Perspectives on cerebral palsy in Africa: Exploring the literature through the lens of the International Classification of Functioning, Disability and Health

Sandra Abdel Malek\(^1\)\(^2\) | Peter Rosenbaum\(^1\)\(^2\)\(^3\) | Jan Willem Gorter\(^1\)\(^2\)\(^3\)

\(^1\)School of Rehabilitation Science, McMaster University, Hamilton, Ontario, Canada
\(^2\)CanChild Centre for Childhood Disability Research, McMaster University, Hamilton, Ontario, Canada
\(^3\)Department of Paediatrics, McMaster University, Hamilton, Ontario, Canada

Correspondence
Sandra Abdel Malek, School of Rehabilitation Science, McMaster University, Hamilton, ON, Canada.
Email: abdels11@mcmaster.ca

Abstract

**Background:** The prevalence of cerebral palsy (CP) in countries in Africa is suggested to be higher than in Western countries. Research in Western resource-rich contexts has demonstrated the importance of environmental factors in the activities and participation of individuals with CP, as illustrated by the International Classification of Functioning, Disability and Health (ICF). Although the domains of the ICF are often acknowledged in research on CP in western contexts, the extent to which these domains have been explored in resource-limited areas is unknown. The current review aims to describe the nature of the current published literature on CP in African countries and how it aligns with the domains of the ICF.

**Methods:** Key informants familiar with the literature on CP in Africa were consulted, and a literature search was conducted to identify articles on CP originating from countries in Africa. Identified articles were assessed to determine the ICF domains that align with the studies. Themes relating to each domain were identified.

**Results:** Twenty-seven studies were included. The majority of studies were quantitative (70.3%) and focused on children or caregivers. Most studies included a body functions and structures component (70.4%), focusing on impairment rather than functioning. Activities and participation domains were addressed, but often focused solely on mobility. Environmental factors typically related to resources and policies and personal factors were often unexplored.

**Conclusions:** Research on CP in Africa tends to be grounded in a biomedical approach to disability and does not explicitly use the ICF as a guiding framework. Studies exploring body function and structures tend to be descriptive in nature, whereas activities and participation, and environmental and personal factors, represent modifiable factors that may be addressed with interventions. Knowledge translation activities that promote the ideas of the ICF in an African context should therefore be considered.

**KEYWORDS**
Africa, cerebral palsy, disability, ICF

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Cerebral palsy (CP) is a diagnosis that refers to "a group of permanent disorders of the development of movement and posture causing activity limitation, that are attributed to non-progressive disturbances that occurred in the developing fetal or infant brain" (Rosenbaum et al., 2007). The way in which CP manifests varies greatly between individuals and often includes impairments in "sensation, perception, cognition, communication, and behaviour, by epilepsy, and by secondary musculoskeletal problems" (Rosenbaum et al., 2007). CP displays heterogeneity in "presentation, etiology, evolution, severity, medical and rehabilitation needs, comorbidities, and outcomes" (Shevell, Miller, Scherer, Yager, & Fehlings, 2011); however, various medical and rehabilitative interventions throughout development have the potential to help children with CP engage in meaningful life activities. The prevalence of CP in Western developed countries has been estimated to be approximately 2–2.5 per 1,000 children (Oskoui, Joseph, Dagenais, & Shevell, 2013). Despite concerns of underreporting, this prevalence has been suggested to be even higher in developing countries throughout Africa (Donald, Samia, Kakooza-Mwesige, & Bearden, 2014).

Views of childhood disability have typically been grounded in a biomedical approach, with an emphasis on “fixing” impairments and achieving normality (Rosenbaum & Gorter, 2012). This traditional approach has been challenged in recent years, given that this model largely ignores society’s contribution to the construction of disability and often discounts social and legal barriers (Wiart & Darrah, 2002). Disability as a sociopolitical movement has emerged in the Western world over the last several decades with the aim of redefining "disability" in terms of shortcomings of the community resulting in limitations to participation, rather than viewing disability as a problem inherent to the individual, as has usually been done (Wiart & Darrah, 2002). This movement mirrors the transition from the World Health Organization’s (WHO) 1980 International Classification of Impairment, Disability and Handicap (ICIDH) to the 2001 International Classification of Functioning, Disability and Health (ICF). The ICIDH was a hierarchical framework consisting of impairment (organ level), disability (functional abilities), and handicap (disadvantages experienced as a result; Wood, 1980). However, the ICIDH focused on impairments rather than functioning and failed to address factors external to the individual. To address these shortcomings, the WHO released the ICF and its framework for health (Figure 1) in 2001.

