in low and middle income countries with INTERGROWTH-21st standard: Analysis of CHERG datasets. *BMJ*, 358, j3677.
- McCullough, F. S., Northrop-Clewes, C. A., & Thurnham, D. I. (1999). The effect of vitamin A on epithelial integrity. *The Proceedings of the Nutrition Society*, 58(2), 289–293.
- McDonald, S. J., Middleton, P., Dowswell, T., & Morris, P. S. (2013). Effect of timing of umbilical cord clamping of term infants on maternal and neonatal outcomes. *Cochrane Database of Systematic Reviews*, 7, CD004074. https://doi.org/10.1002/14651858.CD004074.pub3
- McKinlay, C. J., Rebello, C., & Tarnow-Mordi, W. (2016). Probiotics in very preterm infants: The PiPS trial. *Lancet*, 388(10045), 655.
- McKinlay, C. J. D., Alsweiler, J. M., Anstice, N. S., Burakevych, N., Chakraborty, A., Chase, J. G., Gamble, G. D., Harris, D. L., Jacobs, R. J., Jiang, Y., Paudel, N., San Diego, R. J., Thompson, B., Wouldes, T. A., & Harding, J. E. (2017). Children with hypoglycemia and their later development (CHYLD) study team. Association of neonatal glycemia with neurodevelopmental outcomes at 4.5 years. JAMA Pediatric, 171(10), 972–983.
- Mercer, J. S., Vohr, B. R., McGrath, M. M., Padbury, J. F., Wallach, M., & Oh, W. (2006). Delayed cord clamping in very preterm infants reduces the incidence of intraventricular hemorrhage and late-onset sepsis: A randomized, controlled trial. *Pediatrics*, 117(4), 1235–1242.
- Mihatsch, W. A. (2011). Curr Opin Clin Nutr Metab Care. What is the power of evidence recommending routine probiotics for necrotizing enterocolitis prevention in preterm infants? *Current Opinion in Clinical Nutrition and Metabolic Care*, 14(3), 302–306.
- Mihatsch, W. A., Braegger, C. P., Decsi, T., Kolacek, S., Lanzinger, H., Mayer, B., Moreno, L. A., Pohlandt, F., Puntis, J., Shamir, R., Stadtmüller, U., Szajewska, H., Turck, D., & van Goudoever, J. B. (2012). Critical systematic review of the level of evidence for routine use of probiotics for reduction of mortality and prevention of necrotizing enterocolitis and sepsis in preterm infants. *Clinical Nutrition*, 31(1), 6–15.
- Millar, M., Wilks, M., & Costeloe, K. (2003). Probiotics for preterm infants? Archives of Disease in Childhood. Fetal and Neonatal Edition, 88, 354–358. https://www.ncbi.nlm.nih.gov/pmc/articles/PMC1721619/ pdf/v088p0F354.pdf
- Ofek Shlomai, N., Deshpande, G., Rao, S., & Patole, S. (2014). Probiotics for preterm neonates: what will it take to change clinical practice? *Neonatology*, 105(1), 64–70.
- Oza, S., Lawn, J. E., Hogan, D. R., Mathers, C., & Cousens, S. N. (2015). Neonatal cause-of-death estimates for the early and late neonatal periods for 194 countries. *Bulletin of the World Health Organization*, 93, 19–28.
- Panigrahi, P., Parida, S., Nanda, N. C., Satpathy, R., Pradhan, L., Chandel, D. S., Baccaglini, L., Mohapatra, A., Mohapatra, S. S., Misra, P. R., Chaudhry, R., Chen, H. H., Johnson, J. A., Morris, J. G., Paneth, N., & Gewolb, I. H. (2017). A randomized synbiotic trial to prevent sepsis among infants in rural India. *Nature Research*, 548(7668), 407–412.
- Patel, R. M., & Denning, P. W. (2015). Intestinal microbiota and its relationship with necrotizing enterocolitis. *Pediatric Research*, 78(3), 232–238. https://doi.org/10.1038/pr.2015.97
- Rao, S. C., Athalye-Jape, G. K., Deshpande, G. C., Simmer, K. N., & Patole, S. K. (2016). Probiotic supplementation and late-onset sepsis in preterm infants: A meta-analysis. *Pediatrics*, 137(3), e20153684. https://doi.org/10.1542/peds.2015-3684
- Review Manager (RevMan) [Computer program]. Version 5.3. The Cochrane Collaboration, 2019.
- Robinson, J. (2014). Cochrane in context: probiotics for prevention of necrotizing enterocolitis in preterm infants. *Evid Based Child Health*, 9(3), 672–674.

