Parenteral Nutrition in the Newborn: Associated Disorders and Nutritional Aspects

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Abstract

Severe prematurity at birth is an indicator of additional attention for the multidisciplinary team, since the newborn in this condition did not have the opportunity to develop organic systems under ideal conditions (intrauterine). Several factors may induce premature birth, malformation of organs or metabolic disruptions, however, additional attention has been given to inborn errors of metabolism (IEM) and deficiency/excess of key nutrients. Although relatively rare, IEM are detected in increasingly varied spectra and its importance in the proper fetal development has grown significantly. Variations in nutrient availability are also relate to the functional development of complex structures at the end of fetal development, playing a key role in critical periods of growth. Recent data indicate that an adequate supply of macro and micronutrients during pregnancy may be crucial to the survival of newborns, especially those subject to IEM. The use of parenteral nutrition (PN) in deep affected patients has grown significantly in recent years, and its presence has reduced intensely the morbidity/mortality levels among affected infants. Therapy with PN, however, entails risks to the health of newborns, by either chronic nutritional deficiencies, organ exhaustion or systemic infections. This review aimed to discuss progresses and issues encountered in the treatment of newborns with premature PN.

Keywords: Parenteral Nutrition; Inborn Errors of Metabolism; Malnutrition; Nutritional Therapy

Introduction

Neonatal complications normally occur due to multifactorial components, which are determinant of the type, gravity and survival rate of the disorder. Currently, premature infant mortality rates are reducing significantly. In the United States, at least 40% of the newborn with 22 or more gestational weeks are able to survive childbirth (years 1996-2000). During the 80’s (years 1981-1985), this rates of survive were reached only after the 26th gestational week. After surviving, however, premature patients require intensive care, once systemic immaturity imposes a high risk for the occurrence of a large array of complications [1].

Extremely premature patients or those affected by gestational complications may need support from parenteral nutrition (PN), a decisive resource to ensure survival. Parenteral therapy, however, is not a risk free approach. As highlighted by Calkins and colleagues (2014), the use of parenteral nutrition is associated to a myriad of symptoms, especially parenteral nutrition-associated liver disease (PNALD) and central venous catheter-related bloodstream infection (CRBSI) [2]. Nutritional deficiencies are also extenuating factors during PN use, once they impose barriers to the correct development of the central nervous system (CNS), an organ with high levels of plasticity and metabolic activity during the first years of newborn life [3].
In 2012, the Australasian Neonatal Parenteral Nutrition Consensus (ANPNC), aiming to reduce costs and attenuate prescription errors, have proposed standardized formulations for parenteral nutrition to intense care newborns. It is possible to highlight some of ANPNC’s main orientations, such as: (1) providing up to 150mL/Kg/day of water, including nutrient-containing fluids; (2) caloric content of 100-120Kcal/Kg/day to allow for optimal protein-energy adequation or at least 50-60Kcal/Kg/day to fulfill minimum energy needs; (3) 2g/Kg/day amino acids; (4) 17g/Kg/day carbohydrates and; (5) at least 0.5-1.0g/Kg/day lipids [4].

Arginine, an essential amino acid for the biosynthesis of molecules and detoxification of ammonia, is one of the most important in the balance between survival and pathogenesis. Brunton, et al. (1999) have shown that its presence is essential for survival in premature animals, and that the gastrointestinal system is an important regulator of arginine systemic bioavailability [5]. According to Wu and coworkers (2004), the deficiency of arginine may cause dysfunctions in the cardiovascular, pulmonary, neurological and gastrointestinal systems. Authors highlight that this deficiency occurs due to the limited expression of genes related to the synthesis of citrulline and arginine in intestinal cells, therefore supplementation of these amino acids in PN of premature newborns is a relevant strategy to optimize the nutritional status of these patients [6].

With the emerging increase in the premature pregnancies survival rates, it is essential to homogenize data from the medical literature, allowing accurate therapeutic and nutritional approaches, favorable to the proper development of pediatric patients. This work aims to discuss about the progress and shortcomings encountered in the treatment of premature newborns receiving PN.

Materials and Methods

To ensure methodological and typographic quality of the collected studies, search procedures were applied according to Freitas, et al. (2015). Search for scientific articles was limited to indexed databases (e.g. PubMed) and high impact journals. The following keywords were selected as search parameters: “Parenteral nutrition”; “Critically ill infants”; “Inborn errors of metabolism”; and “Preterm labor complications”. Experimental works, clinical studies and invited reviews were selected preferentially. References from selected documents were also considered for analysis and subsequent use in this work. Excluded items were stored and re-evaluated according to the pre-established search procedures. References from non-selected works were also read and additional documents were collected for verification [7].

