Being a mother of a child with HIV-related Neurodevelopmental Disorders in the Zimbabwean Context

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**ABSTRACT**

**Background:** There is a growing population of mothers caring for their biological children who are infected with Human Immunodeficiency Virus (HIV), in Zimbabwe. Many of these children present with HIV-related Neuro Developmental Delays (NDDs). The occupation of being a mother is a complex and multifaceted role geared towards caring for and nurturing children. The different ways in which mothers negotiate the unique circumstances linked to the occupation of being a mother to a child with diagnosis of HIV-related NDDs warrants exploration.

**Aim:** The aim of the study was to describe the mother’s experiences of engaging in daily occupations relating to caring for their child with HIV-related NDDs.

**Methodology:** A descriptive qualitative study using a hermeneutic phenomenological approach was used to uncover the mothers’ lived experiences of caring for their child with HIV-related NDDs. Data generated from phenomenological interviews conducted with five mothers were analysed inductively using a simplified version of the Stevick Colaizzi Keen method.

**Findings:** Two major themes, namely ‘Ndozvazviri’ (Resilient Acceptance) and ‘Rekindled hope for the future’ emerged from the findings. These themes revealed that caring for a child with HIV-related NDDs is a difficult and demanding role. Despite this, mothers accepted and found meaning in this caring role. Their meaning was expressed through the opportunity to care for their own child and to observe their progress in occupational development and engagement. These interactions created positive experiences for the mothers and rekindled their hope for the future of their child.

**Discussion and recommendations:** Despite the huge demands associated with being a mother of a child with HIV-related NDDs, mothers were committed to this role and were reluctant to entrust this role to others. The findings of this study encourage occupational therapists designing interventions for families, to carefully consider how the mothering role positively shapes the identities of mothers caring for children with HIV-related NDDs.

**Key words:** mothering, HIV-related NDDs, caregiver

**INTRODUCTION**

The academic discipline of occupational science supports the exploration of occupational roles in order to increase our understanding of identity and the daily activities that are associated with these roles. The occupational role of mothering is defined as a social practice geared toward nurturing and caring for dependent children. It is a complex and dynamic role in which mothers devote their time to caring for children as well as being involved in other occupations geared to meet the demands of being a mother. Caring for children constitutes an opportunity for mothers to generate meaning and construct their individual identity as a mother and carer of their child.

Research in the profession of occupational therapy and the academic discipline of occupational science has focused on describing the lived experiences of mothers caring for their children with various physical and mental disabilities. These studies report that caring for a child with a disability impacts on a mother’s occupation. Despite the huge demands associated with being a mother of a child with HIV-related NDDs, mothers were committed to this role and were reluctant to entrust this role to others. The findings of this study encourage occupational therapists designing interventions for families, to carefully consider how the mothering role positively shapes the identities of mothers caring for children with HIV-related NDDs.

Prior to the onset of medical interventions in the form of Antiretroviral Therapy (ART) / Highly Active Antiretroviral Therapy (HAART) it was common for most carers of HIV infected children in Africa to be either grandmothers, siblings or extended family members because most biological mothers would have died or would have been too ill as a result of their own HIV/AIDS diagnosis. Advances in healthcare practices for HIV/AIDS, linked to the advent of antiretroviral therapy (ART/HAART), have increased the life expectancy and quality of life of HIV infected people. Thus, parents of children with HIV are now able to continue with their role of caring for their children.

The diagnosis of a child as HIV positive almost always means that at least the mother (and possibly, the father) is also infected. This makes caring for an HIV infected child by a biological mother unique because the mother has to deal with her own diagnosis, and the possibility that both the child and their partner/s have the disease as well. Many mothers only learn about their HIV status at the time that they find out that they are pregnant. They deal with feelings of shock, anxiety, fear and guilt linked to this discovery.

