Investigating the contribution of physiotherapy in the quality of life of parents with children and adolescents suffering from cerebral palsy

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Abstract

Introduction: Many researches have focused on the quality of life (QoL) of children with cerebral palsy, but few have been done about the quality of life of parents with children suffering from cerebral palsy. In addition, less attention has been paid to the contribution of physiotherapy to the quality of life of parents with those children. The primary objective of this study was to evaluate the quality of life of parents with children and adolescents diagnosed with cerebral palsy. A secondary aim was to investigate the contribution of physiotherapy to the everyday life of these parents, namely that the effects of physiotherapy on the child automatically affect the quality of life of the parents.

Methods: Questionnaires were distributed to parents (N=30) at the Developmental Pediatrics Center "Apostolos Fokas", which belongs to the 1st Pediatric Clinic of the General Hospital of Thessaloniki "Hippocrates", as well as to a private pediatric physiotherapy center in the Thessaloniki area. The study's measurement tool was the specialized quality of life questionnaire for parents with children and adolescents diagnosed with cerebral palsy, which designed by the first author of this dissertation and includes 3 sections (totally, 32 questions).

Results: The results showed that parents have been affected by their child's disability in different aspects of their everyday lives, such as social and family relationships, economical status and individual well-being. On the other hand, parents are involved in the intervention process and consider that physiotherapy contributes positively to the quality of life by enhancing their child's functionality.

Conclusions: According to the research hypothesis, the condition of the children, directly, affects the quality of life of parents. However, through the physiotherapeutic approach and, therefore, the functional independence of the child, the quality of life of these parents may improve.

Introduction

Cerebral palsy is one of the most common developmental disabilities that begins in early childhood and remains throughout life [1]. A child with cerebral palsy requires multiple and specialized care, which exceeds that of a normal coeval child and therefore requires more attention by parents [1]. This chronic care turns into a chronic weight, with the result that parents are often bound to their care for a lifetime [2]. As Seligman and Darling [3] report, for some families care takes 24 hours a day, 7 days a week and for many years. All of these contribute, causing the family to suffer physically and psychologically.

Most researches have focused on the effects of the care for a child with cerebral palsy on parental mental health, but less attention has been paid to their quality of life [4-9].

Within the field of the quality of life, the World Health Organization (WHO) includes: physical health, mental state, independence, social relations, environment, religion, beliefs, determination and conflicting views.

According to the existing literature, the quality of life about parents with children diagnosed with cerebral palsy is lower than the quality of life of the general population and parents with normal children [4-9]. Studies suggest that mothers burden themselves more than the child care [3]. In particular, parents (more so mothers) are in many cases characterized as socially isolated and limited because of the demands of the child care. Assistants in the chronic care of a disabled child (with less frequent help) appear fathers and sisters. Very little help seems to be given by relatives and friends [2].

Regarding the rehabilitation of children with cerebral palsy, it must be complete [10]. Physiotherapy plays a central role in managing a child with cerebral palsy and focuses on the functionality, active motion and optimal use of the child's potentials [10]. The goals of rehabilitation in children with cerebral palsy are to minimize the effects of physical impairments, to help the child gain independence in society and to improve the quality of life of these children and their families, who play an important role in the whole process of rehabilitation [11].

Previously, physical and occupational therapists used to focus on impairments and limitations of the child [12]. However, the provision of services in pediatric rehabilitation has undergone significant changes in recent decades [13]. The most extensive change that has taken place...
was the review of the role of the family. In about half a century, the role of parents in the care process has evolved into being a target for intervention [14].

Recent literature on pediatric rehabilitation has shown that interventions focusing on children and their families can lead to satisfaction by the care, the well-being of parents and the reduction of parental anxiety [15]. Families involved in the care of their child have the opportunity to learn more about their child and their child’s treatment options. Family members, in turn, are able to share their knowledge with professionals and this provides a holistic picture of the child and, also, is a way to raise their child’s awareness, as part of the family [13].