Experimental studies without explicit presentation of a statement of approval from a responsible ethics committee, or performing inadequate procedures that may impose indiscriminate risk to human and/or animal life, were not considered for the completion of this review. Documents published in non-indexed bases, entertainment magazine articles, blogs, social networks, documents containing unreferenced data and abstracts were automatically excluded. After initial selection, papers were carefully read and classified according to the items and sub-items covered in this study [7]. In “Disorders in neonatology” section, were included works related to inborn errors of metabolism (IEM), respiratory, neurological and electrolyte disorders. In “Nutritional aspects”, were selected documents related to nutritional needs and therapeutic approaches, in addition to nutritional deficiencies common to critical infants. In “Parenteral nutrition”, data on the composition, use and complications of PN were included.

Selection criteria

Articles were excluded if not met the following inclusion criteria: neonatology-associated disorders OR physiological, neurological and metabolic disorders OR nutritional aspects of prematurity AND parenteral nutrition AND newborn or infant individuals, except for studies using in vivo (non-human) and in vitro experimental models, which were also considered for the purposes of this review. This study followed three major stages of document analysis:

1. Search for keywords in indexed databases and printed issues (n > 3000 documents);
2. Analysis title and summary, to exclude unrelated studies (n = 435 documents);
3. Careful selection, followed by individual articles reading (n = 70 documents).

Due to the large number of initial documents, we used software (Zotero Standalone 4.0) to organize and select references. Considering the methodological limitations of this narrative review, we have adapted our search procedures from Freitas, et al. (2015) and from the PRISMA (i.e. Preferring Reporting Items for Systematic Reviews and Meta-Analyses) norms and standards, allowing for systematic search, optimization of analysis and discussion of the data obtained [7]. Figure 1 illustrates the procedures followed for document acquisition, selection and evaluation.
A great variety of symptomatic IEM is caused by the possibility of different levels of expression and activity of several enzymes. As example are the errors associated with the metabolism of L-glutathione (GSH), a tri-peptide made of glutamate, cysteine and glycine residues, which are present at millimolar concentrations in virtually every cell of the body. GSH is involved in several fundamental biological activities, especially working as non-enzymatic antioxidant, intra and extracellularly. Of the six enzymes involved in the metabolism of GSH, five are related to known IEM, the residual enzymatic capacity and viability of other associated enzymes determines the severity of the symptoms presented, such as hemolytic anemia, metabolic acidosis, neuromuscular defects and generalized aminoaciduria [12].

Among the disorders related to amino acids, those involved in sulfur amino acid metabolism should receive additional attention. Defects in the methionine cycle are involved in methionine synthase activity, either by disorders in the synthesis of cobalamin due to abnormalities in the apoenzyme, and consequently a failure in the formation of methyltetrahydrofolate, or by defects in the enzyme methionine synthase. All these errors dramatically increase homocysteine (HCT) levels, toxic to the body in high concentrations. Other disorders can significantly raise HCT levels are: the deficiency of the enzymes cystathionine beta-synthase (CBS); cystathionase; methionine adenosyltransferase I and III (MAT I/III); glycine N-methyltransferase; and S-adenosylhomocysteine hydrolase. Apparently, patients presenting any of these disorders respond positively to a methionine restricted diet, indicating that patients under restricted PN presenting these IEM must receive a different combination of amino acids [13].

It is possible, however, that other EIM are involved in the defective metabolism of HCT. Cobalamin deficiency is associated with increased and accumulation levels of methylmalonic acid and HCT. Although dietary deficiency of this and other organometals is rare, there are altogether twenty IEM that can affect the absorption, transport, uptake or intracellular metabolism of cobalamin. A major dysfunction to be highlighted is the deficiency in the synthesis of intrinsic factor, essential for the proper absorption of cobalamin. Specific symptoms of the disease appear when the patient is between 1-5 years of age, however, there is the possibility that megaloblastic anemia and/or neurological problems appears before this period [14].
Data reinforces the need for the use of efficient diagnostic tools in gestational and early postnatal periods. Various manifestations can be associated with IEM and therefore allow for therapeutic approach toward drug or nutritional correction of the disorder. Patients under PN treatment alone should receive additional attention in this context, as small changes in the levels of nutrients and caloric content can influence the survival and further development of the infant. Table 1 shows various symptoms commonly associated with IEM.

