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Ethical issues with artificial nutrition of children with degenerative

- brain diseases ☆,☆☆
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ABSTRACT

This report highlights viewpoints of the authors and comments from the auditory at a workshop, held during the 20 14th international Congress on neuronal ceroid lipofuscinoses (NCL) in Córdoba, Argentina, on ethical aspects of 21 artificial nutrition in children with degenerative brain diseases. The discussion centers on what constitutes the 22 best interest of a patient whose personality was immature before the onset of the disease, who has become 23 demented during its course and is unable to communicate his/her own positions and desires. There is wide 24 consensus that in a child with advanced disease who cannot be fed naturally, decisions to withhold nutrition 25 or to institute or stop artificial nutrition, should only be made by parents (or their representatives) who are 26 adequately prepared on an intellectual and emotional level. We try to show that such decisions are highly 27 individual but can be made in a rationally and emotionally acceptable way after a careful and prolonged dialogue 28 between families and professionals. A checklist summarizes important considerations. This article is part of a 29 Special Issue entitled: "Current Research on the Neuronal Ceroid Lipofuscinoses (Batten Disease)".

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1. Introduction

The neuronal ceroid lipofuscinoses (NCL) are a group of genetic incurable degenerative brain diseases that constitute a relatively frequent prototype of a condition associated with dementia, complete helplessness and inability of communication at a young age [1]. While dementia, which denotes a loss of multiple higher cortical functions in a previously well-functioning person, is regarded as a public health priority by the World Health Organization [2], there is still little awareness of childhood dementia which occurs in a number or rare and mostly genetic brain diseases. Other disorders that develop problems similar to

patients (Table 1). A frequent problem during the latter course of a degenerative brain 52

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NCL during childhood or adolescence include the mucopolysaccharidoses 46 and mucolipidoses, Niemann-Pick disease, certain leukodystrophies and 47 basal ganglia degenerations. Such conditions raise a number of ethical 48 issues, some of which are similar to those in the end-of-life care of elderly 49 persons [3], while other topics are specifically related to the young age of 50

disease in a child or adolescent is the inability of eating and drinking. 53 With the availability of various methods of artificial nutrition, several 54 questions arise regarding their application in children with degenera- 55 tive brain diseases (Table 2). Compared with the growing knowledge 56 and awareness of such diseases in recent years, little medical literature 57 deals with these questions [4,5], although some principles regarding 58 children with severe brain damage have been expressed many years 59 ago [6]. It is clear that other medical therapies aiming at the prolongation of life in the end stage of an incurable brain disease, such as respi- 61 ratory support, are subject to analogous deliberations.

Professionals caring for children with degenerative brain disease are 63 in need of principles that are applicable in practice when the question of 64 tube feeding arises. Clear concepts should create a feeling of security in 65 caregivers and avoid traumatic irritations through the interfering of 66 persons not directly involved.

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Report of a workshop with participation of patient organizations at the 14th NCL Congress in Córdoba, Argentina, 25th of October, 2014.

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Table 1

Reasons that make some ethical issues specific to demented children.

- The patient's personal autonomy is not yet fully developed.
 - The patient has had no chance yet to conceive and fix a consistent conviction regarding questions of vital importance.
 - As dementia in childhood is rare, relevant knowledge and experience are scarce.

1.1. Origin of presented material

This article is a report of issues discussed at a workshop where the authors presented their views on aspects of life support through artificial nutrition for children in the end stage of a progressive brain disease. Members of NCL patient organizations participated in the discussion. For many critical questions a consensus was reached, mostly on the basis of intensive personal experience of families and medical professionals.

1.2. Issues dealt with at the workshop

Discussions centered on a list of questions where a need was felt for analysis of frequently encountered medical situations and for solutions or guidelines to management (Table 2).

In the following, we treat these questions and offer some practical guidelines for handling such issues in young persons with a degenerative brain disease.

1.3. What are medical indications for an artificial nutrition?

Artificial nutrition and artificial hydration are medical treatments associated with significant problems and should never be instituted without an unambiguous medical indication. Although the decision to start such treatment is ideally based on a dialogue between the patient's family and their physician (see below), it is the latter's duty to formulate the indication and put it down in writing with simple clear words. Such a drastic medical measure as artificial nutrition should not be started as "the normal thing to do" in progressive dementia.

