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Measuring disability experienced by adults living with HIV: assessing construct validity of the HIV Disability Questionnaire using confirmatory factor analysis

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ABSTRACT

Objectives: To assess the construct validity of the HIV Disability Questionnaire (HDQ), a self-administered questionnaire that describes the presence, severity and episodic nature of disability experienced by people living with HIV.

Design: We conducted a confirmatory factor analysis. We hypothesised that domains in the HDQ characterised six dimensions of disability, each represented by HDQ items: physical symptoms and impairments (20 items); cognitive symptoms and impairments (3 items); mental and emotional health symptoms and impairments (11 items); uncertainty (14 items); difficulties with day-to-day activities (9 items) and challenges to social inclusion (12 items). We developed a measurement model to test these hypotheses. We used maximum likelihood methods of estimation to determine model fit. We considered a threshold for the Root Mean Square Error of Approximation (RMSEA) of <0.05 as an indication of overall goodness of model fit. We considered variables with factor loadings of >0.30 as representing a given domain of disability.

Setting: We recruited adults with HIV from hospital clinics, AIDS service organisations and a specialty hospital in Ontario.

Participants: Of the 361 adults with HIV who completed the HDQ, 80% were men, 36% were 50 or older and 77% reported living with at least two concurrent health conditions in addition to HIV.

Outcome Measures: We administered the HDQ followed by a demographic questionnaire.

Results: The model achieved good overall fit as indicated by a RMSEA of 0.030 (90% CI 0.028 to 0.033). All HDQ items represented our hypothesised dimensions of disability (factor loadings >0.30). Factor loadings ranged from 0.34 to 0.90. Domains of disability correlated with each other ranging from r=0.47 (between difficulties with day-to-day activities and uncertainty) to r=0.88 (between mental-emotional health challenges and challenges to social inclusion).

Conclusions: The six domain structure of the HDQ demonstrated construct validity when administered to adults living with HIV.

Introduction

As people living with HIV infection are living longer and aging, they are experiencing the complex health-related challenges of the infection, associated comorbidities and potential adverse effects of treatment.1–3 Health-related challenges, collectively termed disability, can include symptoms and impairments (eg, fatigue, neurocognitive impairments, weakness, pain), difficulties

Strengths and limitations of this study

- We assessed the construct validity of the HIV Disability Questionnaire (HDQ), a 69 item self-reported questionnaire developed to assess the presence, severity and episodic nature of disability experienced by adults living with HIV.

- We conducted a confirmatory factor analysis hypothesising that domains in the HDQ represented six dimensions of disability, each represented by HDQ items: physical symptoms and impairments (20 items); cognitive symptoms and impairments (3 items); mental and emotional health symptoms and impairments (11 items); uncertainty (14 items); difficulties with day-to-day activities (9 items) and challenges to social inclusion (12 items).

- The six domain structure of the HDQ demonstrated construct validity when administered to adults living with HIV. The hypothesised model achieved good overall fit as indicated by a RMSEA of 0.030. All HDQ items represented our hypothesised dimensions of disability (factor loadings >0.30).

- The HDQ is the first known HIV-specific instrument of disability developed from the perspective of adults living with HIV.

- The HDQ can be used to describe the multidimensional nature of disability experienced by adults living with HIV and lay the foundation for more widespread measurement of disability in HIV clinical practice and research.
with day-to-day activities (eg, household chores), challenges to social inclusion (eg, ability to work) and uncertainty or worrying about the future.4–6 Disability may be experienced as episodic in nature, whereby health challenges fluctuate on a daily basis and over the entire course living with HIV.7 Given current treatments for HIV and the long-term survival for successfully treated individuals, measuring disability is critical for determining the impact of the disease, its comorbidities and interventions.

We developed a new self-administered instrument, called the HIV Disability Questionnaire (HDQ) using the Episodic Disability Framework, a conceptual framework that describes disability experienced by adults living with HIV. The Episodic Disability Framework consists of three main components: (1) dimensions of disability that may fluctuate on a daily basis and over the entire course of living with HIV (symptoms and impairments, uncertainty, difficulties with day-to-day activities, and challenges to social inclusion), (2) contextual factors (social support, stigma, living strategies and personal attributes) that may exacerbate or alleviate the dimensions of disability and (3) triggers, defined as life events that mark momentous or major episodes of disability.7,8 The purpose of the HDQ is to describe the presence, severity and episodic nature of disability experienced by adults living with HIV.9 Items in the HDQ were derived from the Episodic Disability Framework.7 The HDQ consists of six domains of disability: physical symptoms and impairments; cognitive symptoms and impairments; mental and emotional health symptoms and impairments; uncertainty; difficulties with day-to-day activities and challenges to social inclusion.10 The HDQ possesses sensibility (face and content validity and ease of use).11 Our aim was to assess the construct validity of the HDQ among adults living with HIV.

