Effect of Age and Level of Education on Neurocognitive Impairment in HIV Positive Zambian Adults

Norma Kabuba
University of Zambia and Norwegian University of Science and Technology

J. Anitha Menon
University of Zambia

Donald R. Franklin Jr.
University of California, San Diego

Stian Lydersen
Norwegian University of Science and Technology

Robert K. Heaton
University of California, San Diego

Knut A. Hestad
Innlandet Hospital Trust, Hamar, Norway, and Hedmark University of Applied Sciences

Objective: Older age and lower education levels are known to be associated with worse neurocognitive (NC) performance in healthy adults, and individuals with HIV infection may experience accelerated brain/cognition aging. However, higher education may possibly protect against HIV-associated neurocognitive disorders (HAND). The aim of the current cross-sectional study was to assess the effect of age and education in an HIV-1 clade C infected adult population in urban Zambia. Method: Demographically corrected Zambian norms on a neuropsychological (NP) test battery were used to correct for normal age and education effects. The study assessed 286 HIV positive (+) males (37.1%) and females (62.9%) with a mean age of 41.35 (SD = 8.56) and mean years of education = 10.16 (SD = 2.18). A comprehensive NP test battery was used to assess cognitive domains frequently affected by HIV: attention/working memory, learning/and delayed recall, executive function, verbal fluency, processing speed, verbal and visual episodic memory, and fine motor skills. Results: In younger HIV+ Zambians, higher education evidenced protective effects against NC impairments overall, and for the specific domains of executive functions, learning and speed of information processing. Impairment scores did not support accelerated overall brain aging although the restricted age range and relative youth of our total sample may have precluded detection of such tendencies. Conclusions: The present study raises the need to investigate factors that could be implicated in the poor neurocognitive performance among the younger, less educated HIV+ individuals in Zambia.

General Scientific Summary
Zambia’s younger, less educated HIV+ individuals are at risk for cognitive impairment. Age and education interaction on cognitive performance was investigated in an HIV+ sample in Zambia. Younger, less educated HIV+ should be empowered in disease management in Zambia.

Keywords: HIV, neurocognitive functioning, age, education

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Antiretroviral therapy (ART) has reduced the worldwide HIV and AIDS-related mortality rates from a peak of 2.2 million in the mid-2000s to 1.8 million in 2010 and 1.1 million in 2015 (Negin, Mills, & Bärnighausen, 2012; UNAIDS, 2016). ART also has helped with a reduction in the prevalence of HIV-related dementia by as much as 50% (Bociaga-Jasik, Lickiewicz, Cieśla, Mach, & Garlicki, 2010; Cholewińska & Szymańska, 2009; Rosca, Rosca, Simu, & Chirileanu, 2012; Vance, Fazeli, Ross, Wadley, & Ball, This article was published Online First March 5, 2018.

Norma Kabuba, Department of Psychology, University of Zambia, and Department of Psychology, Norwegian University of Science and Technology; J. Anitha Menon, Department of Psychology, University of Zambia; Donald R. Franklin Jr., Department of Psychiatry, University of California, San Diego; Stian Lydersen, Regional Centre for Child and Youth Mental Health and Child Welfare-Central Norway, Faculty of Medicine, Norwegian University of Science and Technology; Robert K. Heaton, Department of Psychiatry, University of California, San Diego; Knut A. Hestad, Department of Research, Innlandet Hospital Trust, Hamar, Norway, and Department of Public Health, Hedmark University of Applied Sciences.

Correspondence concerning this article should be addressed to Norma Kabuba, Department of Psychology, University of Zambia, Great East Road Campus, P.O. Box 32379, Lusaka, Zambia. E-mail: ewelanjik@gmail.com
on increased cognitive problems for the older HIV infection (Negin & Cumming, 2010).

Sub-Saharan Africa is also undergoing a demographic shift in terms of the number and age range of people living with HIV and AIDS: Various studies report that since more people have access to ART, mortality rates have begun to drop and HIV-positive people are surviving much longer. The successful scale up of ART in developing countries has moved from negligible levels in 2003 to more than 7.9 million in 2012 (Floyd et al., 2012; Negin & Cumming, 2010; Negin, Mills, & Bärnighausen, 2012; Reniers et al., 2009).

