Improved Outcomes in Paediatric Intestinal Failure with Aggressive Prevention of Liver Disease

D. Sigalet, D. Doctor, M. Robertson, V. Lam, M. Brindle, K. Sarkhosh, L. Driedger, M. Sajedi

1 Alberta Children’s Hospital/University of Calgary, Pediatric General Surgery, Calgary, Canada
2 Alberta Children’s Hospital, Department of Pediatrics, Calgary, Canada
3 Alberta Children’s Hospital, Pediatric Surgery, Calgary, Canada
4 Alberta Children’s Hospital, Pediatric General Surgery, Calgary, Canada

Abstract

Background/Purpose: A protocol-driven care algorithm for the care of intestinal failure (IF) centred on therapies to prevent Parenteral Nutrition Associated Cholestasis (PNAC) was instituted in 2006. We report our results from 2006–2009, and compare them to the outcomes of our previous cohort of patients (1998–2006).

Methods: With regional ethics board approval, we have been prospectively gathering data on patient with IF cared for by our regional surgical unit. IF was defined as a residual bowel length of <40 cm or a requirement for PN for greater than 60 days. With the development of a multidisciplinary care team, a protocol-driven strategy to prevent PNAC was instituted in 2006, with aggressive introduction of enteral feeds, use of prophylactic antibiotics to prevent bacterial overgrowth, lipid reduction and use of a fish oil-derived lipid preparation for cholestasis and Serial Transverse Enteroplasty (STEP) if bowel dilation occurred.

Results: In the era from 1998–2006, 33 patients were identified, with a 72% survival; the direct bilirubin averaged 112 ± 34 μM/L after 3 months of PN. 8/33 (27%) of patients received prophylactic antibiotics, and none received fish oil-based lipids. The most common causes of IF were gastrochisis (30%) and atresia (21%); 31 of 33 patients were infants. Average time to intestinal rehabilitation/death was 4.5 ± 3 months. All deaths were related to sepsis or PN/liver failure. In the era from 2006–2009, 22 patients have been followed, with 100% survival. Average bilirubin after 3 months of PN was 8 ± 2.2 μM/L, 20/22 (90%) received prophylactic antibiotics, and 6/22 (27%) received fish oil-based lipid PN. The common causes of IF were gastrochisis 15/22 (68%) and atresia (27%). 18/22 are weaned from PN, and the average time to intestinal rehabilitation was 2.7 ± 1.3 months, 4 patients underwent STEP procedures, (p < 0.05 by Fischer’s exact or Student’s t-test, data mean ± SD).

Conclusions: The institution of an aggressive protocol of advancing enteric feeds, oral antibiotic prophylaxis for bacterial overgrowth, fish oil-based lipid use, and the STEP procedure for dilated bowel has resulted in an apparent increase in survival and a remarkable improvement in liver function in a paediatric IF population. Further studies to define the relative importance of these therapies are recommended.

Introduction

Intestinal failure is a challenging clinical problem in the paediatric population. The ability to provide appropriate calories via parenteral nutrition provides a safety net for the clinician [1,2]. However in the infant population this is a double-edged sword. Historically, the use of parenteral nutrition for periods greater than 6–8 weeks would predictably result in the onset of parenteral nutrition-associated liver disease (PNALD) [3,4]. In the past, the complication of progressive hepatic fibrosis and liver failure was a constant threat for these patients. There is an inverse relationship between the length of bowel and the frequency of liver failure [1,5–8]. This is further exacerbated by episodes of line or bacterial overgrowth-induced sepsis [9–11]. In this context the ability to provide nutrition by the enteral route is the ultimate treatment for PNALD [12,13]. However, in the drive to provide calories by the enteral route the potential for promoting bacterial overgrowth within the small bowel is a continual threat. Episodes of sepsis definitely increase the rate and the severity of liver disease. Thus the clinical care of these
patients consists of a series of small steps to increase enteral nutrition while continually watching for problems with bacterial overgrowth, or recurrent necrotising enterocolitis.

Certain key observations by clinical groups over the recent years have helped refine the general strategy for the care of these patients. Georgeson observed that the aggressive use of rotating antibiotics appeared to reduce the risk of gut-driven sepsis, and also had a beneficial or protective effect on the progress of liver disease [14]. While other groups have had variable results, this concept is theoretically quite logical. It has been shown that the short gut has increased permeability; factors which increase the growth of bacteria within the intestinal lumen increase the exposure of the liver either to intact bacteria or to their breakdown components, which then may worsen cholestasis [1,9,15].