- Saigal, S., O'Neill, A., Surainder, Y., Chua, L. B., & Usher, R. (1972). Placental transfusion and hyperbilirubinemia in the premature. *Pediatrics*, 49(3), 406–419.
- Singhal, P. K., Singh, M., Paul, V. K., Malhotra, A. K., Deorari, A. K., & Ghorpade, M. D. (1991). A controlled study of sugar-fortified milk feeding for prevention of neonatal hypoglycaemia. *Indian Journal of Medical Research*, 94, 342–345.
- Singhal, P. K., Singh, M., Paul, V. K., Lamba, I. M., Malhotra, A. K., Deorari, A. K., & Ghorpade, M. D. (1992). Prevention of hypoglycemia: A controlled evaluation of sugar fortified milk feeding in small-forgestational age infants. *Indian Pediatrics*, 29(11), 1365–1369.
- Smith, E. R., Muhihi, A., Mshamu, S., Sudfeld, C. R., Noor, R. A., Spiegelman, D., Shapiro, R. L., Masanja, H., & Fawzi, W. (2016). The effect of neonatal vitamin A supplementation on morbidity and mortality at 12 months: A randomized trial. *International Journal of Epidemiology*, 145(6), 2112–2121.
- Stevens, G. A., Bennett, J. E., Hennocq, Q., Lu, Y., De-Regil, L. M., Rogers, L., Danaei, G., Li, G., White, R. A., Flaxman, S. R., Oehrle, S. P., Finucane, M. M., Guerrero, R., Bhutta, Z. A., Then-Paulino, A., Fawzi, W., Black, R. E., & Ezzati, M. (2015). Trends and mortality effects of vitaminA deficiency in children in 138 low-income and middle-income countries between 1991 and 2013: a pooled analysis of population-based surveys. *Lancet Global Health*, 3(9), e528–e536.
- Sutor, A. (1995). Vitamin K deficiency bleeding in infants and children. Seminars in Thrombosis and Hemostasis, 21(3), 317–329. https://doi. org/10.1055/s-2007-1000653
- Thompson-Branch, A., & Havranek, T. (2017). Neonatal hypoglycemia. *Pediatrics in Review*, 38(4), 147–157.
- Thornton, P. S., Stanley, C. A., De Leon, D. D., Harris, D., Haymond, M. W., Hussain, K., Levitsky, L. L., Murad, M. H., Rozance, P. J., Simmons, R. A., Sperling, M. A., Weinstein, D. A., White, N. H., & Wolfsdorf, J. I. Pediatric Endocrine Society (2015). Recommendations from the pediatric endocrine society for evaluation and management of persistent hypoglycemia in neonates, infants, and children. *Journal of Pediatrics*, 167(2), 238–245.
- US Institute of Medicine, Food and Nutrition Board (2000). Standing committee on the scientific evaluation of dietary reference intakes. dietary reference intakes for vitamin A, vitamin K, arsenic, boron, chromium, copper, iodine, iron, manganese, molybdenum, nickel, silicon, vanadium, and zinc. National Academy Press.
- Vallabhaneni, S., Walker, T. A., Lockhart, S. R., Ng, D., Chiller, T., Melchreit, R., Brandt, M. E., & Smith, R. M., Centers for Disease Control and Prevention (CDC). (2014). Notes from the field: Fatal gastrointestinal mucormycosis in a premature infant associated with a contaminated dietary supplement--Connecticut. Morbidity and Mortality Weekly Report, 20(64), 155–156.
- Wang, H., Bhutta, Z. A., Coates, M. M., Coggeshall, M., Dandona, L., Diallo, K., Franca, E. B., Fraser, M., Fullman, N., Gething, P. W., Hay, S. I., Kinfu, Y., Kita, M., Kulikoff, X. R., Larson, H. J., Liang, J., Liang, X., Lim, S. S., Lind, M., ... Damtew, S. A. (2015). Global, regional, national, and selected subnational levels of stillbirths, neonatal, infant, and under-5 mortality, 1980-2015: A systematic analysis for the Global Burden of Disease Study. *Lancet (London, England)*, 388(10053), 1725–1774.
- Wells, GA, Shea, B, O'Connell, D, Peterson, J, Welch, V, Losos, M, & Tugwell, P. (2017). The Newcastle-Ottawa Scale (NOS) for assessing the quality for nonrandomised studies in meta-analyses. Department of Epidemiology and Community Medicine, University of Ottawa, Canada. http://www.ohri.ca/ programs/clinical_epidemiology/oxford.asp
- West, K. P., Wu, L. S., Ali, H., Klemm, R. D. W., Edmond, K. M., Hurt, L., Kirkwood, B., Newton, S., Shannon, C., Taneja, S., Mazumder, S., Bhatia, K., Bhandari, N., Katz, J., Tielsch, J. M., Humphrey, J., Agoestina, T., Soofi, S. B., Ariff, S., ... Yoshida, S. (2019). Early neonatal vitamin A supplementation and infant mortality: An individual



