Assessing the Relationship between Caregivers Burden and Availability of Support for Family Caregivers’ of HIV/AIDS Patients in Calabar, South East Nigeria

Ekaete Francis Asuquo¹*, Josephine B. Etowa², Prisca Adejumo³

¹Department of Nursing Science, University of Calabar, Calabar, Nigeria; ²School of Nursing, Faculty of Health Sciences, University of Ottawa, Ottawa, Canada; ³Department of Nursing Science, University of Ibadan, Ibadan, Nigeria.

Email: *ekaetefasuquo@yahoo.com

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ABSTRACT

Purpose: This study examined the level of burden and the extent of support on family caregivers of people living with AIDS (PLWHA) in Calabar, South East Nigeria. Methods: A mixed method with cross sectional approach was used. Purposive sampling technique guided the recruitment process and data collection methods included, semi-structured questionnaires and focusing group discussion. 260 respondents participated in the study. The quantitative data were mined with the aid of SPSS and the qualitative data were analysed with the aid of NVivo8 using thematic analysis. Results: Results indicated high level of burden with limited support to caregivers. A Chi-square value of 25.1 was obtained at P < 0.05, suggesting a significant relationship between availability of support and caregivers burden. This relationship was supported by the themes of physical, social, emotional and financial burden for the caregivers. Similarly, information on coping skills, emotional support, financial assistance and help with caregiving themes emerged for social support. Conclusion: In Nigeria, the burden of caring for HIV/AIDS patients has a remarkable impact on family caregivers. This calls for the development of policies that can systematically address the needs of family caregivers in order to ameliorate the negative consequences of caregiving for PLWHA.

Keywords: Family Caregivers; Caregivers’ Burden; Caregivers’ Support; People Living with HIV/AIDS (PLWHA); Nigeria

1. Introduction

Providing care for a sick family member is a normative role embedded in African culture. It is an age-long tradition of kindness, love and loyalty, which binds family members together [1,2]. Family members therefore become the principal source of informal long-term care to an ill or disabled family member [3,4]. Culturally in Africa, the criteria for selecting primary caregivers are linked to sex, generation and geography [4] and near kinsman or close relation from the nuclear family. These roles are assumed without questioning and may be laden with adverse consequences [5-7]. HIV epidemic has become a major public health challenge of our times and its impact engulfs all sectors of our life including homes [8-11]. Global effort has been directed to tackle this pandemic and to reduce its prevalence in which it is no longer a public health issue. While this effort has led to advancement in medical treatment and improvement in life expectancy, increasing numbers of people are living with chronic illness and disabilities. And this has a detrimental effect on family members who assume caregiving role and bore the brunt of taking care of those with chronic conditions including the HIV epidemic. This is particularly problematic in sub-Saharan Africa where they are 22 million cases (69%) [12]. A synergistic relationship exists between high prevalence of HIV and the demand for health care services and also has a burden on both formal and informal caregivers [13]. Hence HIV’s burden transcends the health care system to families and friends who are without appropriate sensitization and educational preparation to take up the caregiving role to meet the health need of their family members [5,6,14-16].
This is a significant contribution by the informal caregivers who form the backbone of health care system round the world [17,18]. Most often, they remain hidden and their contribution to care receivers is unrecognized. Studies revealed that caregivers experience negative physiological, psychological, behavioral as well as financial consequences of caregiving [5,6].