Since its first reported incidence of HIV in the 1980s, Zambia, a sub-Saharan African country, has been making strides to alleviate the effects of HIV and AIDS on its population. To that end, the Zambian government introduced ART free of charge at the two major hospitals since 2002. Free ART is currently offered at various primary health care centers and all government clinics in Zambia (Stringer et al., 2006).

However, as increasingly more people infected with HIV live longer, it is postulated that there will be a corresponding increase in milder forms of HAND. These forms have been named asymptomatic neurocognitive impairment, and mild neurocognitive disorders, which, according to some estimates, are affecting 30–60% of people living with HIV (Cholewińska & Szymańska, 2009; Heaton et al., 2010; Vance et al., 2012).

The aforementioned HIV-associated neurocognitive disorders affect various aspects of cognitive functioning including impairments in attention, concentration, learning, memory, psychomotor ability and speed of information processing (Kabuba, Menon, Franklin, Heaton, & Hestad, 2017; Negin & Cumming, 2010).

It is well documented that increasing age in itself is associated with declines in cognitive functioning. Equally, HIV infection is also associated with a risk of cognitive decline. Therefore, with the steady increase in the number of older adults living with HIV, there is need to understand the interaction of age and HIV-1 associated cognitive impairment (Hardy & Vance, 2009; Kissel, Pukay-Martin, & Bornstein, 2005; Valcour et al., 2004; Vance, 2009; Vance, McDougall, Wilson, Debsiai, & Cody, 2014).

The combined risks of HIV infection and age are reported to increase the prevalence or severity of impairment beyond effects of either risk alone (Milanini & Valcour, 2017; Saylor et al., 2016). Previous studies have revealed that older HIV+ individuals are more likely to develop cognitive problems and a decline in functional ability (Farooqui & Farooqui, 2009; Sheppard et al., 2015). Neurocognitive impairment within HIV+ is said to significantly correlate with older age (Sheppard et al., 2015). Some scholars have postulated that this could be more directly related to ART toxicities and not HIV, however, comparative studies between drug-conservative groups and those on aggressive treatment demonstrate that uncontrolled HIV is essentially more detrimental than ART toxicities (Deeks & Phillips, 2009; Pathai, Bajililan, Landay, & High, 2014).

Additionally, concerns have been raised to the effect that HIV may precipitate an NP process similar to that observed in Alzheimer’s disease (Iudicello et al., 2012; Kuhlmann, Minihan, Huebce, Nebel, & Rimbach, 2010; Valcour et al., 2004). This evidence thus further potentially poses an increased risk of developing cognitive problems for the older HIV+ individuals.

All these factors make it imperative to understand the increased risk of HIV infection and age especially in sub-Saharan Africa were the number of people aging with HIV is increasing (Joska et al., 2011, 2012; Negin & Cumming, 2010). Another key component related to cognitive functioning and HIV infection is education. There are currently conflicting results regarding aspects of education that are implicated in cognitive functioning in HIV-infected persons. Some studies involving HIV-1 positive subjects suggest that although educational attainment is an important factor in cognitive performance, years of education does not necessarily account for quality of education (Ryan et al., 2005). On the contrary, other studies have shown that level of education itself has an effect on cognitive functioning in infected groups (De Ronchi et al., 2002; Satz et al., 1993).

It is well established that level of education is an important element not only in normal people, but also in those with brain damage, affecting cognitive performance in various domains, especially for verbal performance tests. Higher education groups have been reported to present slower cognitive decline as a result of normal aging than the individuals with lower educational levels. This is because education may have a protective effect on cognitive functioning. Increased cognitive capacity, referred to as cognitive reserve, is typically used to explain the delay in cognitive and functional expression of neurodegenerative illnesses such as HIV (Ardila, 1998; Ardila, Ostrosky-Solis, Rosselli, & Gómez, 2000; Bornstein & Suga, 1988; Le Carret et al., 2003; Lezak, 1995; Ostrosky-Solis, Ardila, Rosselli, Lopez-Arango, & Urriel-Mendoza, 1998; Stern, 2002; Tombaugh, 2004; Welch, Doineau, Johnson, & King, 1996).

In normal populations, there are significant differences in cognitive functioning associated with levels of education. The effect of education is not linear but rather represents a negatively accelerated curve which tends to plateau. This is because the ceiling in NP tests are typically low (Ardila, 1998).