The stigma linked to a diagnosis of HIV/AIDS adds an additional layer to the burden of care, compelling mothers to deal with the tension between the preference for secrecy surrounding the disease and the openness required for them to access health...
and educational services for their child and seek and accept social support\(^1\). In Burkina Faso, Hejoaka\(^2\) explored the life experiences of being a mother of a child with HIV/AIDS. The study findings revealed that often HIV-positive mothers tend to silence their own needs and attend to the needs of their children. In most instances, the mothers ignore their physical, psychological and social needs and/or seek services and support late, thus negatively impacting on their own general health and, consequently, the health of their child. These unique challenges, related to the caregiving experience, motivate the need for an inquiry into the lived experiences of mothers caring for their own children with HIV-related NDDs. This paper aims to explore how personal and contextual factors shaped the occupation of caring for a child with HIV. The research is located in Zimbabwe\(^2,6,8\), the home country of the researcher, where 9.4% of children contracting HIV from their mothers are at risk of developing NDD in Zimbabwe\(^9\).

**METHODOLOGY**

A descriptive qualitative study using a hermeneutic phenomenological approach was employed in this study as it allows the researcher to elicit and understand how the phenomenon of being a caregiver of a child with a HIV-related neurodevelopmental disorder is lived and experienced\(^4\). Qualitative research provides individuals with a voice to relay their unique experiences in a way that they can be well represented and understood\(^5\). Mothers were recruited from the Harare Hospital Children’s Rehabilitation Unit (CRU), the country’s largest referral unit for children with disabilities in Zimbabwe. Through the assistance of the administrator, children whose biological mothers were registered as primary caregivers were identified and listed and using purposive sampling, a list of fourteen mothers who met the selection criteria was obtained. Using the maximum variation criteria, five mothers who had different marital statuses, economic backgrounds and had different numbers of children were selected for the study. See Table I below for profiles of the participants. All the mothers agreed to the use of pseudonyms during the study. On the initial appointment, the researcher explained the details of the study to mothers. This allowed the mother to make an informed choice as to whether or not to consent to participate in the study.

All the selected and contacted mothers agreed to take part in the study. For each mother, three indepth interviews of 60-90 minute durations were used to capture the experiences and the feelings of mothers from their point of view. Interviews were guided by open-ended questions, linked to the study objectives and information gained from the literature review.

Mothers were given the option to be interviewed in the language they were most comfortable in. All mothers chose their home language Shona (the researcher is also fluent in this language). The first interview transcripts were translated to English and analysed before the second interviews took place. This allowed the researcher to gradually uncover provisional themes within the phenomenological interviews and guided the direction of the second interviews\(^9\). The third interview sessions were carried out for member checking purposes to increase credibility, clarification and further explanations from participants\(^7\). During the process of member checking, a combination of analysed data from both interview one and two were brought to the mothers, who in turn reviewed it to ensure that the findings were a true reflection of what they had said. All the mothers agreed that the data were true reflections of their caring experience of coping with a HIV positive NDD child.

Data analysis was guided by a simplified version of the Stevick-ColaiuzzKeen method detailed in Creswell\(^18\) and Moustakas\(^19\). Steps for data analysis included bracketing, familiarisation, horizontalisation, textual description, structural description and the overall essence of the caring experience. The data analysis procedure followed an iterative process whereby the researcher moved between the different steps of the analysis to eventually describe the mothers’ experiences of caring for their HIV positive child with NDD.