In addition, it has been suggested that the provision of calories in the form of fat may also contribute to the development of liver disease. This has led to suggestions of using a lipid sparing or lipid reduction strategy, if cholestasis is apparent. Georgeson observed that the aggressive use of rotating antibiotics appeared to reduce the risk of gut-driven sepsis, and also had a beneficial or protective effect on the progress of liver disease [14]. While other groups have had variable results, this concept is theoretically quite logical. It has been shown that the short gut has increased permeability; factors which increase the growth of bacteria within the intestinal lumen increase the exposure of the liver either to intact bacteria or to their breakdown components, which then may worsen cholestasis [1,9,15].

Finally, in terms of positive therapies to improve the tolerance of enteral feeds, the use of a serial transverse enteroplasty (STEP) procedure to increase the useful surface area, while coincidentally reducing the tendency to bacterial overgrowth, has shown benefits in selected patients [20,21].

Our surgical unit has had a long-standing interest in the treatment of Short Bowel Syndrome. In the years between 1998 and 2006, despite this interest, there was only limited coordination of the care of infants following intestinal resection or gastrochisis repair. Nutritional support was provided, with attention focused on the provision of adequate calories, primarily by PN, and the institution of enteral feeds was primarily at the discretion of the attending surgeon. Coincident with a move of the hospital in 2006, we were able to institute a cohesive multidisciplinary team approach (CHIRP) to care for patients with intestinal failure [6,22]. This occurred in October 2006. All infants undergoing intestinal resection or repair of gastrochisis were under the care of the CHIRP team, as well as older children on long-term PN (longer than 60 days).

At that time, a more aggressive policy of increasing enteral feeds and the use of rotating antibiotics with a promotility agent was instituted (www.ucalgary/pediatricsurgery/CHIRP protocol). In brief, nutrition was initially provided as TPN, following conventional guidelines for total calories and the allocation of fat, protein and carbohydrates [6,23]. This included the provision of calories appropriate for age and weight, following a “balanced” PN approach with fat prescribed at a rate of 2–3 g/kg per day given as a conventional 20% Intralipid emulsion (Fresenius Kabi Canada, Toronto, ON, Canada). If biochemical signs of liver failure ensued (conjugated bilirubin over 50 μmol/L in infants older than 2 months of age), then an aggressive lipid reduction strategy was started, with a reduction in intravenous lipids (Intralipid, 1.0 g/kg/day, Fresenius Kabi, Toronto, ON, Canada). If bilirubin climbed above 60 μmol/L, then Intralipid was discontinued, and a regimen using a fish oil-based lipid preparation (Omegevan, Fresenius Kabi, by special access, Health Canada) was instituted (1.0 g/kg/day), with monthly monitoring of serum essential fatty acid levels.

Patients underwent routine imaging of the small intestine with an upper GI barium study and follow through, every 6 months for the first 2 years, and annually thereafter, to monitor for the development of distended “overadapted” segments. Imaging was also done if there was a loss of tolerance of enteral feeds or an episode of sepsis with intestinal organisms. If areas of dilated bowel were detected in the setting of a loss of tolerance of enteral feeds, then a STEP procedure was recommended to families. In all cases where this was suggested, the families agreed to surgery.

This report details our experiences in the years 1998–2006 as one distinct phase, in which a “conventional” approach to PN and enteral therapy was used. This is contrasted with the second phase from 2006 to early 2009, when the more aggressive, team-based approach to management was begun, with a focus on increasing enteral feeds, and preventing cholestasis and liver disease. The change in strategy forms a natural break point for comparisons.

Methods

Study Population

Infants referred for surgical and nutritional care to the Alberta Children’s Hospital between 1 January 1998 and September 2006 form the first cohort in the study, while infants cared for during the time from October 2006 to January 2009 form the second cohort. Data on the original patient demographics, disease type, therapy and progress of feeding was gathered prospectively. Informed consent was obtained from the subject’s parents. The protocol for gathering data was approved by the Conjoined Regional Health Ethics Board (protocol 16506).

