Food intake, nutritional status and gastrointestinal symptoms in children with cerebral palsy

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ABSTRACT — Background — Cerebral palsy may be associated with comorbidities such as undernutrition, impaired growth and gastrointestinal symptoms. Children with cerebral palsy exhibit eating problems due to the effect on the anatomical and functional structures involved in the eating function resulting in malnutrition. Objective — The aim of this study was to investigate the association between food intake, nutritional status and gastrointestinal symptoms in children with cerebral palsy. Methods — Cross-sectional study that included 40 children with cerebral palsy (35 with spastic tetraparetic form and 5 with non-spastic choreoathetoid form of cerebral palsy, all requiring wheelchairs or bedridden) aged from 4 to 10 years. The dietary assessment with the parents was performed using the usual household food intake inquiry. Anthropometric data were collected. Gastrointestinal symptoms associated with deglutition disorders, gastroesophageal reflux and chronic constipation were also recorded. Results — The median of height-for-age Z-score (-4.05) was lower (P<0.05) than the median of weight-for-age (-3.29) and weight-for-height (-0.94). There was no statistical difference between weight-for-age and weight-for-height Z-scores. Three patients with cerebral palsy (7.5%) exhibited mild anemia, with normal ferritin levels in two. Symptoms of dysphagia, gastroesophageal reflux, and constipation were found in 82.5% (n=33), 40.0% (n=16), and 60.0% (n=24) of the sample, respectively. The patients with symptoms of dysphagia exhibited lower daily energy (1280.2±454.8 Kcal vs 1890.3±847.1 Kcal, P=0.009), carbohydrate (median: 170.9 g vs 234.5 g, P=0.023) and fluid intake (483.1±294.9 mL vs 992.9±292.2 mL, P=0.001). The patients with symptoms of gastrointestinal reflux exhibited greater daily fluid intake (720.0±362.9 mL) than the patients without symptoms of gastroesophageal reflux (483.7±320.0 mL, P=0.042) and a greater height-for-age deficit (Z-score: -4.9±1.7 vs 3.7±1.5, P=0.033). The patients with symptoms of constipation exhibited lower daily dietary fiber (9.2±4.3 g vs 12.3±4.3 g, P=0.031) and fluid intake (456.5±283.1 mL vs 741.1±379.2 mL, P=0.013) intake. Conclusion — Children with cerebral palsy exhibited wide variability in food intake which may partially account for their severe impaired growth and malnutrition. Symptoms of dysphagia, gastroesophageal reflux, and constipation are associated with different food intake patterns. Therefore, nutritional intervention should be tailored considering the gastrointestinal symptoms and nutritional status.


INTRODUCTION

Cerebral palsy is a chronic non-progressive encephalopathy that is caused by various agents and presents with heterogeneous clinical manifestations. It may be associated with comorbidities such as undernutrition (46% to 90% of the patients), impaired growth, mental retardation, epileptic seizures, communication disorders, visual and auditory defects, and gastrointestinal symptoms, including dysphagia, gastroesophageal reflux, and constipation. Food intake is one of the factors that determine malnutrition. Oropharyngeal dysphagia due to motor dysfunction may reduce the food intake of patients, with consequent malnutrition, pulmonary aspiration, respiratory infection, and chronic lung disease. Gastroesophageal reflux in cerebral palsy is associated with severe complications, such as esophagitis and esophageal dysphagia, and also with reduced food intake. As for constipation, it is believed that alterations in the neural modulation are associated with reduced colonic motility. Other factors may also contribute to the development of constipation, such as severe skeletal deformities, spasm, use of anticonvulsants, low levels of physical activity, and a low-fiber diet.

Children with cerebral palsy exhibit eating problems due to the effect on the anatomical and functional structures involved in the eating function, resulting in reduced energy and nutrient intake and in consequent malnutrition. As a function of malnutrition, the body fat reserves become depleted, the muscle mass is reduced, and immune dysfunction occurs, with consequent increased risk of respiratory and urinary tract infections.

No study published up to the present time has sought to associate nutritional status with food intake and the presence of gastrointestinal symptoms in cerebral palsy. Therefore, the aim of this study was to investigate the correlation of food intake and nutritional status with the presence of gastrointestinal symptoms in children with severe cerebral palsy.

Case series and methods

This cross-sectional study was approved by the Ethics Committee of the São Paulo School of Medicine, Federal University of São Paulo. The study was approved by the Ethics Committee of the São Paulo School of Medicine, Federal University of São Paulo. No funding was received for this study and all authors declare no conflict of interest.

