A comparison of self-report and antiretroviral detection to inform estimates of antiretroviral therapy coverage, viral load suppression and HIV incidence in Kwazulu-Natal, South Africa

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Abstract

Background: Accurately identifying individuals who are on antiretroviral therapy (ART) is important to determine ART coverage and proportion on ART who are virally suppressed. ART is also included in recent infection testing algorithms used to estimate incidence. We compared estimates of ART coverage, viral load suppression rates and HIV incidence using ART self-report and detection of antiretroviral (ARV) drugs and we identified factors associated with discordance between the methods.

Methods: Cross-sectional population-based survey in KwaZulu-Natal, South Africa. Individuals 15–59 years were eligible. Interviews included questions about ARV use. Rapid HIV testing was performed at the participants’ home. Blood specimens were collected for ARV detection, LAg-Avidity HIV incidence testing and viral load quantification in HIV-positive individuals. Multivariate logistic regression models were used to identify socio-demographic covariates associated with discordance between self-reported ART and ARV detection.

Results: Of the 5649 individuals surveyed, 1423 were HIV-positive. Median age was 34 years and 76.3% were women. ART coverage was estimated at 51.4% (95%CI:48.5–54.3), 53.1% (95%CI:50.2–55.9) and 56.1% (95%CI:53.5–58.8) using self-reported ART, ARV detection and both methods combined (classified as ART exposed if ARV detected and/or ART reported) respectively. ART coverage estimates using the 3 methods were fairly similar within sex and age categories except in individuals aged 15–19 years: 33.3% (95%CI:23.3–45.2), 33.8% (95%CI:23.9–45.4%) and 44.3% (95%CI:39.3–46.7) using self-reported ART, ARV detection and both methods combined. Viral suppression below 1000cp/mL in individuals on ART was estimated at 89.8% (95%CI:87.3–91.9), 93.1% (95%CI:91.0–94.8) and 88.7% (95%CI:86.2–90.7) using self-reported ART, ARV detection and both methods combined respectively. HIV incidence was estimated at 1.4 (95%CI:0.8–2.0) new cases/100 person-years when employing no measure of ARV use, 1.1/100PY (95%CI:0.6–1.7) using self-reported ART, and 1.2/100PY (95%CI:0.7–1.7) using ARV detection. In multivariate analyses, individuals aged 15–19 years had a higher risk of discordance on measures of ARV exposure (aOR:9.4; 95%CI:3.9–22.8), while migrants had a lower risk (aOR:0.3; 95%CI:0.1–0.6).

Conclusions: In KwaZulu-Natal, the method of identifying ARV use had little impact on estimates of ART coverage, viral suppression rate and HIV incidence. However, discordant results were more common in younger individuals. This may skew estimates of ART coverage and viral suppression, particularly in adolescent surveys.

Keywords: HIV, ART, Self-report, ARV detection, Antiretroviral coverage, HIV incidence, Viral suppression

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Background
Accurately identifying individuals who are on antiretroviral therapy (ART) in HIV surveys is important to determine ART coverage and the proportion of those on ART who are virally suppressed. In addition, antiretroviral (ARV) use is often included in recent infection testing algorithms (RITAs) used to estimate incidence, to ensure that patients on ART are not misclassified as recently infected.

ART exposure can be identified by including questions on prior and current ART use in survey questionnaires (self-report), or by detection of antiretroviral drugs (ARVs) in blood. Self-report has commonly been used in surveys to identify individuals on ART [1–5] but accuracy cannot be verified without an objective measure. ARV detection in dried blood spots (DBS) is an objective method to identify patients with recent ART exposure, but may not identify individuals on ART who have poor or irregular adherence to ART.

We used data from a population-based HIV survey conducted in Mbongolwane and Eshowe, KwaZulu-Natal (KZN), South Africa [6]. In this analysis we compared estimates of ART coverage and viral load suppression rates when we used self-report and/or ARV detection to identify individuals on ART. We assessed the agreement between the two methods of identifying individuals on ART and the socio-demographic covariates associated with discordance between ART self-report and ARV detection. Finally, we determined the impact of including ART use, as ascertained by these two methods, on HIV incidence estimation.

Methods
Design and population
This study is part of a cross-sectional population-based survey conducted from July to October 2013 that aimed to assess HIV prevalence, incidence and the HIV cascade of care in Mbongolwane and Eshowe, KZN, [6]. Individuals aged 15–59 years old living in the area were eligible for inclusion in the survey.