The multidimensional ICF framework conceptualizes functioning and disability across four interconnected domains: body functions and structures, activities and participation, environmental factors, and personal factors. The domains of the ICF emphasize the biopsychosocial nature of functioning, providing an alternative way of viewing disability (WHO, 2001) and forming a valuable framework for the assessment and management of children and youth with CP (Rosenbaum & Stewart, 2004). For example, when considering treatment outcomes, parents, medical professionals, and youth with CP discussed several concepts that align with all domains of the ICF (Vargus-Adams & Martin, 2011). The diversity of these goals across the ICF may be expected, given that the domains of the ICF have been suggested to be interdependent, with relationships between domains likely to be influenced by other factors within the framework. Environmental factors have specifically been thought to play a role in the relationship between the domains

**FIGURE 1** International Classification of Functioning, Disability and Health (World Health Organization, 2001)
between body function and structure and activities and participation (Wright, Rosenbaum, Goldsmith, Law, & Fehlings, 2008).

The importance of environmental factors in the activities and participation of individuals with CP has been demonstrated in a variety of contexts, including at home, school, and within the community (Anaby et al., 2014; Law et al., 2013). Research has generally focused on the role of environmental factors in Western resource-rich contexts, such as Canada, the United States, the United Kingdom, Australia, and Sweden (Anaby et al., 2013; Imms, 2008). However, environments in resource-limited contexts differ markedly from those of the Western world. There is a paucity of literature regarding the extent to which environmental factors, activities, and participation have been explored in resource-limited African countries, despite the likelihood of a higher prevalence of CP in these regions. A systematic review on CP in Africa revealed that research typically focuses on prevalence, aetiology, classification, treatment and management, and comorbidities (Donald et al., 2014). That review also revealed many gaps in the literature such as a lack of qualitative and mixed methods studies exploring individual and family understanding of CP, a lack of studies addressing access to resources and barriers to care, and a lack of longitudinal studies assessing outcomes (Donald et al., 2014). Additionally, although the ICF has been available since 2001, the acknowledgement and application of its domains across cultural contexts, particularly in African countries in both research and practice, remains to be explored.

The purpose of this literature review is therefore to explore the current published literature about CP in African countries. Specifically, this review aims to answer the following questions:

1) What is the nature and extent of the current published literature on cerebral palsy in countries in Africa?

2) How does the research focus of the published literature align with the domains of the ICF?

Applying the ICF framework to the current research will provide information regarding the recognition and endorsement of the domains of the ICF in varying cultural contexts and the need for knowledge translation activities to promote these ideas of functioning and disability.

2 | METHODS

A literature review was conducted in order to explore the scope of the literature regarding CP in Africa and to categorize the literature according to the corresponding ICF domains. To identify relevant studies, key informants who are familiar with the literature in Africa were contacted and provided the names of several prominent researchers who focus on CP in an African context. An author search was conducted through Google Scholar and the PubMed database to identify relevant articles. Additionally, a search of the PubMed database was conducted using medical subject heading terms. Specifically, “cerebral palsy” and “Africa, northern” OR “Africa, south of the Sahara” were used. These terms encompass studies pertaining to cerebral palsy across all African countries.

Reference lists were hand searched to identify other relevant articles.

2.1 | Inclusion/exclusion criteria

Title and abstracts of articles were screened, and relevant articles underwent a full-text review. Articles were included if they were peer-reviewed and published within the last 5 years (2014–present) and if the full text was available. Articles published prior to 2014 were excluded given that the only systematic review (to the author's knowledge) on CP in Africa was published in 2014 and encompassed research from several databases, African and international disability organizations, commercial search engines, and local journals. The present literature review therefore provides an update to the information presented in 2014. Only articles that explicitly reported on CP (rather than other neurodisabilities or developmental disorders in general) in any setting (e.g., hospitals, schools, out-patient centres) were included. Additionally, only articles that reported results for an African country or region were included; articles that reported on “low-resource settings” or “low-income countries” without explicitly providing information relating to an African context were not included. Correspondences, commentaries, and editorials were not included, and no restrictions were placed on language.

2.2 | Data extraction

The purpose and outcomes of each included study were assessed to determine the ICF domains that align with the study. The ICF domains were determined according to the perspective of the individual with CP, rather than the participants of the study (e.g., caregivers’ mental health was reported as the individual with CP’s environment rather than as the caregiver’s body function and structure). The results were analysed, and themes relating to each ICF domain were identified.

3 | RESULTS

Using the inclusion criteria described above, 34 studies were identified. Following title and abstract screening, removal of duplicates, and the inclusion of articles found through the reference search, 27 full-text articles were included in the review (Table 1). Of the included studies, three were published in 2014, seven in 2015, six in 2016, seven in 2017, and four in 2018. All included articles were in English, with no studies meeting inclusion criteria found in other languages.