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Paulo (Universidade Federal de São Paulo). Data were collected at the outpatient clinic of Our Home Spiritualist Center – André Luiz Houses (Centro Espírita Nosso Lar – Casas André Luiz), Guarulhos, and at Saint Francis Children’s Home (Lar Escola São Francisco), São Paulo, Brazil. The children’s guardians and institutional representatives were asked to sign an informed consent form.

The sample of this study was of convenience, which comprised 40 children with cerebral palsy, 23 (57.5%) male and 17 (42.5%) female. The patients socioeconomic level was established based on the classification formulated by the Brazilian Association of Market Research Companies (Associação Brasileira de Empresas de Pesquisa – ABEP)(17).

All of the patients had severe cerebral palsy, requiring wheelchairs or bedridden. The age varied from 4 to 10 years that was the selected range for inclusion in the study. All of the patients who met the inclusion criteria and had medical appointments on the days when the study was conducted were invited to participate in the study. According to the Oxford Feeding Study II classification(18), 87.5% (35/40) of the patients had the spastic tetraparetic cerebral palsy and 12.5% (5/40) had the non-spastic choreoathetoid form of cerebral palsy.

Information was registered in a form that included personal, socioeconomic, and anthropometric data, clinical history, gastrointestinal symptoms, feeding route, characteristics of oral motor function, and dietary survey.

The dietary assessment with the parents was performed using the usual household food intake inquiry. The dietary data was collected by one of the authors (Caramico-Favero DCO) who is an experienced nutritionist. Utensils and utensils album images have been used and sipped over the table (homemade measures). Posteriorly homemade measuring was converted to grams and milliliters. Hydric ingestion was calculated from the liquids intake of the diet (water, coconut water, juices, milk, carbonated drinks, jellies and teas). The water used for cleaning the enteral tube or gastrostomy was also computed. The type of enteral diet was also registered (homemade blended, commercial formula or both homemade and commercial). Information about noncommercial enteral feeding was collected. Nutrition calculations were performed using the NutWin Software for Support for Decision-Making in Nutrition version 2.5 by the Federal University of São Paulo(19).

The food intake values were compared to the references in Dietary Reference Intake (DRI) for the following parameters: Estimated Energy Requirements (EER); Estimated Average Requirement (EAR) of energy; Recommended Dietary Allowance (RDA) and Tolerable Upper Intake Level (UL) of carbohydrates and proteins(20). Fiber intake was assessed based on the American Health Foundation’s recommendation, according to which the minimum daily fiber intake (in grams) is equal to the child’s age (in years) plus five(21).

To assess the participants’ nutritional status, their body weight was measured, and their height was estimated. The body weight was assessed using a Toledo® (São Bernardo do Campo, São Paulo, Brazil) 500-kg capacity scale with a 100-g precision platform. This scale allowed the weighing of children on wheelchairs as needed. The children’s current weight was calculated by subtracting the wheelchair weight from the combined child-wheelchair weight; the children were weighed wearing a minimum of clothing and without diapers. In the remaining cases, as the sample characteristically comprised children who cannot stand up by themselves, they were held by an adult and weighed, and their current weight was calculated by subtracting the adult’s weight from the combined child-adult weight. The children’s height was estimated based on the tibia length(22).

The anthropometric parameters weight-for-age, height-for-age, and weight-for-height were analyzed based on Z-scores relative to the reference values established by the World Health Organization (WHO). Values two standard deviations below the corresponding Z-score were considered to be indicative of nutritional deficits(23,24).

To assess the children’s hemoglobin and ferritin levels, blood samples were collected by puncture of a forearm vein, which was performed by a duly trained professional at America Diagnosis Laboratory (Laboratório Diagnósticos da América, Barueri, São Paulo, Brazil). Sample analysis was performed using a Pentra ABX automated analyzer, Horiba Medical (Kyoto, Japan). The diagnosis of anemia and iron deficiency was based on comparing the measured blood hemoglobin and ferritin levels to the reference values formulated by WHO(25).

Symptoms of dysphagia, gastroesophageal reflux, or constipation were recorded in the patients’ individual forms.

The symptoms of dysphagia were evaluated by speech therapists based on reported occurrence of cough, drooling, or choking; orofacial motor features registered during the study in ad hoc form; utensils used for feeding; and consistency of the food consumed(26).

