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Human breast milk-based nutritherapy: A blueprint for pediatric healthcare

Pravin Shende*, Bhakti Khanolkar

Shobhaben Pratapbhai Patel School of Pharmacy and Technology Management, SVKM’s NMIMS, V.L. Mehta Road, Vile Parle (W), Mumbai, India

Abstract

Human Breast Milk (HBM) is a storehouse of micronutrients, macronutrients, immune factors, microbiota and numerous other bioactive macromolecules. Fulfillment of optimum nutritional requirements of more than 240 million malnourished infants worldwide is possible via adequate amount (570-900 mL/d) of breast milk administration to infants in first few years of life. Technological advancements enable study of multiple components of HBM like stem cells, bioactive proteins, micro RNAs, immunoglobulins and epithelial cells to understand their role in enhancement of nutritional value of HBM. Furthermore, immunological and protective functions of HBM against various illnesses like diabetes, anemia, respiratory and cardiovascular abnormalities, otitis media and gastrointestinal diseases prove superiority of HBM over artificial milk. Presence of major macronutrients like fatty acids, sphingomyelins, proteins, peptides, lactoferrin, lactalbumins, lysozymes, mucins, growth factors, oligosaccharides and cytokines increase nutritive value of HBM. In the future, HBM can serve as a carrier for delivery of drugs, vaccines and genes to infants and offer novel therapeutic applications to stimulate effective health, growth and development of infants. The review article highlights multimodal nutritional benefits of HBM, provides insight into preclinical and clinical studies of HBM-based therapeutics and encourages further research on HBM therapy to suffice nutritional needs of infants.

Keywords: Clinical studies, Human breast milk, Nutritherapy, Pediatric nutrition, Therapeutic applications

1. Introduction

Adequate nutrition (60-95 g/d carbohydrates, 9.1-11 g/d proteins, 30-31 g/d fats 200-1000 IU, 10 μg-10 mg/d vitamins and 0.27-40 mg/d iron) for infants is essential for their optimal growth and development, whereas poor nutrition enhances risk of multiple diseases and disorders like throat infections, bronchitis, skin diseases, bacterial sinusitis, etc. [1]. Globally, malnutrition is the cause of 45% of deaths in children (<5 years-of-age) and its severe social, economic and health effects pose long-lasting negative impact on many communities and countries. World Health Organization (WHO) recommends prioritization of breastfeeding (administration of HBM) to infants (<6 months-of-age), followed by complementary foods (for infants who are 6-23 months-of-age) to overcome malnutrition [2]. In addition to supply of nutrition, breast milk (BM) shows potential to save lives of 820,000 infants (<5 years-of-age) per year, worldwide [3]. HBM consists of a spectacular array of molecular and cellular components to facilitate optimum nutrition, growth and development of infants [4]. Colostrum renders multitude of nutritional, bioactive and immune factors to provide well-balanced nutrition and protect infants against pathogens. Full-term HBM (THBM) provides components like fats (stimulate cognitive functions), peptides such as ghrelin (controls appetite for long-time), sugars such as oligosaccharides (act against pathogens and increase the production of beneficial bacteria) and...
more than 900 types of proteins (for protective functions) to promote adequate nutrition in infants [5]. HBM is a dynamic fluid consisting of multiple vitamins (A, D, B12, B2, B6 and iodine), minerals, microbiota, micro RNAs (miRNAs), immunoglobulins, antibodies, enzymes, bioactive molecules and viable cells (stem cells, immune cells and milk-producing cells) to suffice nutritional requirements of infants [6]. The advancement in drug, vaccine and gene delivery technologies for administration of medicines to infants via HBM cells [7] will ensure specific therapeutic effect at desirable sites of action with lesser adverse events and more biocompatible nature. The present review article focuses on manifold benefits of HBM-based nutritherapy in the treatment of pediatric illnesses and explores various possibilities for modulation of breast milk components for HBM-based drug delivery to infants.