**Burden of Care**

The Nigerian health care system is based on the implicit assumption that, family members are available to provide care in the homes to the dependent ill, the elderly, physically and mentally disabled member. Previous report indicates that caregivers are increasingly being required to perform complex tasks similar to those carried out by paid health or social service providers [3]. Therefore the extent of care provided by these care givers covers minimal assistance with activities of daily living to complete care of a debilitated patient. These care are complicated with the chronic pattern of exacerbation and remission associated with HIV/AIDS, which depends on the caregivers appraisal of the experience that may produce high level of burden [15,19,20]. Caregiving stress model identified the stress process including the background, stressors, mediators and outcome [21]. Agreeing with these assertions, other authors [22,23] declared that factors which influence burden include the characteristics of caregiving context such as duration of care, extent of social support and quality of family relationship, characteristic of the caregiver and characteristic of the care receiver. This perceived stress was identified as caregiver’s burden [24] in reference to inconvenience associated with caring for mentally ill relatives and later conceptualized as the degree to which the emotional, physical health, social life, and financial status of a caregiver has been impaired as a result of caregiving [25]. This conceptualization is used in this study as the degree to which life’s tension is expressed to depict the physical, emotional, financial and social challenges associated with caring for PLWHA.

The high level of burden experienced by the caregivers may notably have produced negative consequences which are manifested as symptoms of anxiety and depression and induced changes that could lead to heart disease, hypertension, psychological worries, loneliness, increasing rate of physiological illness and suppressed immune responses. Others include reduced engagement in preventive health behaviors, social problems, etc. [4,5,15,16,26,27]. It is important to note that the task of caregiving is not only stressful to the family care giver, and it has a serious impact on the patient quality of life. A correlation has been found existing between the level of burden and quality of life of the care receiver [5,7].

**2. Support to Caregivers**

The pattern of HIV occurrence has been classified into four phases namely onset, course, outcome and degree of incapacitation [28]. Utilizing this typology, HIV has a gradual onset with a long term stress on the patient, the course is progressive with symptoms which increases in severity. As disabilities become progressive with less periods of relief, the patients and caregivers are rendered psychologically, socially and physically disabled [28,29]. The level of burden experienced by caregivers differ depending on degree of incapacitation of the recipient [28,30].

The need to support caregivers of PLWHA for continual adaptation to task changes become imperative since the perceived goal is to reduce burden of care on caregivers. The conceptualization of social support is viewed as coping assistance or the active involvement of significant others in the caregivers ability to manage stress [31]. This is the concept utilized in this study. The significant others may be family, community members or friends, and occasionally professional institutions’ help may be required from time to time [28]. The help rendered by significant others enhances coping ability as it satisfied the need for attachment, relieves stress, and boosts a sense of self-worth, trust and life-direction [32-34]. The burden associated with caring for PLWHA differs from caring for someone with cancer or other debilitating diseases, since HIV involves social isolation, contagion and stigmatization, hence caregivers are emotionally laden with high level of burden. Adequate support is needed to encourage and prepare caregivers to express hurt, anger and fear before taking up caregiving role. Burden of care can be ameliorated by factors such as support from other family members, the ability to use problem-focused coping strategies and availability of support from the community [32]. Other factors include the provision of information and advice, participation in social activities and help from caregiving and emotional support [35-37]. Provision of information for coping with stress and dealing with emotions are important areas where caregivers may require help [33,38,39]. On the other hand burden can be worsened when the people involved are socially isolated, less knowledgeable about the condition or have limited interpersonal skills. This study therefore assesses the burden of caregivers, availability of support and proffers the appropriate intervention measures as a panacea for delivering efficient quality care to the patients and wellbeing of caregivers.

**3. Materials and Methods**

**Location of the Study**

The study was carried out in Calabar Municipality of
Cross River State of Nigeria. Calabar is located at the extreme end of southern eastern Nigeria between latitude 4°58′ North and longitude 8°17′ East. Calabar Municipality host the University of Calabar Teaching Hospitals (a tertiary institution) which acts as referral center to primary and secondary health care facilities in the state. It has been selected by the Federal Government of Nigeria as one of the centres for the management of people living with HIV/AIDS. The study was informed by the high HIV prevalence of 10.4% [40] at the study site.

