Caregiver satisfaction with a multidisciplinary community-based rehabilitation programme for children with cerebral palsy in South Africa

Magugu Ngubane* and Verusia Chetty**

*Discipline of Physiotherapy, School of Health Sciences, University of KwaZulu-Natal, Durban, South Africa
**Corresponding author, email: chettyve@ukzn.ac.za

This study sought to determine the levels of satisfaction and experiences of caregivers with a multidisciplinary rehabilitation programme. A total of 26 caregivers were recruited at 3 health settings in South Africa. A survey based on the Multidimensional Assessment of Parental Satisfaction for Children with Special Needs and focus-group discussions were conducted at the end of programme. The majority of the respondents were mothers and unemployed. The participants were generally satisfied with the rehabilitation. However, participants held on to myths, stereotypes, and beliefs about the cause of disability and explained the impact of stigma on having a disabled child in their communities. Caregivers perceived the benefit of rehabilitation but believed that lack of communication and consultation with health professionals limited the care.

Keywords: caregivers, cerebral palsy, multidisciplinary team, rehabilitation satisfaction

Introduction

Cerebral palsy (CP) is the commonest cause of disability in children with a global prevalence of 1.5 to more than 4 children per 1000 children living with this condition. CP is classified as a group of disorders with motor impairments as the hallmark feature ranging from minor to major dysfunction. Commonly associated with comorbid cognitive and sensory disorders resulting from disturbance in the development of an infant or foetal brain, which remain permanent in nature. These impairments result in activity limitations and inhibited participation in daily activities as well as social recreational interaction. Children with CP require a comprehensive multidisciplinary rehabilitation approach to maintain and improve function and prevent secondary complications. The multidisciplinary team (MDT) managing children with CP comprises various health professionals, including doctors (neurologists and paediatricians), nurses, physiotherapists, occupational therapists, speech and language therapists, psychologists, and social workers. Each professional has a varied role in the team but need to collaborate in order to achieve optimal care. Roles and expectations of professionals span early detection, acute care, physical and cognitive rehabilitation intervention, timely referral and psychosocial support with the aim of integrating children back into their communities.

The MDT in a South African public domain manages children with CP from primary care settings through to tertiary settings. There is, however, a shift from centrally situated institutions of health care toward the primary health care model (clinics within the community) as it is fundamental for South African healthcare to become accessible to the larger population. Zimbabwe shares a similar governing infrastructure within its health care systems. A study by Dambi and Jelsma in 2014 compared a community-based model and hospital-based intervention for children with CP in Zimbabwe, which shows evidence of preferred outcomes within the community-based intervention. While the shift is still in its development and implementation stages, principles of practice for the MDT requires that the issues of access in such resource-limited settings be addressed. One of the strategies that has been identified in rehabilitation research within countries with such political climates is task shifting. Task shifting is the training of laypersons, family and community members in skills of rehabilitative care in order to address the need within the communities and the lack of care due to shortage of staff and financial constraints among other barriers to rehabilitation.

Compliance with rehabilitation is directly related to levels of satisfaction and influences commitment to the intervention. In the rehabilitative care of children with CP, the involvement of the caregivers and their level of satisfaction is of importance as their participation has an impact on the rehabilitation goals. Furthermore, satisfaction with rehabilitation intervention is an imperative outcome that is believed to influence the emotional well-being of carers and impacts on a child's psychology. Scarcity in caregiver satisfaction and perception of rehabilitation is evident in resource-poor settings such as South Africa, thus this study is crucial in informing research for rehabilitation professionals involved in the management of children with CP.

This study's emphasis was focused on caregivers of children with CP who were involved in rehabilitation within a task-shifting approach. The study offered a rehabilitation intervention programme addressing the needs of children with CP by capacitating the caregivers; thereafter the caregivers’ level of satisfaction with the intervention programme and perceptions relating to rehabilitation practice were investigated.