2. Health burden associated with pediatric nutrition

Malnutrition in infants is a major contributor to mortality of infants as they are more susceptible to infections, immune-related disorders (hypersensitivity reactions, neoplasia, eczema, contact dermatitis) and other respiratory and gastrointestinal illnesses [2]. Since the complications of chronic malnutrition in infants are irreparable, addressing intergenerational transmission of malnutrition within first two years (after birth) is essential. Triple burden of malnutrition (undernutrition, overweight and hidden hunger) is harmful to children as well as mothers [3]. Undernutrition includes stunting and wasting which results in poor growth and cognition, susceptibility to infections, perinatal complications, chronic diseases and low birth weight of infants. The overweight and obese infants are prone to diabetes, cardiovascular problems, metabolic disorders and obstetric complications [1,3]. Deficiency of micronutrients is the cause for hidden hunger that manifests neural tube abnormalities in neonates, maternal morbidity and mortality, impaired cognition, poor immunity and improper tissue and organ development [8]. According to United Nations Children's Fund [3] around 149 million children (<5 years-of-age) are stunted, 50 million wasted and 40 million are overweight, where more than 1 in 3 children experience improper growth due to inadequate nutrition. Globally, 21.3% (<5 years-of-age) children suffer from stunting, 6.9% from wasting and severe wasting and 5.6% from obesity [9].

Children with short height endure weight loss in early childhood and fail to attain adequate size (Body Mass Index: 18.5-24.9) as adults, resulting in lesser physical capacity to carry out day-to-day activities, lower IQ (<90 score) and higher susceptibility to infections as compared to normal children (who receive adequate neonatal nutrition) [10]. Breastfeeding exhibits numerous benefits for infants especially in the first hour after birth, therefore, such infants show lower mortality and morbidity rates in comparison to non-breastfed and late-breastfed infants [11]. Despite such prominent benefits of administration of HBM, only 44% (two out of five) infants are breast-fed in the first hour (after birth) and 42% infants (<6 months-of-age) are exclusively breastfed [9]. Increase in promotion of breast milk substitutes (BMS), the presence of cultural and social taboos against breastfeeding, lack of balance between work and childcare by mothers and insufficient support for breastfeeding are the main reasons for lower rate of breastfeeding to infants universally [12]. Low birth weight (below 2.5 kg body weight), very low birth weight (1-1.5 kg) and extremely low birth weight (below 1 kg) infants suffer from serious nutritional deficiencies due to immature development of tissue and organ systems. Nutrient absorption from gastrointestinal tract (GIT) and digestion and absorption of fat-soluble vitamins, trace elements and fats are primarily damaged in such infants [13]. Brain, kidney and liver are vulnerable to detrimental effects of insufficient nutrition, followed by impairment of overall growth and development of infants. Energy requirements are more in immature infants as their needs of incidental cold stress, resting metabolic rates and losses of unabsorbed nutrients are higher [14]. Insufficient amount and lower quality of proteins administered to low birth weight (LBW) infants can impair their renal functions. Disruption in maintenance of water and energy balance, insufficient supply of optimum amounts of proteins, vitamins and minerals and dysregulation of calcium-phosphorus homeostasis are some deleterious effects of inadequate administration of macronutrients (in form of HBM) to infants [15].

3. Therapeutic applications of HBM

HBM is classified into three types: 1. Colostrum, 2. Transitional milk and 3. Mature milk, where each type possesses distinct characteristics (colostrum: composed of minerals, proteins, vitamins and
immunoglobulins, transitional milk: contains proteins and calories and mature milk: consists of 90% water and 10% proteins, fats and carbohydrates. The composition of HBM changes during the course of lactation period to facilitate effective nutritional requirements for infants at all ages (Fig. 1) [16]. Around 87% water, 7% lactose, 4% fat, 1% proteins and small amounts of carbohydrates and vitamins are nutritional components of HBM, whereas its non-nutritional constituents are hormones and growth factors [15]. In addition to supply of nutrition, milk therapy using HBM demonstrates immunological and protective properties to protect infants from *H. pylori* infections, immunodeficiency-related diarrhoea, anemia, irregular tissue growth, respiratory and cardiovascular abnormalities, enteral nervous system and autoimmune disorders, inadequate intestinal maturation and gastrointestinal diseases. Furthermore, HBM plays an important role where infants cannot acquire sufficient nutrition such as in low, middle-income and developing countries [17]. Multiple bioactive and immunomodulatory elements (antibodies and immunoglobulins) of HBM like miRNAs, cellular components (macrophages, T-cells and leukocytes), progenitor cells, growth-regulating hormones (somatostatin and calcitonin) epithelial cells and growth factors like vascular endothelial growth factor (VEGF), brain-derived neurotrophic factor (BDNF), glial cell-line-derived neurotrophic factor (GDNF), etc. show significant nutritional benefits in preclinical studies for adequate growth and development of infants [18,19]. Fats (long, short and medium-chain fatty acids (FA) and sphingomyelins), proteins (casein, osteopontin, peptides, cytokines, amylase, growth factors and haptocorrin), whey (alpha-lactalbumin, lactoferrin, lipases, lysozyme, sIgA and mucins) and carbohydrates (oligosaccharides and lactose) in HBM (Table 1) supplement energy, modulate immune system, neutralize toxins, absorb vitamins, stimulate brain and neural development, promote absorption of calcium, iron and phosphorus, inhibit pathogen attack and protect infants from antimicrobial and antiviral infections [6]. The supply of HBM to infants via breastfeeding is endorsed by WHO to render optimum nutrition for health and well-being of very low birth weight (VLBW) and preterm infants. Administration of HBM reduces hospitalization cost as it protects infants against healthcare-associated and community-related diseases like necrotizing enterocolitis (NEC), otitis media, neonatal sepsis, gastroenteritis and respiratory infections [15].