participant data meta-analysis of randomised controlled trials. Archives of Disease in Childhood, 104(3), 217–226.

- Weston, P. J., Harris, D. L., Battin, M., Brown, J., Hegarty, J. E., & Harding, J. E. (2016). Oral dextrose gel for the treatment of hypoglycaemia in newborn infants. *Cochrane Database Systematic Reviews*, 5, CD011027. https://doi. org/10.1002/14651858.CD011027.pub2
- World Health Organization (2009a). Global prevalence of vitaminA deficiency in populations at risk 1995-2005. WHO global database on vitamin A deficiency.
- WHO (2009b). Technical consultation on neonatal vitamin A supplementation research priorities: Meeting report. Geneva, World Health Organization.
 2009. http://www.who.int/nutrition/publications/micronutrients/ vitamin_a_deficieny/NVAS_report.pdf
- WHO (2015). WHO recommendations on interventions to improve preterm birth outcomes. http://apps.who.int/iris/bitstream/handle/10665/ 183037/9789241508988_eng.pdf?sequence=1
- WHO (2017a). Newborn: reducing mortality. WHO 2017. http://www.who. int/mediacenter/factsheets/fs333/en
- WHO (2017b). Newborn care at birth. WHO 2017. http://www.who.int/ maternal_child_adolescent/newborns/care_at_birth/en/
- Wickström, R., Skiöld, B., Petersson, G., Stephansson, O., & Altman, M. (2018). Moderate neonatal hypoglycemia and adverse neurological development at 2-6 years of age. *European Journal of Epidemiology*, 33(10), 1011–1020.
- Williams, A. F. (1997). Hypoglycaemia of the newborn: A review. Bulletin of the World Health Organization, 75(3), 261–290.
- Wolbach, S. B. (1933). Epithelial repair in recovery from vitamin A deficiency: An experimental study. *Journal of Experimental Medicine*, 57(3), 511–526. https://doi.org/10.1084/jem.57.3.511
- World Bank. (2017). Low and middle income countries. Data: https://data. worldbank.org/income-level/low-and-middle-income

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APPENDIX A

Literature Search Strategy

Medline Strategy using PubMed

Vitamin A

(((("Vitamin A"[Mesh]) OR (Vitamin A[tiab] OR Aquasol A[tiab] OR Retinol[tiab] OR All Trans Retinol[tiab] OR All-Trans-Retinol[tiab] OR Vitamin A1[tiab] OR Vitamin A 1[tiab] OR 11-cis-Retinol[tiab] OR 11 cis Retinol[tiab] OR Tretinoin[tiab]) AND Supplement*[tiab])) AND (("Infant"[Mesh] OR "Premature Birth"[Mesh]) OR (Neonat*[tiab]) OR neo nat*[tiab]) OR (newborn* OR new Born*[tiab] OR newly born* [tiab]) OR (preterm[tiab] OR preterms[tiab] OR pre term[tiab] OR pre terms[tiab]) OR (premature*[tiab] AND (birth*[tiab] OR born[tiab] OR deliver*[tiab])) OR (low[tiab] AND (birthweight*[tiab]) OR birth weight*[tiab])) OR (lbw[tiab] OR vlbw[tiab] OR elbw[tiab]) OR infant* [tiab] OR (baby[tiab] OR babies[tiab])))) NOT ("Animals"[Mesh] NOT ("Animals"[Mesh] AND "Humans"[Mesh]))