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<table>
<thead>
<tr>
<th>Mothers</th>
<th>2 (Tinashe)</th>
<th>3 (Memory)</th>
<th>4 (Nomsa)</th>
<th>5 (Kudzi)</th>
<th>6 (Zivai)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age of mother</td>
<td>35</td>
<td>33</td>
<td>40</td>
<td>25</td>
<td>30</td>
</tr>
<tr>
<td>Marital status</td>
<td>Married</td>
<td>Married</td>
<td>Married</td>
<td>Single (separated)</td>
<td>Married</td>
</tr>
<tr>
<td>Level of education</td>
<td>Form 4</td>
<td>Form 2</td>
<td>Grade 7</td>
<td>Form 4</td>
<td>Form 4</td>
</tr>
<tr>
<td>Occupation/ employment/ work</td>
<td>Not working</td>
<td>Not working</td>
<td>Not working</td>
<td>Buying and selling of goods</td>
<td>Buying and selling of meat</td>
</tr>
<tr>
<td>Economic status</td>
<td>Good (husband is an engineer)</td>
<td>Poor (both parents do not work)</td>
<td>Average (husband is a welder, his job is not wellpaying)</td>
<td>Poor (vending business is not generating enough income for her)</td>
<td>Average (husband works in a supermarket)</td>
</tr>
<tr>
<td>Number of children</td>
<td>3</td>
<td>2</td>
<td>2</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Ages of the children</td>
<td>9; 5; 3</td>
<td>6; 4</td>
<td>16; 2</td>
<td>2</td>
<td>5; 3</td>
</tr>
<tr>
<td>HIV Status of the children</td>
<td>First two are negative and only last born is positive</td>
<td>Both are positive</td>
<td>First born is negative and second is positive</td>
<td>Child is positive</td>
<td>First child is negative and the second is positive</td>
</tr>
</tbody>
</table>

**Table I: Profiles of the participants**

**Children with NDD: type of disability**

<table>
<thead>
<tr>
<th>Disability</th>
<th>Delayed developmental milestones</th>
<th>Epileptic/left hemiparesis</th>
<th>Cerebral Palsy</th>
<th>Cerebral Palsy</th>
<th>Delayed developmental milestones</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>3</td>
<td>4</td>
<td>2</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Gender</td>
<td>Male</td>
<td>Male</td>
<td>Female</td>
<td>Female</td>
<td>Male</td>
</tr>
</tbody>
</table>

\(^{**}\) Zimbabwe is a landlocked country which has recently emerged from some marked political and socioeconomic challenges. Zimbabwe is in southern Africa; Zambia and Botswana lie to the north, Namibia to the west, South Africa to the south and Mozambique to the east. Its population is 11.4 million. The capital city is Harare, which has a population of 1.6 million\(^12\).
collection commenced. In Zimbabwe, where the data were collected, the Medical Research Council also reviewed the study and approval was granted (MRCZ/B/501). The permission to carry out the study at Harare Central Hospital (CRU) was sought from the hospital administrative offices. Records of children with HIV-related neurodevelopmental disorders were sourced from the Harare Central Hospital Children's Rehabilitation Unit’s administration office.

**FINDINGS**

Two major themes emerged: Resilient Acceptance ‘Ndozvazviri’ and “Rekindled hope for the future”. (See Table 2 below for the themes and subthemes). These two themes will be presented below drawing on the sub-themes to support the description of each theme.

<table>
<thead>
<tr>
<th>Theme</th>
<th>Subthemes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Resilient Acceptance ‘Ndozvazviri’</td>
<td>Nothing I can change; you just have to accept it. God’s case: no appeal. Meeting cultural expectations.</td>
</tr>
<tr>
<td>Rekindled hope for the future</td>
<td>Excited by progress. Privilege of caring.</td>
</tr>
</tbody>
</table>

**Theme One: ‘Ndozvazviri’ (Resilient acceptance)**

The first theme ‘Ndozvazviri’ describes how the mothers were consistent in adapting to the multiple and significant sources of stress they encountered during the process of firstly finding out their own positive HIV status and then the HIV positive status of the child; followed soon thereafter by a growing realisation that their child was not developing typically because of an HIV-related NDD, and then also facing the prospect of the long term dependence of their child on them.

The mothers were devastated and described in detail their traumatic experiences of receiving the news of their own HIV positive status and that of their children,

I was so devastated...I was just a worrying mother. (Zivai)

I was so devastated; such issues are worrisome in life. (Kudzi)

The sub-theme Nothing I can change; you just have to accept it, however captures their commitment to accepting the responsibility they held for the care of their child.