HBM offers several benefits to mothers like prevention of postpartum hemorrhage, amelioration of birth spacing and reduction in risk of ovarian and breast cancers [17]. Preterm HBM (PHBM) and THBM-derived exosomes show presence of peptides (lactadherin and lactotransferrin) and oligosaccharide 2’-fucosyllactose that protect infants against gastrointestinal diseases. HBM exosomes exhibit multiple beneficial effects like reduction in obesity and exacerbations of asthma, minimization of interleukin (IL)-2 and Tumor Necrotic Factor (TNF-α) production and activation of T-regulatory cells [20]. Lactoferrin (glycoprotein) present in HBM possesses antimicrobial and immunomodulatory characteristics that protect infants against multiple infections by three mechanisms: 1. Modulation of growth of bacteria in GIT, 2. Promotion of

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**Fig. 1. Variability of nutritional components in: a. Colostrum, b. Transitional HBM and c. Mature/Term HBM. [sIgA: secretory Immunoglobulin A, EGF: Epidermal Growth Factor, VEGF: Vascular Endothelial Growth Factor, IGF: Insulin-like Growth Factor].**
proliferation of intestinal cells and 3. Differentiation, maturation and regulation of host-cell responses against pathogens [21]. HBM contains large quantities of immunoglobulin A (IgA) antibodies that demonstrate preventive effects against immune-mediated diseases and bacterial skin infections by inhibiting the harmful microorganisms from penetrating the tissues and organs of infants [22]. Two categories of growth factors of HBM: 1. Transforming Growth Factors (TGF-α and TGF-β) and 2. Insulin-like Growth Factors (IGF-1 and IGF-2) stimulate wound healing and repair of cartilage and muscles. TGF-α and TGF-β promote activities like cell proliferation, tissue and muscle repair, embryonic development and musculoskeletal growth, whereas IGF-1 induces anabolism, reduces catabolism and augments wound-healing properties [23].  

Table 1. Functions of fats, proteins, whey and carbohydrates in HBM.