Taking into account the deficits of existing literature, the purpose of the present study was to investigate the contribution of physiotherapy to the quality of life of parents with children and adolescents diagnosed with cerebral palsy. It is widespread that the aim of physiotherapy in children and adolescents with cerebral palsy is to improve functionality, by enhancing the gross motor function and learning functional activities. Therefore, through the specially formulated quality of life questionnaire for parents with children and adolescents suffering from cerebral palsy, we assume that the functionality of the person with cerebral palsy is inextricably linked to the quality of life of their parents and guardians. Thus, if we thoroughly record the quality of life of this category of parents, we will contribute to a more holistic and sufficient physiotherapeutic intervention.

Methods

Participants

In the present study, 30 parents (20 mothers and 10 fathers) of children and adolescents (aged 3-17 years) diagnosed with cerebral palsy took part. The selection criteria pertain to parents of children and adolescents diagnosed with any form of cerebral palsy and any classification (I-V) at the Gross Motor Function Classification System (GMFCS) and age range 3-17 years. The exclusion criteria refer to mentally handicapped parents, inability to read and write the Greek language and, generally, the inability to complete this specific questionnaire. In addition, a customized written acquiescence and briefing, concerning the participation of the parents in the present dissertation, is provided, as well as their ability to be informed about the results of this study. Furthermore, the Research Ethics Committee at the Technological Educational Institute of Thessaloniki approved the study protocol.

Study design

Questionnaires were distributed to parents at the Developmental Pediatrics Center “Apostolos Fokas”, which belongs to the 1st Pediatric Clinic of the General Hospital of Thessaloniki “Hippocratio”, as well as to a private pediatric physiotherapy center in the Thessaloniki area. Parents filled in the questionnaires following their oral and written briefing.

Assessment of QOL

The study’s measurement instrument was the specialized quality of life questionnaire for parents with children and adolescents diagnosed with cerebral palsy, which designed and made by the first author of this dissertation and includes 3 sections (totally, 32 questions):

1st section: Demographic information that includes each child’s data, such as age and disability, as well as parent’s data, such as occupation, educational level, living area and family structure.

2nd section: General quality of life questions from the parent questionnaire for children with disabilities. This section includes general questions concerning parents’ feelings during diagnosis, condition of the family at present, and how much disability has affected the routine of parents, and, finally, parents’ opinion about physiotherapists.

3rd section: Specific questions concerning parent’s opinion about physiotherapy and how it helps to improve the child’s functionality and, thus, to improve the quality of life of parents.

Statistical analysis

Data was analyzed using descriptive statistics of PSPP version 0.8.5 and Microsoft Office Excel 2010 programmes. We did not proceed to inferential statistics (t-test) due to the limited sample size (N=30) and the existence of qualitative variables.

Results

2nd section-general questions

Parents’ feelings during diagnosis: The feelings that the question included was anger, disappointment, lamentation, loneliness and shock. As for the feeling of anger, the majority of parents, in particular, the 50% answered “enough”, for the feeling of disappointment the 60% of parents answered “enough”, for the feeling of lamentation the 36.67% of parents answered “enough”, for the feeling of loneliness the 50% answered “moderately” and for the sense of shock the 90% of parents answered “enough”. The emotion of shock is the first stage that parents experience, when a child with disabilities is born and is manifested by emotional disorganization and confusion [2].

Time to realize the possible consequences of diagnosis: The 40% of parents realized the possible consequences of diagnosis “after two months”, the 26.67% of parents “immediately”, the 20% “in less than one year”, while the 13.33% “after one year”.

Time that parents spend in activities of everyday life: Of all parents, the 43.33% spends “not at all” time on themselves, the 43.33% “not enough” and the 13.33% devotes “enough” time to themselves (Figure 1). For the time about the relationship with the partner, as a couple, the 70% of the parents responded “not enough”. For the time with the family the 50% of the parents responded “not enough” and the remaining 50% “enough”. For the child’s rehabilitation activities, the 80% of parents dedicate “enough” time. In addition, the 46.67% of parents dedicate “enough” time to play with their child. As for the time about friends, the 63.33% of parents responded “not enough”. Finally, the 50% of parents devotes “not at all” time to their entertainment.

