Transition From Parenteral to Enteral Nutrition and Postnatal Growth in Very Preterm Infants During Their First 28 Days of Life

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Background: Nutrition practices for preterm infants during the first few weeks of life can be divided into three phases: the parenteral nutrition (PN), enteral nutrition (EN), and transition (TN) phases; the TN phase includes both PN and EN. Our purpose was to analyze nutrition practices for very preterm infants during the TN phase and their association with the infants’ growth during the first 28 days of life.

Methods: Data from 268 very preterm infants < 32 weeks old from six neonatal intensive care units were analyzed retrospectively. The TN phase was defined as enteral feedings of 30-120 ml/kg/d. Postnatal growth failure (PGF) was defined as a 28-day growth velocity < 15 g/kg/d. Differences in protein and energy intake between the PGF and non-PGF groups during the TN phase were calculated, and risk factors for PGF were identified using multivariate regression analysis.

Results: The total protein (parenteral + enteral) intake during the TN was 3.16 (2.89, 3.47) g/kg/d, which gradually decreased as the enteral feeding volume increased in the TN phase. The total energy (parenteral + enteral) intake during the TN phase was 115.72 (106.98, 122.60) kcal/kg/d. The PGF group had a lower total protein intake (parenteral + enteral) than the non-PGF group had [3.09 (2.85, 3.38) g/kg/d vs. 3.27 (3.06, 3.57) g/kg/d, P = 0.007, respectively]. No significant difference was found in energy intake during the TN phase. The variables associated with PGF included a lower total protein intake (parenteral + enteral) intake, a smaller day of age at the end of the TN phase, and a higher birth weight z-score.

Conclusion: Increasing the total protein intake (parenteral + enteral) during the TN could reduce the incidence of PGF.

Keywords: enteral nutrition, parenteral nutrition, premature [MeSH], growth, first 28 days of life
INTRODUCTION

Postnatal growth failure (PGF) is prevalent among very preterm infants (1, 2), and the risk of developing adverse metabolic and neurodevelopmental outcomes later in life is high (3). In a longitudinal cohort study of the long-term growth of preterm infants, Andrews et al. (4) concluded that early PGF can be prevented. Hence, the importance of early nutritional management for the prevention of early growth failure among preterm infants has been gradually recognized (1, 5). The implementation of an enhanced parenteral nutrition (PN) protocol was found to promote growth, but it was also associated with an increased prevalence of severe hyperglycemia and higher mortality (6–9). Appropriate postnatal growth is more likely to be followed by better long-term outcomes (10, 11); therefore, it is particularly important for preterm infants to achieve appropriate nutrition intake and growth during their first few weeks of life.

Actual nutrition practices for preterm infants during their first few weeks of life can be divided into three phases: the parental nutrition (PN), enteral nutrition (EN), and transition (TN) phases, with the TN including both the PN and EN (12). Nutrition during the TN phase is a new challenge for neonatal physicians because it involves the transformation of parenteral amino acid (AA) to enteral protein (13–17). Liotto et al. (14) found that intensified nutritional management during the TN improved the postnatal growth of preterm infants. Moreover, a high incidence of poor growth during the TN predicted growth failure at discharge. However, poor growth during the PN phase was not significantly associated with growth failure at discharge (18).

Martin et al. (19) reported improved growth during the neonatal period by optimizing the early nutrition of preterm infants, but the relationship between nutrition during the TN phase and growth during the neonatal period remains unknown. The purpose of our study was to assess in the discharge. However, poor growth during the PN phase was not significantly associated with growth failure at discharge (18).

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Nutrition Management

Infants received PN within 24 h after birth, and their initial glucose dose of 4–8 mg/kg/min was increased by 1–2 mg/kg/min, up to 11–14 mg/kg/min. Their AA intake was increased from 1.5–2.0 to 3.5–4.0 g/kg/d; lipids were started at 1.0 g/kg/d and increased by 0.5–1.0 g/kg/d, up to 3 g/kg/d.

Infants without contraindications to EN support should receive minimal enteral nutrition as early as possible after birth, with human milk or formula (10–20 ml/kg/d for 3–5 days). Full EN is achieved when the EN calorie intake is at least 80 kcal/kg/d. The target caloric value of full PN is 80–100 kcal/kg/d, and the target caloric value of full EN is 110–135 kcal/kg/d (20).

Data Collection and Definitions

The TN phase began with the reduction of PN, i.e., an EN volume of 30–120 ml/kg/d (≥30 ml/kg/d, ≤120 ml/kg/d). The PN phase began with the PN included a minimal volume of EN and a PN-only phase. Full EN was defined as an EN volume > 120 ml/kg/d or the absence of a PN infusion.