The presence of nausea, vomiting, and regurgitation was considered symptoms of gastroesophageal reflux(27). Constipation was defined as pain and/or strain to pass stools combined with fragmented and hard or cylindrical stools with diameter larger than a hotdog sausage with cracks in the surface, and/or two or fewer defecations per week. Having only two or fewer isolated bowel movements per week was considered to be indicative of constipation. Patients who did not meet any of those criteria but used laxatives were also characterized as having constipation. This definition was adapted from previous studies that assessed constipation in children with neurological diseases(10,28,29). The symptoms of gastroesophageal reflux and constipation were assessed by a gastroenterologist.

The results were analyzed by means of parametric and non-parametric statistical tests depending on the distribution of the variables and are described together with the results. Analysis was performed using the software SigmaStat version 3.5. (Systat Software, San Jose, California, USA)(30) and Epi-Info version 3.2.2 (Atlanta, GA, USA)(31). The value set to reject the null hypothesis was 0.05 or 5%.

RESULTS

The sample comprised 40 children with cerebral palsy with mean age of 6.7±2.4 years old, of whom 33 (82.5%) were fed by the oral route, five (12.5%) via a gastrostomy tube, and two (5.0%) by the oral route combined with a nasogastric or gastrostomy tube. According to Friedman’s test followed by Dunn’s multiple comparison test, the height-for-age Z-score of the 40 children with cerebral palsy (median = -4.05; 25th and 75th percentiles: -5.30 and -2.89) exhibited a greater deficit (P<0.05) than weight-for-age (median = -3.29; 25th and 75th percentiles: -3.95 and -2.28) and weight-for-height (median = -0.94; 25th and 75th percentiles: -2.06 and 0.12). Weight-for-age and weight-for-height did not reach statistical significant difference (P>0.05).

TABLE 1 describes the data on daily carbohydrate, protein intake, distributed according to the RDA and EAR for age and gender. The protein and carbohydrate intake was above the RDA in 92.5% (37/40) and 85.0% (34/40) of the participants, respectively.
TABLE 1. Carbohydrate and protein intake according to Estimated Average Requirement (EAR), Recommended Dietary Allowance (RDA), and Tolerable Upper Intake Level (UL) of children with cerebral palsy.

<table>
<thead>
<tr>
<th>Nutrient</th>
<th>&lt; 2 SD of EAR</th>
<th>From -2 SD of EAR to EAR</th>
<th>Within EAR and RDA</th>
<th>&gt; RDA</th>
<th>&gt; UL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbohydrates* (g/day)</td>
<td>1 (2.5%)</td>
<td>1 (2.5%)</td>
<td>4 (10.0%)</td>
<td>34 (85.0%)</td>
<td>–</td>
</tr>
<tr>
<td>Protein* (g/day)</td>
<td>–</td>
<td>–</td>
<td>3 (7.5%)</td>
<td>37 (92.5%)</td>
<td>–</td>
</tr>
</tbody>
</table>

*Mean intake (standard-deviation) of carbohydrates and protein were, respectively, 199.6±85.3 and 48.6±17.9 g.

TABLE 2 compares the anthropometric and dietetic data, nutritional indexes and the intake of energy, macronutrients, fiber, and fluids as a function of the presence or absence of symptoms of dysphagia. The mean energy intake was significantly lower among the children with clinical signs of dysphagia (P=0.009). In regard to the macronutrients, the children with dysphagia exhibited a greater intake of carbohydrates and protein and lower intake of fat, but only the carbohydrate intake exhibited a significant difference (P=0.023). Fluid intake was significantly lower among the children with dysphagia (P=0.001).

TABLE 3 compares the anthropometric and dietary indicators as a function of the presence or absence of clinical manifestations compatible with gastroesophageal reflux disease. The children with signs and symptoms suggestive of gastroesophageal reflux disease exhibited greater fluid intake (P=0.042).

TABLE 4 compares the nutritional and dietary data as a function of the presence or absence of clinical evidences of constipation. The children with symptoms of constipation exhibited significantly lower daily intake of fiber (P=0.031) and fluids (P=0.013).

As TABLES 2, 3 and 4 show, the anthropometric parameters did not differ between the children with or without gastrointestinal symptoms, except, for patients with gastroesophageal symptoms who presented lower height-for-age Z-score.