The included articles originated from 10 African countries, whereas one paper looked at Africa as a continent. Of studies looking at specific countries, one was conducted in each of the following: Ghana (3.7%), Zambia (3.7%), Malawi (3.7%), and Ethiopia (3.7%). Three studies were conducted in Egypt (11.1%), Nigeria (11.1%), and Zimbabwe (11.1%). Four studies were conducted in Botswana (14.8%) and South Africa (14.8%), and five in Uganda (18.5%). Of the included
### Table 1: Articles meeting inclusion criteria and included in review

| Authors | Title                                                                 | Year | Journal                                                      | Country |
|---------|------------------------------------------------------------------------|------|--------------------------------------------------------------|---------|
| Abd El-Kafy, E. M. | The clinical impact of orthotic correction of lower limb rotational deformities in children with cerebral palsy: A randomized controlled trial. | 2014 | Clinical Rehabilitation                                      | Egypt   |
| Adepoju, F., Hamzat, T., & Akinyinka, O. | Comparative efficacy of progressive resistance exercise and biomechanical ankle platform system on functional indices of children with cerebral palsy. | 2017 | Ethiopian Journal of Health Sciences                         | Nigeria |
| Adolfsson, M., Johnson, E., & Nilsson, S. | Pain management for children with cerebral palsy in school settings in two cultures: Action and reaction approaches | 2018 | Disability and Rehabilitation                               | South Africa |
| Bearden, D. R., Monokwane, B., Khurana, E., Baier, J., Baranov, E., Westmoreland, K., & Steenhoff, A. P. | Pediatric cerebral palsy in Botswana: etiology, outcomes, and comorbidities | 2016 | Pediatric Neurology                                          | Botswana |
| Chikwanha, T. M., Chidhakwa, S., & Dangarembizi, N. | Occupational therapy needs of adolescents and young adults with cerebral palsy in Zimbabwe: Caregivers' perspectives. | 2015 | Central African Journal of Medicine                          | Zimbabwe |
| Chiluba, B. C., & Moyo, G. | Caring for a cerebral palsy child: A caregivers perspective at the University Teaching Hospital, Zambia. | 2017 | BMC Research Notes                                          | Zambia |
| Coker-Bolt, P., DeLuca, S. C., & Ramey, S. L. | Training paediatric therapists to deliver constraint-induced movement therapy (CIMT) in sub-Saharan Africa. | 2015 | Occupational Therapy International                          | Ethiopia |
| Conchar, L., Bantjes, J., Swartz, L., & Derman, W. | Barriers and facilitators to participation in physical activity: The experiences of a group of South African adolescents with cerebral palsy | 2016 | Journal of Health Psychology                                 | South Africa |
| Dambi, J. M., & Jelsma, J. | The impact of hospital-based and community based models of cerebral palsy rehabilitation: a quasi-experimental study. | 2014 | BMC Pediatrics                                               | Zimbabwe |
| Dambi, J. M., Jelsma, J., & Mlambo, T. | Caring for a child with Cerebral Palsy: The experience of Zimbabwean mothers. | 2015 | African Journal of Disability                               | Zimbabwe |
| Dambi, J. M., Mandizvidza, C., Chiwaridzo, M., Nhuwzi, C., & Tadyanemhandu, C. | Does an educational workshop have an impact on caregivers' levels of knowledge about cerebral palsy? A comparative, descriptive cross-sectional survey of Zimbabwean caregivers. | 2016 | Malawi Medical Journal                                       | Zimbabwe |
| Donald, K. A., Kakooza, A. M., Wammanda, R. D., Mallewa, M., Samia, P., Babakir, H., & Chugani, H. | Pediatric cerebral palsy in Africa: Where are we? | 2015 | Journal of Child Neurology                                  | Africa |
| El-Tallawy, H. N., Farghaly, W. M., Shehata, G. A., Rageh, T. A., Metwally, N. A., Badry, R., & Kandil, M. R. | Cerebral palsy in Al-Quseir City, Egypt: Prevalence, subtypes, and risk factors. | 2014 | Neuropsychiatric Disease and Treatment                     | Egypt |
| Johnson, A., Gambah-Sampaney, C., Khurana, E., Baier, J., Baranov, E., Monokwane, B., & Bearden, D. R. | Risk factors for malnutrition among children with cerebral palsy in Botswana. | 2017 | Pediatric Neurology                                         | Botswana |
| Johnson, E., Nilsson, S., & Adolfsson, M. | Eina! Ouch! Eish! Professionals' perceptions of how children with | 2015 | Augmentative and Alternative Communication                  | South Africa |
| (Continues) | | | | |


**Studies and Participants:**

- One was a comparative study between South Africa and Sweden, and one was a cross-cultural study in Malawi, Russia, Poland, and Brazil. This cross-cultural study was the only one that explicitly used the ICF.
- The study by Schiariti et al. (2018) in Malawi used the International Classification of Functioning, Disability, and Health (ICF) to assess the implementation of its core sets for children and youth with cerebral palsy.