HIV Disability Questionnaire

The HDQ consists of 69 items that describe the presence, severity and episodic nature of disability experienced by adults living with HIV. Each item consists of a statement about a health-related challenge (eg, ‘I have trouble remembering things like appointments and when to take my medications’) and has both a seven-point ordinal response scale asking the respondent to rate the challenge on the day of administration (from 0 – ‘None at all’ to 6 – ‘Extreme trouble’) and a nominal response scale asking whether the challenge fluctuated (or changed) over the past week (‘Yes’ or ‘No’).

All data were entered into a database and 20% of cases were independently checked for accuracy. We removed any cases with >10% of item responses missing. We calculated disability presence, severity and episodic scores for each domain and for the total HDQ scale. Disability presence scores were calculated by summing the number of challenges (≥1 response option on the seven-point scale) for a total disability presence score (ranging from 0–69). Disability severity scores were calculated by summing individual item scores (ranging from 0 to 6) for each domain. Disability episodic scores were calculated by summing the number of challenges identified as episodic (‘Yes’ responses; ranging from 0–69). All presence, severity and episodic scores were linearly transformed to a score ranging from 0 to 100. Higher scores on each scale indicated a greater degree or episodic nature of disability.

Analysis

We conducted a confirmatory factor analysis to assess the construct validity of the domains of disability in the HDQ. We hypothesised that domains in the HDQ represented six dimensions of disability, each represented by the following HDQ items: physical symptoms and impairments (represented by 20 items); cognitive symptoms and impairments (3 items); mental and emotional health symptoms and impairments (11 items); uncertainty (14 items); difficulties with day-to-day activities (9 items) and challenges to social inclusion (12 items; figure 1).

We assessed convergent construct validity by determining the extent to which disability severity scores in the HDQ represented a hypothesised domain of disability with factor loadings >0.30. We assessed divergent construct validity by determining the extent to which domains of disability were distinct constructs that together comprised the larger construct of disability. We considered correlations between latent variables <0.80 as signifying distinct dimensions of disability.12

We used the maximum likelihood methods of estimation, which is preferred for non-normally distributed data.13 Prior to our analysis, we conducted mean imputation for missing data in order to maximise the sample size for analysis.12 We estimated and reported standardised parameter estimates (or factor loadings) for each item. We defined factor loadings >0.30 as indicating a
relationship between an HDQ item and a given domain; these items were considered as ‘loading’ on that domain.

We used a combination of approaches to evaluate the overall goodness of fit of the confirmatory factor analysis solution. If the solution demonstrated adequate goodness of fit we considered this as constituting evidence in favour of construct validity of the domains of the HDQ. We considered a Root Mean Square Error of Approximation (RMSEA) <0.05, Comparative Fit Index (CFI) >0.95 and Tucker Lewis Index (TLI) >0.95 to indicate good model fit. The RMSEA is a population-based index that assesses the extent to which a model fits reasonably well in the population by evaluating the discrepancy between the hypothesised model, with optimally chosen parameter estimates, and the population covariance matrix. The RMSEA ranges from 0 to 1, with smaller values indicating better model fit. The CFI and TLI assess model fit by examining the discrepancy between the data and the hypothesised model while adjusting for sample size. CFI and TLI values range from 0 to 1 with higher values indicating better fit. We considered the RMSEA as the primary statistic for overall goodness of model fit because it is less sensitive to sample size and is recommended for confirmatory factor analysis. We estimated our sample size using the rule that a minimum of five participants per item are required for factor analysis. With 69 disability severity items in the HDQ, we required at least 345 participants. We used SPSS (V.19.0) for computation of HDQ scores and Mplus (V.7.0) for the confirmatory factor analysis.

RESULTS

Three-hundred and sixty-one participants were recruited from AIDS service organisations in Toronto (51%), word of mouth (28%), AIDS service organisations in Hamilton, Niagara and Durham regions (7%), a specialty hospital in Toronto (3%) and hospital clinics in Toronto and other areas of southern, Ontario (3%). The majority of participants were men (80%), the median age was 46 years, most were taking antiretroviral medications (83%), and many (77%) were living with two or more health conditions in addition to HIV (table 1). The most common self-reported concurrent health conditions included mental health challenges such as anxiety and depression (40%), arthritis (27%), osteoporosis or osteopenia (16%) and cardiovascular disease (15%).