To determine the combined effects of HIV serostatus and education level on cognitive abnormalities, Satz et al. (1993), employed five NP tests (grooved pegboard, verbal fluency, symbol digit modalities, and Rey auditory verbal learning). There were abnormalities (38%) in those with less than 12 years of education compared with 17% cognitive abnormalities that were found in those with more than 12 years of education. They report that the interaction between education level and serostatus was evident even after possible confounding factors such as age, ethnicity and CD4 count level were controlled for. Their study had the strength of comprising a large sample size of 888 HIV+ and 855 HIV− participants. These results point to the potential interaction between level of education and HIV+ status on cognitive function.

In Zambia, the formal education system is broadly structured as basic education (1–9 years of education) and higher education...
(10–12 years of education). Basic education is a three-tiered system with lower basic (1–4 years), middle basic (5–7 years), and upper basic (8–9). The first two tiers are primary education while upper basic (8–9 years) is the first phase of secondary education and this is followed by high education (10–12 years). Students who complete 12 years of education are thereafter expected to pursue tertiary education offered in colleges and universities (UNESCO, 2011).

According to the Zambia National Education Profile (2014), the net enrollment rate for primary school is at 94% and the primary completion rate is 91%. The enrollment rate for lower secondary education rate is 68% while the transition rate to higher education is 56%.

In light of the foregoing, the current study sought to investigate whether age and education has an effect on the cognitive functioning of the HIV+ adult population in Zambia. The current study had the strength of employing a comprehensive NP test battery measuring seven cognitive domains. Normal effects of age and education were corrected for using a previously collected large sample of healthy HIV− adult Zambians (Hestad et al., 2016).

The main aim of the current study was to establish how neurocognitive functioning is affected by age and level of education in people infected with HIV-1 clade C. Specifically, we wanted to find out the possible neurocognitive differences between HIV+ young adults and HIV+ older adults. Furthermore, the study sought to determine if there are neurocognitive differences between HIV+ participants with more years of formal education versus those with fewer years of formal education.

We hypothesized that there would be a main effect for age and education on NP performance in our HIV+ sample. We expect that the older HIV+ participants will have poorer results on demographically corrected NP test scores (from published norms), compared with the younger HIV+ participants and, furthermore, we expect that the HIV+ participants with higher levels of education would have better scores than the HIV+ participants who had lower levels of education.

Method

Participants

The sample was drawn from six urban clinics in the Zambian capital city of Lusaka. The clinics sampled were: Chilenje, Chipata, Kabwata, Kalingalinga, Matero Main, and Matero Referral clinic. All of the clinics sampled are under the management of the Lusaka District Health Management Team in Zambia. These clinics were chosen because of the presence of the antiretroviral centers, which routinely provides HIV counseling and testing services as well as providing treatment.

Participants were eligible to participate in the study if they were: HIV sero-positive and on antiretroviral treatment. Information about HIV status was based on the participants’ medical files. They also were required to have a minimum of 5 years of formal education. English is the primary language of instruction in the Zambian school system, and the testing was performed in English (Hestad et al., 2016). Participants were also required to be between 20 and 65 years of age (to conform to the age range of the HIV− controls who participated in the Zambian NP norming study; Hestad et al., 2016).

Potential participants were excluded from the study if they had a history of non-HIV related neurological disorders such as epilepsy or closed head injury. This information was obtained by means of the Neurobehavioural Medical Screen Form, which assesses past medical and neurological histories (Kabuba et al., 2017). Participants with a history of drug abuse were also excluded from the study. This information was obtained using a structured interview (Heaton, Miller, Taylor, & Grant, 2004; Kabuba, Menon, & Hestad, 2011). Furthermore, Individuals with obvious physical disabilities were excluded from the study to minimize the possibility of test performance requiring motor dexterity being impaired due to the handicap.

The target sample size was 324 participants, but only 286 HIV+ participants were included in the current study because 38 participants with pulmonary tuberculosis were excluded due to the possible effect of tuberculosis on NP functioning (Robertson, Liner, & Heaton, 2009).

Procedure

The recruitment process was carried out with the assistance of nurses at each clinic who identified participants who routinely seek ART meeting the inclusion criteria. Once informed consent was obtained, the participants were referred to one of 10 neuropsychology master of science students from University of Zambia, who had received extensive training in interviewing, as well as administration and scoring of the NP tests. The testing process for each participant took approximately 2 hr 30 min (Kabuba et al., 2017).