4. Methods

Study participants constituted a convenient sample of 260 caregivers who were primary caregivers to PLWHA between June and December 2009. Participants were identified through HIV clinic of the University of Calabar Teaching Hospital and visitation list of a voluntary caregiver of PLWHA, an NGO (Positive Development Foundation, Calabar, Nigeria). Inclusive criteria were: being a caregiver for more than one month and also caring for PLWHA patient with some functional abilities as a family member. A total of 299 potential caregivers met the inclusive criteria of being primary caregivers, but only 260 completely filled their questionnaires. 11 caregivers did not know the diagnosis of ailment of their care receivers. Twelve (12) refused to participate while 16 did not complete their questionnaires. Two sessions of focus group discussions were also used to encourage self-disclosure among the participants [41] and to complement findings from the questionnaires. Being a primary caregiver involved participants who provided unpaid physical support such as helping in activities of daily living, shopping, food preparation, helping in administering medication, overseeing medical appointment, financial, and emotional support to PLWHA. The study was submitted to the University of Calabar Teaching Hospital Ethics and Research Committee, who gave the approval for conducting the study. Consent was also obtained from the Managing Director of Positive Development Foundation. Informed consent was also obtained from study participants following full description of the aims and objectives of study.

Caregivers Burden. The Zarit Burden Interview scale questionnaire, semi structured questionnaire on social support and focus group discussion were the main instrument for data collection. Zarit Burden Interview (ZBI) scale made up of 22 items was used to measure the level of burden. This scale reflects the degree of emotional, physical health and social impact of caregiving on caregivers while caring for PLWHA [42]. The respondents indicated the discomfort they experienced of particular items on a 5-point Likert scale ranging from 0 - 4. The total score ranged from 0 to 88 and a high score correlated with higher level of burden. The psychometric properties of Zarit Burden Interview scale had been examined in many studies with an estimated internal consistency reliability of Chronbach’s alpha ranges of 0.88 to 0.92 [43]. The social support questionnaire consisted of 12 items on a 4-point Likert scale assessing the availability of support on Information, financial help, help with caregiving and emotional support available to caregivers. The questionnaire was translated into Efik (the native language of the people) and back-translated into English by experts to ensure that there was no lost of meaning. The Efik version was administered to caregivers with little or no education. The NGO (Positive Development Foundation) provided the forum for three focus group discussions with about 8 to 10 participants at each session. A total of 27 caregivers participated in the focus group discussion after having completed the ZBI and social support questionnaire. Focus group discussion as an assessment tool involves an organised discussion with a selected group of individuals to gain information about their views and experiences on a topic [44]. It has the advantage of creating greater room for spontaneity and participants feel supported and empowered within the acuity of the group and its cohesiveness [45].

Data Analysis

Data collected were analyzed using the Statistical Package for the Social Sciences (SPSS 16.0) software to generate the frequency and percentages of the values. By the application of inferential statistics (Chi-Square test), the level of significance was also determined at the P-value of 0.05. For level of burden, scores ranged from 0 to 88. Zero to twenty (0 - 20) represented little or no burden, 21 - 40, mild to moderate burden; 41 - 60, moderate to severe burden and 61 - 88, severe burden [46]. Thematic analysis was used for assessing the focus group discussion sessions [47]. Sixty five (65) codes emerged which were developed into 4 basic themes namely physical, social, emotional and financial burden, which provided impetus for understanding the level of burden experienced by the caregivers. Four basic themes namely providing information on practical and coping skills, financial assistance, help with caregiving and emotional support emerged as caregivers support needs.

5. Results

In this study, the demographic variables and level of caregivers burden are presented as frequency, percentages and ranges. Table 1 shows the socio-demographic characteristics of the respondents. The majority of the respondents were females 199 (76.5%) while 61 (23.5%) respondents were males who voluntarily took up the care...
Table 1. The Socio-demographic data of respondents (N = 260) in Calabar Municipality, Nigeria.