The indication may be easy to determine in patients with swallowing problems but who are otherwise in relatively good condition. Insufficient nutrition in such patients may lead to avoidable additional morbidity and unnecessary suffering. The indication is much less clear in far advanced stages of cerebral decline, as improvement of the quality of life through artificial nutrition may not be convincing at this stage. Quality of life, however, is a very variable notion that only can be assessed regarding the individual situation. As a rule this is only possible for parents and for physicians thoroughly acquainted with the medical history.

1.4. When is prolonging life by artificial nutrition in the best interest of the patient?

There is little doubt that decisions of vital importance should be made in the best interest of the patient, but difficulties may arise as to what constitutes the patient's best interest, particularly in a mentally incapable, non-communicative child with severe brain damage [7].

Table 2

Questions related to artificial nutrition in children with degenerative brain disease.

When a child in the later stage of disease has become unable to be fed naturally:

- What are medical indications for an artificial nutrition?
- When is prolonging life by artificial nutrition in the best interest of the patient?
- Can artificial nutrition be withheld or terminated in certain situations?
- · If artificial nutrition is indicated, what methods should be used?
- By whom and how should decisions be made?
- \bullet What are the psychological, social and legal implications of such decisions?

Sometimes, arguments outside the realm of empathy and the presumed 107 well-being of the child are put forward, such as "sanctity of life" or other 108 concepts colored by religious, cultural, and racial or ethnic perspectives. 109 Such arguments are rarely emphasized by persons who have been intensively confronted with the situation of such children. There seems 111 to be a wide consensus that ethical questions of the nature discussed 112 here should be settled on a highly individual basis and not according 113 to rules or directions of outsiders.

It must be emphasized here that feeding against the patient's wishes 115 is widely regarded as an assault on personal integrity. In this respect, 116 the presumed wish of a demented patient (or the wish reasonably 117 projected on the patient) not to be fed artificially is particularly weighty 118 and has been subject of intense discussion in demented adults [8].

1.5. Can artificial nutrition be withheld or terminated in certain situations? 120

When, for whatever motive, artificial nutrition has been instituted, 121 a situation may evolve with good reasons for termination. After 122 prolonged observation of the patient, parents and physicians may 123 come to recognize that continuation of artificial nutrition will be with- 124 out benefit to the patient and will prolong life unnaturally. In this situation, parents, who are the only ones responsible for their child's welfare, 126 may be under psychological pressure towards continuation of artificial 127 nutrition.

If a decision to withdraw or withhold treatment that prolongs life is 129 considered to be in the best interest of the patient, it does not contradict 130 prevailing laws in most countries. This implicates that it can be in the 131 patient's best interest to accept death as a consequence of the decision. 132 These deliberations should be touched upon and be adequately docunamented already at the very first consultation where questions of nutrition come up.

It should be noted that the concept of artificial nutrition comprises 136 the supply of both, food as well as water. Medical experts expressed 137 concern that withholding food but continuing the supply of water 138 would lead to a particularly poor physical condition and undesirable 139 prolongation of life. In a patient dying from a cerebral condition as 140 discussed here, keeping the oral mucous membranes moist is thought 141 to be adequate palliative care.

1.6. If artificial nutrition is indicated, what methods should be used?

Clinical judgment must be used to determine if nutritional support 144 will improve the patient's well-being. Evaluating growth and the 145 gaining of weight of a severely handicapped child on the basis of the 146 usual pediatric charts is not useful. In immobile adolescents, obesity 147 provoked by relative over-feeding sometimes creates unnecessary 148 difficulties of handling the patient.

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Once a clear decision has been made for tube feeding, the help of a 150 gastroenterologist with pediatric experience must be sought. Percuta- 151 neous endoscopic gastrostomy is a good solution and feasible in many 152 parts of the world [9]. If this service is not available, naso-gastric tubes 153 are an alternative that frequently works for prolonged periods of time 154 without serious complications.

The type of food used for artificial nutrition via tubes deserves critical examination. As these neurological patients usually do not have an
intestinal disease, food composition should be close to a regular diet.
Food can be prepared at home using mechanical homogenization. This
is not only economical but also avoids much discomfort that is associated with industrial preparations. Erroneously, unphysiological highcalorie preparations are frequently given to patients with muscular
atrophy due not to malnutrition but to motor inactivity. Caloric requirements should not be prescribed on the basis of healthy children
of the same age. Other details, such as the schedule of feeding in individual patients, are beyond the scope of this discussion.

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