The influence of disability on different aspects of parental life: This question related to the degree of which disability affects social relationships, occupation, life with the partner as a couple, level of stress, family relationships, economic status and individual welfare of parents (Figure 2). For the social relationships, the 36.67% of parents responded that the disability of the child has affected them “enough”. As for the occupation, the 40% of parents has been affected “moderately” and for the life with the partner as a couple, the 53.33% of parents responded “moderately”. Regarding stress level, the 70% of parents responded “enough”. As for family relationships, disability has affected 53.33% of parents “moderately”, while 46.67% of the parents have been affected “enough”. Finally, with regard to individual well-being, the 53.33% of parents have been affected “moderately”.

Characteristics attributed by physiotherapists to parents: In this question, the 53.33% of parents answered “enough” about the efficiency...
Chasidou AE (2017) Investigating the contribution of physiotherapy in the quality of life of parents with children and adolescents suffering from cerebral palsy

Figure 1. Time devoted by parents to people and activities of everyday life

Figure 2. The influence of disability on different aspects of parental life
of physiotherapists, the 73.33% responded “moderately” to the protection afforded by physiotherapists, while for the physiotherapists’ ability about care the 60% of the parents responded “moderately”. In addition, for the question, in which, physiotherapists attribute stress to parents, the 53.33% responded “moderately”, for anguish, the 50% of parents responded “moderately”, and, finally, for stiffness, the 66.67% answered “not at all”.

How qualified parents consider physiotherapists are about pathology, comprehension of family needs, prevention of stress and communicating skills: Regarding pathology, the 56.67% of parents responded that physiotherapists are “enough” qualified. In respect to comprehension of the family needs, the 56.67% responded that physiotherapists are “moderate” qualified. In terms of stress prevention, the 60% of parents responded “moderately”. Finally, in regard of communication skills, the 56.67% of parents responded that they are “moderate” qualified.

3rd section-specific questions

Age of child’s physiotherapy initiation: The majority of parents, namely 30%, replied that their child started physiotherapy at the age of two months and all parents responded that they started physiotherapy immediately after the referral (Figure 3).

Parent’s involvement during the physiotherapeutic session: Of the total population of parents, the majority, namely 57%, does not participate in the physiotherapy session, while the 43% participates.

Parents who like the process of attending the physiotherapeutic session: Of the parents participating in the physiotherapeutic session, the 60% enjoys it, while the 40% does not like it.

Preference for parents to take advantage of the physiotherapeutic session’s time with something else: The 67% of parents do not want to take advantage of the physiotherapeutic session’s time, with something else.

Encouragement by the physiotherapist about the parent’s active participation in the physiotherapeutic session: According to the 70% of parents’ population, the physiotherapist encourages their active participation in the session, while only the 30% says, that the physiotherapist does not encourage their participation.

Opinion of the parents: a) on the physiotherapy and the functional development of the child, b) on further occupation at home, and c) on the combination of physiotherapy with an alternative therapy (e.g., hippotherapy, hydrotherapy): The 90% of parents believe that only physiotherapy is not enough for the child’s functional development, and all parents believe that they need further occupation at home. Finally, in regard to the question of whether physiotherapy should be combined with alternative treatment, 80% of parents responded “yes”.

Application, by parents, at home, of specialized exercises, position changes, training of activities, under the instructions of the physiotherapist, and how many hours they apply them per week: As for the question of home exercises, under the instructions of the physiotherapist, the 87% of parents responded “yes”. The 36.67% of parents apply them “4 hours per week”, the 26.67% of parents “7 hours per week” and the 23.33% “6-7 hours per week”.