<table>
<thead>
<tr>
<th>Variables</th>
<th>Frequency</th>
<th>Percentage (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>61</td>
<td>23.5</td>
</tr>
<tr>
<td>Female</td>
<td>199</td>
<td>76.5</td>
</tr>
<tr>
<td>Age (yrs)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10 - 20</td>
<td>22</td>
<td>8.5</td>
</tr>
<tr>
<td>21 - 30</td>
<td>30</td>
<td>11.5</td>
</tr>
<tr>
<td>31 - 40</td>
<td>51</td>
<td>19.6</td>
</tr>
<tr>
<td>41 - 50</td>
<td>90</td>
<td>34.6</td>
</tr>
<tr>
<td>51 and above</td>
<td>67</td>
<td>25.8</td>
</tr>
<tr>
<td>Educational Qualification</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Primary</td>
<td>15</td>
<td>5.8</td>
</tr>
<tr>
<td>Secondary</td>
<td>112</td>
<td>43.1</td>
</tr>
<tr>
<td>Tertiary</td>
<td>111</td>
<td>42.7</td>
</tr>
<tr>
<td>Never been to school</td>
<td>22</td>
<td>8.5</td>
</tr>
<tr>
<td>Household Income Level US$*/Day</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Less than $6.00</td>
<td>130</td>
<td>50.0</td>
</tr>
<tr>
<td>$7.00 - 8.00</td>
<td>59</td>
<td>22.7</td>
</tr>
<tr>
<td>$9.00 - 10.00</td>
<td>25</td>
<td>9.6</td>
</tr>
<tr>
<td>$11.00 - 12.00</td>
<td>18</td>
<td>6.9</td>
</tr>
<tr>
<td>$13.00 - 14.00</td>
<td>13</td>
<td>5.0</td>
</tr>
<tr>
<td>$14.00 and above</td>
<td>15</td>
<td>5.8</td>
</tr>
<tr>
<td>Number of People in the Household</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 - 3</td>
<td>19</td>
<td>7.3</td>
</tr>
<tr>
<td>3 - 6</td>
<td>153</td>
<td>58.9</td>
</tr>
<tr>
<td>7 and above</td>
<td>88</td>
<td>33.8</td>
</tr>
<tr>
<td>Duration (Hrs/day)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 - 8 Hrs/day</td>
<td>52</td>
<td>20</td>
</tr>
<tr>
<td>9 - 12 Hrs/day</td>
<td>162</td>
<td>62.3</td>
</tr>
<tr>
<td>13 - 24 Hrs/day</td>
<td>46</td>
<td>17.7</td>
</tr>
<tr>
<td>Duration (Yrs)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Less than 1 year</td>
<td>101</td>
<td>38.9</td>
</tr>
<tr>
<td>Between 1 - 2 years</td>
<td>109</td>
<td>41.9</td>
</tr>
<tr>
<td>Between 2 - 3 years</td>
<td>38</td>
<td>14.6</td>
</tr>
<tr>
<td>Above 3 years</td>
<td>12</td>
<td>4.6</td>
</tr>
</tbody>
</table>

6. Level of Burden on Caregiving

Generally, the results revealed a level of burden ranging from 14 - 71 on ZBI Scale (Table 2). 49 (18.8%) experienced no burden while providing care, 67 (25.8%) respondents experienced mild to moderate level of burden, 93 (35.8%) experienced moderate to severe level of burden while 51 (19.6%) respondents experienced severe level of burden in providing care to PLWHA.

Focus Group: The extent of burden expressed following discharge from health care institution was categorized into four basic themes namely physical, social, emotional and financial burden. Carers generally perceived caring for PLWHA to be very stressful as they reported that their tasks ranged from simple to complex and covering all the “activities of daily living” such as putting care recipient to bed and getting up, bathing, dressing, toileting and coping with incontinence worsened during period of exacerbation, preparing food, giving medications, providing financial assistance, and emotional support like keeping company especially when care recipient is depressed. These activities left them with aches and pains, and fatigue and exhaustion were regular experiences in their life.