HIV Disability Questionnaire

Participants took a median of 14 min to complete the HDQ (1st–3rd quartile: 10–20 min). Almost all HDQ items (n=66) demonstrated a floor effect with >15% of responses rated ‘0’ indicating no disability. Floor effects

Figure 1  A priori measurement model for confirmatory factor analysis of the HIV Disability Questionnaire.
were most common in items that referred to symptoms and impairments or difficulties with day-to-day activities. Ten HDQ items demonstrated a ceiling effect with >15% of responses rated ‘6’ indicating the highest severity of disability. Ceiling effects were most common in items pertaining to uncertainty or worrying about the future. Missing responses were <5% across all HDQ disability severity and episodic items.

Median HDQ presence scores (ranging from 0 to 100) ranged from 44 in the difficulties with day-to-day activities domain (1st–3rd quartile: 11–78) to 100 in the cognitive health challenges domain (1st–3rd quartile: 67–100). Median HDQ severity scores were highest in the uncertainty domain (45; 1st–3rd quartile: 27–67) followed by challenges to social inclusion (33; 18–51), cognitive (28; 11–50), mental-emotional (27; 14–53) and physical symptoms and impairments (25; 14–38). Median HDQ episodic scores (number of challenges that fluctuated within the week) were greatest in the physical (35; 8–54) and cognitive symptoms and impairments domains (33; 0–67) followed by the mental-emotional symptoms and impairments domain (18; 0–64), difficulties with day-to-day activities (11; 0–33), uncertainty (7; 0–50) and challenges to social inclusion (0; 0–33). The most common health challenges that were reported as episodic spanned the physical and mental-emotional symptoms and impairments domains and included fatigue (50%), feeling sad, down or depressed (48%), aches and pains (46%), headaches (42%), feeling anxious (41%), having trouble sleeping (40%) and feeling weakness in muscles (40%). Eighty-two per cent of participants completed the HDQ on what they considered a ‘good day’ living with HIV.

Confirmatory factor analysis results

Results of the confirmatory factor analysis are presented in table 2. Correlation matrices are available on request. The RMSEA was 0.030 (90% CI 0.028 to 0.033), satisfying our primary goodness of fit criterion. The χ² value was 3020.981 (p<0.001). CFI (0.812) and TLI (0.805) statistics did not meet our prespecified criteria for goodness of fit. All HDQ items represented our hypothesised domains of disability (factor loadings >0.30). To interpret the first factor loading in table 2, one standard deviation (SD) increase in physical symptoms and impairments is associated with a 0.64 SD increase in loss of energy. Equivalently, the value of the Pearson correlation coefficient between loss of energy and physical symptoms and impairments is 0.64.

Factor loadings ranged from 0.34 (item 2: ‘I have diarrhoea’, loading on the physical symptoms and impairments domain) to 0.90 (item 22: ‘I have trouble thinking clearly’, loading on the cognitive symptoms and impairments domain). Domains of disability correlated with each other ranging from r=0.47 (between difficulties with day-to-day activities and uncertainty) to r=0.88 (between mental-emotional symptoms and impairments and challenges to social inclusion; table 3).

DISCUSSION

This is the first study to assess the construct validity of the HDQ, the only HIV-specific measure of disability. Results of our confirmatory factor analysis demonstrated good overall model fit of items with the domain structure, supporting the validity of the six domains of disability in the HDQ.

Floor and ceiling effects were evident across the HDQ. We believe that the floor effect, primarily seen in items that addressed day-to-day activities, likely reflected the way in which participants were sampled; most were living independently in the community and faced few challenges to mobility or self-care activities. Ceiling or floor

| Table 1 Characteristics of participants (n=361) |
|-----------------------------------------------|
| **Characteristic** | **Number (%)** |
| Gender | |
| Men | 289 (80) |
| Women | 66 (18) |
| Transgendered | 6 (2) |
| Age (median; 1st–3rd quartile) | 46 years (40–52 years) |
| 50 years or older | 130 (36) |
| Year of HIV diagnosis | 1998 (1991–2005) |
| (median; 1st–3rd quartile) | |
| Diagnosed prior to 1996 | 139 (38) |
| Taking antiretroviral therapy | 301 (83) |
| Nadir CD4 count | 167 (46) |
| (<200 cells/mm³) | |
| Undetectable viral load | 196 (76) |
| Earning between | 192 (53) |
| $20 000–20 000 per year | |
| Currently working for pay | 72 (20) |
| Living alone | 227 (63) |
| Have children | 117 (32) |
| Live with children | 26/117 (22) |
| Self-rated health status | |
| Poor | 22 (6) |
| Fair | 75 (21) |
| Good | 138 (38) |
| Very good | 82 (23) |
| Excellent | 42 (12) |
| Living with two or more concurrent health conditions | 279 (77) |
| Common concurrent health conditions | |
| Mental health (eg, anxiety or depression) | 143 (40) |
| Arthritis | 97 (27) |
| Osteoporosis or osteopenia | 59 (16) |
| Cardiovascular disease (eg, heart attack or stroke) | 55 (15) |
| Hepatitis C | 51 (14) |
| Cancer | 40 (11) |
| Diabetes | 39 (11) |
| Neurocognitive decline | 36 (10) |
| Liver disease | 36 (10) |