The hemoglobin and ferritin levels shown that three male parients (7.5%) exhibited mild anemia, with normal ferritin levels in two. No child exhibited ferritin reduction not accompanied by a decreased hemoglobin level. The mean iron intake of the sample was 11.0±6.0 mg/daily.
TABLE 4. Comparison of anthropometric nutritional parameters and daily energy, macronutrient, fiber and fluid intake between children with cerebral palsy with or without constipation.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Constipation</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Yes (n=24)</td>
<td>No (n=16)</td>
</tr>
<tr>
<td>Height-for-age Z-score*</td>
<td>-4.4 ± 1.6</td>
<td>-3.9 ± 1.7</td>
</tr>
<tr>
<td>Weight-for-height Z-score*</td>
<td>-0.9 ± 1.5</td>
<td>-1.0 ± 2.1</td>
</tr>
<tr>
<td>Weight-for-age Z-score*</td>
<td>-3.2 ± 1.0</td>
<td>-3.1 ± 1.2</td>
</tr>
<tr>
<td>Intake</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Energy (kcal/day)*</td>
<td>1488.8 ± 656.1</td>
<td>1234.3 ± 412.4</td>
</tr>
<tr>
<td>Carbohydrates (g/day)**</td>
<td>195.6 (158.6; 250.8)</td>
<td>166.6 (131.8; 197.0)</td>
</tr>
<tr>
<td>Protein (g/day)**</td>
<td>35.6 (34.9; 57.3)</td>
<td>34.9 (27.5; 45.9)</td>
</tr>
<tr>
<td>Fat (g/day)**</td>
<td>46.8 (29.5; 58.0)</td>
<td>33.9 (27.5; 45.9)</td>
</tr>
<tr>
<td>Fiber (g/day)*</td>
<td>9.2 ± 4.3</td>
<td>12.3 ± 4.3</td>
</tr>
<tr>
<td>Iron</td>
<td>8.0 (5.6; 12.9)</td>
<td>11.9 (8.1; 15.0)</td>
</tr>
<tr>
<td>Fluids (mL/day)*</td>
<td>456.5 ± 283.1</td>
<td>741.1 ± 379.2</td>
</tr>
</tbody>
</table>

*Mean and standard deviation, Student's t-test; **Median and 25th and 75th percentiles, Mann-Whitney test.

DISCUSSION

The results of this study show that children with cerebral palsy often exhibit inadequate food intake, severe anthropometric deficits, and a high frequency of gastrointestinal symptoms associated with certain diet characteristics.

Malnutrition and impaired growth are a common in children with cerebral palsy due to several factors, some of which are related to diet[3,12,32,13]. Although inadequate intake of energy, protein, essential fatty acids, vitamins, and minerals is considered to be the main cause of these conditions[35,34], few studies have assessed the food intake of children with cerebral palsy. A study conducted in Greece compared the energy intake of 16 children with cerebral palsy and 16 children without neurologic abnormalities (control group) and found it to be inadequate in both groups[35]. In addition to inadequate energy intake, muscle tone, level of physical activity, and the presence of involuntary motions may contribute to the incidence of malnutrition found in children with cerebral palsy by increasing their daily energy requirements[34].

A study conducted in Norway on 221 children with mental deficiency found that the presence of orofacial dysfunction is associated with reduced daily energy intake[35]. It should be observed that the reference values used in this study were based on the nutrient intake of children who did not present cerebral palsy and thus may not be fully appropriate for children with cerebral palsy. The EAR values correspond to the median distribution of the nutrient requirements of healthy individuals of the same gender and age range and meet the needs of 50% of the corresponding population[36]. In children with spastic paralysis, the muscles are hypertonic, which increases their energy requirements[36]. In addition, in such children, orofacial motor dysfunction interferes with sucking, chewing, and swallowing. The prevalence of oropharyngeal dysphagia varies from 16 to 99%[32]. Affection of the oral phase of swallowing is characterized by inability to control the food in the mouth due to problems with sealing the lips, loss of oral reflexes and of the motion of the anterior and dorsal parts of the tongue, and difficulties in chewing. Individuals with cerebral palsy have difficulty closing the lips while swallowing, which contributes to the food bolus escaping the mouth, aggravating malnutrition, and hinders the assessment of effective food intake[6,32]. In this study, clinical symptoms of dysphagia were found in most of the participants (33/40) who exhibited lower energy intake (TABLE 2) than the ones without evidence of dysphagia. A Brazilian study carried out in Santos (São Paulo, Brazil) included 90 children with quadriplegic cerebral palsy aged between 2 and 13 years showed high prevalence difficult to chewing (41%) and swallow (12.8%)[36].