Definition of Terms

Intestinal failure was defined to have occurred in any child undergoing a laparotomy who 1) was left with less than 40 cm small bowel length, or 2) required parenteral nutrition after

laparotomy for more than 60 days. This follows our original definition proposed for surgical cases, but increases the time required for PN support as suggested in recent NIH Paediatric Intestinal Failure Consortium Guidelines (PIF-Con) [4]. Patients with major cardiac or chromosomal anomalies that precluded initial surgical management or prevented an approach with conventional care were excluded. Patients dying in the acute phase of NEC (before considering the start of enteral feeds) were also excluded.

Data Collection
In the first cohort, from 1998 to 2000, patients were identified by a review of the parenteral nutrition records of the pharmacy, and demographic and feeding data were collected retrospectively. From 2000–2005, patients were identified and followed up prospectively; some elements of data collection were done prospectively, and some elements (lab values and feeding protocols) were collected prospectively. In the recent cohort, data was collected prospectively from the weekly patient chart reviews and from hospital records. Data was entered on a separate spreadsheet and analyzed using Prism statistical software (Prism, La Jolla, CA, USA). Outcomes were calculated from the time of the original surgery.

Data collection included initial patient demographics, gestational age, birth weight, and the primary pathology. The surgical procedure was recorded and the length of bowel measured. The percentage of measured bowel length compared with the predicted bowel length based on a published algorithm was calculated [24]. In cases of gastroschisis, an accurate measure of bowel length could not be performed, and so the pre-resection length was assumed to be equivalent to the norms for gestational age and weight.

Data relating to the ongoing clinical status, including tolerance of enteral feeds, were recorded weekly. Biochemical parameters were also collected weekly as part of the routine of parental nutritional monitoring, and included conventional blood counts, electrolytes and liver function tests.

The use of medications including promotility agents, rotating antibiotics, and other medications was recorded. Operative details regarding the STEP procedure, if performed, were also recorded.

Results

The majority of patients treated over both time periods were infants, presenting with intestinal failure in the neonatal period (Table 1). There were no significant differences in the patients’ average age, weight or gestational age at surgery. There was a difference in the number of patients presenting with gastroschisis, which increased in the second phase, with a corresponding reduction in the number of babies having necrotising enterocolitis.

The changes in treatments used, according to era, are listed in Table 2. There was a clearly significant increase in the use of rotating antibiotics, as well as of the strategies invoked to preserve liver function, that is lipid reduction, and the use of a fish oil-based lipid preparation in the recent cohort. Finally, use of the STEP procedure has increased in the recent era.

Outcomes are listed in Table 3. The most striking change is in survival, which improved significantly in the recent cohort (2006–2009: survival was 22 out of 22 vs. 1998–2005: 24 out of 33, Fig. 1). Although the length of follow up was necessarily longer in the first cohort, most deaths occurred within the first 15 months of follow up. The cause of death in all cases was end-stage liver failure and sepsis. In the only older child in the 1998–2006 cohort who died after 3 years of PN, liver failure was attributed to the soybean-derived lipid preparation.

The development of liver failure also differed significantly between the two groups (Fig. 2). In the treatment era from 1998–2006, the average direct bilirubin after 3 months of PN was 112 ± 34 μM/L; in the recent era it was 8 ± 2 μM/L (p < 0.01 by Student’s t-test). No patients in the recent cohort developed direct bilirubin which stayed above 100 μM/L, while in the 1998–2006 cohort roughly 25% of all patients on PN for longer than 25 weeks developed this biochemical marker of liver failure. This difference was clearly related to the change in nutritional support and PN treatment in the recent cohort, as the groups were similar in age and diagnosis. No other changes with regard to surgical or medical aspects of therapy were made dur-
ing this time frame. The improvement in liver preservation does not appear to be due to an increased ability to induce intestinal adaptation and the corresponding ability to stop PN therapy. Although there was a reduction in the time required to achieving independence from PN in the recent cohort, the reduction was not statistically significantly different from the earlier time period (Fig. 3).

Finally, given the historical nature of the controls the average type of follow up is much longer in the 1998–2006 treatment group. However it is important to note that there was only one case of liver failure in this group which did not present in the group. However it is important to note that there was only one type of follow up is much longer in the 1998–2006 treatment period.