Not all characteristics add to the total n due to missing responses.
Table 2: Confirmatory factor analysis results—domains of disability in the HIV Disability Questionnaire (HDQ; n=361)

| Domain of disability (latent variables) | HDQ items (indicator variables) | Item no | Standardised factor loadings | 95% CI |
|-----------------------------------------|---------------------------------|---------|-----------------------------|-------|
| **Physical symptoms and impairments (20 items)** | | 1 | I feel too tired to do my usual activities (loss of energy) | 0.64 | 0.59 to 0.70 |
| | | 2 | I have diarrhoea | 0.34 | 0.25 to 0.42 |
| | | 3 | I feel nauseous | 0.66 | 0.60 to 0.71 |
| | | 4 | I have headaches | 0.60 | 0.54 to 0.67 |
| | | 5 | I have numbness or tingling in my hands | 0.60 | 0.54 to 0.67 |
| | | 6 | I have numbness or tingling in my feet | 0.58 | 0.52 to 0.64 |
| | | 7 | I have aches or pains | 0.68 | 0.64 to 0.73 |
| | | 8 | I have trouble swallowing food | 0.52 | 0.44 to 0.59 |
| | | 9 | I have less desire to have sex (decreased libido) | 0.46 | 0.39 to 0.53 |
| | | 10 | I have shortness of breath | 0.59 | 0.53 to 0.66 |
| | | 11 | I have fever, chills or sweats | 0.57 | 0.50 to 0.64 |
| | | 12 | I feel weakness in my muscles | 0.73 | 0.69 to 0.78 |
| | | 13 | I have muscle cramps | 0.67 | 0.62 to 0.72 |
| | | 14 | I have stomach cramps | 0.63 | 0.57 to 0.69 |
| | | 15 | I am losing weight | 0.42 | 0.34 to 0.50 |
| | | 16 | I lack an appetite for food | 0.46 | 0.39 to 0.53 |
| | | 17 | I have trouble sleeping | 0.50 | 0.43 to 0.57 |
| | | 18 | I have problems with my vision | 0.57 | 0.50 to 0.64 |
| | | 19 | I have problems with my hearing | 0.42 | 0.34 to 0.50 |
| | | 20 | I feel dizzy | 0.70 | 0.65 to 0.76 |
| **Cognitive symptoms and impairments (3 items)** | | 21 | I have trouble remembering things like appointments and when to take my medication | 0.72 | 0.67 to 0.78 |
| | | 22 | I have trouble thinking clearly | 0.90 | 0.87 to 0.92 |
| | | 23 | I have trouble concentrating | 0.87 | 0.84 to 0.90 |
| **Mental health symptoms and impairments (11 items)** | | 24 | I feel anxious | 0.67 | 0.62 to 0.71 |
| | | 25 | I feel sad, down or depressed | 0.77 | 0.73 to 0.80 |
| | | 26 | I am afraid for my future | 0.77 | 0.74 to 0.81 |
| | | 27 | I lack confidence around others | 0.76 | 0.72 to 0.81 |
| | | 28 | I am uncomfortable with how my body looks | 0.62 | 0.57 to 0.67 |
| | | 29 | I feel isolated even when I am around others. In other words, I feel that I do not belong | 0.81 | 0.78 to 0.84 |
| | | 30 | I feel embarrassed around others | 0.82 | 0.79 to 0.85 |
| | | 31 | I feel guilty | 0.76 | 0.72 to 0.81 |
| | | 32 | I feel lonely | 0.75 | 0.71 to 0.79 |
| | | 33 | I feel discouraged about my future life options | 0.81 | 0.78 to 0.84 |
| | | 34 | I feel ‘shut out’ by my friends or family | 0.66 | 0.60 to 0.72 |
| **Uncertainty or worry about the future (14 items)** | | 35 | I worry about my future health living with HIV | 0.84 | 0.81 to 0.87 |
| | | 36 | I worry about my lab test results such as my CD4 count and viral load | 0.76 | 0.72 to 0.80 |
| | | 37 | I worry about having a serious illness. | 0.86 | 0.84 to 0.88 |
| | | 38 | I worry about what the outcome of my next episode of illness might be | 0.87 | 0.85 to 0.89 |
| | | 39 | I worry about the side effects of HIV treatments | 0.70 | 0.66 to 0.75 |
| | | 40 | I worry about my income or financial security living with HIV | 0.68 | 0.64 to 0.72 |
| | | 41 | I worry what might happen to my family and friends if I have an episode of illness | 0.68 | 0.63 to 0.73 |