| Components of HBM | Functions |
|-------------------|-----------|
| Fats              | Long-chain FA 1. Visual and neural development of infants 2. Modulation of immune system 3. Antiprotozoal and antiviral activities |
|                   | Short-chain FA 1. Source of energy 2. Maturation of GIT |
|                   | Medium-chain FA 1. Promotes peripheral glucose utilization 2. Source of energy |
|                   | Sphingomyelins 1. Myelination of central nervous system 2. Visual development of infants |
| Whey              | Lactoferrin 1. Protects against iron-dependent pathogens 2. Antimicrobial activity |
|                   | Secretory IgA 1. Antipathogenic effect 2. Neutralizes toxins and viruses |
|                   | Lysozyme 1. Supports preterm growth of infants 2. Bactericidal and bacteriostatic effects 3. Promotes growth of commensal bacteria |
|                   | α-Lactalbumin 1. Immunoprotective properties 2. Gut maturation and development 3. Promotes absorption of amino acids 4. Lactose synthesis |
|                   | Bile salt-stimulated lipase 1. Infant growth 2. Digestion of fats |
|                   | Mucins Inhibit binding of pathogens |
| Carbohydrates     | Oligosaccharides 1. Antimicrobial, anti-infective and anti-adhesive properties 2. Prebiotic for gut colonization 3. Brain development in infants |
|                   | Lactose 1. Major source of energy 2. Prebiotic for gut colonization 3. Calcium absorption |
|                   | Proteins Casein 1. Source of calcium and phosphorous 2. Provide faster gastric transit than artificial formula milk |
|                   | Peptides Antihypertensive, antimicrobial, Antithrombotic and immunomodulatory activities |
|                   | Osteopontin 1. Immunomodulatory effects 2. Gut barrier functions |
|                   | Amylase 1. Antibacterial activity 2. Digestion of polysaccharides |
|                   | Cytokines 1. Anti-inflammatory effects 2. Anti-infective functions |
|                   | Growth factors 1. Promote intestinal growth and maturation of intestinal mucosa of infants 2. Anti-inflammatory effects 3. Stimulate growth of cells 4. Regulate development of multiple tissues and organs 5. Reduce effects of hypoxia, NEC, hemorrhagic shock, etc. 6. Develop neuronal system and enhance gut peristalsis 7. Enhance hematocrit level and stimulate erythropoiesis |

Cells of HBM are majorly divided into two classes: 1. Probiotic bacteria and 2. BM cells. Breast-derived cells (myoepithelial cells, lactocytes, stem cells and...
progenitor cells) and blood-derived cells (immunological and hemopoietic stem cells) are two major subclasses of BM cells (Fig. 2) [24]. Around $10^7 - 10^8$ bacterial cells in 800 mL of HBM are ingested by infants that help in development of their strong immune system with effective resistance against infectious diseases. Probiotic bacteria (*Bifidobacteria* and *Lactobacilli*) in HBM facilitate establishment of microbiome in infants, enhance nutritive value of HBM and enable use of HBM as an alternative to antibacterials and other antibiotics for the treatment of infections (mastitis and atopic dermatitis) in infants [25,26]. HBM is identified as a source for differentiation of neural cell lineage, thus indicating its involvement in the development of enteric nervous system which regulates the functions of GIT in infants. Human Breast Milk Stem Cells (hBSCs) exhibit application in regenerative medicine and assist in epigenetic regulation and development of tissues in infants. Easy and accessible harvesting of hBSCs via non-invasive techniques and negligible tumorigenic potential indicate their future use in autologous transplantation of tissues and organs in infants [27]. Useful constituents in HBM like probiotic and lactic acid bacteria, *Bifidobacteria*, Human Alpha-Lactalbumin Made Lethal To Tumour (HAMLET) cells, commensal bacteria and stem cells render natural antibacterial, anti-inflammatory and immunomodulatory properties to treat manifold skin problems (cuts, wounds, lesions and scrapes), skin irritations and allergies in infants [17].

Among nutrients present in HBM, triacylglycerides (TAGs) show highest amount of variability throughout lactation period (8 d post-partum to 6 months). Palmitin-Olein-Olein (POO) and Palmitin-Olein-Linolein (POL) are two significant TAGs (>49%) found in colostrum, wherein maternal factors like age, weight, gestational period, etc. do not affect concentrations of TAGs in HBM [16]. HBM reduces risk of sepsis, retinopathy and urinary tract infections, strengthens visual development in infants, promotes proliferation of diverse and well-balanced microbiota and suppresses immune-mediated complications like type 1 diabetes, inflammatory bowel disease and asthma [28]. Defensins and lactoferrin (in HBM) block the entry of harmful pathogens, preserve microbiota composition and promote action of good bacteria (*Lactobacillus, Bacteroides, and Clostridia*). The good bacteria are imperative in activation of neonatal immunologic functions like tolerance, maintenance of T-helper cell balance, mucus production, development of mucosal barrier homeostasis and promotion of tight junction expression [29]. IgA and IgG produced by B-cells in HBM exhibit antimicrobial protection against number of pathogens and post-natal virus acquisitions. HBM depicted significant neutralization potency for monoclonal