Methodology

This study is phased within a larger project in three purposively selected healthcare settings within semi-rural communities in the province of KwaZulu-Natal, South Africa. A mixed-methods approach using both qualitative and quantitative paradigms was used to investigate and explore a multidisciplinary six-week rehabilitation intervention programme for children with CP and their caregivers. The sessions were conducted on a weekly basis at each of the sites in a large seminar room/gym. A team of rehabilitation professionals planned the structured six-week programme. An occupational therapist, a speech and language therapist and a physiotherapist formed the core team. The physiotherapist, who was skilled in neurodevelopmental training...
as a postgraduate specialty, was responsible for executing the intervention. Table 1 summarises the intervention programme for the six weeks. The educational session always initiated the day’s events and on some days the caregivers had individual contact with the therapist. The educational sessions were based on evidence that emerged from managing children living with CP. Therapists delved into basic understanding of the condition and the importance of rehabilitation, as well as addressing challenges when caring for children with CP. The group therapy included mat-work, practical sessions on positioning and joint protection, and play therapy as well as toy-making.

This study addressed the level of satisfaction following the six-week programme and experiences of the caregivers regarding rehabilitation at the various sites.

Design
For this phase of the study the authors used a cross-sectional survey with closed-ended questions as well as feedback focus-group discussions with all the caregivers following the six-week intervention. This triangulation of positivist and naturalistic paradigms allows for strengthening of findings in a field with a paucity of research.12

Participant recruitment
Once approval was obtained from the University of KwaZulu-Natal, permission was obtained from the relevant health and institutional authorities governing the three public healthcare sites in KwaZulu-Natal. Thereafter children with caregivers were recruited during their scheduled sessions at the respective sites. All caregivers consented voluntarily to participate in the study and no incentives were offered at any point in the study.

Data collection
A survey was conducted with caregivers at the completion of the six-week programme and a focus-group discussion was held with each caregiver. The researchers agreed on the thematic representation of the data. The focus-group discussions were conducted in the same space as the sessions after the six-week intervention programme. A focus-group guide was used to inform discussions and included experiences of rehabilitation services and the six-week programme, as well recommendations for implementation of such programmes. Data were recorded using a Dictaphone, OLYMPUS VN-425PC Digital Voice Recorder and transcribed verbatim following the focus-group discussions.

Data analysis
The quantitative data from the survey were analysed using the Statistical Package for Social Sciences (SPSS®) version 21.0 (IBM Corp, Armonk, NY, USA) and was interpreted and reflected using descriptive statistics. Nvivo 10® (QSR International (UK) Ltd, Daresbury, UK) was used for thematic analysis of the narratives arising from the discussions following the six-week programme.

Results
Three sites within semi-rural communities in the province of KwaZulu-Natal, South Africa were purposively selected as they offered rehabilitation to children with CP. Twenty-six caregivers, who were all female and of African ethnicity, consented to participate in the study. They were conveniently recruited at the respective sites when they attended their healthcare visit. The preferred language medium was isiZulu.

Most of the caregivers accompanying the children were mothers (88%, n = 23) while 11% (n = 3) were grandmothers. The majority were unemployed (85%, n = 22) and 22 of the caregivers were unmarried women.

The focus-group discussions were conducted using Nvivo 10® and the themes that emerged were read and re-read until the researchers agreed on the thematic representation of the data.
The themes, sub-themes and illustrative quotes are reported in Table 3. Although focus-group discussions centred on perception of the six-week rehabilitation programme, participants shared their experience of caring for children with special needs as well as their views of rehabilitative services.

Discussion
Rehabilitation for children with CP is crucial to maximise functional ability and improve quality of life for those affected. Often it is the caregivers of the children who are critical to the success of any type of rehabilitation intervention. They adopt the dominant role of executing the necessary intervention and are responsible for the continuity of such programmes, as in the intervention programme used in this study.6 Assessing caregivers’ satisfaction with the six-week intervention programme compared with an institutional based health setting improved the outcomes of a rehabilitation intervention 3, providing home and community support. Supported by another South African study advocating that community-based interventions provide psychosocial support, support relationships and improve social functioning and self-esteem.17 In another study in Turkey looking at the quality of life of mothers of children with CP as primary caregivers, researchers corroborate that lack of resources and transport limitations pose a challenge to caregivers’ primary role. Barnes and Good18 highlighted the advantages of community-based rehabilitation as improving these limitations through family involvement, decreasing costs and providing home and community support.