Physical Burden: Caregivers generally express that they experience extreme fatigue as they indulge in day to day caregiving. Caregiving to them is very challenging and produces ill effect on their physical health. The physical ill effect of caregiving as perceived by the caregivers include fatigue and exhaustion “you keep doing one task to another and no rest period, even some nights...”

Table 2. The distribution of caregivers burden on Zarit Burden Interview scale (ZBI).

<table>
<thead>
<tr>
<th>Level of Burdens</th>
<th>n = 210</th>
<th>%</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Burden</td>
<td>49</td>
<td>18.8</td>
<td>0 - 21</td>
</tr>
<tr>
<td>Mild to Moderate Burden</td>
<td>67</td>
<td>25.8</td>
<td>21 - 40</td>
</tr>
<tr>
<td>Moderate to Severe</td>
<td>93</td>
<td>35.8</td>
<td>41 - 60</td>
</tr>
<tr>
<td>Severe Burden</td>
<td>51</td>
<td>19.6</td>
<td>61 - 88</td>
</tr>
<tr>
<td>Total</td>
<td>260</td>
<td>100</td>
<td>0 - 88</td>
</tr>
</tbody>
</table>

\[^1\text{US$} = \text{N 164.00 (Nigerian Naira).}\]
become day because it is pack filled with activities”. Some expressed the feeling of being trap to responsibility, “I just can’t ignore him as other members of our household do, it becomes my responsibility and mine alone as the wife”. The stress experienced enhances discouragement and helplessness. Reports of physical ailments such as continuous waist pain, “I have been lifting him for bathing, but as the waist pain persist if no help is available I bath him in bed”. Other physical effects include, weight loss, headaches and many lack the energy to face another day. With reference to their perceived health status, the discussants generally agreed that their health had suffered as noted “I used to be plum, but since I started caring for her, I’ve lost weight and I don’t have sufficient time to care for myself, and some friends ask me, are you sick? I really know that this work has affected me.” Another reported that she is now diagnosed of having hypertension which she attributed to excessive worries due to caregiving. Most discussants revealed that burden of care increases during period of exacerbation. One said “I suffer most when he has diarrhea, the wrappers (native African wear around the waist down to the knee) are not enough, you keep washing and reusing, he is weak and frail, I am caring for him and many other things I have to do to make ends meet. At the end of the day I feel drained, tired and worn out”. Some narrated that burden also increased when the care receiver exhibited some abnormal behavioral symptoms, a situation which demanded continuous supervision coupled with providing care. “I can cope with bathing and feeding or anything else but when he starts talking to air, becoming very suspicious of everybody and acting abnormal, I feared for my life and that of others in the house”.

Emotional Burden: During the FGD sessions, participants raised some issues which impacted them emotionally such as being unduly irritable, fear of the unknown, guilt and fear of being susceptible to HIV through caregiving. Most of the discussants revealed that the burden has influence their emotional wellbeing as they become unduly irritable and used aggressive words when provoked, which were later regretted. This was portrayed by the words of a female caregiver who said “I slapped my child who insisted that I should be at the school’s parent teachers association, there was no reason for that for God sake no reason. I felt disappointed by my action and I started crying”. Another reported, “I woke up some days feeling depressed and afraid of what the day holds, and on such days, tears flow freely from my eyes”. Another reported, “sometimes I get annoyed and upset when others are laughing, and I often wondered what is good about life that thrills their heart”. Fear of the unknown has been a serious trait to caregivers, as most of them harbor anticipatory grief about the imminent death of their sick relative especially knowing that HIV has no cure. This fear is common especially when the care receiver is a bread winner. A woman said “when my husband is dead (sick relative) and the pension stop coming in, what will become of the family”. Some felt already susceptible to HIV through caregiving and some felt stigmatized as HIV positive. Caregivers were generally committed to the care of their relatives with some strong emotional ties which would produce guilt if they should leave the caregiving role, “We’ve been married for 30 years although the last eight years was nothing to write home about, but he is my husband anyhow” another added “he is my only son I will always love and care for him”.