Glucose

((((("Glucose"[Mesh]) OR (Dextrose OR Glucose[tiab]) AND supplement*)))) AND (("Infant"[Mesh] OR "Premature Birth"[Mesh]) OR (Neonat*[tiab] OR neo nat*[tiab]) OR (newborn* OR new Born* [tiab] OR newly born*[tiab]) OR (preterm[tiab] OR preterms[tiab] OR pre term[tiab] OR pre terms[tiab]) OR (premature*[tiab] AND (birth* [tiab] OR born[tiab] OR deliver*[tiab])) OR (low[tiab] AND (birth* [tiab] OR born[tiab] OR deliver*[tiab])) OR (low[tiab] OR vlbw[tiab] OR weight*[tiab] OR birth weight*[tiab])) OR (lbw[tiab] OR vlbw[tiab] OR elbw[tiab]) OR infant*[tiab] OR (baby[tiab] OR babies[tiab])))) NOT ("Animals"[Mesh] NOT ("Animals"[Mesh] AND "Humans"[Mesh])) **Probiotics**

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((((("Probiotics"[Mesh] OR "Prebiotics"[Mesh] OR "Synbiotics"[Mesh]) OR (Probiotic*[tiab] OR prebiotic*[tiab] OR synbiotic* [tiab]))) AND (("Infant"[Mesh]OR "Premature Birth"[Mesh]) OR (Neonat*[tiab] OR neo nat*[tiab]) OR (newborn* OR new Born*[tiab] OR newly born*[tiab]) OR (preterm[tiab] OR preterms[tiab] OR pre term[tiab] OR pre terms[tiab]) OR (premature*[tiab] AND (birth* [tiab] OR born[tiab] OR deliver*[tiab])) OR (low[tiab] AND (birth* [tiab] OR born[tiab] OR deliver*[tiab])) OR (low[tiab] OR vlbw[tiab] OR elbw[tiab]) OR infant*[tiab] OR (baby[tiab] OR babies[tiab])))) NOT ("Animals"[Mesh] NOT ("Animals"[Mesh] AND "Humans"[Mesh]))

CINAHL Strategies

Vitamin A

(MH"Vitamin A") OR TI ("Vitamin A" OR "Aquasol A" OR Retinol OR "All Trans Retinol" OR "All-Trans-Retinol" OR "Vitamin A1" OR "Vitamin A 1" OR "11-cis-Retinol" OR "11 cis Retinol" OR Tretinoin) OR AB ("Vitamin A" OR "Aquasol A" OR Retinol OR "All Trans Retinol" OR "All-Trans-Retinol" OR "Vitamin A1" OR "Vitamin A 1" OR "11-cis-Retinol" OR "11 cis retinol" OR Tretinoin)

AND

TI (Supplement*) OR AB (Supplement*) OR MH "Dietary Supplementation" OR MH "Dietary Supplements"

AND

(MH "Infant" OR MH "Infant, Premature" OR MH "Infant, Newborn") OR TI ((Neonat* OR neo nat*) OR (newborn* OR new Born* OR newly born*) OR (preterm OR preterms OR pre term OR pre terms) OR (premature* AND (birth* OR born OR deliver*)) OR (low AND (birthweight* OR birth weight*)) OR (lbw OR vlbw OR elbw) OR infant* OR (baby OR babies)) OR AB ((Neonat* OR neo nat*) OR (newborn* OR new Born* OR newly born*) OR (preterm OR preterms OR pre term OR pre terms) OR (premature* AND (birth* OR born OR deliver*)) OR (low AND (birthweight* OR birth weight*)) OR (lbw OR vlbw OR elbw) OR infant* OR (baby OR babies))

NOT

(MH "Animals" NOT (MH "Animals" AND MH "Humans")) Limiter: Exclude MEDLINE records

Glucose

(MH "Glucose") OR TI (Dextrose OR Glucose) OR AB (Dextrose OR Glucose)

AND

AND

TI (Supplement*) OR AB (Supplement*) OR MH "Dietary Supplementation" OR MH "Dietary Supplements"

(MH "Infant" OR MH "Infant, Premature" OR MH "Infant, Newborn") OR TI ((Neonat* OR neo nat*) OR (newborn* OR new Born* OR newly born*) OR (preterm OR preterms OR pre term OR pre terms) OR (premature* AND (birth* OR born OR deliver*)) OR (low AND (birthweight* OR birth weight*)) OR (lbw OR vlbw OR elbw) OR infant* OR (baby OR babies)) OR AB ((Neonat* OR neo nat*) OR (newborn* OR new Born* OR newly born*) OR (preterm OR preterms OR pre term OR pre terms) OR (premature* AND (birth* OR born OR deliver*)) OR (low AND (birthweight* OR birth weight*)) OR (lbw OR vlbw OR elbw) OR infant* OR (baby OR babies))