The perceptions of caregivers of children with cerebral palsy toward rehabilitation and health care in this paper included untimely referral, scarcity of information from healthcare staff, poor consultation with the healthcare team and lack of communication. Studies by Dambis and Jelsma and Ones et al concur that a lack of information hinders caring for children with CP optimally. Likewise, in a Norwegian study with carers of children with CP, the mothers and sometimes grandmother or aunt predominantly adopted the role of primary caregiver.2,4,6 Fathers became the source of income, relegating the primary caring to the female.8 In this South African setting, however, a large majority were single women. The role of a single mother caring for a child with special needs adds a different paradigm and will need further investigation.

Augmenting the psychosocial challenges experienced by caregivers were the lack of community support, lack of information, financial constraints and transport barriers that hindered optimal rehabilitative care. In an analogous resource-limited African setting a study revealed that caregivers of children with CP withdraw from social participation due to their limited African setting. A study revealed that caregivers of children with CP withdraw from social participation due to their physical stressors.8,16 Ones et al. believed that carers of children with CP withdraw from social participation due to their responsibilities and this impacts on their quality of life. Raina et al. concurred that caring for children with CP does affect psychosocial well-being and Pfeifer et al. felt that carers are not merely affected physically and socially but financially as well. The majority of caregivers in this study were unemployed and the belief is that the duty to the child is limiting them from obtaining employment.

The primary caregivers were female and included mothers and a few grandmothers. Similarly, in other studies centred on caregivers of children with CP, the mothers and sometimes grandmother or aunt predominantly adopted the role of primary caregiver.2,4,6 Fathers became the source of income, relegating the primary caring to the female.8 In this South African setting, however, a large majority were single women. The role of a single mother caring for a child with special needs adds a different paradigm and will need further investigation.

The myths and beliefs held by caregivers regarding cerebral palsy spanned the belief that the condition is due to witchcraft or a curse on the family, the stress and strain undergone by the mother as well as her behaviour and the chronic medication that the mother administered. This perception inadvertently leads to stigmatisation of the already burdened carer, adding psychosocial stressors. Caregivers of children with CP in this setting further believed that their lack of understanding, psychosocial barriers and physical stressors challenged them as carers. Other studies reflect that primary carers often experience psychological and physical stressors.8,16 Ones et al. believed that carers of children with CP withdraw from social participation due to their responsibilities and this impacts on their quality of life. Raina et al. concurred that caring for children with CP does affect psychosocial well-being and Pfeifer et al. felt that carers are not merely affected physically and socially but financially as well. The majority of caregivers in this study were unemployed and the belief is that the duty to the child is limiting them from obtaining employment.

Table 2: Caregivers’ level of satisfaction with the six-week rehabilitation intervention

| Question | Poor | Neutral | Rating excellent |
|----------|------|---------|------------------|
| How would you rate your level of satisfaction with the quality of the six-week intervention | 0% | 15% | 85% |
| How well did the staff deal with your needs | 0% | 23% | 77% |
| What did you think of the quality of care you received | 0% | 15% | 85% |
| How did you find the explanation of procedure/treatment provided | 0% | 8% | 92% |
| How did you find the explanation of medical condition and treatment | 0% | 15% | 85% |
| How would you rate your level of satisfaction with the referral to other healthcare professionals | 0% | 10% | 90% |
| How would you rate your level of satisfaction with the explanation of discharge and follow-up care instructions | 0% | 46% | 54% |
| How did you find the courtesy and concern you received from the rehabilitation healthcare worker who treated your child | 0% | 38% | 62% |
| How did you feel about the rehabilitation healthcare worker’s concern about your confidentiality and privacy | 0% | 62% | 38% |
| How did you feel about your involvement in treatment sessions | 0% | 23% | 77% |
| What do you think about the material given about the services, content and understanding | 0% | 27% | 73% |
| How would you rate the exercises | 0% | 23% | 77% |