Continued
| Domain of disability (latent variables) | Item no | HDQ items (indicator variables)                                                                 | Standardised factor loadings | 95% CI |
|----------------------------------------|---------|-----------------------------------------------------------------------------------------------|-----------------------------|--------|
|                                        |         | I worry about being able to remain in the workforce or return to the workforce                  | 0.56                        | 0.50 to 0.62 |
|                                        |         | I worry about dying                                                                          | 0.64                        | 0.58 to 0.70 |
|                                        |         | I worry about my bodily appearance                                                           | 0.67                        | 0.62 to 0.71 |
|                                        |         | I worry about the legal issues of telling others about my HIV status                          | 0.58                        | 0.52 to 0.63 |
|                                        |         | I worry about what others would think of me if they knew I was HIV positive                  | 0.59                        | 0.54 to 0.64 |
|                                        |         | I worry about transmitting HIV to others                                                     | 0.39                        | 0.31 to 0.46 |
|                                        |         | I have put certain life decisions on hold (such as buying a house, returning to work or school, or starting a family) because of my uncertainty living with HIV | 0.58                        | 0.53 to 0.64 |
| Difficulties with day-to-day activities (9 items) | 49      | I am unsteady on my feet                                                                     | 0.69                        | 0.64 to 0.74 |
|                                        | 50      | I have trouble walking                                                                       | 0.75                        | 0.70 to 0.79 |
|                                        | 51      | I have trouble climbing stairs                                                               | 0.73                        | 0.69 to 0.77 |
|                                        | 52      | I have trouble with daily activities such as eating, bathing, grooming, or dressing           | 0.71                        | 0.66 to 0.77 |
|                                        | 53      | I have trouble doing household chores such as cleaning, doing dishes, laundry, and cooking   | 0.78                        | 0.74 to 0.82 |
|                                        | 54      | I have trouble taking part in leisure or recreation, such as exercise or dancing              | 0.78                        | 0.74 to 0.82 |
|                                        | 55      | I have trouble getting out to do errands such as grocery shopping, banking, or doctor’s appointments | 0.85                        | 0.82 to 0.88 |
|                                        | 56      | I have trouble keeping track of my finances                                                  | 0.53                        | 0.47 to 0.60 |
|                                        | 57      | I have trouble getting around, such as driving or taking public transportation                | 0.74                        | 0.68 to 0.79 |
| Challenges to social inclusion (12 items) | 58      | I find it hard to meet the needs of those I care for (such as children, parents, grandparents, partners, or pets) | 0.59                        | 0.52 to 0.66 |
|                                        | 59      | I find it hard to fulfil my role as a family or community member living with HIV              | 0.72                        | 0.68 to 0.76 |
|                                        | 60      | I feel cut off from my friends, networks, ethnic or religious communities                    | 0.71                        | 0.66 to 0.76 |
|                                        | 61      | My illness prevents me from doing volunteer or paid work or going to school                   | 0.60                        | 0.54 to 0.66 |
|                                        | 62      | I feel that my work performance is limited because of my illness                             | 0.64                        | 0.59 to 0.70 |
|                                        | 63      | I struggle to maintain safe and stable housing living with HIV                                | 0.48                        | 0.41 to 0.56 |
|                                        | 64      | I find it hard to talk with others about my illness, even my family and friends              | 0.56                        | 0.51 to 0.62 |
|                                        | 65      | I find it hard to ask others for help when I go through an episode of illness                 | 0.67                        | 0.62 to 0.72 |
|                                        | 66      | I find it hard to start new friendships living with HIV                                       | 0.70                        | 0.66 to 0.75 |
|                                        | 67      | I find it hard to start new, intimate, sexual relationships living with HIV                  | 0.54                        | 0.48 to 0.60 |
|                                        | 68      | I tend to isolate myself from others because I am HIV positive                               | 0.76                        | 0.72 to 0.80 |
|                                        | 69      | I find it hard to take part in leisure or recreational things like going to the movies, out to dinner, or on vacation because I cannot afford it | 0.48                        | 0.41 to 0.55 |