**Participants:**

- In the majority of the studies (12; 44.4%), participants were children ranging from 0 to 18 years of age. Six (22.2%) of the studies used caregivers of individuals with CP as participants—five (18.5%) of these studies were interested in those caregivers' perspectives on pain management.

### Table 1 (Continued)

| Authors | Title | Year | Journal | Country |
|---------|-------|------|---------|---------|
| Kakooza-Mwesige, A., Andrews, C., Peterson, S., Mangen, F. W., Eliasson, A. C., & Forssberg, H. | Prevalence of cerebral palsy in Uganda: A population-based study. | 2017 | The Lancet Global Health | Uganda |
| Kakooza-Mwesige, A., Forssberg, H., Eliasson, A. C., & Tumwine, J. K. | Cerebral palsy in children in Kampala, Uganda: Clinical subtypes, motor function and co-morbidities. | 2015 | BMC Research Notes | Uganda |
| Kakooza-Mwesige, A., Byanyima, R. K., Tumwine, J. K., Eliasson, A. C., Forssberg, H., & Flodmark, O. | Grey matter brain injuries are common in Ugandan children with cerebral palsy suggesting a perinatal aetiology in full-term infants. | 2016 | Acta Paediatrica | Uganda |
| Kakooza-Mwesige, A., Tumwine, J. K., Eliasson, A. C., Namusoke, H. K., & Forssberg, H. | Malnutrition is common in Ugandan children with cerebral palsy, particularly those over the age of five and those who had neonatal complications. | 2015 | Acta Paediatrica | Uganda |

| Monokwane, B., Johnson, A., Gambrah-Sampaney, C., Khurana, E., Baier, J., Baranov, E., ... & Bearden, D. R. | Risk factors for cerebral palsy in children in Botswana. | 2017 | Pediatric Neurology | Botswana |

| Munyuru, K., Idro, R., Abbo, C., Kadamukasa, M., Katabira, E., Mupere, E., & Kakooza-Mwesige, A. | Prevalence and factors associated with sleep disorders among children with cerebral palsy in Uganda; a cross-sectional study. | 2018 | BMC Pediatrics | Uganda |

| Nilsson, S., Johnson, E., & Adolfsson, M. | Professionals' perceptions about the need for pain management interventions for children with cerebral palsy in South African school settings. | 2016 | Pain Management Nursing | South Africa |

| Ogoke, C. C., & Iloeje, S. O. | Severity of motor dysfunction in children with cerebral palsy seen in Enugu, Nigeria. | 2017 | Pan African Medical Journal | Nigeria |

| Patel, P., Baier, J., Baranov, E., Khurana, E., Gambrah-Sampaney, C., Johnson, A., ... & Bearden, D. R. | Health beliefs regarding pediatric cerebral palsy among caregivers in Botswana: A qualitative study. | 2017 | Child: Care, Health and Development | Botswana |

| Polack, S., Adams, M., O`banion, D., Baltussen, M., Asante, S., Kerac, M., ... & Zuurmond, M. | Children with cerebral palsy in Ghana: malnutrition, feeding challenges, and caregiver quality of life. | 2018 | Developmental Medicine & Child Neurology | Ghana |

| Schiariti, V., Longo, E., Shoshmin, A., Kozhushko, L., Besstrashnova, Y., Król, M., ... & Mwale, M. | Implementation of the International Classification of Functioning, Disability, and Health (ICF) core sets for children and youth with cerebral palsy: Global initiatives promoting optimal functioning. | 2018 | International Journal of Environmental Research and Public Health | Malawi |

| Shehata, A. I., Hassanein, F. I., & Abdul-Ghani, R. | Seroprevalence of Toxoplasma gondii infection among patients with non-schizophrenic neurodevelopmental disorders in Alexandria, Egypt. | 2016 | Acta Tropica | Egypt |
providing care to children, whereas one study (3.7%) explored the perspectives of caregivers of adolescents. Five studies (18.5%) were conducted with professionals (teachers, clinicians, etc.), and two (7.4%) used an adult population. One study (3.7%) used both caregivers and children with CP as participants, whereas another study recruited individuals of all ages with CP.