![Cells of Human Breast Milk](image_url)
antibodies (present in colostrum), anti-HIV-1 activity and inhibition of HIV transmission across mucosal barrier [30]. Free Secretory Component (FSC), an element of secretory IgA (slgA) found in HBM, performs immune exclusion activity against pathogens to treat neonatal infections. FSC and slgA inhibit attachment of pathogens on intestinal epithelial cells and neutralize their toxic effects in neonatal gut. Development of innate and adaptive immunities in infants, maintenance of integrity of intestinal epithelial barrier, regulation of mucosal homeostasis and inhibition of adhesion of Enteropathogenic Escherichia coli (EPEC) to epithelial cells suggest possible therapeutic effects of FSC for treatment of diarrhoea in infants [31]. HBM improves enteral feeding of preterm neonates, lowers incidences of vomiting and abdominal discomfort and shortens gastric residual time in comparison with formula-fed infants [17].

MicroRNAs are short (22 nucleotides) and fragile strands present in exosomes (extracellular vesicles) to regulate gene expression in humans. Researchers identified more than 1,400 miRNAs in HBM and suspected that they play a key role in modulating prominent aspects of child development like nutritional requirements and immune functions [18,32]. Evaluation of composition of PHBM revealed presence of higher concentration of metabolic miRNAs and macronutrients (protein and fats), suggesting that constitution of miRNA in HBM changes to help infants grow rapidly [33]. miR-335, miR-155, miR-223, miR-375 and several other types of miRNAs are present abundantly in HBM and are responsible for epigenetic growth, development of immune and neuronal systems, reduction in allergies and prevention of autoimmune diseases in infants [17]. HBM miRNA (miR-148a-3p) acts as a biomarker for the preparation of artificial infant formulae, examination of breast aberrations, standardization and quality control of raw milk processing in dairy industry and evaluation of health status of lactating mother [34]. HBM miRNAs can be administered to infants via uptake of BM cells, exosomes and microvesicles or as free molecules in skim milk. Absorption of miRNAs will occur via gastrointestinal epithelial cells, followed by transport to tissues, organs and systemic circulation to perform immunoprotective and developmental programming functions [35]. Antioxidant capacity of HBM exhibited by bioactive proteins like lactoferrin, adiponectin, leptin and lysozyme indicate that it is an ideal source of nutrition for preterm infants. Antioxidant properties of HBM help to reduce oxidative stress-induced complications and reinforce immature antioxidant defense system of infants [4].

3.1. Pre-clinical studies

P-glycoprotein (P<sub>gp</sub>) deficiency causes NEC in newborn mice, whereas P<sub>gp</sub> induction protects epithelial cells from apoptosis mediated by pathogens. In a preclinical study, administration of HBM to newborn mice and rats resulted in increase of intestinal P<sub>gp</sub> levels on days 1-5 and 3-7, respectively. HBM stimulated P<sub>gp</sub>-inducing factors like Epidermal Growth Factor (EGF), oligosaccharides, lactoferrin and heparin-binding EGF for potentiation of P<sub>gp</sub> expression in intestine and enterocyte cells of neonatal rodents to protect them from NEC [36]. In another preclinical study by Dai et al. [37] Human Milk Growth Factor III (HMGF III) derived from HBM demonstrated similar biological and chromatographic properties like EGF-urogastrene and reduced severity, duration and number of incidences of duodenal ulcer formation in CD-1 mice. A dose of 32 mg/kg of HMGF III resulted in substantial reduction of ulcer index (2.77 to 0.68), incidence of ulcers (62.50% to 18.18%) and severity score (1.52 ± 0.33 to 0.32 ± 0.228).