**Confirmatory Factor Analysis—Goodness of Fit Statistics.**

$\chi^2=3020.981$ (p value <0.0001).
Degrees of freedom (df)=2262.
Comparative Fit Index (CFI)=0.812 (ideal is ≥0.95).
Tucker-Lewis Index (TLI)=0.805 (ideal is ≥0.95).
Root Mean Square Error of Approximation (RMSEA)=0.030 (good fit indicated by <0.05).

**Results**

All standardised factor loadings were statistically significant (p<0.0001).
All factor loadings were >0.30 which indicate the HDQ items ‘load’ on a given domain of disability.
The prevalence of disability including physical impairments, activity limitations and social participation restrictions among people living with HIV have been documented using measures based on the International Classification of Functioning, Disability and Health (ICF).^{1, 5–6, 23} The ICF (and the measures derived from the ICF) do not take into account the domain of uncertainty, nor the episodic nature of HIV. The HDQ was developed from the Episodic Disability Framework, a conceptual framework specifically derived and empirically validated with, adults living with HIV.^{7–8, 24} Results showed the highest median HDQ severity score was in the uncertainty domain, highlighting the importance of uncertainty as a key component of disability for adults living with HIV. The Episodic Disability Framework also has been used to inform qualitative approaches to exploring experiences of older men who self-identify as having HIV-associated neurocognitive challenges and has been considered an approach to conceptualise disability among people living with HIV internationally.^{25–26}

However, this is the first known study to use the Episodic Disability Framework to inform the development and validation of a new quantitative measure of disability for adults living with HIV. With episodic health and disability identified as a key research priority in the HIV and rehabilitation field,^{27} the HDQ will be integral to accurately and consistently describe the health-related consequences of HIV, aging and related comorbidities with adults living with HIV.

The highest median HDQ presence score was in the cognitive symptoms and impairments domain. In other studies, self-reported cognitive symptoms have been associated with depression.^{28} We confirmed this association in our analysis; the correlation between cognitive and mental-emotional domains was 0.65. While subjectively measured components of mental health correlate with each other, treatment strategies to address mental-emotional and cognitive health symptoms can differ, suggesting these are distinct clinical concepts^{29–30} as represented in the HDQ.

Compared with other HDQ domains, symptoms and impairments tended to fluctuate more on a daily basis with median HDQ episodic scores greatest in the physical (35 challenges that fluctuated within the week) cognitive (33 challenges), and mental-emotional (18 challenges) symptoms and impairments domains, demonstrating the potential episodic nature of disability. This was expected given items related to symptoms and impairments such as fatigue, weakness and trouble concentrating may fluctuate more readily than those associated with social inclusion such as the ability to engage or re-engage in the workforce. Specific symptoms and impairments that fluctuated the most included fatigue, feeling sad, down or depressed, aches and pains, headaches, feeling anxious, having trouble sleeping and feeling weakness in muscles. Despite low episodic domain scores, participants demonstrated a range of episodic scale of the HDQ.

Further work is needed to assess the properties of the episodic scale of the HDQ.

Lastly, the majority of participants (82%) reported completing the HDQ on a ‘good day’ living with HIV.

### Table 3

| HDQ domain of disability | HDQ domain correlated with… | Factor loading (correlation) (range 0–1) |
|--------------------------|-----------------------------|-----------------------------------------|
| Cognitive symptoms and impairments | Physical | 0.70 |
| Mental-emotional symptoms and impairments | Physical | 0.64 |
| Mental-emotional | Cognition | 0.65 |
| Uncertainty | Physical | 0.57 |
| Mental-emotional | Cognition | 0.51 |
| Difficulties with day-to-day activities | Physical | 0.78 |
| Mental-emotional | Cognition | 0.80 |
| Mental-emotional | Uncertainty | 0.59 |
| Challenges to social inclusion | Physical | 0.55 |
| Mental-emotional | Uncertainty | 0.47 |
| Mental-emotional | Day | 0.47 |
| Mental-emotional | Physical | 0.68 |
| Mental-emotional | Cognition | 0.64 |
| Mental-emotional | Uncertainty | 0.88 |
| Mental-emotional | Day | 0.67 |