Parent or relative who applies physiotherapy at home, prescribed by the physiotherapist: Of the total respondent parents, the 63.33% responded that only “mother” is concerned with physiotherapy at home prescribed by the physiotherapist, the 33.33% answered “both parents”, while only the 3.33% replied “no”.

Parent’s opinion on physiotherapy and if it affects his free time: The 67% of parents believe that physiotherapy affects “enough” positively their free time, the 23% responded “moderately”, while only the 10% reported “not at all”.

Contribution of physiotherapy to the improvement of child’s functionality, according to the parents’ opinion: The 97% of parental population believe that physiotherapy helps “enough” in the improvement of their child’s functionality, while the 3% believes that it helps “moderately”.

According to parents’ opinion, how physiotherapy, by improving child’s functionality, facilitates, by extension, their daily routine: The 97% of parents believe that physiotherapy helps to improve child’s functionality “enough” and, thus, makes easier their daily routine, while only the 3% of parents believes that physiotherapy facilitates their daily routine “moderately”, by improving their child’s functionality.

Discussion of the parent with physiotherapists, on what concerns them, in regard to their child’s progression: The 63% of parents responded that they discuss “enough” with their physiotherapist, on what it concerns them, in regard to their child’s progression.

Confidence of parents in the opinion and advice of the physiotherapist: The 80% of parents trusts the opinion and advice of their physiotherapist “enough”, while the 20% of parents trusts them “moderately”.

Parents’ estimation about physiotherapists in respect to the comprehension of difficulties faced by parents, in their daily life: The 53% of parents believes that physiotherapists understand the difficulties that parents face in everyday life “moderately”, while the 47% believes that physiotherapists understand “enough” parents’ difficulties.

Discussion

The primary objective of this study was to investigate the quality of life of parents with children, aged 3-17 years and diagnosed with cerebral palsy. A secondary objective was to investigate the contribution of physiotherapy to the quality of life of these parents, specifically whether physiotherapy, by enhancing the child’s functionality, also improves the quality of life of parents, as well as the parents’ opinion on physiotherapy and physiotherapists and, finally, whether the last ones support the model of family-centered therapeutic approach.

In terms of parents’ quality of life, questions were asked about the condition of the family at present, namely the time that parents spend...
on people and activities of everyday life, and how disability affects different aspects of their everyday lives. From the results it turned out that parents do not spend enough time on themselves or even on the relationship with their partner, as a couple. As for their family, half of the parents answered that they spend enough time, while the remaining half do not spend enough. In regard of child’s rehabilitation activities, the majority of parents spend enough time. The majority of them, also, spend enough time, playing with their child. Finally, most parents do not spend enough time with their friends and the majority of parents do not spend any time at all about their entertainment.

In addition, the child’s disability seems to affect the daily routine of the parents. In particular, it greatly affects parents’ social relationships, stress levels and economic conditions, while moderately affects occupation, life with the partner, family relationships and individual welfare of parents.

As confirmed by literature, the solicitude for a child with cerebral palsy may positively affect a parent’s ability to create new social support networks, but the condition of such a child’s care may adversely affect parents’ physical health, social relationships, freedom and independence, family relationships and economic stability [4].

From the knowledge, regarding parents’ opinion about physiotherapists, it seems that the last ones are highly competent, in terms of efficiency, they attribute moderate protection and ability, in terms of solicitude, to parents, little anxiety and anguish, while they do not express any stiffness at all.

Also, according to parents’ opinion, physiotherapists are highly trained in pathology, while they are themselves moderately satisfied with regard to the comprehension of family needs, the prevention of stress, as well as communicating skills. However, most parents discuss with their physiotherapists what they are concerned about their child’s development, and they trust their opinion and advice. The role of parents in the health care process and the attitudes of healthcare professionals, are essential elements in understanding care processes [15].

With the family unit as the primary framework for the promotion of children’s health and well-being, the family-centered approach has been shown to contribute to the emotional well-being of parents [16-19]. For this reason, in the questionnaire of this dissertation questions about the involvement of parents in physiotherapeutic sessions included.