Data on the nutrient intake of all intravenous and enteral products during the TN phase and daily body weights were collected retrospectively to evaluate the intake of intestinal protein and energy, parenteral AA and energy, and total protein and energy. An EN volume of 30 ml/kg/d, which was considered equivalent to 30–39 ml/kg/d, was fed to the infants. When the feeding amount was the same for more than 1 day, the daily nutrient intake on these days was divided by the infant’s body weight of that day, and the average of these nutrients in the feeding volume was calculated as the average nutrient composition of the feeding volume.

Growth indicators included birth weight, gestational age, time to regain birth weight, and growth velocity (GV). Being small for gestational age was defined as a birth weight less than the 10th percentile for gestational age (21). The birth weight z-score was calculated using an online preterm growth calculator (available at www.peditools.org) (22), and the formula was: z-score = (the individual value – mean value)/standard deviation. The GV (from birth to the 28th day of life) was based on an exponential model of regaining birth weight. The formula was: $ GV = [1,000^*\ln (W_{28}/W)]/(28-D); $ where $ W_{28} = $ weight on the 28th day of life, $ W = $ birth weight, and $ D = $ the day (of age) that birth weight was regained (23). The term PGF was defined as a 28-day GV < 15 g/kg/d. The term non-PGF was defined as a 28-day GV ≥ 15 g/kg/d (24).

Feeding types included human milk and formula; human milk included mother’s own milk (MOM) and donated human milk (DHM). When the volume of human milk reached 50–100 ml/kg/d, human milk fortifiers (HMFs) were added as supplements (20). The formula included: formula for preterm infants, extensively hydrolyzed infant formula and amino acid-based formula. The protein and energy components of the human milk were calculated in accordance with the respective nutrient reference values of transitional milk, mature milk, and donated milk (25, 26). The nutrient composition of the formula and HMFs were calculated according to the products’ instructions (Table 1).
Energy calculations of parenteral nutrition were as follows: glucose = 4 kcal/g (glucose injection); AA = 4 kcal/g (6% pediatric compound amino acid injection 18AA-I/19AA-I); and lipids = 9 kcal/g (20% medium and long-chain fat emulsion injection, Cs−24Ve). During the PN-dominant TN phase, the amino acids in the PN were expressed as protein (1 g protein = 1.13 g AA) to calculate the total protein.

**Clinical Factors**

The clinical factors associated with growth outcomes included bronchopulmonary dysplasia (BPD), with continuing oxygen requirements at 36 weeks corrected gestational age (27), necrotizing enterocolitis ( NEC) ≥ grade 2 (28), intraventricular hemorrhage (IVH) ≥ grade 3/periventricular leukomalacia (29), confirmed sepsis (30) and invasive mechanical ventilation.

**Statistical Analysis**

Continuous variables were expressed as median and interquartile range (IQR), and categorical variables were expressed as frequencies and percentages. The Mann–Whitney U and Chi-square tests or Fisher’s exact test were used to analyze differences between groups. Variables with statistically significant differences in the univariate analysis were entered into the multivariate logistic regression model to determine the factors predicting PGF. P < 0.05 was considered statistically significant. All analyses were performed using SPSS statistical software, version 26 (IBM Corp. Armonk, NY).

**RESULTS**

**Study Population**

In 2019, 742 preterm infants were treated in the six NICUs; 474 were excluded from the study: 294 because their length of stay was <28 days, 10 because complete data were not available, 2 because their admission to the NICU was >24 h after birth, and 168 because they had not completed the TN by the 28th day after birth. Thus, data from the remaining 268 infants were included in this study.

Participants’ demographic characteristics and clinical outcomes are compared in Table 2. Birth weight and birth weight z-scores were significantly higher in the PGF group than those in the non-PGF group (P = 0.001, P = 0.013, respectively), but none of the clinical outcomes was significantly different.

**Comparisons of the Nutritional Data**

Infants with PGF had a longer start time for the parenteral AA (P = 0.029), a smaller day of age at the end of the TN (P = 0.045), a lower AA intake during the PN phase (P = 0.01), and a lower total protein intake (PN + EN) during the TN phase (P = 0.007) than those without PGF. No significant difference was found in feeding types or other nutritional data during the TN phase (Table 3).

During the TN phase, with the increase in the enteral feeding volume, the energy and protein provided by the EN gradually decreased, and the energy and protein provided by the EN gradually increased, as shown in Figure 1. When the enteral feeding volume reached 80 ml/kg/d, the protein and energy provided by the PN was equivalent to that provided by the EN. Total protein (parenteral + enteral) during the TN phase was 3.16 (2.89, 3.47) g/kg/d, which gradually decreased with the increase in the enteral feeding volume. The total energy (parenteral + enteral) intake during the TN phase was 115.72 (106.98, 122.60) kcal/kg/d.