The first part of the testing process involved obtaining the participants’ demographic characteristics, medical, and psychiatric information based on self-report. Medical details were confirmed by the patients’ medical records provided by the medical personnel. Administration of NP tests was carried out in the same order for all participants, and they were compensated the Zambian Kwacha equivalent of $5 (USD) for transport and refreshment allowance at the end of the testing process (Kabuba et al., 2017).

Measures

Cognitive functioning. Cognitive functioning was measured using an NP test battery that measures cognition across the seven ability domains that have been identified as frequently affected in HIV-associated neurocognitive disorders (Antinori et al., 2007): Executive functioning (Stroop Color–Word Interference trial, Category Test errors, WI Card Sorting Test–64 total errors, and Color Trails 2), working memory/attention (Paced Auditory Serial Addition Test–50 and Wechsler Memory Scale–III Spatial Span Test), speed of information processing (Wechsler Adult Intelligence Scale–III [WAIS–III], Digit Symbol, WAIS–III Symbol Search, Trails A, Color Trails 1, Stroop Color Naming and Stroop Word Naming), verbal fluency (letter fluency, animal fluency, and action fluency), learning (Hopkins Verbal Learning Test—Revised [HVLT-R] and Brief Visuospatial Learning Test—Revised [BVMT-R], delayed recall (HVLT-R delay and BVMT-R delay), and complex motor function (grooved pegboard [dominant and nondominant hands])). The test battery is appropriate for the current study because it measures the domains typically affected by HIV. It is also an internationally well-recognized NP assessment tool that has been translated into multiple languages around the world.
females, with an average age of 38.5 (healthy sample comprised 157 (48.5%) males and 167 (51.5%)
data (Heaton, Miller, et al., 2004). The raw scores were corrected
process was similar to that used for the United States normative
information were considered in the category of “low education level,”
upper quartile distribution of the data; less than 9 years of education
were the education levels at which major examinations are taken in
Zambia to qualify for high school and tertiary education, respec-
tively (UNESCO, 2011). This approach uses all the available data,
focusses on conceptually relevant age and education contrasts, and
allows for testing significant effects for age, education, and their interaction. Also, these education lengths are the lower and upper quartile in our data set. Education and age were analyzed as continuous covariates in the current study instead of dichotomizing these for two main reasons: first, dichotomizing implicitly means that the effect is assumed constant within each side of the cutpoint, and makes a discrete “jump” at the cutpoint. This is practically never realistic. Second, dichotomizing implies loss of statistical power compared with using the variable as it is.
Here, p values <0.05 are considered significant. However, due
to multiple hypotheses, p values between 0.01 and 0.05 should be
interpreted with caution. Analyses were carried out in SPSS Version 22.

Results

Demographics and Disease Characteristics

Demographics and disease characteristics are shown in Table 1. The HIV+ sample (n = 286) had a mean age of 41.35 (SD = 8.56), and a mean number of years of education of 10.16 (SD = 2.18). There were 37.1% males and 62.9% females. In our sample, 56 persons (19.6%) had less than 9 years of education, 128 (44.8%) had 9 to 11 years, and 102 (35.7%) had 12 or more years of education.

The "normal" effects of age, education, and sex on the NP test performance were controlled for by using demographically corrected standard scores (T scores) generated with a previously collected normative sample of 324 HIV-negative participants (Hestad et al., 2016).

Table 2 shows the estimated means of NP performance across the seven domains using T score means. Results are illustrated for the lower and upper quartiles of age and education: for younger and older age levels estimated for ages 35 and 47 years, respectively, and for low and high education levels, 9 and 12 years of education, respectively. In these analyses, years of education, age, and their interaction were included as continuous covariates.

As seen in the table, there is a statistically significant effect of age and/or education for learning, recall, and motor functions. This is illustrated in Figure 1. In what follows, we report p values for age and education for these.

For learning, young but not old age had a significant (p = .003) effect of education, with higher education showing relatively preserved/normal abilities. There is only a negligible and nonsignificant effect of education.

For recall, at the young age level there is a significant (p = .026) effect of education, with low education being associated with poorer performance. An at older age level, again higher education was marginally with better performance associated with higher education.

For motor functions, all mean estimated scores were high and normal, regardless of age and education with lower education actually being associated with better motor performance.