There is nothing I can change! His pregnancy came without being planned. I thought I was gaining weight until my husband advised me to go to the clinic. I went there and the reports from scan showed that I had a three months old pregnancy. I walked away from that place and I said to myself whatever…. (Tinashe)

The word ‘whatever’ here is a close translation of the Shona phrase ‘chero zvazvaita’ which portrays a sense of despair and confusion. It describes the essence of Tinashe’s helplessness, of not knowing how to handle this situation, or what to do.

In all the cases, the whole experience of having a child with HIV-related NDD was never imagined or thought of.

My son just woke up one day with a hot temperature. I did not understand what it was. I took him to hospital and the doctors did their stuff assuming that it was meningitis and what they were saying I do not know much about it. I don’t know much about that. So I stayed with him in hospital then was later discharged after 3 weeks. He could not sit, he could not walk or do anything and he had fits, he is now epileptic. And to think of it that he is also HIV positive… I just didn’t know what else to do. (Memory)

The doublebarrelled condition of the child being HIV positive, as well having a NDD, resulted in the mothers experiencing mothering as a role consisting of all-consuming, difficult and demanding tasks. The mothers spoke of how the caring role for the child with NDDs was different from that of their other children because of their increased levels of dependence on their mothers for a wider array of activities than the other siblings.

Caring for him is different from other children, it is much more demanding because my child cannot do most of the things alone like other four year olds whom you can send to go and fetch water and they go. (Tinashe)

She (his sister) could go to the toilet but he (the child) cannot. He needs me all the time. (Zivai)

The mothers however demonstrated resilience in the face of these multiple negative insults. They described how they needed to come to terms with the fact that their child had HIV/AIDS. This was a situation that was not going to change, and thus it was best to accept their situations and move on with life. Their resilience, reflected in their ability to accept, adapt and respond to these many demands is evident in the quotes below:

I did not panic, because already I knew my child was HIV positive and anything could happen. I just told myself that this is the way it just supposed to be (Zvazviri ndozvazviri). (Memory)

It was difficult for me to accept it. What else can you do? But you just have to accept it. (Nomsa)

There is nothing I can change…ndozvazviri…just have to accept it. (Zivai)

The term ‘ndozvazviri’ is widely used in the Shona language. It signifies a very difficult situation that one encounters and has no power to change. In response, the individual has to accept the situation as it is, as nothing can be done to change it.

All the mothers who participated in this study were Christian. Their belief in God contributed to their acceptance of the condition of the child and their caring role. This is represented in the sub-theme God’s case, no appeal. Mothers believed that whatever God would have done was final and no one had the power to change it. Having a child with HIV-related NDD was in accordance with God’s plan. The mothers thus felt this was reason enough to accept their situation as is.

Nomsa who, after spending 12 years trying to have a baby, eventually gave birth to a child who developed an HIV-related neurodevelopmental disorder. She believed that this was God’s will:

I used to cry that why was I given one child, should I stay with only one. If only God was to see and bless me with another child and it happened … Aaah …am happy with that, I have to be, God gave, to whom could it have been done to.

Most of the mothers also held the belief that a child is a blessing from God. Because the mothers have been blessed with a child, no matter his/her condition, they could not complain or reject a blessing from God. Mothers drew their strength and energy from spirituality. In some difficult situations mothers narrated how they would ask for God’s intervention; which no man could change or go against. The personal connection with a superior being brought comfort, hope and strength to continue with the occupation of caring for a child with HIV-related NDD.

She (her child) got admitted to hospital and doctor said an operation has to be done because her intestines are not functioning very well. I prayed to God. I was praying every day and God heard me. (Tinashe)

Although the mothers had a high regard for the power of their God, some mothers highlighted that they did not have trust in the church. Mothers felt that disclosing their status and that of their children would make them vulnerable to stigma from family, friends and the community.

The church is just the same as in the location (suburb), like at church it depends if pastor’s wife can keep a secret because you can tell her and she discloses to people. (Kudzi)
Mothers who participated in this study shared the Shona culture, a culture which gives the responsibility of caring for others, especially children, to the mothers.