NOT

(MH "Animals" NOT (MH "Animals" AND MH "Humans")) Limiter: Exclude MEDLINE records

Probiotics

`(MH "Probiotics") OR (MH "Prebiotics") OR TI (probiotic* OR prebiotic* OR synbiotic*) OR AB (probiotic* OR prebiotic* OR synbiotic*) AND

(MH "Infant" OR MH "Infant, Premature" OR MH "Infant, Newborn") OR TI ((Neonat* OR neo nat*) OR (newborn* OR new Born* OR newly born*) OR (preterm OR preterms OR pre term OR pre terms) OR (premature* AND (birth* OR born OR deliver*)) OR (low AND (birthweight* OR birth weight*)) OR (lbw OR vlbw OR elbw) OR infant* OR (baby OR babies)) OR AB ((Neonat* OR neo nat*) OR (newborn* OR new Born* OR newly born*) OR (preterm OR preterms OR pre term OR pre terms) OR (premature* AND (birth* OR born OR deliver*)) OR (low AND (birthweight* OR birth weight*)) OR (lbw OR vlbw OR elbw) OR infant* OR (baby OR babies))

NOT

(MH "Animals" NOT (MH "Animals" AND MH "Humans")) Limiter: Exclude MEDLINE records

Scopus Strategies

Vitamin A

(TITLE-ABS("Vitamin A" OR "Aquasol A" OR retinol OR "All Trans Retinol" OR "Vitamin A1" OR "11-cis-Retinol" OR tretinoin)) AND (TITLE-ABS(Supplement*)) AND (TITLE-ABS ((neonat* OR "neo nat*") OR (newborn* OR "new born*" OR "newly born*") OR (preterm OR preterms OR "pre term" OR "pre terms") OR (premature*) AND (birth* OR born OR deliver*) OR (low AND (birthweight* OR "birth weight*")) OR (lbw OR vlbw OR elbow) OR infant* OR (baby OR babies))) AND NOT INDEX(medline)

Glucose

TITLE-ABS (Glucose OR Dextrose) AND TITLE-ABS (supplement*) AND TITLE-ABS ((neonat* OR "neo nat*") OR (newborn* OR "new born*" OR "newly born*") OR (preterm OR preterms OR "pre term" OR "pre terms") OR (premature*) AND (birth* OR born OR deliver*) OR (low AND (birthweight* OR "birth weight*")) OR (lbw OR vlbw OR elbow) OR infant* OR (baby OR babies)) AND NOT INDEX (medline)

Probiotics

TITLE-ABS (Probiotic* OR Prebiotic* OR Synbiotic*) AND TITLE-ABS ((neonat* OR "neo nat*") OR (newborn* OR "new born*" OR "newly born*") OR (preterm OR preterms OR "pre term" OR "pre terms") OR (premature*) AND (birth* OR born OR deliver*) OR (low AND (birthweight* OR "birth weight*")) OR (lbw OR vlbw OR elbow) OR infant* OR (baby OR babies)) AND NOT INDEX (medline)

Vitamin A

CENTRAL

- 1. 1 MeSH descriptor: [infant] explode all trees
- 2. MeSH descriptor: [Premature Birth] explode all trees
- 3. (Neonat*:ti,ab OR neo nat*:ti,ab OR (newborn*:ti,ab OR new Born*:ti,ab OR newly born*:ti,ab) OR (preterm:ti,ab OR preterms:ti,ab OR pre term:ti,ab OR pre terms:ti,ab) OR (premature*:ti,ab AND (birth*:ti,ab OR born:ti,ab OR deliver*:ti,ab)) OR (low:ti,ab AND (birthweight*:ti,ab OR birth weight*:ti,ab)) OR (lbw:ti,ab OR vlbw:ti,ab OR elbw:ti,ab) OR infant*:ti,ab OR (baby:ti,ab OR babies:ti,ab)
- 4. #1 OR #2 OR #3
- 5. MeSH descriptor: [Animals] explode all trees
- 6. MeSH descriptor: [Humans] explode all trees
- 7. (#5 NOT (#5 AND #6))
- 8. supplement*:ti,ab
- 9. MeSH descriptor: [Vitamin A] explode all trees
- "Vitamin A":ti,ab OR "Aquasol A":ti,ab OR Retinol:ti,ab OR "All Trans Retinol":ti,ab OR "All-Trans-Retinol":ti,ab OR "Vitamin A1":ti,ab OR "Vitamin A 1":ti,ab OR "11 cis Retinol":ti,ab OR "11-cis-Retinol":ti,ab OR Tretinoin:ti,ab
- 11. #9 OR #10
- 12. #11 AND #8
- 13. #12 AND #4 NOT #7
- 14. "accession number" near pubmed
- 15. #13 NOT #14