This was the unusual case of liver failure caused by macrophage activation syndrome acquired in infancy, the time to successful adaptation typically takes 16 weeks, and only occurs after complete discontinuation of PN [13]. However, there may be some improvement in the rate of adaptation in the recent cohort of patients, as shown by the Kaplan-Meier curve of the time to reaching intestinal autonomy (Fig. 3; p < 0.06).

In examining each of the elements of therapy in the modern era, the fundamental reasons for the improved outcome are not evident, and likely is to be due to a combination of factors. The use of continual drip feeds is likely important. Anecdotally we have observed that in situations where infants are managed with intermittent oral feeds or oral plus NG top up, as the volumes are observed that in situations where infants are managed with intermittent oral feeds or oral plus NG top up, as the volumes are reached. The reasons for this are not clear, however it is interesting to note that observations in both the lab setting and in our human patients show that continuous feeds appear to result in a maximal stimulation of GLP-2 levels [27–29]. In turn, continuous elevation of GLP-2 levels have been shown to maximally induce adaptation [28,29]. To preserve oral feeding skills, we routinely start to encourage oral feeds when the infant is held for 2 h, and this volume is given by bottle with a gavage top up, twice daily. This maintains oral skills, and facilitates transitioning once the infant tolerates near full feeds Secondly, the use

These results support the utility of aggressive multidisciplinary treatment of intestinal failure with attention to bacterial overgrowth, and the modification of conventional parenteral nutrition using a lipid reduction/fish oil-based strategy if hyperbilirubinaemia occurs.

The observed improvement in long-term survival, as well as the improved preservation of liver function are the most important findings of this study (Figs. 1, 2). The ability to preserve normal liver function over the long term has a number of critically important benefits for this patient population. Firstly, the cause of death in all the historical cohort patients was related to liver failure and sepsis. As noted, in one patient the development of liver failure was associated with macrophage activation syndrome, but the ultimate cause of death was liver failure and sepsis. All other cases had straightforward hepatic fibrosis and cirrhosis. This resulted in the end-stage liver failure with ascites, poor hepatic synthetic function, and repeated cycles of bacterial and fungal sepsis. Several patients were listed for intestinal or combined intestinal-liver transplantation but did not receive grafts because of the lack of appropriate donors [9,22,25].

The ability to maintain liver function in this patient population is critical to their survival, and quality of life. The time required for intestinal adaptation in these patients ranged from 4.5 months in the initial cohort to 2.7 months in the more recent cohort; these results are very similar to other recent reports in similar populations with mixed causes of post-surgical intestinal failure [10,11,26]. In the most severe cases of short bowel syndrome acquired in infancy, the time to successful adaptation averaged 2.6 years [8]. In our historical cohort, all patients in this category had succumbed to liver-related disease long before this adaptation would have been expected to occur (Table 3).

The improved outcome in the modern era was not related to the simple increase in the speed of adaptation; all of the 6 patients who were started on the lipid reduction/omegaven therapy had rapid normalisation of liver function (within 10 weeks in all cases), despite the fact that only 2 of them came off PN in that time. Historically, resolution of PN-associated hyperbilirubinaemia typically takes 16 weeks, and only occurs after complete discontinuation of PN [13]. However, there may be some improvement in the rate of adaptation in the recent cohort of patients, as shown by the Kaplan-Meier curve of the time to reaching intestinal autonomy (Fig. 3; p < 0.06).

In examining each of the elements of therapy in the modern era, the fundamental reasons for the improved outcome is not evident, and likely is to be due to a combination of factors. The use of continual drip feeds is likely important. Anecdotally we have observed that in situations where infants are managed with intermittent oral feeds or oral plus NG top up, as the volumes are slowly increased, the speed of adaptation seems to be somewhat slower than those managed with continuous NG feeds until full feeds are reached. The reasons for this are not clear, however it is interesting to note that observations in both the lab setting and in our human patients show that continuous feeds appear to result in a maximal stimulation of GLP-2 levels [27–29]. In turn, continuous elevation of GLP-2 levels have been shown to maximally induce adaptation [28,29]. To preserve oral feeding skills, we routinely start to encourage oral feeds when the infant is receiving more than 10cc/h of EBM or formula, the feeds are held for 2 h, and this volume is given by bottle with a gavage top up, twice daily. This maintains oral skills, and facilitates transitioning once the infant tolerates near full feeds Secondly, the use

**Discussion**

This study shows a distinct change in the incidence of liver failure and an improvement in survival in paediatric patients on long term parenteral nutrition from the initial cohort (1998–2006) and patients treated in the recent era, from 2006–2009.
of rotating antibiotics has long been suggested to be useful in preventing PNALD [14]. The mechanism underlying this is not clear, however it has been postulated that ongoing bacterial overgrowth affects liver function via systemic, and especially portal, release of cytokines, this is turn may worsen the progress of liver disease [13,18,29]. This appears to have the greatest impact in infants younger than six months of age; older infants appear much less susceptible to this induced tendency to liver failure.