### Table 1: Symptoms associated to inborn errors of metabolism

<table>
<thead>
<tr>
<th>Disorder</th>
<th>Symptoms</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lactase deficiency</td>
<td>Mitochondrial disorders*</td>
</tr>
<tr>
<td>Abetalipoproteinemia</td>
<td>Enteropeptidase deficiency</td>
</tr>
<tr>
<td>Lysinuric protein intolerance</td>
<td>Sucrase/isonaltase deficiency</td>
</tr>
<tr>
<td>Fatty acids oxidation disorders</td>
<td>Mitochondrial disorders*</td>
</tr>
<tr>
<td>Glycogenolysis disorders</td>
<td>Myoadenylate deaminase deficiency</td>
</tr>
<tr>
<td>5,10-methylenetetrahydrofolate reductase deficiency Familial hypercholesterolemia</td>
<td>Fabry disease</td>
</tr>
<tr>
<td>Homocystinuria</td>
<td>Multiple deficiency of carboxylase</td>
</tr>
<tr>
<td>Metachromatic leukodystrophy</td>
<td>HHH syndrome</td>
</tr>
<tr>
<td>Mitochondrial disorders*</td>
<td>Peroxisomal disorders</td>
</tr>
<tr>
<td>Metachromatic leukodystrophy</td>
<td>Genetic disorders of glycosylation</td>
</tr>
<tr>
<td>Galactosemia</td>
<td>3-oxothiolase deficiency</td>
</tr>
<tr>
<td>d-2-hydroxyglutaric aciduria</td>
<td>Mitochondrial disorders*</td>
</tr>
<tr>
<td>Type I glycogenosis</td>
<td>Type I and IV hyperlipoproteinemia</td>
</tr>
<tr>
<td>Lipoprotein lipase deficiency</td>
<td>Lysinuric protein intolerance</td>
</tr>
<tr>
<td>Mitochondrial disorders*</td>
<td>Type C Niemann-Pick disease</td>
</tr>
</tbody>
</table>

*Disorders of mitochondrial etiology may be related to respiratory chain components, oxidation of lipids and various other components of the organelle. Adapted from [16]

In large populations, the fatality rate related to respiratory disorders is relatively low, however, there is a wide variety of affections capable to impose health risks to preterm infants or newborns from mothers whose age is inadequate for pregnancy. Rubaltelli, et al. (1998) investigated a population of approximately 17,000 Italian neonates, of these, about 2.8% has developed some signal of respiratory distress, and 15.88% were cases of fatality. The fatality rate by respiratory disorders, in any population observed, was 0.45% [17].

According to Levine, et al. (2001), the incidence of pulmonary hypertension in neonates is about five times greater in individuals born by cesarean compared to the traditional delivery procedure [18]. Several authors have investigated the consequences of a cesarean delivery, such as the occurrence of uterine rupture [19]. In addition, recent data suggest that this mode of delivery can significantly affect the body mass index (BMI) of these subjects when adults (23-25 years), without, however, the presence of other metabolic risk markers [20].

Performing of cesarean delivery have increased significantly over the years, about 37% of US births are performed this way [21]. When the procedure (i.e. cesarean) is the mother’s choice, a responsible medical team has to inform the patient about the risks and benefits of the procedure. In fact, recent results indicate that the mother’s reproductive plans are a key factor in the successful development of the gestational process, whatever the mode of delivery selected by the family [22].
Genetic and epigenetic disorders can affect respiratory function in the postnatal period; such disorders are greatly influenced by behavioral factors and family history. As expected, infants requiring intensive care often present a varied array of hereditary dysfunctions. Data about inaccuracies in the expression of alleles of the surfactant protein A are heterogeneous, but indicate that this protein is related to adequate protection against viral and bacterial infections. Surfactant proteins C and D, in contrast, appear to be involved in the development of chronic affections. Loss of function in alleles expressing the surfactant protein B seems to have greater impact on postnatal affections, potentially generating irreversible acute or chronic effects in susceptible infants [23].

Primary ciliary dyskinesia (PCD) is an autosomal recessive hereditary manifestation resulting in inappropriate mucociliary transport, characterized by airway obstruction and recurrent infections of the lung, middle ear and paranasal sinuses. Individuals with PCD normally live satisfactorily if treated with the appropriate periodicity, however, there is certain degree of unpredictability in the progression of the symptoms, which can result in severe pulmonary dysfunction and, ultimately, to respiratory failure and death [24].