To further investigate the effects of age and education on NP performance across the seven domains, we conducted analyses adjusted for the following potential confounders simultaneously: nadir CD4, current CD4, AIDS status, duration on ART, viral load detection, BMI, and gender. These results remained substantially the same when adjusting for the potential confounders; data not shown.

Table 3 shows the estimated means of NP deficit scores across the seven domains and globally for ages 35 and 47, and 9 and 12 years of education. In these analyses, years of education, age, and their interaction were included as continuous covariates.

As seen in the table, there is a statistically significant effect of age and/or education for learning, executive function, motor functions, and the global deficit score and a trend for processing speed. This is illustrated in Figure 2. Although the overall model for motor functions was statistically significant, age and education on their own had no statistically significant effect in this domain. In what follows, we report p values for age and education for these.

For learning, young but not old age had a significant (p = .01) effect on education. The younger, less educated had poorer performance compared with the younger, more educated group.

For executive functions, at the younger age level there is a significant (p = .05) effect of education. The younger, less educated had poorer performance compared with the younger, more educated group. At the older age there is only a negligible and nonsignificant effect of less age or less education.

For speed of information processing, the younger, less educated had poorer performance (p = .03) compared with the younger, more educated group. The older age groups showed nonsignificant effects of both age and education.

For global deficit scores, the young age and less educated group had poorer performance (p = .03) compared with the younger, more educated group. The older age group exhibited negligible and nonsignificant effect of age and education effects.

To further investigate the effects of age and education on NP performance across the seven domains, we conducted analyses adjusted for the following potential confounders simultaneously: nadir CD4, current CD4, AIDS status, duration on ART, viral load detection, BMI, and gender. The results remained substantially the same when adjusting for the potential confounders; data not shown.

Table 1

<table>
<thead>
<tr>
<th>Demographic and Disease/Treatment Characteristics of HIV+ Sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demographic and disease characteristics</td>
</tr>
<tr>
<td>------------------------------------------</td>
</tr>
<tr>
<td>Age</td>
</tr>
<tr>
<td>Education</td>
</tr>
<tr>
<td>Female gender</td>
</tr>
<tr>
<td>Nadir CD4</td>
</tr>
<tr>
<td>Current CD4</td>
</tr>
<tr>
<td>Undetectable viral load</td>
</tr>
<tr>
<td>Duration on ART</td>
</tr>
<tr>
<td>With AIDS</td>
</tr>
<tr>
<td>BMI</td>
</tr>
</tbody>
</table>

Note. The sample sizes (n) for disease characteristics vary due to missing data. Neuropsychological performance based on age and education in the HIV+ sample. ART = antiretroviral therapy; BMI = body mass index.
Age and Neurocognitive Functioning in the HIV+ Sample

We hypothesized that the older HIV+ participants would have poorer cognitive performance on the NP tests compared with the younger HIV+ participants. Advancing age is typically reported to be associated with a decline in cognitive functioning (Hardy & Vance, 2009; Kissel, Pukay-Martin, & Bornstein, 2005; Milanini & Valcour, 2017; Vance, 2009; Vance et al., 2014). In the current study, the effects of normal aging and education levels in healthy individuals were controlled by the use of demographically corrected NP test norms. Thus our explanation was that HIV infection would exacerbate the process of normal cognitive aging.

Contrary to our hypothesis, advancing age in our HIV+ sample did not negatively affect cognitive performance on the NP tests. The statistically significant effect of age yielded from the current study showed that the younger HIV+ group with less education performed worse on learning and recall on the mean T scores than the younger HIV+ group with more education. A similar trend was observed on the impairment scores in which the younger, less educated performed more poorly on the learning domain scores and global deficit scores than the younger, more educated group. The results obtained in the current study are counterintuitive although similar to those reported by Wilkie et al., 2003, who reported that certain cognitive processes may be more impaired in the younger than the older HIV-1 infected adults. They reported that learning and delayed recall were more impaired in the younger HIV+ compared with the older HIV+ group. Caveat the NP performance differences observed in the current study we assume, were not due to age effects but rather due to education effects/cognitive reserve. Although the current study findings may appear to be counterintuitive, we postulate that, age effects (young vs. old) were likely masked because the sample in the current study was relatively young compared with the previous studies that reveal that individuals over the age of 60 typically demonstrate negative cognitive results owing to age on NP evaluation (Ardila, 1998; Becker, Lopez, Dew, & Aizenstein, 2004; De Ronchi et al., 2002; Milanini & Valcour, 2017; Wilkie et al., 2003).