Four studies (14.8%) were conducted in rehabilitation centres, four (14.8%) in school settings, and four (14.8%) in the community. Six (22.2%) of the studies occurred in a hospital (inpatient and outpatient sites) and nine (33.3%) took place in a referral centre (inpatient and outpatient, paediatric neurology clinics, and tertiary referral centres).

Nineteen (70.3%) of the studies were quantitative in nature, two (7.4%) used mixed methods, and six (22.2%) were qualitative studies. Of the six qualitative studies, one explored the perspectives of adolescents with CP, one explored caregiver perspectives, three explored perspectives of professionals (teachers, occupational therapists, physical therapists, social workers) in a school setting, and one described a meeting of physicians across Africa related to the evaluation and management of children with CP.

### 3.1 | ICF domains

In terms of ICF domains, some studies focused solely on one domain. For example, a study investigating neuroimaging patterns, specifically grey matter brain injuries, of children with CP aligns solely with the body functions and structures domain (Kakooza-Mwesige et al., 2016). However, the majority explored multiple domains, such as a study exploring the clinical subtypes, motor function, and comorbidities of children with CP (Kakooza-Mwesige, Forssberg, Eliasson, & Tumwine, 2015). This study aligns with body functions and structures (clinical subtype; speech, visual, and hearing impairments as comorbidities) as well as activities (motor function; learning impairment as a comorbidity). Table 2 outlines the ICF domains addressed by each included study.

Most studies (20; 74.1%) included a body functions and structures component. Studies that addressed body functions and structures were found to fall into three categories: those exploring comorbidities, those exploring risk factors, and those that are descriptive in nature. Activities were addressed by 11 (40.7%) studies, with the majority (9; 81.8%) focusing solely on mobility and the remaining two addressing self-care and learning. Participation was the least addressed of the domains, with only one of five studies addressing more than one area of participation. Participation focused on recreation and leisure, religion, education, and interpersonal interactions. Environmental factors were explored by 16 (59%) of the included studies. Environmental factors were related to the resources and policies in different settings, the support available to the individual, and attitudes of family and society in general. Three studies addressed products/technology, and three explored aspects of the natural environment. Personal factors were explored in two of the included studies. Table 3 indicates concepts addressed within each ICF domain by the included studies.

### 4 | DISCUSSION

This review reveals the nature of the current literature about CP in African countries and how the focus of the published research aligns with the domains of the ICF. The literature appears to be grounded in a biomedical approach to disability, with the majority of studies exploring some aspect of body functions and structures and a minority exploring activities and participation beyond mobility.

Studies included in this review indicate that the published research is not representative of all African countries, given that the included studies originate from 10 countries out of a possible 54. This may be due to a lack of resources in many African countries to support this kind of research. According to the WHO, the African Region (excluding countries in the Eastern Mediterranean Region such as Egypt and Morocco) accounts for 24% of the international burden of disease while having access to only 1% of the world’s financial resources for health (WHO, n.d.-a). Additionally, resources are more often allocated to curative services that tend to be high-cost, rather than to prevention and health promotion measures (WHO, n.d.-a). These services are often focused on infectious diseases rather than noncommunicable diseases, including disabilities. This reality is likely attributable to a variety of factors, such as weak health systems, inadequate training for healthcare providers, lack of financial resources, and a perception that infectious diseases should be the primary health focus of the region (WHO, n.d.-a). This may explain, at least in part, some of the cross-continental underrepresentation of conditions like CP in the literature. Additionally, differences in education levels may play a role in which countries are represented in the literature. The 10 countries from which research was identified have an average literacy rate of 72.2%, with the majority of the unrepresented countries having a lower percentage, ranging as low as 22% in Chad and 14% in Niger (United Nations, 2018). Lower resources may therefore be a barrier to conducting research in these areas, posing a challenge in painting an accurate picture of the understanding of CP across Africa.