Note: n = 26.
| Sub-theme                                           | Quotes                                                                                                                                                                                                 |
|---------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| **Myths and beliefs about cerebral palsy**         | *Witchcraft or curse*<br>‘Our ancestors were angry and caused the child to be sick’ (Caregiver 3)<br>‘Her father took her to his family before they performed necessary rituals for her. That made the ancestors angry’ (Caregiver 8)<br>**Mother’s stress caused cerebral palsy**<br>‘My daughter was abused by the child's father when she was pregnant that is why he is like this’ (Caregiver 7)<br>**Mother’s medication as a cause of cerebral palsy**<br>‘It is the tablets that I take when I was pregnant that caused the child to be disabled’ (Caregiver 10)<br>**Behaviour as a cause**<br>‘An arrogant or rude mother can cause the child to be sick’ (Caregiver 1)<br>‘If we being shouted at a lot when mom was pregnant’ (Caregiver 2) |
| **Perceptions of caregivers of children with cerebral palsy toward rehabilitation and health care** | *Untimely referral*<br>‘I get help but they ordered wrong shoes for him [physiotherapy referral to orthotist] and it took a long time to do so’ (Caregiver 4)<br>‘I tell them there at physio to give him a chair, they take long to get it from the other therapist [Referring to OT]’ (Caregiver 6)<br>**Scarcity of information from healthcare staff**<br>‘To me they said that there was blood in her brain. But did not explain how it got there or what challenges I will face with her. I only heard at physio that she had cerebral palsy’ (Caregiver 11)<br>‘For me they said she had cerebral palsy, marked with cerebral atrophy and small head. I can see the small head but do not understand what the other stuff means’ (Caregiver 8)<br>**Poor consultation with healthcare team**<br>‘Depends on person attending you. Some are nice and help, others don’t explain to us. They just look at the child and you don’t know what they are doing. They would not even explain like you did here [referring to 6-week programme] and just give you your next appointment’ (Caregiver 20)<br>**Lack of communication from rehabilitation team**<br>‘They just tell us that the child has delayed milestone. But what does it mean, At least here you tell us more [referring to 6-week programme], later the child presents this way’ (Caregiver 19)<br>**Rehabilitation benefits children**<br>‘Attending physiotherapy clinic helps. My child was tight now it is better. She can now walk for short distances, exercises helped a lot’ (Caregiver 14)<br>‘Exercises and advice was to sit him for long to strengthen muscles of his body and head. That helped me’ (Caregiver 7)<br>‘What help me and my child was physiotherapy. My child could not turn and laugh but now can’ (Caregiver 10)<br>**Intrinsic inhibitors of caregivers**<br>‘I just don’t know what to do when he does not want to eat and when he chokes on drinking and during feeding. No one tells us what to do’ (Caregiver 10)<br>**Psychosocial barriers**<br>‘I don’t have friends because they laughed at me and my child because she was disabled and the father left us as soon as the child was sick. My mom is my friend now’ (Caregiver 8)<br>‘I just worry when he is sick with flu or when get a fits because I know after that he will get sicker and they might think I don’t take care of him. Sometimes I don’t even attend therapy when he is not well’ (Caregiver 23)<br>**Physical stressors**<br>‘I stress a lot when I think about him and his future. I just don’t know what will happen to him when I die because we are both sick and no one supports us’ (Caregiver 10)<br>**Extrinsic barriers to caring for children with cerebral palsy**<br>‘People laugh at you when you walking with your disabled child. People laugh all the time. It’s very sad’ (Caregiver 19)<br>‘Some people in the community would look at you and think that you deserved having a child like this more especially when you are young like us. They say we did not listen to our parents and we are cursed’ (Caregiver 10)<br>‘People still don’t know that our kids with disability are a gift like other kids. Some mothers I know don’t walk with them but they get paid for disability grant. I walk with my child and I’m not ashamed’ (Caregiver 18)<br>‘Finances are always a challenge for me. I sometimes do not have money to take him to hospital for his physiotherapy appointment because I need money for three people to be able to ride the taxi’ (Caregiver 14)<br>‘My family does not support me. Being alone with no one to support is hard [no support from the father and also financially]’ (Caregiver 10) |
Concerning services were discussed. Myrhaug et al., in their work on family-centred services, believed that parental involvement is essential to rehabilitation, as well as adequate information and the sharing of in-service delivery. The lack of information as perceived in this study could be due to the lack of staff and high patient volume. The rehabilitation staff shortage is recognised in South Africa as a barrier to optimal rehabilitation. Dambi and Jelsma’s study also factored high patient volumes versus staff complement as a possible barrier to sharing necessary information. The South African Health care system is premised on 11 Batho Pele Principles, which advocate adequate information being shared when caring for patients, including children with CP and caregivers alike, which is contradicted in the delivery of care as perceived by caregivers in the present study. Hence, further research is critical to explore how to translate these principles into practice in a South African context.