The sub-theme Meeting cultural expectations captures how cultural expectations contributed to their committed approach to caring for their child.

“It is not easy but I am the mother. I have to do it.” (Nomsa)

The experience of getting to know about their HIV status as well as the NDD condition of the child was difficult for the mothers and there was a feeling of resigned acceptance ‘ndozvazviri’ (That is the way it is). The belief in God as a spiritual being and the anchoring foundation of their culture played a role in guiding mothers to this resigned acceptance.

Theme Two: ‘Rekindled hope for the future’

The second theme of ‘Rekindled hope for a future’, unpacks the positive and gratifying aspects of the mothers’ experiences of their caregiving occupation. These included instances where mothers, while caring for and playing with their children, noted improvements in the developmental milestones and overall health of the child. Mothers expressed that taking care of their children was something very important and meaningful to them and they would want to continue doing it.

The sub-theme Privilege of caring reflects the mothers’ positive attitudes towards their caregiving role despite this being a role associated with many challenges.

What I value most is caring for my child as well as planning for his future. Like any other child, you would want to be able to take good care of him, see him grow, go to school and have a better life. (Ziva)

Because it is just so difficult to take care of such a child in such a condition and being able (to do so) is so important to me. (Nomsa)

Mothers valued the times and opportunities for engagement and participation in play of the child. It was important for the mothers to set time aside to play with their child. Mothers eagerly shared their experiences of engaging in play with their child. Two of the mothers reported that they felt happy to watch the child play safely with siblings or other children in the community. Watching their children being included in play with other children brought a sense of happiness and joy to the mothers. This could be linked to the sense of belonging the mothers experienced because their child was doing what other children in the community were doing, despite the disability.

I was happy to watch my child play with others. He can now safely play with others. You see that he has a lot of potential and it is beginning to show. (Memory)

Excited by progress

The privilege to care for their children allowed them to note the progress of the child and the positive changes happening in the child, which brought a sense of joy, hope and relief to the mothers. This applied even more when the mothers had experienced situations where they had been told their child would not be able to do anything in life or that the child’s life expectancy would not be long. As the child showed progress, emotions of grief, sorrow and despair were complemented with hope for the future.

I am just happy with the improvements I am noticing. Yes I feel so happy because you know if you are in closed doors and you hear doctors saying this and that and on top of that, he (the child) got tested and found to be HIV positive. (I) … would say now he is an invalid cabbage. he won’t do anything in life. So now with the improvements I feel so happy. (Memory)

Some of the mothers had heard of various myths and beliefs that a child with HIV and disability had a short life expectancy. These beliefs worried the mothers and made them uncertain of the future of their children. In one of the interviews, Nomsa mentioned that she had been told that children infected with HIV die before the age of seven years, but when she had come to the CRU and saw other children infected with HIV who were older than seven years, she was relieved. She felt better as she learnt of the progress other children with HIV, who were achieving their developmental milestones and maintaining good general health. Improvements in the general health of the child rekindled hope in the mothers. Progress of the child was described in terms of general health, increase in body weight and attainment of developmental milestones.

These days I am celebrating the fact that my child is able to say a word mama…huya (mom… come) what an improvement! (Memory)

I am excited! At first he could not do anything but now he is able to do this (bringing hands together) so now he can hold things. (Nomsa)

Their child’s improved ability to initiate and do activities alone meant a lot to the mothers especially when a child had not been able to do anything before. Noticing the child achieving a new ‘doing’ – no matter how big or small, good or bad – these mothers cherished every ‘doing’ of their child with happiness. These small tasks included the child bringing two hands together, eating alone, playing, saying a word or even choosing what they wanted to wear. Also, engagement and participation of the child in play was a key factor that brought relief and excitement to the mothers.

This could have been attributed to the fact that, engaging in play, for both children and adults, is identified in the Shona culture as a significant measure of health and wellbeing in children. If a sick child was still able to engage in play, then nature of illness was not viewed as serious. Shona nurses also use play as a yardstick to determine the severity of illness and often nurses ask mothers if the child is playing when they take case histories. Engagement in play for a child with NDD signified a sense of health and well-being, thus, mothers were excited when their children engaged in play.