3.2. Clinical studies

Bio-active proteins in HBM show antioxidant properties that are beneficial for supply of adequate nutrition to preterm-born infants. The findings from clinical study conducted by Mehta et al. revealed that lactoferrin, leptin, lysozyme and adiponectin levels affect the Total Antioxidant Capacity (TAC) of HBM. Analysis of 60 HBM samples from 15 women (21-43 years-of-age) showed significant effect of the proteins on HBM TAC (R² = 0.635 ± 0.102, p < 0.0001). Furthermore, higher concentrations of slgA, adiponectin and lysozyme in transitional HBM and PHBM demonstrated their superior antibacterial and anti-inflammatory activities in comparison to THBM [38]. Another clinical study revealed that PHBM rendered more nutritional and antimicrobial benefits (higher quantities of immunoglobulin A, sodium, potassium, chloride, lysozyme, proteins and lower quantities of fats, potassium and carbohydrates than THBM after first month of parturition. Levels of OPG, leptin and lactoferrin were higher in THBM (p < 0.05-0.0001) as compared to PHBM, indicating that THBM is imperative for neuroendocrine regulation [39]. Study of HBM-administered (n = 24) vs formula-fed infants (n = 15) depicted that IgA, lactoferrin and lysozyme content of HBM renders anti-infective and protective properties to infants. At 1 month (p < 0.025) and 7 months (p < 0.025) after discharge, infants who received HBM exhibited upper respiratory symptoms for fewer days in comparison to formula-fed
infants [40]. Analysis of HBM (collected from 35 mothers) using filter-based mid-infrared (MIR) spectroscopy rendered quantification of macronutrient composition (amount of fats, lactose and proteins) of HBM. In comparison to Kjeldahl, Mojonnier and High-Performance Liquid Chromatography (HPLC) methods used for determination of proteins, fats and lactose, respectively, MIR spectroscopy emerged as easy and rapid point-of-care test (POCT) for evaluation of nutrients in HBM [41]. A clinical trial involving 1738 infants indicated that length of treatment and re-hospitalization rate for infants suffering from neonatal abstinence syndrome (NAS) were significantly decreased with administration of HBM (n = 430) in comparison to formula-fed infants (n = 1308) [42]. In another clinical study, administration of at least 50% HBM to VLBW infants (n = 202) for first two weeks (after birth) resulted in six-fold reduction of NEC ((OR = 0.17, 95% CI: 0.04 to 0.68), p = 0.01) [43]. Deep sequencing technology-based study of 602 miRNAs originating from pre-miRNAs (precursors of miRNAs) revealed that about 67.82% of immune-related miRNAs are expressed in HBM exosomes. The miRNAs are transferred from HBM to GIT of infants and perform critical role in development of strong immune system. Unsuitable conditions like RNase-mediated digestion, prolonged incubation of miRNAs at room temperature and exposure to multiple freeze-thaw cycles did not affect stability and functions of the miRNAs [44]. DNA samples were extracted from HBM and subjected to amplification to examine clinical use of DNA in pharmacogenetic studies. The experiment revealed that unpasteurized HBM served as an alternative to blood for pharmacogenetic analysis and suggested enhancement in administration of HBM to infants for improving their nutritional requirements [45]. Data of clinical and preclinical studies using HBM is illustrated in Table 2.

4. Future implications of pediatric drug delivery via HBM

Drug delivery to pediatric population demonstrates considerable variability in age, weight,

| Sr. No. | Objective | Study design | Result | Interpretation | Reference |
|---------|-----------|--------------|--------|----------------|-----------|
| Pre-clinical studies |
| 1. | To understand HBM-induced Pgp expression for protection against NEC in neonates | Postpartum Sprague-Dawley rats and newborn rats were studied. 1. HBM was obtained from healthy volunteers and centrifuged to remove insoluble matter and lipids | 1. Low Pgp levels were identified as cause of NEC in newborn rodents. 2. Intestinal epithelium of rodents was protected against bacterial colonization and NEC due to supply of HBM. | Administration of HBM decreases the risk of NEC in newborns and protects against NEC. | [36] |
| 2. | To study effect of HBM-derived HMGF-III on duodenal ulcers induced in mice | Duodenal ulcers in male CD-1 mice were induced by cysteamine HCl, followed by treatment with HMGF-III | Number of incidences, ulcer index and severity score of duodenal ulcers in CD-1 mice were reduced | HMGF-III exhibited protective effect on cysteamine-induced duodenal ulcers in CD-1 mice. | [37] |
| Clinical Trials |
| 1. | To study the effect of bio-active proteins on TAC of HBM | 1. PHBM samples: 60 2. THBM samples: 20 | 1. Content of bio-active proteins affects the TAC of HBM (R² = 0.635 ± 0.102, p = 0.001) 2. Higher levels of adiponectin, IgA and lysozyme in PHBM 3. Higher level of OPG, leptin and lactoferrin in THBM | 1. Antioxidant activity of HBM is enhanced due to presence of bio-active proteins, thus rendering HBM as an ideal nutritive supplement for infants 2. PHBM exhibited higher levels of proteins (adiponectin, IgA and lysozyme) that show anti-inflammatory and antibacterial activities 3. PHBM showed lower level of leptin indicating improper neuroendocrine regulation | [38] |