Overall, these results support the use of an aggressive pattern of rotating antibiotics in order to suppress small intestine bacterial overgrowth. This is particularly relevant because of the intentionally high enteral load of nutrients, in a condition of congenitally dilated, post-surgical alterations in motility, all of which will encourage the constitutive increase of small intestinal flora.

Fundamentally, the most likely cause of the improved outcomes was the use of lipid reduction/fish oil-based therapy. It is notable that while lipid reduction was used in infants whose direct bilirubin was increasing over 34μmol/L, this was not in itself adequate in the majority of cases [19]. Accordingly, if the bilirubin had not dropped significantly within 2–3 weeks, Omegaven therapy at 1 g/kg/day was initiated [19]. The average time to resolution of hyperbilirubinemia in these patients was 6.2 weeks; in 4 cases we were not able to advance enteral nutrition beyond 20% of total calories during that same time period. Thus we feel the benefit is primarily due to the use of the fish oil-based lipid preparation.

Importantly, we monitored the fatty acid profile using the serum triene:tetraene ratios. In 2 out of 3 long-term PN patients maintained exclusively with Omegaven for longer than 6 months, lipid profile abnormalities and clinical rashes suggestive of essential fatty acid deficiency, were detected. Once this was detected, the patients were supplemented with Intralipid as 50% of total IV fat and both the biochemical and clinical evidence of fatty acid deficiency improved; in the future we plan to switch to a combination of Intralipid/Omegaven after 6 months of PN. It is also important to note that while the cohort of patients treated with Omegaven had a dramatic normalisation of hepatic biochemistry and synthetic parameters, those patients who required long-term PN had persistent fibrosis on serial liver biopsies [30]. This has also been seen in other groups, and indicates the importance of continuing to aggressively increase enteral feeds, even in very short bowel syndrome patients; ultimately the only true “cure” for PNALD is discontinuation of PN [9,17,18].

Finally, the STEP procedure was used in 4 patients in the recent era. This low incidence makes comments difficult; however, our experience is similar to that of other groups [20,21,31]. In these cases, each STEP procedure resulted in a significant improvement in the tolerance of enteral feeds; however, complete intestinal autonomy was only possible in 1 patient. It is important to note that in 2 of these patients a repeat STEP procedure was contemplated because of persistent recurrent dilation of the bowel. Thus we view the STEP procedure as a useful adjunct in the care of these patients. However, the fundamental drive to increase crypt cell proliferation may continue to cause intestinal dilation until the absorptive capacity is sufficient to maintain nutrition enterally [32].

Limitations

This study suffers from the limitation of using historical cohorts. As shown in the demographic data, there has been a clear increase in patients presenting for intestinal failure as a result of gastrochisis (Table 1). This reflects the worldwide increase in the number of babies born with gastrochisis [33,34]. In our population the prevalence of gastrochisis is 1 per 2400 live births. Importantly, the causes of intestinal failure, size, gestational age, and survival of this patient population are very similar to those in a number of recent reports [1,8,10,11,17,35]. Based on this, we suggest that the survival of the previous cohort of patients was similar to that seen in many other contemporary series; the finding of improved outcomes after the combination of a protocol-driven increase in enteral feeds, the use of rotating antibiotics, lipid reduction/fish oil-based lipid preparations and the STEP procedure may be a new standard for the care of this patient population. The results seen in this initial cohort are encouraging and justify continued evaluation.

Conflict of Interest: None

References

1 Goutet O, Ruemmele F. Causes and management of intestinal failure in children. Gastroenterology 2006; 130: S16–S28
9 Kelly DA. Intestinal failure-associated liver disease: What do we know today? Gastroenterology 2006; 130: 570–577