Table 2

<table>
<thead>
<tr>
<th>Domain</th>
<th>9 years edu. EM (SE)</th>
<th>12 years edu. EM (SE)</th>
<th>9 years edu. EM (SE)</th>
<th>12 years edu. EM (SE)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Executive</td>
<td>46.05 (.56)</td>
<td>47.38 (.66)</td>
<td>47.11 (.53)</td>
<td>47.23 (.57)</td>
<td>.211</td>
</tr>
<tr>
<td>Fluency</td>
<td>46.39 (.65)</td>
<td>46.59 (.76)</td>
<td>47.53 (.61)</td>
<td>47.12 (.65)</td>
<td>.483</td>
</tr>
<tr>
<td>Working memory</td>
<td>44.80 (.72)</td>
<td>44.09 (.84)</td>
<td>45.46 (.68)</td>
<td>44.68 (.73)</td>
<td>.580</td>
</tr>
<tr>
<td>Learning</td>
<td>43.05 (.66)</td>
<td>45.59 (.77)</td>
<td>45.60 (.62)</td>
<td>45.76 (.66)</td>
<td>.002</td>
</tr>
<tr>
<td>Recall</td>
<td>44.02 (.69)</td>
<td>46.02 (.80)</td>
<td>45.99 (.65)</td>
<td>47.00 (.69)</td>
<td>.014</td>
</tr>
<tr>
<td>Motor</td>
<td>54.04 (.90)</td>
<td>51.69 (1.05)</td>
<td>52.05 (.85)</td>
<td>49.86 (.91)</td>
<td>.008</td>
</tr>
<tr>
<td>Speed of information processing</td>
<td>45.83 (.67)</td>
<td>46.53 (.72)</td>
<td>47.17 (.58)</td>
<td>46.81 (.62)</td>
<td>.292</td>
</tr>
<tr>
<td>Global mean T</td>
<td>46.18 (.48)</td>
<td>46.80 (.56)</td>
<td>47.27 (.45)</td>
<td>46.98 (.48)</td>
<td>.249</td>
</tr>
</tbody>
</table>

Note. p value for the combined effect of these covariates (F test with 3 degrees of freedom). Bold values are meant to highlight the statistically significant results. Results are presented as estimated means with associated standard errors for ages 35 and 47 years (lower and upper quartile in our data set), and for 9 years and 12 years of education years. See the online supplemental materials for tables showing beta estimates and confidence intervals. NP = neuropsychological; EM = estimated mean; SE = standard error.

Discussion

Age and Neurocognitive Functioning in the HIV+ Sample

Figure 1. Estimated mean T scores for learning, recall, and motor functions, from linear regression with age, education, and their interaction as covariates. See the online article for the color version of this figure.
tive functions (Hardy et al., 1999; Van Gorp et al., 1994; Wilkie et al., 2003).

Education and Neurocognitive Functioning in the HIV + Sample

We had postulated that within the context of HIV infection, participants with higher levels of education would show less cognitive impairment than those with lower levels of education. This was confirmed in the current study but only at the lower age range (see Tables 2 and 3). The younger and less educated in the current study were found to perform more poorly on learning, executive function, processing speed, and on the global deficit scores. The results yielded in the present study are in agreement with previous studies (Satz et al., 1993; De Ronchi et al., 2002), which show that level of education has an effect on cognitive functioning in the HIV +. This result shows that the effect of cognitive reserve was more apparent in the young age level than in