Social Burden: Social groups provide companionship, support and a feeling of togetherness which has a positive effect on caregivers’ health and well-being. Some discussants feel isolated from their social group due to lack of time for such functions or due to poor self-esteem. Some reported reduced time spent on self, spouse and children. A discussant reported feeling uncomfortable especially when they had a known visitor. “I just felt they may be thinking I am also HIV positive”. Another retorted, “It’s like falling from grace to grass, and your self esteem is gone”. Another remarked “My boy friend called it quit when he found out I was caring for my brother who had HIV”. Some wished they could leave the caregiving role to someone else, one caregiver complained “but who do you leave it to”.

Financial Burden: The financial burden of caregiving is a significant aspect that worsens caregiving experience especially in low and middle income countries such as study setting where most individuals live below one dollar per day [47]. Most carers revealed that this aspect is an issue as it influences the administration of drugs and other activities such as school fees for children and general family upkeep. Sometimes children drop out of school taking to hawking as a way out. Even with those who chose hawking as a remedy, discrimination and stigmatization takes its toll on their ability to sell their farm products where they are known. A discussant narrated “my children started hawking to make ends meet because I couldn’t pay for their secondary education any more”. Another said, “I had to hawk my farm products away from home because neighbors won’t buy from us”. Another discussant reported, “We sometimes suspend giving drugs until when food is available, since he cannot take drugs with empty stomach. Although we are aware that this is not right but what can we do when there is no money to buy the food”. Generally, they agreed that lack of money was a serious constraint to caregiving although some agreed to some level of financial support from Church and NGOs.
6.1. Availability of Support to Caregivers

The availability of support was categorized into four sections namely: providing information on practical and coping skills, financial assistance, help with caregiving and emotional support (Figure 1). Among the 260 respondents, 152 (58.5%) agreed to have received some form of support while 108 (41.5%) received no form of support at all. The form of support received include: 6 (0.2%) information on practical and coping skills, 36 (13.9%) financial assistance, 72 (27.7%) help with caregiving and 38 (14.6%) emotional support. Most caregivers perceived their role to be extremely stressful with adverse consequences on their health, which some form of support from individuals or government could have helped to ameliorate.

Providing Information on Practical and Coping Skills: Majority of discussants revealed that they had no prior information from hospital staff on how to care for recipients. Their knowledge and skills on care came from trial and error and past experiences based on previous life situations. Many wished they had training to assure themselves they were providing the right care “I wish I was trained, nevertheless I just do all I can for him, if this did not work, I try something else”. The caregivers generally wished they had training to assist them in decision making and problem solving relating to their roles. The fear of stigmatization also prevented some from seeking advice from some support staff. The forum provoked negative feeling when some learnt about the foundation that supported PLWHA and available services. Although the foundation had their own challenges such as number of people that volunteered to provide care for PLWHA, information regarding recipient health conditions and their implications coupled with how to prevent contagion through caregiving was also identified as areas that required attention.

6.2. Financial Assistance

All caregivers unanimously express the need for financial support as recipient disease condition had impinged on the family resources. Even caregivers who had some form of financial support felt it was not enough. A caregiver expressed “not just giving money from time to time, I wish the government can employ one of my children, it will help make life easy for us”. Coping strategies varied among the caregivers which included children dropping out of school to hawking, begging and other subservient means of survival as a way out.

Help with Caregiving: Most care givers expressed that they needed a break occasionally from providing care and a relief period would have cushioned the effects from both physical and mental stress of caregiving. It would also foster social relationship and alleviate feelings of isolation. “I know if I had the time I would be going to church more often and something would have been said to help me. But it seems this work is my sole responsibility, no time for what I want to do for myself”.

Emotional Support: Throughout the focus group sessions, group members expressed anticipatory grief about the sad end of their stressful endeavor. Always probing to know if early death was still inevitable with drugs. Fear of susceptibility to HIV was also raised which compounded caregiving role. The need for counseling services was identified to help in eradicating fear and anxiety associated with caregiving.