Correlations between the latent variables ranged from r=0.47 to r=0.88 (Table 3). A high correlation between mental-emotional challenges and challenges to social inclusion (r=0.88) suggested that these latent variables may not be empirically distinct.^{12} However, these domains of disability were represented by items with similar wording, such as ‘I feel…’ (mental-emotional domain) and ‘I find it hard to…’ (social inclusion domain) which could explain the high correlation and obscure the discriminant nature between these two domains. The correlation between mental and emotional health challenges and challenges to social inclusion also may reflect the influence of mental health on aspects of social inclusion such as employment among people living with HIV.^{21–22} Overall, we concluded six domains of the HDQ represent the dimensions that could explain the high correlation and associations among people living with HIV internationally.^{25–26}

Table 3: Correlations between domains in the HIV Disability Questionnaire (HDQ)

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Implications for practice, research and policy

The HDQ is the first known HIV-specific Disability Questionnaire developed to assess the multidimensional nature of disability experienced by adults living with HIV. The HDQ has the potential to be used in clinical research, practice and policy. Patient-reported outcome measures are important for their ability to detect disability, monitor disease progression and facilitate patient–clinician communication. Further psychometric assessment including test–retest reliability, interpretability and responsiveness, will enable researchers to use the HDQ to document the presence, severity and episodic nature of disability experienced by adults living with HIV. The HDQ may be considered for use by clinicians and AIDS service organisations to assess the profile of disability experienced by their clients. This may help to identify areas to implement programmes, services and interventions to reduce disability experienced by clients with HIV. Universal measurement of disability with the HDQ may facilitate more broad and ongoing tracking of episodic disability trends and evaluation of interventions to inform resource allocation, as well as income and employment programme and policies to ensure optimal care and social inclusion for people living with HIV.

Strengths of our analytical approach included our adherence to the COSMIN guidelines for assessing validity and hypothesis testing, such as providing a clear description of the a priori hypothesised measurement model and goodness of fit criteria; providing details on the distribution of HDQ scores and missingness sample; and describing the correlations between domains. Our study has limitations. First, the HDQ was developed and validated primarily with men in their 40 s who were taking antiretroviral therapy, living with concurrent health conditions and not currently working. Generalisability of these findings to other people living with HIV has not yet been demonstrated. Similarly, the validity of the HDQ in other cultural and developmental contexts is unknown. Second, our goal was to validate the domain structure of the HDQ (not to measure disability) so HDQ scores should be interpreted cautiously. Third, our a priori goodness of fit criteria only was met for the RMSEA. We considered the RMSEA as the primary statistic for overall goodness of model fit because it is recommended for confirmatory factor analysis. Fourth, while our results indicate that dimensions of disability are correlated to comprise the larger construct of disability, the direct relationships between the domains of disability in the HDQ are unknown. Our results provide a measurement model which can serve as the foundation for future structural equation model analyses to determine the relationships between the domains of disability.

Our analysis focused on assessing the construct validity of the domains of the disability scale of the HDQ. Future research will assess the test–retest reliability, interpretability and responsiveness of the HDQ. Additionally, we will consider the measurement properties of the episodic scale. Longitudinally exploring the episodic nature of disability experienced by adults living with HIV is important to consider in future research.

CONCLUSIONS

Results from this confirmatory factor analysis support the validity of the domains of disability in the HDQ when administered to adults living with HIV in Ontario. This is the first known HIV-specific instrument of disability developed from the perspective of adults living with HIV. The HDQ can be used to describe the multidimensional nature of disability experienced by adults living with HIV and lay the foundation for more widespread measurement of disability in HIV clinical practice and research.

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REFERENCES

1. Gaidhane AM, Zahiruddin QS, Waghmare L, et al. Assessing self-care component of activities and participation domain of the international classification of functioning, disability and health (ICF) among people living with HIV/AIDS. AIDS Care 2008;20:1096–104.

2. Weiss JJ, Osorio G, Ryan E, et al. Prevalence and patient awareness of medical comorbidities in an urban AIDS clinic. AIDS Patient Care STDs 2010;24:39–48.

3. Willard S, Holzemer WL, Wantland DJ, et al. Does “asymptomatic” mean without symptoms for those living with HIV infection? AIDS Care 2009;21:322–8.

4. Dray-Spira R, Persoz A, Boufassa F, et al. Exploring disability from the perspective of adults living with HIV/AIDS: development of a conceptual framework. Health Qual Life Outcomes 2004;2:46.

5. Myezwa H, Stewart A, Musenge E, et al. Assessment of HIV-positive in-patients using the International Classification of Functioning, Disability and Health (ICF) at Chris Han Baragwanath Hospital, Johannesburg. Afr J AIDS Res 2009;8:93–105.