Pharmacological approaches at the end of pregnancy (34-36 weeks) and in the postpartum period are apparently ineffective in mitigating the incidence of inflammatory or dysplastic airway diseases. Porto, Coutinho and Amorim (2011), after administration of 12 mg intramuscular betamethasone in pregnant women with imminent risk of having premature delivery, found no reduction in the incidence of respiratory disorders in neonates [25]. Kinsella and colleagues (2006) also found no reduction in the incidence of bronchopulmonary dysplasia in infants with birth weight > 1000 g after the administration of nitric oxide (NO). Their data, however, indicate that the use of NO was able prevent overall risk of brain damage [26].

Extremely premature infants may undergo major loss of water and electrolytes, this is mainly due to disproportion between the relatively large body surface area and the immaturity of their skin. This relationship allows insensible water loss, which can be exhausted in the presence of other factors such as radiant heat, tachypnea, omphalocele, gastrochisis and exposure to dry conditions. According Bhatia (2006), six strategies may be used for proper maintenance of fluid and electrolytes in premature infants with body weight under 1000g: (1) effective parenteral delivery of amino acids; (2) proper fluid replacement; (3) continuous measurement of electrolytes; (4) progressive increase in amino acid supply; (5) adjustment of lipids supply and; (6) monitor weight, vital signs and plasma urea daily, providing sodium (2-3mEq/kg/day) only when measured under 130mEq/L, plus 2-3/kg/day of potassium [27].

In addition to the regular continued attention needed to maintain the health of the newborn, diseases and dysfunctions of hereditary nature may induce electrolyte imbalance in premature and at term infants. Acute renal failure (ARF) may occur due to a congenital etiology, as in the case of polycystic kidney disease. This disorder can also occur in the postnatal period as a result of ischemic hypoxia or severe toxicity induced by chemical contaminants or IEM/drugs [28]. Gouyon and Guignard (2000) discuss procedures needed for the application of intense care in neonates with ARF. According to authors, the initial treatment of complications related to ARF should aim to correct the hypotensive crisis, acidosis and hypoxemia, allowing reduced renal vasoconstriction and improved tissue perfusion [29].

Infants with less than 3 months of age, in which the formation of the urinary tract was inadequate, in concomitance to pyelonephritis, are at risk of transitional pseudoaldosteronism. Thies, et al. (1999) report the case of newborn males, 17 days of life, which presented with pseudoaldosteronism as consequence of pyelonephritis and vesicoureteral reflux. The initial diagnosis, however, was congenital adrenal hyperplasia, characteristic of the symptoms of insensitivity to aldosterone and excessive loss of sodium [30]. Martinerie, et al. (2009), however, describing physiological mechanisms of interaction between aldosterone and excretion of electrolytes in the postpartum period, point out that the emergence of partial resistance to aldosterone action is common in the first moments of life [31], indicating that additional markers such as intense metabolic acidosis and dehydration should be added to the diagnostic process.

According to recent data from Diogo, et al. (2015), metabolic and electrolyte disorders in newborns cause damage to the CNS, preferably originating in the deep region of the gray matter, with less involvement of the white matter and cortical regions. These lesions are typically bilateral and symmetrical, with the exception of non-keto hyperglycemia, which is typically unilateral [32]. Neurological disorders in newborns will be briefly discussed next.

The evaluation of neurodevelopmental status is one of the most critical remarks to be held in the first days after birth [33]. Since the middle of the ninth century, authors have recognized the need to evaluate physical parameters, such as the signals for the adequate development of the nervous system. During this period, aspects such as quality of hair, skin and plantar creases served as clinical evidence for neurological maturity [34]. Recent studies indicate that even discrete neurological signs serve as a parameter for assessing the presence of cognitive and psychiatric disorders in newborns, especially those with low birth weight [35]. It is necessary, therefore, that grading standards are clear about the etiology and severity of the presented neurological dysfunction, allowing emergency therapeutic procedures to be more effective [36].
In addition to intrinsic factors, environmental factors and maternal quality of life during pregnancy can result in observable neurological disorders. In a prospective observational study, Smith and colleagues (2000) evaluated the neuropsychological performance of newborns born from pregnant women with subclinical hypo or hyperthyroidism. Their data indicate that in the first half of pregnancy, the presence of subclinical hypothyroidism was associated with lower mental development index during the first year of life [37]. Chiriboga, et al. (1999) investigated the neurological effects of fetal exposure to cocaine in about 100 newborns whose mothers were regular users of different doses of the substance. Drug-exposed children have high rates of growth retardation, smaller head circumference and neurological abnormalities, such as global hypertension, tremors and stiffness in the extensor muscle. Using linear regression analysis, the authors also showed that the extent of neurologic damage occurred in a dose-dependent manner [38].