<table>
<thead>
<tr>
<th>Domain</th>
<th>9 years edu.</th>
<th>12 years edu.</th>
<th>9 years edu.</th>
<th>12 years edu.</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>EM (SE)</td>
<td>EM (SE)</td>
<td>EM (SE)</td>
<td>EM (SE)</td>
<td></td>
</tr>
<tr>
<td>Executive</td>
<td>.44 (.04)</td>
<td>.28 (.05)</td>
<td>.36 (.04)</td>
<td>.34 (.04)</td>
<td>.032</td>
</tr>
<tr>
<td>Fluency</td>
<td>.52 (.06)</td>
<td>.45 (.07)</td>
<td>.43 (.05)</td>
<td>.44 (.06)</td>
<td>.589</td>
</tr>
<tr>
<td>Working memory</td>
<td>.67 (.07)</td>
<td>.59 (.08)</td>
<td>.60 (.07)</td>
<td>.63 (.07)</td>
<td>.679</td>
</tr>
<tr>
<td>Learning</td>
<td>.66 (.06)</td>
<td>.46 (.07)</td>
<td>.40 (.06)</td>
<td>.46 (.06)</td>
<td>.001</td>
</tr>
<tr>
<td>Recall</td>
<td>.56 (.06)</td>
<td>.43 (.07)</td>
<td>.46 (.06)</td>
<td>.37 (.06)</td>
<td>.133</td>
</tr>
<tr>
<td>Motor</td>
<td>.14 (.05)</td>
<td>.17 (.06)</td>
<td>.25 (.05)</td>
<td>.35 (.05)</td>
<td>.009</td>
</tr>
<tr>
<td>Speed of information processing</td>
<td>.49 (.05)</td>
<td>.39 (.06)</td>
<td>.37 (.05)</td>
<td>.43 (.05)</td>
<td>.056</td>
</tr>
<tr>
<td>Global deficit score</td>
<td>.49 (.04)</td>
<td>.39 (.04)</td>
<td>.40 (.04)</td>
<td>.43 (.04)</td>
<td>.035</td>
</tr>
</tbody>
</table>

*Note. p value for the combined effect of these covariates (F test with 3 degrees of freedom). Bold entries meant to highlight the statistically significant results. Results are presented as estimated means with associated standard errors for ages 35 and 47 years (lower and upper quartile in our data set), and for 9 years and 12 years of education years. See the online supplemental materials for tables showing beta estimates and confidence intervals. NP = neuropsychological; EM = estimated mean; SE = standard error.*

Figure 2. Estimated mean deficit scores for executive functions domain deficit score (DDS), Learning DDS, global deficit scores and speed of information processing DDS from linear regression with age, education and their interaction as covariates. See the online article for the color version of this figure.
the older age level participants with more years of education. Previous studies have shown that due to cognitive reserve, higher education groups tend to have a slower decline on cognitive tests than those with lower educational levels (Ardila, 1998; Ardila et al., 2000; Bornstein & Suga, 1988; Le Carret et al., 2003; Lezak, 1995; Ostrosky-Solis et al., 1998; Stern, 2002; Tombaugh, 2004; Welch et al., 1996).

In addition, the current study yielded peculiar results regarding education and performance on motor functions. Interestingly, although the scores for all age and education groups regarding motor functions were within the normal range; the older, more educated group did not perform as well as the less educated, younger group. This result is consistent with results obtained by Robertson et al. (2007) in Uganda and Heaton et al. (2011), who report that grooved pegboard tasks were less affected in the HIV+ in Combination antiretroviral therapy (CART) era. This result could be an indication that motor functions are not adversely affected in HIV+ individuals on ART in Zambia.

On the contrary, this result could possibly be an indicator that the younger sample with less education might be more involved in manual work that demands motor dexterity as opposed to the older sample. Presumably, since the older participants in the current study are engaged in cognitively demanding jobs, their performance on neuropsychology tests is better and more homogeneous due to the positive neuroplasticity resulting from the cognitively demanding tasks they perform. Furthermore, the assumption would be that the younger group in our sample is presumably engaged in different work conditions where the younger, less educated are involved in manual work and the younger, more educated in cognitively demanding jobs, which could account for the differences observed between the two groups. The younger, more educated benefit from the cognitive reserve as a result of education and continued use of cognitive skills and thus perform better on the NP tests (Mahmcke, Bronstone, & Merzenich, 2006; Vance & Burrage, 2006).

The implication of this is that cognitive performance is worse for HIV+ people who are younger and less educated in Zambia. There is, however, a need to carry out further research to ascertain why this could be the case. It could also be necessary to carry out a longitudinal study that would better take into account the differences that could account for the effects of age and education on neurocognitive impairment in HIV+ adults in Zambia.

Conclusion

The findings obtained from the current study show that there was an effect of level of education on neurocognition in the HIV+ younger sample. The cognitive reserve properties associated with higher education enabled the younger, more educated participants to perform better than the younger, less educated group on neurocognitive tests. The current study also revealed that motor functions are preserved in the HIV+ on CART in Zambia. This could be an indication that the younger, less educated need to be empowered in disease management.

References


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