Participants in the included studies were generally children with CP or individuals interacting with children with CP, such as professionals or caregivers. Very few studies explored CP in adolescence or adulthood. This may be due to the lower life expectancy observed across Africa (average 61 years) compared with the global life expectancy (72 years; WHO, n.d.-b). In Western contexts, children with CP are likely to have a life expectancy similar to that of the general population, provided that appropriate healthcare is available (Koman, Smith, & Shlitt, 2004; Strauss & Shavelle, 1998). However, this may not be the case in African countries where healthcare resources are often limited (Donald et al., 2015). Infants are at a higher risk of prematurity, low birth weight, and complications such as birth asphyxia (WHO Regional Office for Africa, 2014), all of which may contribute to an increased risk of CP (Odding, Roebroeck, & Stam, 2006). Although infant mortality rates have been decreasing (WHO Regional Office for Africa, 2014), countries in sub-Saharan Africa have the lowest improvements in child survival (WHO, n.d.-a), which may contribute to CP being perceived as a childhood disorder and thus being researched as such. However,
| Authors | Title | Body functions/structures | Activities | Participation | Environment |
|---------|-------|---------------------------|------------|---------------|-------------|
| Abd El-Kafy, E. M. | The clinical impact of orthotic correction of lower limb rotational deformities in children with cerebral palsy: A randomized controlled trial. | x | | | |
| Adepoju, F., Hamzat, T., & Akinyinka, O. | Comparative efficacy of progressive resistance exercise and biomechanical ankle platform system on functional indices of children with cerebral palsy. | | x | | |
| Adolfsson, M., Johnson, E., & Nilsson, S. | Pain management for children with cerebral palsy in school settings in two cultures: Action and reaction approaches | x | x | x | |
| Bearden, D. R., Monokwane, B., Khurana, E., Baier, J., Baranov, E., Westmoreland, K., ... & Steenhoff, A. P. | Pediatric cerebral palsy in Botswana: Etiology, outcomes, and comorbidities | x | | x | |
| Chikwanha, T. M., Chidhakwa, S., & Dangarembizi, N. | Occupational therapy needs of adolescents and young adults with cerebral palsy in Zimbabwe: Caregivers’ perspectives. | | x | x | x |
| Chiluba, B. C., & Moyo, G. | Caring for a cerebral palsy child: A caregiver’s perspective at the University Teaching Hospital, Zambia. | | | | x |
| Coker-Bolt, P., DeLuca, S. C., & Ramey, S. L. | Training paediatric therapists to deliver constraint-induced movement therapy (CIMT) in sub-Saharan Africa. | x | | | |
| Conchar, L., Bantjes, J., Swartz, L., & Derman, W. | Barriers and facilitators to participation in physical activity: The experiences of a group of South African adolescents with cerebral palsy | x | x | | |
| Dambi, J. M., & Jelsma, J. | The impact of hospital-based and community based models of cerebral palsy rehabilitation: A quasi-experimental study. | | x | x | |
| Dambi, J. M., Jelsma, J., & Mlambo, T. | Caring for a child with Cerebral Palsy: The experience of Zimbabwean mothers. | | | | |
| Dambi, J. M., Mandizvidza, C., Chwiridzo, M., Nnumzi, C., & Tdyanemhandu, C. | Does an educational workshop have an impact on caregivers’ levels of knowledge about cerebral palsy? A comparative, descriptive cross-sectional survey of Zimbabwean caregivers. | | | | x |
| Donald, K. A., Kakooza, A. M., Wammanda, R. D., Mallewa, M., Samia, P., Babakir, H., ... & Chugani, H. | Pediatric cerebral palsy in Africa: Where are we? | | | | x |
| El-Tallawy, H. N., Farghaly, W. M., Shehata, G. A., rageh, T. A., Metwally, N. A., Badry, R., ... & Kandill, M. R. | Cerebral palsy in Al-Quseir City, Egypt: Prevalence, subtypes, and risk factors. | | | x | |
| Johnson, A., Gambrah-Sampaney, C., Khurana, E., Baier, J., Baranov, E., Monokwane, B., & Bearden, D. R. | Risk factors for malnutrition among children with cerebral palsy in Botswana. | x | x | | |
| Johnson, E., Nilsson, S., & Adolfsson, M. | Eina! Ouch! Eish! Professional’ perceptions of how children with cerebral palsy communicate about pain in South African school settings: Implications for the use of AAC. | | x | | |

(Continues)
studies in Western contexts have shown the importance of addressing the needs of adolescents with CP (Gorter, Stewart, & Woodbury-Smith, 2011), particularly because participation seems to become more restricted as children transition into adolescence and adulthood (King et al., 2003), which may have a negative influence on health and well-being (Shikako-Thomas, Kolehmainen, Ketelaar, Bult, & Law, 2014). Therefore, future studies should aim to explore adolescents’ experiences and perspectives in an African context to promote well-being and functioning according to the ICF domains.