Infants in the PGF group had a lower parenteral AA intake throughout the TN phase than those in the non-PGF group. Before the enteral feeding volume reached 80 ml/kg/d, the infants in the PGF group had a lower parenteral AA intake [2.85 (2.44, 3.20) g/kg/d vs. 3.07 (2.60, 3.46) g/kg/d, P = 0.025, respectively] and a lower enteral protein intake [0.82 (0.77, 0.89) g/kg/d vs. 0.84 (0.81, 0.92) g/kg/d, P = 0.03, respectively]. After the enteral feeding volume reached 80 ml/kg/d, infants in the PGF group had a lower parental AA intake [1.30 (1.01, 1.69) g/kg/d vs. 1.40 (1.20, 1.68) g/kg/d, P = 0.048, respectively]. No significant difference was found in the parenteral or enteral energy intake before or after the enteral feeding volume reached 80 ml/kg/d (P > 0.05).

**Predictors of Postnatal Growth Failure in the First 28 Days of Life**

We performed a multivariate logistic regression analysis of the nutritional predictors of PGF during the first 28 days of life (Table 4). Three variables were kept in the model (P < 0.001):
the total protein intake (PN + EN) during the TN (P = 0.005), the day of age at the end of the TN (P = 0.006), and the birth weight z-score (P = 0.017). For every 0.1 g/kg/d increase in the total protein intake during the early TN phase, the risk for 28-day growth failure decreased by 8.3%.

**DISCUSSION**

This study analyzed the nutrition data of very preterm infants from six NICUs, excluding data from preterm infants who were still in the TN phase at their 28th day of life. Analysis of the relationship between nutrition intake during the TN phase and the growth of very preterm infants during the first 4 weeks of life showed that lack of protein during the TN phase was associated with growth failure during the first 28 days. This finding differs from those of previous studies on the nutrition and growth of preterm infants (1, 10, 19, 26). Previous studies have reported that achieving adequate nutritional intake during the first few weeks of life is challenging for preterm infants, and a potential strategy to solve this problem is to start parenteral nutrition early, i.e., shortly after birth (31, 32). Our study found that preterm infants without PGF started parenteral nutrition earlier than infants with PGF did, and their amino acid intake during the PN phase was higher; however, neither factor predicted growth failure during the neonatal period. In our study, the nutrition of preterm infants was divided by nutritional phase rather than by postnatal period. We found that protein deficiency was most likely to occur in the TN phase. Therefore, total protein intake (PN + EN) during the TN can predict growth failure during the neonatal period, while total energy intake has no obvious correlation with growth during the neonatal period.

The median duration of the TN phase was 10 days, which is consistent with data from previous studies (33). The older the day of age at the end of the TN phase, the lower the risk of 28-day growth failure, which may be related to lack of enteral protein intake during the TN phase, resulting in a longer period of PN intake. Furthermore, our study found that the birth weight z-score was negatively correlated with the GV because the preterm infants with a lower birth weight z-score often had a higher parenteral amino acid intake during the PN-only and TN phases. This finding is consistent with the negative correlation between birth weight z-score and changes in the birth weight z-scores reported by Izquierdo et al. (2).

The introduction and promotion of the enteral feeding of preterm infants with NEC is a major modifiable risk factor (34), which undoubtedly affects the outcomes of preterm infants with NEC (35). Despite newer and gentler modes of ventilation, BPD remains a major cause of neonatal mortality and morbidity (36). Preterm infants with BPD often show better postnatal growth because of continued PN (37). Therefore, NEC, BPD, and the other clinical factors analyzed in this study may be related to the growth of preterm infants. Our study population included very preterm infants who had completed the TN phase during the neonatal period and achieved full enteral feeding. The inclusion criteria precluded neonates with severe clinical outcomes during the neonatal period; therefore, no adverse outcomes (e.g., NEC or BPD) were identified as risk factors in the multivariate analysis of 28-day growth failure.

In a systematic review of GV calculations, the most commonly used measurement for calculating growth was g/kg/d (38). In our study, we found that birth weight was an important variable affecting PGF. However, the GV was based on an exponential calculation combined with the time to regain birth weight (39). The formula itself contained two important variables: birth weight and the time to regain birth weight, which were beneficial for our growth evaluations (40, 41). Therefore, it is more accurate and reasonable to conclude that insufficient protein intake during the TN phase can predict growth failure during the neonatal period.