According to Dammann, et al. (2015), damage to the lungs and brain are the most common affections found in immature neonates. In cortical regions, there was extensive damage to white matter, including focal necrosis and diffuse astrocytosis. Ultrasound methods (USM) are typically used for detection of large histological injury areas, however, only about 8% of infants born with very low birth weight (< 1500 g) show observable signs of tissue injury. Methods such as magnetic resonance imaging (MRI), however, are able to detect indicators of morphological abnormalities in about 50% of individuals born before 34 gestational weeks [39]. Data corroborate concisely about the accuracy of MRI methods [40]. In addition, epidemiological studies indicate that preterm (< 32 weeks) and extremely low birth weight (< 1500 g) individuals exhibit high levels of cognitive impairment and have great difficulty in becoming completely independent of familial care [39].

In a recent study, Miller and colleagues (2007) showed that congenital cardiovascular diseases are correlated with abnormal morphological brain development. Their data suggest similarity with results from preterm infants, born with extremely low birth weight, presenting significant increase in tissue diffusivity and loss of white matter [41]. Recent data suggest that treatment with low doses of recombinant human erythropoietin is capable of reducing risk of morphological and functional disorder in neonates with moderate hypoxic-ischemic encephalopathy [42]. These data indicate that severe hypoxemia and vascular disorders produce potentially irreversible morphologic damage in premature neonates, stressing the need for effective and sensitive diagnostic tools during pregnancy.

As previously discussed, there is a need to develop effective tools for diagnosing diseases initiated during pregnancy. Due to very specific mechanisms and phenotypes expressed, however, researchers are struggling to find effectively predictive methods for unusual neurological diseases [43]. Currently, authors investigate the role of mitochondrial dynamics in neuroplasticity and development of neurological diseases [44], searching, through the use of stable clonal lineages of neural stem cells, new methods in regenerative medicine for the central tissue [45].

**Nutritional aspects**

After birth, dramatic changes in energy metabolism occur in order to ensure proper body development, especially the essential consolidation of CNS structures, which takes many years to reach some level of stability. In premature newborns, these nutritional requirements not only override, but also create a differentiated demand for protein, total calories and key minerals, such as calcium. Recommendations for caloric intake to low birth weight infants (i.e. 120Kcal/kg/day) have remained unchanged for 30 years [4,46]. Authors have, however, revalued guidelines on protein requirements, considering this nutrient a limiting factor for the proper development of premature newborns. Rigo and Senterre (2006) highlighting the role of protein on weight gain, growth stimulus and accumulation of lean mass, recommends about 3.8-4.4g of protein/kg/day and protein/energy ratio of about 3.0-3.3g/100Kcal/day for preterm infants with 26-30 weeks of age [47].

The use of breast milk (BM) is essential for proper development of the newborn, and recent data suggest that preterm infants may benefit from approaches where the provision of BM in complemented with nutritional therapy. Authors point out that the use of BM promotes gastrointestinal maturation processes, improvements in neurobehavioral tests, optimized coordination of sucking/swallowing, oxygen saturation and body temperature, indicating that feeding premature infants with BM comprises a physiological recovery strategy [48]. The use of BM has also been explored as a way to counter approaches with strictly unnatural food, that despite the ability to promote improvements in growth and calorie intake, does not optimize survival rates or decrease the length of stay under hospital care [49]. Therefore, it's of great importance to highlight the need for early observation of malnutrition and prematurity signs, once this practice may shorten the demand for permanence under hospital care, PN therapy and even invasive procedures.

