POSITIVE BENEFITS OF BLENDED DIET: WEIGHING IN ON GASTROINTESTINAL DYSTONIA

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Introduction Children with severe neurodisability and tone disorders can present with debilitating symptoms and declining growth. Gastrointestinal dystonia (GID) describes the clinical manifestations of: pain behaviour; hypertonicity; retching; vomiting; vagal phenomena; abdominal distension and straining attributable to the GI tract in the context of severe neurodisability. Blended diet (BD) has an emerging role in the nutritional management of patients with neurodisability, however we are unaware of data pertaining to BD specifically in patients with GID.

Aim A tertiary centre review of outcomes of patients with GID receiving BD.

Method Patients who commenced BD between 07/2017 and 02/2020 were identified from prospectively gathered complex enteral nutrition (CEN) and specialist dietetic databases. BD was initiated with specialist paediatric dietetic support within the CEN clinic or in other clinics. Data gathered included: demographics; primary diagnosis; enteral feeding plan; fundoplication; weight standard deviation z score at 0, 6, 12 and 18 months prior to and from commencing BD; medications at 0 and 6, 12, 18 months; reasons if BD discontinued; parenteral nutrition (PN) requirement.

Results 29 children met criteria for GID and commenced BD. 18 were male. 14 had a fundoplication. Feeding method prior to BD: gastric bolus 17; gastric continuous 8; jejunal 4. Mean age BD commenced was 7 years. Follow up ranged 8 to 40 months. 25 patients continue BD to date. 4 discontinued BD within 1–2 months citing increased GI symptoms or device blockage.

Of the 25 who continue BD, median weight z scores declined from -1.77 at 12 months prior to BD, -1.86 at 6 months prior, -1.94 at 0 months, then rose to -1.54 at 12 months, and -1.40 at 18 months. Mean weight z scores were maintained at -1.97 (CI -0.85 to -3.09) at 0 months, -1.97 (CI -0.27 to -3.02) at 12 months and -1.79 (CI -0.34 to -3.25) at 18 months [figure 1]. This trend was more significant in a subset of 6 patients, z score -2.06 (CI -1.46 to -2.67) at 0 months, -1.13 (CI -0.59 to -1.67) at 12 months and -0.84 (CI -0.46 to -1.21) at 18 months (p = 0.003) [figure 2].

9 of the 25 were able to discontinue one GI medication, 4 discontinued two GI medications. Tone medications: 3 reduced; 20 unchanged; 2 increased. No patients received PN during the study.

Conclusion GID represents the severest end point of gastrointestinal symptoms in neurodisability, with progressive decline often a feature. Our data show children with GID receiving BD continue to track their weight trajectory, whilst some experience significantly improved growth. No patients required trial on PN during the study. The authors advocate for BD’s role in minimizing the need for invasive treatments in GID whilst addressing symptoms and maintaining nutritional status and growth superiorly to formula feeds alone. Improvements in symptomatology and quality of life will be better described by prospective survey of patients commencing BD for GID. The authors are currently gathering this data.

REFERENCES
2. Hay J, et al. (presented BSPGHAN 2020)