In our study, the TN phase was defined as 30 ml/kg/d ≤ the enteral feeding volume ≤ 120 ml/kg/d, and could be divided.

### Table 2: Comparisons of baseline characteristics and clinical outcomes.

| Characteristic                      | Total (n = 268) | GV ≥ 15 g/kg/d (n = 71) | GV < 15 g/kg/d (n = 197) | P-value |
|------------------------------------|----------------|-------------------------|--------------------------|---------|
| Birth weight z-score               | 0.42 (–0.08, 0.89) | 0.29 (–0.39, 0.70) | 0.46 (0.03, 0.95) | 0.013*  |
| Birth weight, median (IQR), g      | 1,290 (1,160,1,490) | 1,210 (1,110,1,380) | 1,330 (1,170,1,525) | 0.001a  |
| Gestational age, median (IQR), weeks | 29 (28,30)   | 29 (28,30)   | 29 (28,30)   | 0.145a  |
| Male sex, n (%)                    | 133 (49.6)    | 44 (61.9)    | 89 (45.1)    | 0.015b  |
| Small for gestational age, n (%)   | 2 (0.75)      | 1 (1.41)     | 1 (0.51)     | 0.460c  |
| Time to regain birth weight, median (IQR), days | 9 (7, 11) | 8 (7, 11) | 9 (7, 11) | 0.518c  |
| NEC (≥ grade 2), n (%)             | 18 (6.72)     | 6 (8.45)     | 12 (6.09)    | 0.497c  |
| IVH (≥ grade 3)/PVL, n (%)         | 17 (6.34)     | 3 (4.23)     | 14 (7.11)    | 0.394c  |
| Confirmed sepsis, n (%)            | 20 (7.46)     | 8 (11.27)    | 12 (6.09)    | 0.156c  |
| Invasive mechanical ventilation, n (%) | 59 (22.01) | 15 (21.13) | 44 (22.34) | 0.835c  |

IQR, interquartile range; BPD, bronchopulmonary dysplasia; NEC, necrotizing enterocolitis; IVH, intraventricular hemorrhage; PVL, periventricular leukomalacia; GV, growth velocity; PGF, postnatal growth failure.

*Mann-Whitney U-test; b Fisher’s exact test; c Chi-square test.

P < 0.05, significant difference between groups.
The phase before the volume of enteral feeding that reached 30 ml/kg/d was the PN phase. In our study, the median of the amino acid intake during the PN phase was above 3.5 g/kg/d, which met the CSPEN Guideline’s recommendation (20); however, the intake of the non-PGF group was significantly higher than that of the PGF group during this phase. Hence, a higher amino acid infusion may be beneficial to the growth of the newborn during the PN phase.

During the following TN phase, the median total protein intake gradually decreased; it was <3.5 g/kg/d. Moreover, the PGF group had a lower parenteral AA intake throughout the TN than the non-PGF group. Thus, the common phenomenon of insufficient protein intake during the TN phase predicted the 28-day growth failure, which further explains the association of insufficient protein intake during the TN with poor growth during the neonatal period. Our study found the cut-off point between the early and late stages of the TN phase (i.e., when the amount of enteral feeding was 80 ml/kg/d and the protein...
intake during the PN phase was greater than that of the EN phase, and after the volume of enteral feeding was 90 ml/kg/d and the protein intake of the EN phase gradually exceeded that of the PN phase) reflected the current situation of insufficient protein intake throughout the TN for very preterm infants. Therefore, during the TN phase, the total protein intake should reach 3.5 g/kg/d to promote better growth among preterm infants. Although human milk feeding has a lower protein content...
than formula feeding, human milk can promote the deposition of fat-free substances, resulting in better neurodevelopment. Therefore, human milk is still the preferred choice for preterm infants during the late TN phase (42). In order to increase the total protein intake while considering the individual differences of preterm infants, human milk fortifiers can be used as supplements, as appropriate, when the human milk volume reaches 50-100 ml/kg/d (14). At the same time, regulating the composition of the standardized PN solutions during the TN phase, including carbohydrates, fats, and amino acids, can achieve the goal of optimal nutrient intake to improve the growth of preterm infants during the neonatal period (13, 14). Moreover, considering the high mortality rate caused by hyperglycemia in extremely low-birth-weight infants, the optimal composition of the early PN phase to avoid PGF must be balanced against the risk for hyperglycemia (9).