In newborns receiving enteral or PN, attention should be paid to the adequate supply of essential nutrients and the recovery from symptoms of congenital affections, such as IEM. VanderVeen, et al. (2013) show that limited supply of essential nutrients, especially lipids and, therefore, total calories, may exhaust symptoms of retinopathy of prematurity (ROP). Data indicate that weight gain is a key approach to reduce symptoms of ROP in extremely premature infants [50]. In fact, recent data indicate that parenterally-provided lipids to premature infants with extremely low birth weight is related to weight gain and possibly optimize the nutritional status during the intensive care period [51]. With the advent of new technologies in the production of concentrated/isolated food compounds for nutritional therapy, investigators seek to highlight the participation of specific lipids in the recovery of several bouts. Recent evidence indicates that long-chain polyunsaturated fatty acids play an important role in the treatment of IEM [52] and morphological development of the CNS [53].
A problem faced by nutritional approaches in pre or postnatal periods is that, in many cases, damage or delay in the development and differentiation of cells in the CNS makes it impossible to restore proper cognitive state during childhood and adulthood. Gestational periods where proliferation of neural cells is critical seem to be crucial in the morphofunctional consolidation of brain structures and thus prevents drastic recovery or induction of plasticity only by nutritional approaches during or after birth [54]. Due to the prematurity of certain disorders, the use of PN can act only as a stopgap measure, losing its essentiality or ability to promote improvements in the framework of profoundly affected newborns. These cases require deep ethical reflection, in order to provide parents or guardians with the best guidance about the non-effectiveness of PN and therefore the interruption of its use [55].

Nutritional deficiencies during pregnancy may impose profound impact on the proper fetal development, and also affect the emergence of symptoms related to weight disorders in adulthood. Triunfo and Lanzone (2015) highlight aspects of malnutrition in the prenatal period and its direct consequences at birth. According to authors, cases of gestational iron-deficiency anemia can induce premature labor, prolonged labor, postpartum hemorrhage and also fetal growth restriction. Vitamin A deficiency, in turn, can promote episodes of maternal night blindness, low birth weight, reduced growth rate and raise the mortality/morbidity levels of infants [56]. Evidence suggests that, while some of the damage may occur prior to birth, supplementation with micronutrients through PN would promote total or partial recovery from symptoms or nutrient deficiencies (see Table 3).

Zinc, a key micronutrient for essential cellular functions, seems to be an important factor in preventing premature births. It’s also common to observe the development of cretinism in the offspring of women presenting severe iodine deficiency. This disorder, however, can be eliminated with supplementation during the first trimester of pregnancy or during the prenatal period. The lack of folate may prevent proper formation of the neural tube, with serious impacts on morbidity/mortality. Finally, calcium deficiency was related to hypertensive disorders during pregnancy, mortality, premature birth, fetal growth restriction and also those related disorders of vitamin D metabolism [56].

In a recent study, Sonne, et al. (2012) developed neonate mice with mutations in the OCTN2 carnitine transporter, which plays an important role in carnitine resorption, accounting for over 90% of the carnitine transport capacity to body tissues. Data show severe morphological changes in the intestinal epithelium, infiltration of lymphocytes and macrophages, generalized atrophy, pro-apoptosis and inflammatory responses [57]. Failure of expression or deficiency of carriers for certain nutrients is a causal factor for many disorders in newborns, especially in premature ones. Deficiencies in the expression of an intestinal zinc transporter (i.e. ZIP4), for example, leads to severe zinc malabsorption, which performs its biological function at picomolar (pM) concentrations. This deficiency, when chronic, causes acrodermatitis enteropathica, a recessive inherited disease that imposes severe damage to epithelia and other systems [58].

In an epidemiological study by McLimore and colleagues (2013), multiple risk factors (intrinsic and environmental) proved to be proportionally correlated to the adequacy of iron in the postnatal period, indicating that nutritional deficiencies can be influenced by factors other than those traditionally observed [59]. Maldonado-Cedillo and colleagues (2015), in turn, have recently demonstrated that malnutrition may extenuate other phenomena responsible for inducing fetal developmental disorders. In their study on rodents, the use of a malnutrition protocol with lead poisoning led to reduced body and brain weight from the fourth week of pregnancy, significant increases in lipid peroxidation and increased stress-like activity in cerebellar regions [60].

Malnutrition during critical periods of gestation can also negatively regulate the expression of genes related to development and child growth. Ojha, et al. (2013), in a model of sheep, showed that the supply of sub-optimal feeding to pregnant animals led to lower expression of genes related to the activity of the brown adipose tissue, which is intrinsically involved with substrate metabolism during the initial periods of mammal development [61]. Current trends in nutrition of premature infants indicate that it is imperative that mothers aim to provide BM as the main nutrient source, with or without additional nutritional therapy, unless it’s necessary to guarantee the required substrates for postnatal growth [62]. Authors also highlight the importance of observing the specificity in the supply of key nutrients, ensuring that the minimum needs for polyunsaturated chain fatty acids are adequate, and that the iron supply, necessary to balance oxygen perfusion systems and nutrients throughout the body, is preserved [63].