(continued on next page)
disease condition, health status, degree of physiological and psychological development, etc. [3]. Differences in transportation and metabolism of drugs and lack of safer and biocompatible drug-carrier systems for pediatric patients generate the need for HBM-based drug carriers. Bovine milk-derived exosomes serve as carriers for lipophilic and hydrophilic drugs (withaferin, anthocyanidin, docetaxel, curcumin and paclitaxel), anti-cancer agents and small molecules for efficacious activity.

Table 2. (continued)

| Sr. No. | Objective | Study design | Result | Interpretation | Reference |
|---------|-----------|--------------|--------|----------------|-----------|
| 2.      | To analyze effect of content of bioactive proteins in PHBM and THBM | 1. PHBM samples: 60 2. THBM samples: 20 | 1. Reduction in levels of OPG, lysozyme, adiponectin and lactoferrin, adiponectin was observed but no change in SgA content was reported in PHBM and THBM in the first month of lactation. 2. Levels of OPG, leptin and lactoferrin were higher in THBM ($p < 0.05$-$0.0001$) as compared to PHBM | PHBM renders better antimicrobial and anti-inflammatory properties than THBM, whereas PHBM helps in neuroendocrine regulation of infants | [39] |
| 3.      | To examine effect of HBM in reduction of symptoms of infections | 24 infants: Administered HBM 15 infants: Administered milk formula | Decrease in duration of upper respiratory in infants after one month post discharge ($p < 0.025$) and at seven months ($p < 0.025$) | HBM reduces symptoms of upper respiratory infection in infants in first year after birth | [40] |
| 4.      | To validate MIR spectroscopy for analysis of macronutrients in HBM | HBM samples collected from 35 mothers to assess quantities of fats, lactose and proteins | ICC for fats: 0.997, lactose: 0.776 and proteins: 0.839 | MIR spectroscopy is an effective method for analysis of macronutrients in HBM | [41] |
| 5.      | To assess benefit of HBM (administered to infants) in shortening duration of pharmacological treatment for NAS and length of hospitalization of infants | 1738 HBM-fed infants analyzed. Infants were divided into 3 main groups: 1. No HBM group, 2. HBM group and 3. Formula-fed/any BM group | Median duration of treatment and hospitalization was found to be lower in HBM-fed infants than no HBM-fed infants | HBM decreased duration of pharmacological treatment for NAS and hospitalization stay | [42] |
| 6.      | To test whether administration of HBM (≥50%) for first 14 days after birth is protective against NEC in VLBW infants | 222 VLBW infants divided into 2 groups: 1. <50% HBM administration (n = 46) 2. ≥50% HBM administration (n = 156) | 1. 5/46 (10.6%) infants affected with NEC in first group 2. 5/156 (3.2%) infants affected with NEC in second group | ≥50% administration of HBM for first 14 d after birth resulted in sixfold reduction of risk of NEC | [43] |
| 7.      | To understand expression of immune-related miRNAs in HBM exosomes | 602 miRNAs from 452 pre-miRNAs were studied from 20-30 mL HBM samples | 59 out of 87 (67.82%) immune-related pre-miRNAs contain HBM exosomes ($p < 10$-$16$) | 1. HBM exosomal miRNAs are transported to infants via GIT 2. These exosomes are resistant harsh extracellular environmental conditions DNA extraction from HBM is possible and shows ability to serve as source of DNA for pharmacogenetic studies | [44] |
| 8.      | To assess possibility of DNA extraction from HBM for pharmacogenetic studies | HBM samples (74) obtained from 37 mothers, of which some were subjected to pasteurization. Amplification of DNA performed using PCR technique | Mean DNA concentration in: 1. Pasteurized HBM samples: 2.0 ± 1.8 ng/μL 2. Unpasteurized HBM samples: 2.6 ± 2.0 ng/μL | DNA extraction from HBM is possible and shows ability to serve as source of DNA for pharmacogenetic studies | [45] |