The results also showed that the majority of parents did not participate during the physiotherapy session. However, parents who participate in the physiotherapy session often enjoy the process of attendance, while very few of them would prefer to use this time with something else. According to the opinion of parents, the majority of themselves do not participate in the physiotherapy session, although the physiotherapist encourages their active participation.

Parents’ beliefs, early on, in terms of their child’s treatment, may have a significant impact on the attendance during therapeutic session and, consequently, on the therapeutically outcomes [20]. In addition, parents have been shown to be more satisfied with family-centered interventions [16,17].

Regarding the question about physiotherapy and whether suffices for the functional development of the child, the majority of parents believes that is not sufficient in itself, whereas all parents believe that further work at the home with physiotherapeutic exercises is needed. Most of the parents, also, believe that physiotherapy should be combined with an alternative therapy.

Most parents implement specialized exercises, position changes and activities training, under the guidance of the physiotherapist, at home. However, the majority of parents spends less than four hours per week, proving either that they do not have enough time, due to daily duties, or because of the psychological and physical weariness they experience both of the parents.

Finally, with regard to the contribution of physiotherapy to the quality of life of parents, it seems, initially, that physiotherapy, according to the parents’ opinion, helps to improve the child’s functionality. In addition, by improving child’s functionality, the majority of parents believes that it facilitates, as a result, their daily routine. Many of the parents, also, believe that physiotherapy, positively, affects their free time.

However, in this dissertation, there were some limitations. Initially, the sample was limited (n=30), as many parents refused to complete the questionnaire. After that, the questionnaire did not cover all the dimensions of quality of life. In the sequel, the study consisted of children with a large age range (3-17 years), which affects the children’s demands in terms of their parents care and, hence, their quality of life. Therefore, studies with a more specific age range need to be conducted. The study, also, included all forms of cerebral palsy and all levels of GMFCS (Gross Motor Function Classification System), which may again affect the quality of life of parents. Finally, research bibliography at the databases (scopus.com; sciencedirect.com; pubmed.com), which concerned with the quality of life of parents with children and adolescents diagnosed with cerebral palsy, was deficient [21-25].

Conclusions

According to the results of the present study, which are perfectly consistent with the initial hypothesis, physiotherapy seems to contribute positively to the quality of life of parents with children and adolescents suffering from cerebral palsy. This is achieved by increasing the child’s functionality, involving parents in the intervention process, and implementing family-centered therapeutic approach, where parents learn their child better and their care is facilitated. However, further researches, in regard to the contribution of physiotherapy to the quality of life of parents, with a larger sample of subjects, needs to be carried out.

References

1. Guillamón N, Nieto R, Pousada M, Redolar D, Muñoz E (2012) Quality of life and mental health among parents of children with cerebral palsy: the influence of self-efficacy and coping strategies. J Clin Nurs 22: 1579–1590. [Crossref]
2. Tsimpidaki A (2013) Child with special needs, family and school. A relationship in interaction, pp. 72, 73, 84.
3. Seligman M, Darling B (2007) Ordinary families, special children. (3rd ed.) New York: The Guilford Press.
4. Davis E, Shelly A, Waters E, Boyd R, Cook K, et al. (2010) The impact of caring for a child with cerebral palsy: quality of life for mothers and fathers. Child Care Health Dev 36: 63-73. [Crossref]
5. Kaya K, Unsal-Delialiloglu S, Kutay Ordu-Gokkaya N, Ozisler Z, Ergun N, et al. (2010) Musculoskeletal pain, quality of life and depression in mothers of children with cerebral palsy. Disabil Rehabil 32: 1666-1672. [Crossref]
6. Prudente COM, Barbosa MA, Porto CC (2010) Relation between quality of life of mothers of children with cerebral palsy and the children’s motor functioning, after ten months of rehabilitation. Revista Latino-Americana de Enfermagem 18: 149–155.
7. Romeo DM, Cioni M, Distefano A, Battaglia LR, Costanzo L, et al. (2010) Quality of life in parents of children with cerebral palsy: is it influenced by the child’s behaviour? Neuropediatrics 41: 121-126.
8. Okarowska-Zawada B, Kulak W, Wójcikowska J, Sienkiewicz D, Pazuko-Patej G (2011) Quality of life of parents of children with cerebral palsy. Progress in Health Sciences1: 116-123.
9. Ström H, Kreuter M, Rosberg S (2011) Quality of life in parents/caretakers of children with cerebral palsy in Kampong Cham, Cambodia. *J Trop Pediatr* 58(4): 303-306.