Parenteral Nutrition
The supply of PN is, in first instance, a mandatory procedure to balance nutrient losses and imbalances caused by diseases capable of impairing oral feeding. Even if PN is not able to fully replace the nutritional needs of premature newborns, its use prevents morbidity and mortality in several infants in critical condition. IEM patients, for instance, have a mandatory need for specific amino acids, lipids and other derivatives to be supplemented or definitively excluded from their diet, therefore PN may be used to provide with essential nutrients, absent in a restrictive diet. Infants with extreme low birth weight presenting with respiratory complications, however, may demand greater caloric input, which is frequently impaired due to the prematurity of digestive mechanisms, intestinal epithelium and liver. Premature newborns at risk of developing neurological disorders, such as cognitive impairment and/or extensive neural/glial death may receive parenteral support to provide polyunsaturated fatty acids, micronutrients and energy necessary to allow for neuronal growth, survival and differentiation. Finally, preventing loss or excessive accumulation of electrolytes is one of the most essential uses of PN, once it’s able to prevent CNS injuries and compensate for irregular reabsorption/excretion of several micronutrients. Table 2 presents nutrient recommendations (via PN) for high-risk newborns.
<table>
<thead>
<tr>
<th>Nutrient</th>
<th>Estimated needs</th>
</tr>
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<tbody>
<tr>
<td>Calorie (kcal/kg)</td>
<td>80–90</td>
</tr>
<tr>
<td>Carbohydrates (g/kg)</td>
<td>16</td>
</tr>
<tr>
<td>Proteins (g/kg)</td>
<td>3–4*</td>
</tr>
<tr>
<td>Lipids (g/kg)</td>
<td>3–4</td>
</tr>
<tr>
<td>Sodium (mEq/kg)</td>
<td>3–4*</td>
</tr>
<tr>
<td>Potassium (mEq/kg)</td>
<td>2–4</td>
</tr>
<tr>
<td>Calcium (mg/kg)</td>
<td>80–120</td>
</tr>
<tr>
<td>Phosphorus (mg/kg)</td>
<td>60–90</td>
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<tr>
<td>Magnesium (mg/kg)</td>
<td>9–10</td>
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<tr>
<td>Copper (g/kg)</td>
<td>65</td>
</tr>
<tr>
<td>Zinc (g/kg)</td>
<td>350–450*</td>
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<tr>
<td>Vitamin A (IU/kg)</td>
<td>500</td>
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<tr>
<td>Vitamin D (IU/kg)</td>
<td>160</td>
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<td>Vitamin E (IU/kg)</td>
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<td>Vitamin K (IU/kg)</td>
<td>80</td>
</tr>
<tr>
<td>Vitamin B1 (mg/kg)</td>
<td>150</td>
</tr>
<tr>
<td>Vitamin B12 (g/kg)</td>
<td>0.4</td>
</tr>
<tr>
<td>Vitamin C (mg/kg)</td>
<td>32</td>
</tr>
</tbody>
</table>

*Ileostomy, surgical drain and other procedures may result in increase in demand. Adapted from [64]

Table 2: Nutritional recommendations for parenteral nutrition in high-risk newborns

Prior to the availability of PN, premature children and high-risk pregnancy had lower rates of survival. With its emergence and subsequent adaptation for neonates, there was a drastic increase in survival and clinical evolution, even in high-risk infants. On should, however, to consider the use of NP as an emergency feature, which imposes significant side effects to the neonate and also involves significant hospital costs. PN standardized solutions usually consist of dextrose, as a source of carbohydrate; essential and non-essential amino acids, thus mimicking protein intake orally; fat emulsions, providing essential polyunsaturated lipids and fatty acids as an energy source; carnitine to promote the oxidation of long chain fatty acids; sodium, calcium and phosphorus, essential electrolytes; vitamins and trace elements, which can vary according to symptoms or procedures to which the infant is subjected [64]. Table 3 presents a brief example of neonate progression through PN and possible modifications according to specific needs and disorders.

*Inborn errors of metabolism. **Central nervous system. Adapted from [30], [46], [47], [63] and [64]

Table 3: Variable requirements in neonate progression through parenteral nutrition (PN)
Several authors discuss the need for using individualized PN prescriptions, considering the availability of novel automated systems, that do not require daily evaluation and can automatically modulate the availability of nutrients needed for critical patients. According to Beecroft, Puntis and Martin (1999), Yeung, et al. (2003) and a review from Riskin, Shiff and Shamir (2006), the use of standardized prescriptions apparently doesn't induce significant changes in biochemical standards, and promotes a significant reduction in error rates, also attenuating total expenses with therapeutic procedures. Authors also point out that the use of computerized systems for the administration of nutrients may optimize this procedure [65-68].