In regard to macronutrient distribution, the intake of carbohydrates and protein was over the RDA in 85.0% (34/40) and 92.5% (37/40) of the sample, respectively, as shown in TABLE 1. The carbohydrate intake was significantly higher among the children without dysphagia (TABLE 2). A study conducted in Greece[35] with children with cerebral palsy up to 10 years old found that carbohydrates represented 47%, fat 36%, and protein 17% of the energy intake, and that this distribution was adequate in the case of carbohydrates and proteins but slightly above the recommended intake in the case of fat according to the Acceptable Macronutrient Distribution Range (AMDR), which indicates the ideal distribution of energy provided by each macronutrient relative to the total energy intake[36]. A previous study in Brazil found low consumption of carbohydrates (52%), adequate intake of protein (53%) and high intake of lipids (43%)[36]. In the Queensland, Australia, a study verified that non-ambulant, tube-fed cerebral palsy patients had significantly lower protein intakes compared to orally fed children[37]. There were no other differences in macronutrient intake between children with cerebral palsy and the control group.

The low prevalence of iron deficiency anemia found in this study is noteworthy. Only a single child with cerebral palsy had reduced hemoglobin and ferritin. The two patients presenting reduced hemoglobin did not have low serum ferritin levels.

Low fluid intake combined with insufficient dietary fiber intake may contribute to the development of constipation[6,7]. Approximately 60% of the sample in this study exhibited constipation, which agrees with the reports in the literature[3,12,38]. The average daily dietary fiber intake was lower among the participants with constipation (TABLE 4). A similar association was found with fluid intake, i.e., the amount of water consumed by the children with constipation was lower than the amount consumed by the children without constipation (TABLE 4). The reason for the lower dietary fiber intake exhibited by individuals with cerebral palsy may be that they consume foods with a low degree of consistency[12,13]. A Brazilian group of children with cerebral palsy also had low daily intake of fruits and vegetables (sources of dietary fiber) and liquids (less than three cups of 200 mL/daily). In this study, 14 of 39 children with quadriplegic cerebral palsy presented less than 3 evacuations per week[38]. In addition to the dietary factors, cerebral palsy is associated with intestinal motility disorders characterized by increased transit time in the transverse colon and rectum[7,38], as well as with changes in recto-anal function that result in a longer duration of the anal inhibitory reflex[39].

Gastrointestinal symptoms of gastroesophageal reflux disease were found in 40.0% of the participants in this study. The children with evidence of gastroesophageal reflux exhibited lower height-for-age in addition to greater fluid intake, most likely because softer foods are less uncomfortable to swallow (TABLE 3).
The sample size might represent a limitation of this study. However, the frequency of gastrointestinal symptoms and the dietary data obtained were so clinically relevant that the number of patients was sufficient to obtain statistically significant differences in several parameters. A second limitation might be the decision to evaluate study only gastrointestinal symptoms. However, the Rome III diagnostic criteria for gastrointestinal functional disorders in patients without neurologic abnormalities recommends that the diagnosis should be established taking into account the gastrointestinal symptoms. In addition, we considered unethical to perform diagnostic tests for dysphagia, gastrointestinal reflux disease/esophagitis (exposing the participants to radiation and sedation) specifically for this study.

CONCLUSION

This study found a wide variation in the results of the dietary survey. Whereas the protein intake was excessive in a large number of cases. The participants exhibited significant anthropometric deficits, which may be partially related to dietary factors. However, it is important to note that the nutritional deficit exhibited by patients with cerebral palsy is multifactorial. Gastrointestinal symptoms are a frequent occurrence among individuals with cerebral palsy. The patients with constipation exhibit lower fluid and dietary fiber intake; energy consumption is reduced in the patients with symptoms of dysphagia; and the individuals with symptoms of gastroesophageal reflux consume greater amounts of fluids. As a conclusion, within the context of multi-professional care provided to patients with cerebral palsy, nutritional interventions should be individualized as a function of the particular needs of each patient.

ACKNOWLEDGEMENTS

We thank the patients and families who participated in the study as well as the institutions (Casas André Luiz and Lar Escola São Francisco) where the study was conducted. We thank American Journal Experts which provided writing services on behalf of Brazilian National Research Council (Conselho Nacional de Pesquisa).

Authors’ contribution

All the authors participated in the conception and study design, interpretation of results and the writing and revising the final version of the manuscript. Caramico Favero DCO was responsible for data collection. All authors read and approve the final manuscript.
REFERENCES