HMGF III: Human Milk Growth Factor III, ICC: intra-class correlation coefficients, MIR: Mid-InfraRed, PCR: Polymerase Chain Reaction, TAC: Total Antioxidant Capacity.
against lung tumors [46]. The exosomes demonstrated protective effects against inflammatory diseases, boosted immunity in infants, improved targeting of anticancer drugs for tumors and exhibited cross-species tolerance with minimum side effects. Improvisation of oral bioavailability, enhancement of safety and efficacy of drugs and increase in biocompatibility, physical and biological stability, tolerability and cost-effective scalability for large-scale manufacturing were some predominant functions demonstrated by bovine milk-derived exosomes [46]. HBM was found to contain mRNA transcripts embedded in microvesicles that enabled transfer of genomic information from mother to infant during breastfeeding. The microvesicles exhibited reverse transcription activity, followed by integration into genome of infants to permanently rectify clinical manifestations of genetic diseases [47]. HBM microvesicles enhanced acceptance of allografts and showed tolerance to Major Histocompatibility Complex (MHC) antigens transferred from mother to infant after breastfeeding. The microvesicles displayed stability in extracellular environment, maintained their structure and function at low pH in stomach and exhibited resistance to enzymatic degradation by pepsin. Due to such effective functional properties of HBM microvesicles, their use as gene delivery vehicles for treatment of genetic diseases in infants is encouraged [47,48]. Isolation and engineering of HBM cells as potent vehicles for rapid and painless drug and gene delivery can facilitate tissue and organ-specific treatments for diseases in infants. Modulation of immune cells of HBM (for delivery of vaccines), stem cells (for prevention of birth defects) and epithelial cells (for production of specific proteins) in order to alter genetic diseases are some advanced approaches suggested for drug delivery to infants [48]. Collection of milk from mother with sick child, isolation of cells from BM, transfection of these cells with modified DNA, incubation of cells (for their growth and division), transfer of cells back to BM and finally administration of this modified BM to infants will result in fortification of nutritive value delivered to infants (Fig. 3). Epithelial cells of HBM produce large quantities of specific type of RNA that prevents translation of mRNAs into proteins and controls expression of maternal genes. Thus, epithelial cells in BM (also called as “protein-replacement factories”) show capability of integration into tissues and organs of breastfed infants resulting in reinforcement of their growth and development [7]. A four-month-old infant can consume more than 100 million cells of HBM (during breastfeeding) everyday; however, it is extremely difficult to deliver large macromolecules into GIT of infants due to limited intestinal permeability. However, HBM cells with 1000 X larger size

Fig. 3. Engineering HBM cells for development of drug delivery systems for infants.
than macromolecules are efficiently transported to GIT where they remain intact and proliferate without degradation [5]. Stem cells in HBM show potential to transform into any other cell in body of infants, thus, they show potential in development of regenerative medicine and generation of efficient systems to deliver drugs and vaccines for cure and prevention of untreatable diseases in infants [27]. Delivery of HBM-derived cells to specific locations in infant body is predicted to enable manifestation of specific therapeutic effects with lesser toxicity in comparison to other synthetic drug delivery systems [48,49].

5. Conclusions

Nutritional deficiencies in pediatrics increase risk of respiratory and gastrointestinal infections, brain and neural impairment, improper growth and immunological disorders. HBM is a natural biofluid composed of manifold bioactive macromolecules that fulfill nutritional requirements of infants, strengthen their growth and development processes and protect them from infectious diseases. Colostrum demonstrates more benefits than term breast milk and protects infants in initial days of life as it contains higher amounts of growth and immune factors. Exosomes derived from HBM show significant anti-infective activity, immunomodulatory properties and anti-cancer effects with possibility to function as carriers for drug delivery to infants. Several types of miRNAs are considered as key components of HBM due to their ability to exhibit both therapeutic and diagnostic properties. Development of regenerative medicine and use of drug, vaccine and gene delivery systems for infants to render specific treatment of untreatable diseases is possible via effective manipulations in stem cells, immune cells and epithelial cells of HBM. In the future, greater knowledge and advancement in research on HBM will aid in conceptualization of novel therapeutic applications to reinforce overall growth and development of infants.

Conflict of interest

The authors declare that there are no conflicts of interest.

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