What excites me is that my child can play, he enjoys playing even with other children. (Ziva)

Despite the negative moments described by the mothers about caring for a child with HIV-related NDDs, they also reported experiencing some positive moments. The positive experiences mainly occurred when mother noticed progress in the child as well as when playing with the child. Progress in the child rekindled hope in the mother and it brought some sense of joy and excitement. Also, instances when the child was able to engage in play and do as other children would be doing add to the positive experience of the mothers.

DISCUSSION

Caregiving occupations presented challenges and opportunities which mothers used to create and transform their lives, pursue aims, overcome barriers, learn new ways of achieving health and happiness and also discover their purpose in life20. Wilcock21 supports the notion that people engage in occupations that are meaningful. However, there can be situations where mothers engaged in the caring occupation, not because it was meaningful to them, but where the occupation has been forced by circumstance. In this study some mothers reported that the pregnancy was not planned and to those who had planned to have a child, no one hoped for a child with HIV-related NDD. Despite these challenges mothers committed to their role as caregivers and found meaning through engagement in this role over time.

What people do is shaped by the context and the external structures of society that shape the individual’s aspirations, expectations and lifestyle22. This emphasises the need to consider not only the individual but also the impact of the environment when striving to understand the occupational engagement of people23. The caring experience of the mothers could not be described outside of the influence of their contexts. The mothers described the trauma linked to the process of discovering their own HIV status, the HIV status of their child and then the child’s diagnosis of NDD. Despite the impact of various environmental factors (discussed below) on
their ability to deal with this trauma, it also supported their ability to meet challenges of their care giving role.

Culture is an occupational determinant which shapes what people do and how they experience their engagement in meaningful occupations. In this study the mothers’ experiences of caring for their children were guided by the sociocultural norms of the Shona culture, a culture that dictates the implicit rules and expectations of being a good mother. The Shona culture gives the responsibility of caring for children, to mothers. Mothers are expected to be the custodians of children and answerable to health issues of the child. The mothers in this study strongly aligned themselves with the expectations of the Shona culture and took on the responsibility of caring for their child with HIV-related NDD with diligence. Their motivation to care was fuelled by their cultural obligations.

Implications for practice

The study highlighted the transactional occupation of caring for a child with HIV-related NDD as shaped by both personal and contextual factors. Thus, occupational therapy is encouraged to enlarge its scope of practice from focusing on the child with impairment, to include the mother as the primary caregiver who is engaged in the occupation of caring for the child, in relation to both their context and their own personal attributes. There is a need for assessment procedures and interventions that take cognisance of the interplay of personal factors and context for mother and child. Also, the various ways in which these mothers orchestrate their daily occupations to accommodate the care of their child as well as the value they place on the care of their child, warrants further investigation.

CONCLUSION

The study reveals the transactions that happen between the mother and the context (environment) in which the occupation of caring for a child with NDD takes place. The role of the context - which included culture and spirituality - in shaping the experiences of the mothers, should not be underestimated and it would be unjust to look at the mother as an individual outside her context. The findings of this study have also highlighted the interplay and interdependence between the contextual and the personal factors that shape the experience of the occupation of caring for a child with HIV-related NDD. Thus, the findings of this study should encourage occupational therapists to carefully consider the transactional nature of occupation and how the mothering role positively shapes these mothers’ identities, when they design interventions for this population group.

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**AUTHORS’ CONTRIBUTIONS**

Munambah and Gretschel developed the concept and design of the study. Munambah submitted the proposal draft for ethical approval and conducted the data collection. Munambah and Gretschel conducted the data analysis and drafted the manuscript. Sonday played on oversight role during data collection and analysis. All the authors assisted in revising the manuscript for submission to a journal. All authors read and approved the final manuscript and Munambah was the corresponding author.

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