The relationship between availability of support and caregivers’ burden was also evaluated. Out of 152 respondents that had received some form of support, 37 experienced no burden, 44 respondents experienced mild to moderate burden, the majority, 56 respondents experienced moderate to severe burden while 15 respondents experienced severe burden (Table 3).

Out of 108 respondents that received no support, 12...
Assessing the Relationship between Caregivers Burden and Availability of Support for Family Caregivers’ of HIV/AIDS Patients in Calabar, South East Nigeria

Table 3. The relationship between availability of support and caregivers’ burden.

<table>
<thead>
<tr>
<th>Availability of Support</th>
<th>No Burden</th>
<th>Mild to Moderate Burden</th>
<th>Moderate to Severe Burden</th>
<th>Severe Burden</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Received any form of support</td>
<td>37 (28.7)</td>
<td>44 (39.2)</td>
<td>56 (54.4)</td>
<td>15 (29.8)</td>
<td>152</td>
</tr>
<tr>
<td>Received no support</td>
<td>12 (20.4)</td>
<td>23 (27.8)</td>
<td>37 (38.6)</td>
<td>36 (21.2)</td>
<td>108</td>
</tr>
<tr>
<td>Total</td>
<td>49</td>
<td>67</td>
<td>93</td>
<td>51</td>
<td>260</td>
</tr>
</tbody>
</table>

Figures in brackets are expected frequencies. $X^2$ Cal. = 25.1, $X^2$ Tab. = 7.81, DF = 3, N = 260, P > 0.05.

7. Discussion

The improvement in technology and the discovery of anti-retroviral drugs (medical innovations) have increased the longevity of PLWHA and turned HIV into a chronic ailment, which has apparently increased the burden of home caregivers. Many caregivers expressed distress about the consequences of caregiving on their general health as revealed in this study. Majority of the caregiver’s experienced moderate to severe level of burden which suggests partial or complete reliance of most care receivers (PLWHA) on the caregivers for their physical, financial, emotional and social support especially during periods of exacerbation of HIV symptoms. This observation is similar to previous report [28] who asserted that HIV has a gradual onset with a long term stress on the patient, the course is progressive with symptoms which increases in severity, and as disabilities become progressive with less periods of relief, it renders the patient and caregivers psychologically, socially and physically disabled. Therefore the level of dependency is related to the functional abilities of the care recipient. The lower the functional abilities the higher the level of burden perceived by the caregiver. This also corroborates with the findings by others in different locations [5-7].

Further study [48] revealed that informal caregivers experienced a lot of stress while caring for recipient and remains a valuable asset to the recipient for their quality of life. Major caregiving problem emanate from carers ability to cope with the increased needs of the care receiver as a result of physical and/or mental illness [49]. Although this finding was contrary to the situation in Canada [50], PLWHA relied more on similarly disadvantaged peers and formal caregivers for support hence there was no increased level of burden on family members. The differences in this study could also be attributed to cultural expectations especially in Africa, where experienced no burden, 23 respondents experienced mild to moderate burden, the majority 37 respondents experienced moderate to severe burden while 36 respondents experienced severe burden. A Chi-square value of 25.1 was obtained at P < 0.05, which showed a significant relationship between availability of support and caregivers burden (Table 3).

The high prevalence of HIV in sub-Saharan African has depleted financial resources of many who cannot afford funds for health care [51]. This study revealed that the majority of the caregivers daily earning was low (Table 1), in conformity with previous observation by WHO [52]. Though this study did not assess the correlation between burden and quality of care, it is important to know that the quality of life of the care recipient depends on the quality of care provided, and the high level of burden has a detrimental effect on the caregiver and a direct effect on quality of care receiver [5,8].