Glucose

- 1. 1 MeSH descriptor: [infant] explode all trees
- 2. MeSH descriptor: [Premature Birth] explode all trees
- 3. (Neonat*:ti,ab OR neo nat*:ti,ab) OR (newborn*:ti,ab OR new Born*:ti,ab OR newly born*:ti,ab) OR (preterm:ti,ab OR preterm:ti,ab OR preterm:ti,ab OR preterm:ti,ab) OR (premature*:ti,ab AND (birth*:ti,ab OR born:ti,ab OR deliver*:ti,ab)) OR (low:ti,ab AND (birthweight*:ti,ab OR birth weight*:ti,ab)) OR (lbw:ti,ab OR vlbw:ti,ab OR elbw:ti,ab) OR infant*:ti,ab OR (babies:ti,ab)
- 4. #1 OR #2 OR #3
- 5. MeSH descriptor: [Animals] explode all trees
- 6. MeSH descriptor: [Humans] explode all trees
- 7. (#5 NOT (#5 AND #6))
- 8. supplement*:ti,ab
- 9. MeSH descriptor: [Glucose] explode all trees
- 10. Dextrose:ti,ab OR Glucose:ti,ab
- 11. #9 OR #10
- 12. #11 AND #8
- 13. #12 AND #4 NOT #7
- 14. "accession number" near pubmed
- 15. #13 NOT #14

Probiotics

- 1. 1 MeSH descriptor: [infant] explode all trees
- 2. MeSH descriptor: [Premature Birth] explode all trees
- (Neonat*:ti,ab OR neo nat*:ti,ab) OR (newborn*:ti,ab OR new Born*:ti,ab OR newly born*:ti,ab) OR (preterm:ti,ab OR preterms:ti,ab OR pre term:ti,ab OR pre terms:ti,ab) OR (premature*:ti,ab AND (birth*:ti,ab OR born:ti,ab OR deliver*:ti,ab)) OR (low:ti,ab AND (birthweight*:ti,ab OR birth weight*:ti,ab)) OR (lbw:ti,ab OR vlbw:ti,ab OR elbw:ti,ab) OR infant*:ti,ab OR (baby:ti,ab OR babies:ti,ab)

Campbell_WILEY-

- 4. #1 OR #2 OR #3
- 5. MeSH descriptor: [Animals] explode all trees
- 6. MeSH descriptor: [Humans] explode all trees
- 7. (#5 NOT (#5 AND #6))
- 8. MeSH descriptor: [Probiotics] explode all trees
- 9. MeSH descriptor: [Prebiotics] explode all trees
- 10. MeSH descriptor: [Synbiotics] explode all trees
- 11. #8 OR #9 OR #10
- 12. Probiotic*:ti,ab OR prebiotic*:ti,ab OR synbiotic*:ti,ab
- 13. #11 or #12
- 14. #13 AND #4 NOT #7
- 15. "accession number" near pubmed
- 16. #14 NOT #15

LILACS

Vitamin A

(tw:(("Vitamin A"))) OR (ti:(("Aquasol A" OR retinol OR "All Trans Retinol" OR "Vitamin A1" OR "11-cis-Retinol" OR tretinoin))) OR (ab:(("Aquasol A" OR retinol OR "All Trans Retinol" OR "Vitamin A1" OR "11-cis-Retinol" OR tretinoin))) AND (ti:(supplement*)) OR (ab:(supplement*)) AND (tw:(Infant)) OR (tw:("Premature Birth")) OR (ti:(((neonat* OR "neo nat*") OR (newborn* OR "new born*" OR "newly born*") OR (preterm OR preterms OR "pre term" OR "pre terms") OR (premature*) AND (born OR deliver*) OR (low AND (birthweight* OR "birth weight*")) OR (lbw OR vlbw OR elbw) OR (baby OR babies)))) OR (ab:(((neonat* OR "neo nat*") OR (newborn* OR "new born*" OR "newly born*") OR (preterm OR preterms OR "pre term" OR "pre terms") OR (premature*) AND (born OR deliver*) OR (low AND (birthweight* OR "birth weight*")) OR (lbw OR vlbw OR elbw) OR "pre term" OR "pre terms") OR (premature*) AND (born OR deliver*) OR (low AND (birthweight* OR "birth weight*")) OR (lbw OR vlbw OR vlbw OR elbw) OR (baby OR babies)))) AND db:("LILACS")