6. O’Brien KK, Bayoumi AM, Strike C, et al. Exploring disability from the perspective of adults living with HIV/AIDS: development of a conceptual framework. Health Qual Life Outcomes 2008;6:76.

7. O’Brien KK, Davis AM, Strike C, et al. Putting episodic disability into context: a qualitative study exploring factors that influence disability experienced by adults living with HIV/AIDS. J Int AIDS Soc 2009;12:5.

8. O’Brien KK, Bayoumi AM, King K, et al. Community engagement in health status instrument development: experience with the HIV Disability Questionnaire. Progress in community health partnerships: research, education and action. In press.

9. O’Brien KK, Bayoumi AM, Stratford P, et al. Which dimensions of disability does the HIV Disability Questionnaire (HDQ) measure? A factor analysis. Disabil Rehabil Early Online. 2014:1–9.

10. O’Brien KK, Bayoumi AM, Bereket T, et al. Sensibility assessment of the HIV Disability Questionnaire. Disabil Rehabil 2013;35:566–77.

11. Brown TA. Confirmatory factor analysis for applied research. New York, NY: The Guilford Press, 2006.

12. Muthén LK, Muthén BO. Mplus user’s guide 6th edition. Los Angeles, CA, 1998–2010.

13. Hu LT, Bentler PM. Fit indices in covariance structure modeling: sensitivity to underparameterized model misspecification. Psychol Methods 1998;3:424–53.

14. Hu LT, Bentler PM. Cutoff criteria for fit indexes in covariance structure analysis: conventional criteria versus new alternatives. Struct Equation Model 1999:6:1–55.

15. Rigdon EE. CFI versus RMSEA: a comparison of two fit indexes for structural equation modeling. Struct Equation Model 1996:3:389–79.

16. Kline P. An easy guide to factor analysis. New York: Routledge,1994.

17. IBM Corp. SPSS Statistics for Windows, Version 19.0. Armonk, NY: IBM Corp, 2010.

18. Muthén LK, Muthén BO, Mplus Version 7.0. Los Angeles, CA, 2012.

19. Bollen KA,vars KH. Longitudinal structural equations model analysis with nonnormal data. Am Sociol Rev 1981;46:232–9.

20. Rueda S, Raboud J, Rourke SB, et al. Influence of employment and job security on physical and mental health in adults living with HIV: cross-sectional analysis. Open Med 2012;6:e118–26.

21. O’Brien KK, Hanna S, Gardner S, et al. Validation of the Episodic Disability Framework with adults living with HIV. Disabil Rehabil 2014;36:319–29.

22. Hopcroft L, Bester L, Clement D, et al. My body’s a 50 year-old but my brain is definitely an 85 year-old: exploring the experiences of men ageing with HIV-associated neurocognitive challenges. J Int AIDS Soc 2013;16:18506.

23. Hanass-Hancock J, Nixon SA. The fields of HIV and disability: past, present and future. J Int AIDS Soc 2009;2:3.

24. O’Brien KK, Ibáñez-Carrasco F, Solomon P, et al. Advancing research and practice in HIV and rehabilitation: a framework of new and emerging research priorities in HIV, disability and rehabilitation. Can J Infect Dis Med Microbiol 2014;25:41A.

25. Atkins JH, Rubenstein SL, Sota TL, et al. Impact of social support on cognitive symptom burden in HIV/AIDS. AIDS Care 2010;22:793–802.

26. Carlier SL, Rourke SB, Murji S, et al. Cognitive complaints, depression, medical symptoms, and their association with neuropsychological functioning in HIV infection: a structural equation model analysis. Neuropsychology 2003;17:410–19.

27. Oliveira SA, Basset C. Neurocognitive complaints in HIV-infection and their relationship to depressive symptoms and neuropsychological functioning. J Commun Disabil 2006;10:489–97.

28. Valdés JM, Kotzeva A, Espallargues M, et al. The impact of measuring patient-reported outcomes in clinical practice: a systematic review of the literature. Qual Life Res 2008;17:179–93.

29. Mokkink LB, Terwee CB, Knol DL, et al. The COSMIN checklist for assessing the methodological quality of studies on measurement properties: a clarification of its content. BMJ Med Res Methodol 2010;10:22.

30. de Vet HC, Ader HJ, Terwee CB, et al. Are factor analytical techniques used appropriately in the validation of health status questionnaires? A systematic review on the quality of factor analysis of the SF-36. Qual Life Res 2005;14:1203–18; discussion 1219–1221, 1223–1204.