The demographic characteristics of caregivers were also significant in this study as it portrayed the cultural norms of the African society such as women’s role being associated with primary caregiving in the society. Similar finding [26] revealed that within the complex system of long-term care, women’s caregiving is essential in providing a backbone of support to the debilitated patient. In relation to HIV scourge, United Nations study [13] revealed that women are significantly affected as care responsibilities fall to older and married women since a substantial proportion of PLWHA move back to their communities of origin at some stage of the illness to be cared for by family members.

Also significant to the study were the age groups 41 to 50 years that form the majority that participated in caregiving. It would imply that majority of caregivers are wives, mothers, grand-mothers. Previous report [26] indicated that the average caregiver is 46 years old female, married and working outside the home. Majority of caregivers lived in large households with minimal household income, coupled with high financial costs of caregiving. These households face the possibility of moving further into indigence [53].

This study revealed that most caregivers received limited or no support from family members, community or government and they expressed need for support. Caregivers took up the caregiving role without educational preparation. They were in most cases unfamiliar with the type of care they must provide or the extent of care needed. Knowledge on how to assess community re-
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sources was limited. This could be due to failure by the health care workers to assess informal primary caregivers skills before assumption of caregiving role or they were just silent on the continuity of care at home which they were afraid to initiate a follow-up. In most cases, the caregivers are unprepared to provide care, due to inadequate knowledge of health care delivery since they received limited guidance from the formal health care providers [54,55]. It has been suggested that caregivers require some form of training to effectively provide care and that it is unfair to presume that carers have the necessary knowledge and skills to care for an ill relative without some form of support [56-58]. Therefore providing information and training prior to care recipient discharge would ensure caregiver’s preparedness for caring role.

During the study, the need for emotional support was also identified which could be provided through health care providers linking caregivers to existing support system available in the community. Such an assistance could minimize the pressure and burden on carers since reports indicate that caregivers experienced mixture of conflicting emotions such as commitment, frustration, anger, loneliness, exhaustion and depression, guilt, anxiety, love and warmth which could have negative consequences on the caregiver [39,57]. They further revealed that helping the care giver to acknowledge and reconcile with these emotions can enhanced coping with the challenges of being a carer.

Financial assistance was also identified as area caregivers needed support. The scourge of HIV is greater among the low socioeconomic class and tends to sap the meager family resources thereby producing a higher level of burden. Caregiving is time consuming therefore caregivers who provide substantial amounts of care divert their time from other productive chores resulting in financial hardship. This is a common feature in low and middle income countries like Nigeria where many live below one dollar per day. Attempt to balance caregiving with other activities, such as work produced negative consequences and more stress on carer. Report indicated that homes engulfed in poverty and are burdened with financial costs of caregiving may move further into indigence [13]. Such households with a sick family members face the opposing pressures to work fewer hours and spend more time caring.

Assistance with physical care such as in activities of daily living is identified as area of need in this study. It is apparent that most caregivers reach a point when they realized that housekeeping routines and regular errands were accomplished with great difficulty or left undone. It has been suggested that assistance in activities of daily living is also needed to alleviate the burden of care [13] and such help should come from other members of the family, individuals with the same culturally background (community support) or those with similar experience such as HIV support [31].

8. Conclusions and Recommendation

In Nigeria, with 2.98 million people infected with HIV, many people will be needed to function as caregivers even though caregiving is not without consequences. This study revealed that caregivers experienced high level of burden (physical, emotional, social and financial) as a result of the care they rendered to PLWHA. Therefore there is need to relieve this burden by providing necessary support in order to enhance the quality of care of the patients as well as maintain the optimum well-being of the caregiver. There should be training and re-training of caregivers on coping and management strategies before they assume the caregiving role for PLWHA.

The development of policies that can systematically assess the caregivers’ need should be promulgated. Counseling sessions should be organized by government and voluntary organizations regularly to present a forum where caregivers could air their problems and challenges encountered during caregiving services. Government should encourage the formation of HIV community support groups to provide support to caregivers.

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