As regards the micronutrients, small changes in their supply may modulate common disorders of patients undergoing therapy with PN. There is hard debate on whether is needed to fine-tune the standardization of nutritional formulas for PN, however, few clinical evidences are collected on these matter. The provision of chromium (Cr), for example, has been reported as excessive by recent studies, whereas only contaminants in PN solution may increase from 10-100% metal concentrations in the administered compound [69]. Manganese (Mn), in turn, is a trace element found in virtually all tissues, playing roles related to the metabolism of macromolecules/nutrients as an allosteric modulator, acting through a large family of enzymes related to these pathways. Studies indicate that, despite its essentiality, the removal of Mn solutions from PN may benefit patients with cholestatic liver disease (CLD), even in those infant patients who develop symptoms at the expense of PN therapy chronicity [70,71].

Similarly to Mn, copper (Cu) is an essential nutrient, even in low concentrations, which should be present in the food or therapeutic solution in order to avoid symptoms of anemia, neutropenia, skeletal abnormalities, and other clinical manifestations. Evidence point, however, to the need for reducing the concentration of Cu in PN solutions for infants and children presenting CLD [72]. Corrections in the levels of potentially harmful nutrients and contaminants may help to improve the general framework of premature infants, thus avoiding specific symptoms developed during pregnancy. Authors, however, have not yet been able to demonstrate improvements in the general health of affected newborns by providing supplemental nutrients thought to be beneficial. A study of glutamine supplementation was not able to induce significant improvements in neonates and infants undergoing surgery in the digestive tract [73], reinforcing the idea that protocols that prioritize standardization of regulations, adequate supply of needed nutrients and prevention of deficiencies are the best to optimize neonate survival.

One of the main complications related to the use of PN are those associated with the liver, about 40-60% of infant patients undergoing PN for long periods develop PNALD. Among the common liver onsets are: hepatic steatosis, cholestasis, deficiency of nutrients and diseases associated with the gallbladder. Oral decontamination, encouraging enteral feeding, fiber intake and treating catheters with antibiotics are among the most common strategies to prevent or treat PNALD [74], however, these procedures must follow a strict schedule, considering that the period for permanence under PN support may not last for more than few weeks, once a long-term use of PN implies in elevated risk of infection, organ failure and death. In these cases, the family must consider the possibility of either an induced recovery through oral/enteral feeding or an invasive procedure, such as gastrostomy.

Recent evidence, including original articles and case reports point to the use of lipid emulsions containing fish oil as a resource to prevent and even reverse PNALD. De Meijer, et al. (2009) presented data indicating that treatment with emulsions containing fish oil may revert hepatic onsets in a murine model [75], while Gura, et al. (2006) report two cases where infants with intestinal failure were treated with omega-3 fatty acids. One patient could be withdrawn from a transplant list, while the second was completely free of symptoms of cholestasis [76]. These data point to the importance of exploring the effects of lipid emulsions enriched with fatty acids, especially long chain n-3 polyunsaturated fatty acids and its derivatives.

As discussed above, one approach to reduce liver symptoms is the treatment of PN catheters with antibiotics and, in some cases, its periodic replacement. Catheter-related bloodstream infections are common, and must be primarily prevented by proper handling during exchange and maintenance of both catheter and perfusion systems. According to Puntis and Hodge (2002), some procedures may be applied to prevent contamination and related infections of the PN catheter, such as: (1) installation and handling of the catheter by experienced operators; (2) use of catheter as a route passage of nutrient solutions only; (3) use of pre-planned treatment protocols and; (4) continuous training of the critical care team and multidisciplinary professionals [77].

Conclusions

Hereditary or environmental factors may affect proper fetal development and induce prematurity of systems. In those cases, susceptibility to nutritional deficiencies, infections, and other physiological disorders are increased. High risk of prematurity may require treatment with PN to delayed or reverse symptoms, allowing greater chance of survival and quality of life in adulthood. Some disorders, however, occur in critical periods of fetal development and therefore may be irreversible, resulting in ethical conflict regarding the maintenance of PN in patients without positive perspective of progression. The medical team should therefore use technical information and based-on-evidence arguments to provide parents with better advice about the maintenance of PN therapy. PN is a resource able to significantly mitigate the levels of child mortality and morbidity, and has been enhanced with the use of precise technological resources and studies that regulate the real nutritional requirements of deeply affected patients, avoiding shortages and induction of toxicity on fragile systems.
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References


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