4.1 | ICF domains

This literature review revealed that the ICF is not explicitly addressed in research on CP in African countries. However, as the current work

| Table 2 (Continued) |
|---------------------|
| **Authors** | **Title** | **Body functions/structures** | **Activities** | **Participation** | **Environment** |
| Kakooza-Mwesige, A., Andrews, C., Peterson, S., Mangen, F. W., Eliasson, A. C., & Forssberg, H. | Prevalence of cerebral palsy in Uganda: A population-based study. | x | x |
| Kakooza-Mwesige, A., Forssberg, H., Eliasson, A. C., & Tumwine, J. K. | Cerebral palsy in children in Kampala, Uganda: Clinical subtypes, motor function and co-morbidities. | x | x |
| Kakooza-Mwesige, A., Byanyima, R. K., Tumwine, J. K., Eliasson, A. C., Forssberg, H., & Flodmark, O. | Grey matter brain injuries are common in Ugandan children with cerebral palsy suggesting a perinatal aetiology in full-term infants. | x |
| Kakooza-Mwesige, A., Tumwine, J. K., Eliasson, A. C., Namusoke, H. K., & Forssberg, H. | Malnutrition is common in Ugandan children with cerebral palsy, particularly those over the age of five and those who had neonatal complications. | x |
| Monkwane, B., Johnson, A., Gambrah-Sampaney, C., Khurana, E., Baier, J., Baranov, E., ... & Bearden, D. R. | Risk factors for cerebral palsy in children in Botswana. | x | x |
| Munyumu, K., Idro, R., Abbo, C., Kaddumukasa, M., Katabira, E., Mupere, E., & Kakooza-Mwesige, A. | Prevalence and factors associated with sleep disorders among children with cerebral palsy in Uganda; a cross-sectional study. | x |
| Nilsson, S., Johnson, E., & Adolffson, M. | Professionals’ perceptions about the need for pain management interventions for children with cerebral palsy in South African school settings. | x | x | x |
| Ogoke, C. C., & Iloeje, S. O. | Severity of motor dysfunction in children with cerebral palsy seen in Enugu, Nigeria. | x | x |
| Patel, P., Baier, J., Baranov, E., Khurana, E., Gambrah-Sampaney, C., Johnson, A., ... & Bearden, D. R. | Health beliefs regarding pediatric cerebral palsy among caregivers in Botswana: A qualitative study. | x |
| Polack, S., Adams, M., O’banion, D., Baltussen, M., Asante, S., Kerac, M., ... & Zuurmond, M. | Children with cerebral palsy in Ghana: Malnutrition, feeding challenges, and caregiver quality of life. | x | x | x |
| Schiariti, V., Longo, E., Shoshmin, A., Kozhushko, L., Besstrashnova, Y., Król, M., ... & Mwale, M. | Implementation of the International Classification of Functioning, Disability, and Health (ICF) core sets for children and youth with cerebral palsy: Global initiatives promoting optimal functioning. | x | x | x | x |
| Shehata, A. I., Hassanein, F. I., & Abdul-Ghani, R. | Seroprevalence of Toxoplasma gondii infection among patients with non-schizophrenic neurodevelopmental disorders in Alexandria, Egypt. | x | x |

Total number of studies addressing each ICF domain.
illustrates, it is possible implicitly to align the findings of these studies analytically to a variety of concepts within each domain of the ICF. Each domain is discussed in further detail below.

### 4.1.1 | Body functions and structures

The majority of the studies included in this review had a body functions and structures component, with many exploring comorbidities such as malnutrition and visual, hearing, and speech impairments. Risk factors included infections and issues relating to birth, such as prematurity and neonatal infections. Descriptive studies focused on neuromusculoskeletal impairments and clinical subtype. Studies exploring components of body functions and structures tended to focus on impairments rather than functioning, which aligns with the biomedical view of disability (Rosenbaum & Gorter, 2012). However, research suggests that interventions targeting impairments do not necessarily increase functional independence and participation (Law & Darrah, 2014; Novak, 2012; Wright et al., 2008), indicating a need for interventions, and research, to go beyond the scope of physical impairment in order to benefit individuals with CP more effectively.

Additionally, only one study explored the mental health of individuals with CP by investigating depression and anxiety among adolescents in South Africa as it relates to their participation in physical activity (Conchar, Bantjes, Swartz, & Derman, 2016). Children with CP have been suggested to be at a higher risk of experiencing symptoms associated with mental illness (Downs et al., 2018), suggesting the need to investigate the extent to which these disorders affect adolescents and young adults with CP in an African context.