10. Kerem Günel M (2009) Rehabilitation of children with cerebral palsy from a physiotherapist’s perspective. *Acta Orthop Traumatol Turc* 43(2): 173-180. [Crossref]

11. Cusick A, McIntyre S, Novak I, Lamin N, Lowe K (2006) A comparison of goal attainment scaling and the Canadian Occupational Performance Measure for paediatric rehabilitation research. *Pediatr Rehabil* 9: 149-57. [Crossref]

12. Bly L (1991) A historical view and current view of the basis of NDT. *Pediatric Physical Therapy* 3: 131-135.

13. Siebes R, Wijnroks L, Ketelaar M, van Schie PEM, Gorter JW (2006) Parent participation in paediatric rehabilitation treatment centres in the Netherlands: a parents’ viewpoint. *Child: care, health and development* 33(2): 196-205.

14. Kruijsten-Terpstra M, Ketelaar H, Boeije MJ, Jongmans JW, Gorter J, et al. (2013) Verschuren, Parents’ experiences with physical and occupational therapy for their young child with cerebral palsy: a mixed studies review. *Child: care, health and development* 40(6): 787-796.

15. Alsem M, Verhoef JW, Gorter LCM, Langezaal JMA, Visser-Meliny M (2016) Parents’ perceptions of the services provided to children with cerebral palsy in the transition from preschool rehabilitation to school-based services. *Child: care, health and development* 42(4): 455–463.

16. King SM, Rosenbaum PL, King GA (1996a) Parents’ perceptions of caregiving: development and validation of a measure of processes. *Dev Med Child Neurol* 38: 757–772.

17. King GA, King SM, Rosenbaum PL (1996b) Interpersonal aspects of care-giving and client outcomes: a review of the literature. *Ambulatory Child Health* 2: 151-160.

18. Rosenbaum P, King S, Law M, King G, Evans J (1998) Family-centred service: a conceptual framework and research Overview. *Physical and Occupational Therapy in Pediatrics* 18: 1-20, 174.

19. Raina P, O’Donnell M, Rosenbaum P, Brehaut J, Walter SD, et al. (2005) The health and well-being of caregivers of children with cerebral palsy. *Pediatrics* 115: e626-636. [Crossref]

20. Nock MK, Ferriter C, Holmberg E (2007) Parent beliefs about treatment credibility and effectiveness: assessment and relation to subsequent treatment participation. *Journal of Child and Family Studies* 6: 27-38.

21. Dunst C, Trivette C, Cornwell J, Davis M (1998) Enabling and empowering families of children with health impairments. *Children’s Health Care* 17: 71-81.

22. Ireys HT, Perry JJ (1999) Development and evaluation of a satisfaction scale for parents of children with special health care needs. *Pediatrics* 104: 1182-1191. [Crossref]

23. King GA, Law M, King SM, Rosenbaum PL (1998) Parents’ and service providers’ perceptions of the family-centredness of children’s rehabilitation services. *Physical and Occupational Therapy in Pediatrics* 18: 21–40.

24. King G, King S, Rosenbaum P, Goffin R (1999) Family-centered caregiving and well-being of parents of children with disabilities: linking process with outcome. *Journal of Pediatric Psychology* 24: 41–53.

25. Letourneau NL, Elliott MR (1996) Pediatric health care professionals’ perceptions and practices of family-centered care. *Children’s Health Care* 25: 157.