Glucose

((tw:(glucose)) OR (ti:(dextrose)) OR (ab:(dextrose)) AND (ti:(supplement*)) OR (ab:(supplement*))) AND ((tw:(infant)) OR (tw:("premature birth")) OR (ti:((neonat* OR "neo nat*") OR (newborn* OR "new born*" OR "newly born*") OR (preterm OR preterms OR "pre term" OR "pre terms") OR (premature*))) AND (ti:((born OR deliver*) OR (low AND (birthweight* OR "birth weight*")) OR (lbw OR vlbw OR elbw) OR (baby OR babies))) OR (ab:((neonat* OR "neo nat*") OR (newborn* OR "new born*" OR "newly born*") OR (preterm OR preterms OR "pre term" OR "pre terms") OR (premature*))) AND (ab:((born OR deliver*) OR (low AND (birthweight* OR "birth weight*")) OR (lbw OR vlbw OR elbw) OR (baby OR babies)))) AND (instance:"regional") AND (db:("LILACS"))

Probiotics

((tw:(probiotics OR prebiotics OR synbiotics)) OR (ti:(probiotic* OR prebiotic* OR synbiotic*)) OR (ab:(probiotic* OR prebiotic* OR synbiotic*))) AND ((tw:(infant)) OR (tw:("premature birth")) OR (ti:((neonat* OR "neo nat*") OR (newborn* OR "new born*" OR "newly born*") OR (preterm OR preterms OR "pre term" OR "pre terms") OR (premature*))) AND (ti:((born OR deliver*) OR (low AND (birthweight* OR "birth weight*")) OR (lbw OR vlbw OR elbw) OR (baby OR babies))) OR (ab:((neonat* OR "neo nat*") OR (newborn* OR "new born*" OR "newly born*") OR (preterm OR preterms OR "pre term" OR "pre terms") OR (premature*))) AND (ab:((born OR deliver*) OR (low AND (birthweight* OR "birth weight*")) OR (lbw OR vlbw OR vlbw OR elbw) OR (baby OR babies)))) AND (instance:"regional") AND (db:("LILACS"))

EMBASE

Vitamin A

- 1 'retinol'/exp OR 'retinol palmitate'/exp OR '11 cis retinol'/exp OR 'retinoic acid'/exp
- 'vitamin a':ti,ab OR 'aquasol a':ti,ab OR retinol:ti,ab OR 'all trans retinol':ti,ab OR 'all-trans-retinol':ti,ab OR 'vitamin a1':ti,ab OR

- 3. supplement*:ti,ab
- 4. 'supplementation'/exp
- 5. #1 OR #2
- 6. #3 OR #4
- 7. #5 AND #6
- 8. 'infant'/exp OR 'prematurity'/exp OR 'newborn'/exp OR 'low birth weight'/exp OR 'very low birth weight'/exp OR 'extremely low birth weight'/exp OR 'premature labor'/exp
- 9. neonat*:ti,ab OR 'neo nat*:ti,ab OR newborn*:ti,ab OR 'new born*':ti,ab OR 'newly born*':ti,ab OR preterm:ti,ab OR 'pre terms:ti,ab OR 'pre terms':ti,ab OR 'pre terms':ti,ab OR (premature*:ti,ab AND (birth*:ti,ab OR born:ti,ab OR deliver*:ti,ab)) OR (low:ti,ab AND (birthweight*:ti,ab OR 'birth weight*':ti,ab)) OR lbw:ti,ab OR vlbw:ti,ab OR elbw::ti,ab OR infant*:ti,ab OR babies:ti,ab
- 10. #8 OR #9
- 11. #7 AND #10
- 12. #11 NOT ([animals]/lim NOT [humans]/lim)
- 13. #12 NOT [medline]/lim