One of the strengths of our study is its multicenter cohort with a large sample size. In addition, few studies have examined the relationship between TN nutrition, and early growth of preterm infants. We also reported the current situation of insufficient protein intake during the TN phase. However, there are some limitations. First, the growth assessment was based only on the infant's weight and the time to regain birth weight, without considering the infant's length or head circumference. Second, the study's retrospective observational design and lack of randomization limits the interpretation of the results. Furthermore, standardized nutrition guidelines for the TN phase had not been established before the study. Thus, future studies should examine the associations between the infants’ nutrition during the TN phase (including carbohydrates, fats, and amino acids) and their long-term prognoses.

**CONCLUSION**

The TN from PN to EN is a critical period for maintaining adequate growth among preterm infants. Our study showed that protein intake during the TN phase could predict growth during the neonatal period, suggesting that neonatal physicians should pay more attention to infants’ total protein intake (PN + EN) during the TN phase, especially parenteral amino acid intake.

**REFERENCES**

1. McKenzie BL, Edmonds L, Thomson R, Haszard JJ, Houghton LA. Nutrition practices and predictors of postnatal growth in preterm infants during hospitalization: a longitudinal study. *J Pediatr Gastroenterol Nutr.* (2018) 66:312–7. doi: 10.1097/MPG.0000000000001747.
2. Izquierdo Renau M, Aldecoca-Bilbao V, Balcells Esponera C, Del Rey Hurtado de Mendoza B, Iriondo Sanz M, Iglesias-Platas L. Applying methods for postnatal growth assessment in the clinical setting: evaluation in a longitudinal cohort of very preterm infants. *Nutrients.* (2019) 11:E2772. doi: 10.3390/nu11127772.
3. Luu TM, Rehman Mian MO, Nuyt AM. Long-term impact of preterm birth: neurodevelopmental and physical health outcomes. *Clin Perinatol.* (2017) 44:305–14. doi: 10.1016/j.clp.2017.01.003.
4. Andrews ET, Ashton JJ, Pearson F, Beattie RM, Johnson MJ. Early postnatal growth failure in preterm infants is not inevitable. *Arch Dis Child Fetal Neonatal Ed.* (2019) 104:F235–41. doi: 10.1136/Archdischild-2018-315082.
5. Maas C, Franz AR, von Krogh S, Arand J, Poets CF. Growth and morbidity of extremely preterm infants after early full enteral nutrition. *Arch Dis Child Fetal Neonatal Ed.* (2018) 103:F79–81. doi: 10.1136/archdischild-2017-312917.
6. Thoene M, Anderson-Berry A. Early enteral feeding in preterm infants: a narrative review of the nutritional, metabolic, and developmental benefits. *Nutrients.* (2021) 13:2289. doi: 10.3390/nu13072289.
7. Törer B, Hanta D, Özdemir Z, Çetinkaya B, Gülcan H. An aggressive parenteral nutrition protocol improves growth in preterm infants. *Turk J Pediatr.* (2015) 57:236–41.
8. Roelants JA, Joosten K, Van D, Huist JM, Reiss I, Vermeulen MJ. First week weight dip and reaching growth targets in early life in preterm infants. *Clin Nutr.* (2018) 37:1526–33. doi: 10.1016/j.clnu.2017.08.023.
9. Stensvold HJ, Strommen K, Lang AM, Abrahamsen TG, Steen EK, Pripp AH, et al. Early enhanced parenteral nutrition, hyperglycemia, and death among extremely low-birth-weight infants. *JAMA Pediatr.* (2015) 169:1003–10. doi: 10.1001/jamapediatrics.2015.1667.
10. Stephens BE, Walden RV, Gargus RA, Tucker R, McKinley L, Mance M, et al. First-week protein and energy intakes are associated with 18-month developmental outcomes in extremely low birth weight infants. *Pediatrics.* (2009) 123:1337–43. doi: 10.1542/peds.2008-0211.
11. Sammallahiti S, Kajantie E, Matinlinna HM, Pyhältö R, Lahti J, Heinonen K, et al. Nutrition after preterm birth and adult neurocognitive outcomes. *PLoS ONE.* (2017) 12:e0185632. doi: 10.1371/journal.pone.0185632.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/supplementary material, further inquiries can be directed to the corresponding author/s.

**ETHICS STATEMENT**

This study was approved by the Research Ethics Committee of the Women's Hospital of Nanjing Medical University (Ethics Number: Ning Fulun [2016] No. 73). Written informed consent from the participants’ legal guardians was not required for participation in this study, in accordance with national legislation and the institutions’ requirements.

**AUTHOR CONTRIBUTIONS**

NW, ZY, and JZ contributed to conception and design of the study. NW and SH contributed to the acquisition and analysis of the data. BW, RC, LG, XL, HW, YG, and WH performed the statistical analysis. NW wrote the first draft of the manuscript. All authors contributed to manuscript revision, read, and approved the submitted version.

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