Personal enablers in coping with caring for a child with cerebral palsy in this study were spiritual and religious support. Additionally, community-awareness strategies and support groups were recommended by caregivers to improve the challenges experienced with rehabilitation. Studies with caregivers of children with CP showed that caregivers seek support from others who are in similar situations and often meet in therapeutic environments. These support groups for caregivers have been affirmed as vital and are believed to assist with emotional and informational support. Pfeifer et al. argued, though, that social support provided by extended family, friends and neighbours was a secondary mechanism to the immediate family nucleus. Moreover, it is evident that focusing on the family improves rehabilitation outcomes and family support for a child living with a disability. However, researchers believe that added social support improves psychosocial well-being. Involving the community is also believed to enhance caregivers’ satisfaction and rehabilitation outcomes in providing holistic care to children with special needs.

Conclusion
A holistic approach to individualised assessment, intervention planning and rehabilitation of children with CP should be implemented for quality patient-centred care. The utilisation of a nonconventional contextualised rehabilitation programme as in this study may prove beneficial in resource-limited settings such as South Africa. The aim of this community-based programme was to provide a support structure and education as well as to improve access for the caregivers. Consequently findings revealed high levels of satisfaction with its implementation. However, accessing such services is still plagued by poverty. Financial instability, transport barriers and other environmental barriers inhibit carers accessing such care even though it was placed within the community. This study did, however, aim to equip the caregivers with knowledge and skills that could be translated into a home-based rehabilitation programme. This task-shifting approach attempts to bridge the gap and enable continuity of the rehabilitation programme.

Limitations
The long-term effects of capacitating caregivers need to be evaluated in order to understand whether this approach is appropriate for a South African context. The study is focused on one of the nine provinces in SA and thus has its own restrictions in generalisability of findings.

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Table 3: (Continued)

| Sub-theme | Quotes |
|-----------|--------|
| Transport barriers | ‘Some taxis would stop for you, but when they see that you have a disabled child on a buggy they would just leave you and rush say they are rushing.' (Caregiver 17) |
| Personal enablers to coping with caring for child with cerebral palsy | ‘Prayer helped me. I feel less sad as I pray’ (Caregiver 7) |
| Spiritual and religious support | ‘I think I would tell the community to take their kids to the clinics if they see that their growth is slower or they are later than the other kids same age to advise about caring for their children’ (Caregiver 8) |
| Recommendations by caregivers | ‘Meet with other mothers to encourage them to take step to help their kids. For example, I attend clinics and I continue with their therapy programmes because at a long run they help. I’ll tell them about my child’ (Caregiver 14) |
| Community-awareness strategies | ‘I think I would tell the community to take their kids to the clinics if they see that their growth is slower or they are later than the other kids same age to advise about caring for their children’ (Caregiver 8) |
| Support groups | ‘Meet with other mothers to encourage them to take step to help their kids. For example, I attend clinics and I continue with their therapy programmes because at a long run they help. I’ll tell them about my child’ (Caregiver 14) |
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Received: 22-07-2016 Accepted: 24-10-2016