### 4.1.2 | Activities and participation

Mobility was the most reported activity and was addressed in 11 out of 14 studies that aligned with the activities domain, indicating a narrow focus of the current research. The majority of these studies used the Gross Motor Function Classification System (Palisano, Rosenbaum, Bartlett, & Livingston, 2008) to conceptualize mobility and classify participants according to motor functioning. Participation was the least reported domain—out of a total of five studies addressing components of participation, three were explicit in their aim to investigate this concept. Children and adolescents with CP in resource-rich contexts have been found to engage in a narrower range of activities compared with their typically developing peers, with more activities occurring at home, alone, or with family rather than peers (Engel-Yeger, Jarus, Anaby, & Law, 2009; King, Law, Hurley, Petrenchik, & Schwellnus, 2010; Shikako-Thomas, Majnemer, Law, & Lach, 2008). Groups of international researchers, as well as clinicians and consumers, have therefore identified participation as a priority for research and interventions (McIntyre, Novak, & Cusick, 2010; Schiariti, Masse et al., 2014). Given that environmental factors have been shown to play an important role in meaningful participation (Anaby et al., 2013; Shikako-Thomas et al., 2008), it is important to investigate the participation of individuals with CP in different cultural contexts, particularly in resource-limited African countries, where the environment differs significantly from resource-rich countries from which the majority of the literature on participation originates.

### 4.1.3 | Environmental factors

A scoping review of environmental factors influencing participation of children and youth with disabilities identified social support and geographic location as the most common facilitators of participation, whereas attitudes, physical environment, transportation, policies, and lack of support were the most frequently cited barriers (Anaby et al., 2013). Every category of the environment domain has been found to influence participation in some way, whether it be as a facilitator or as a barrier (Anaby et al., 2013). The main environmental contextual factors that were addressed by the studies included in this review are resources and policies, support and relationships, and attitudes. Resources and policies were described in the context of schools, transportation, and health services, as well as more broadly at the national level. Support was provided by two main sources, caregivers...
and professionals. Familial and societal attitudes were also discussed. Importantly, although contextual factors were discussed in the literature, personal factors were rarely addressed. Although the social and cultural environment influences an individual's participation in daily activities (Law et al., 1999), it has been suggested that the effects of both environmental and personal factors on health and well-being should be equally explored (Rosenbaum & Stewart, 2004).

4.1.4 | Personal factors

Personal factors such as age and gender, as well as likes and interests, have been shown to play a role in the participation of children and youth with CP (Shikako-Thomas et al., 2008; Verschuren, Wiart, Hermans, & Ketelaar, 2012). Of the qualitative studies addressing environmental factors, only one considered the perspective of adolescents with CP; the remaining qualitative/mixed methods studies reported the perspectives of caregivers and professionals. The importance of considering the perspectives of both children and their caregivers has been described in the literature (Garth & Aroni, 2003), particularly because children and caregivers often have differing perspectives on the same topics (Schiralti, Suave et al., 2014). Individuals with CP may therefore perceive, experience, or value things differently than their caregivers, offering unique input in research and in the development of interventions. Future studies should therefore consider the perspectives of individuals with CP, and particularly their personal factors, and ways in which these factors can be incorporated in clinical practice.

This literature review has some potential limitations. First, only one database was searched by the first author (S. A.), so articles from other databases or from African journals not included in the database may have been missed. Additionally, articles published in other languages and not included in PubMed may have also been missed. However, this literature review was meant to be exploratory in nature in order to provide a sense of the current literature on CP in Africa, rather than trying to be exhaustive. Next steps therefore include creating a comprehensive search strategy for multiple databases and searching the grey literature in order to assess whether there may be a more accurate state of knowledge on CP in the African continent.

5 | CONCLUSION

The current published literature on CP in Africa originates from a handful of countries. The target population of these studies is largely children with CP, with very few studies looking at adolescents and adults. Additionally, the majority of the studies are quantitative in nature and primarily have a body functions and structures component focusing on impairment rather than on functioning. Studies addressing body functions and structures report comorbidities and risk factors and tend to be descriptive in nature. In contrast, studies that align with activities and participation, and environmental and personal factors, provide insight into modifiable factors that may be addressed with interventions. Mobility was the most reported activity, suggesting the need for research regarding other activities. Participation was the least reported domain, although it has been shown to be closely linked to health and well-being. Interventions aiming to increase participation in all activities of life should therefore be considered. Participation may be increased by focusing interventions on environmental modifications. The most reported environmental factors were contextual, relating to resources and policies, support, and attitudes. Policy changes may be necessary to provide more support for caregivers and to reduce barriers to participation in schools and in the larger community. Personal factors should also be addressed when developing interventions by investigating the perspectives of individuals with CP and by exploring ways in which these factors can be applied to clinical practice. This literature review reveals the need for knowledge translation activities to promote the ideas presented by the ICF in an African context.

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ORCID

Sandra Abdel Malek @ https://orcid.org/0000-0002-8230-7879
Peter Rosenbaum @ https://orcid.org/0000-0001-6751-5613
Jan Willem Gorter @ https://orcid.org/0000-0002-3012-2119

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