Procedures
Participants were interviewed at home using a structured questionnaire. Interviews were carried out in an isolated part of the dwelling to preserve confidentiality. We collected socio-demographic information and history of HIV clinical care. The questionnaire included three questions about whether or not the participant was on ART: ‘Have you ever initiated ART, antiretroviral drugs against HIV/AIDS?’, ‘When did you first start antiretroviral drugs?’ ‘Are you still receiving antiretroviral drugs?’ HIV testing was performed at the participants’ home using the Determine Rapid HIV-1/2 Antibody test kit, as a screening test, followed by the Unigold Rapid HIV test kit. HIV-positivity was confirmed by ELISA. Venous blood specimens were collected for single-well limiting antigen avidity enzyme immunoassay (LAg-Avidity EIA) and Nucleic Acid Amplification Testing (NAAT). DBS were prepared at the laboratory to test for the presence of ARVs. We performed qualitative testing for the presence of nevirapine, efavirenz and lopinavir by liquid chromatography tandem mass spectrometry with a limit of quantification of 0.04 micrograms/ml for all drugs. The median time window from last drug intake to a negative detection depends on the drug and ranges from around 24 h for lopinavir up to 7 days for efavirenz and nevirapine [7–9]. Viral load was quantified using a NucliSens EasyQ HIV-1 v2.0 assay from BioMerieux. NAAT testing was performed on 5 member pools using Roche AMPLISCREEN, with positive pools reflexed to individual specimen testing using the Roche CAP/CTM method. HIV-1 genotyping was carried out at the Centre for AIDS Prevention and Research in South Africa (CAPRISA) on samples from participants on ART with viral load higher than 1000 copies/ml, using the TRUGENE HIV-1 Genotyping assay from Siemens Healthcare Diagnostics. Genotypic resistance was interpreted according to the Stanford University algorithm for HIV drug resistance.

Definitions
Self-reported ART: Individuals who reported that they had “ever initiated ART” and were “still receiving ART”.

ARV detected: Nevirapine, efavirenz or lopinavir detected in DBS.

Combined method to identify ART exposure: Categorized as on ART if either ART was self-reported or ARV was detected.

ART discordance: Individuals identified as on ART by one measure but not the other. There were 2 categories of discordance: 1. Individuals who reported being on ART but had no detectable ARV (these likely represent individuals on ART with regular adherence who did not take any ART in the 24 h before, with a longer window for efavirenz). 2. Individuals in whom an ARV was detected who did not report taking ART (these are likely individuals who chose not to disclose being on ART).

ART coverage among all HIV-positive individuals: proportion of HIV-positive individuals on ART among all HIV-positive individuals.

ART coverage among individuals qualifying for ART: proportion of HIV-positive individuals on ART among individuals whether already on ART or qualifying for ART. Criteria for initiation of ART by National Guidelines at the time of the survey were: CD4 < 350cells/μl, pregnant and breastfeeding women.
Viral load suppression rate: viral load below 1000 cp/mL among those identified as being on ART.

Mobility categories: Resident: individuals belonging to the household who had not moved their residency in the previous 10 years. Migrant: individuals belonging to the household who had moved their residency in the previous 10 years. Visitor: individuals not belonging to the household who slept in the household the night before the day of the interview.

Data analyses
Descriptive analyses are presented with 95% confidence intervals (95%CI). Three estimates of ART coverage and viral suppression were calculated using self-reported ART, ARV detection and the combined method to identify ART exposure. HIV incidence was calculated by the method of Kassanjee et al. [10]. The case definition for ‘recent infection’ was: either acute infection (antibody negative but NAAT positive) or recent on the compound rule of LAg-Avidity EIA normalized optical density below 1.5, viral load above 100 cp/mL and not on ART. Three estimates of HIV incidence were calculated: employing no measure of ART use and employing either self-reported ART or ARV detection to identify the individuals not on ART. We estimated Mean Duration of Recent Infection (MDRI) for this RITA at 184 days (95% CI: 159–219), based on a subtype C-specific estimate for the LAg-Avidity EIA and the sensitivity of the screening algorithm [11]. The False Recent Rate (FRR) was estimated in a similar context at 0.2% [12].

Interrater agreement (kappa, $\kappa$) was used to quantify agreement between self-report and ARV detection. Multivariate logistic regressions were used to explore associations between participant characteristics and discordance between self-reported ART and ARV detection with 3 outcomes: (1) ARV detected and not self-reported, (2) ART self-reported and not detected, (3) any ART discordance (ARV detected and not reported or ART reported and not detected). Factors included in the model were: sex (women, men), age (15–19, 20–34, 35–44, 45–59 years), marital status (never married, married/living together, divorced/separated/widowed), level of education (primary or less, secondary or more), area of residence (urban, rural), mobility (resident, migrant, visitor) and employment (employed, not employed). After selecting factors associated with discordance with $p < 0.25$ in the univariate analyses, we applied a step-wise decreasing strategy to select the variables in the multivariate analyses. Sex, age and variables with $p$-value $< 0.10$ were kept in the final model. Data were analyzed using Stata 13 (Stata Corp., College Station, Texas, USA). Incidence analyses were performed using the inctools R package [13].

Results
Description of the survey population
The survey included 5649 individuals of which 1423 were found to be HIV-positive. Median age of HIV positive participants was 34 years (IQR: 27–42) and 1085 (76.3%) were women, 345 (24.3%) lived in couple, 686 (48.2%) had completed at least secondary education, 1142 (80.3%) lived in rural areas, 225 (15.8%) were migrants, 711 (50.0%) were unemployed (Table 1).

ART coverage estimates
ART coverage among all HIV-positive individuals was estimated at 51.4% (95%CI: 48.5–54.3) of HIV-positive individuals using self-reported ART, 53.1% (95%CI: 50.2–55.9) using ARV detection and 56.1% (95%CI: 53.5–58.8) using with both methods combined (Table 2).

Table 1: Socio-demographic characteristics of the HIV-positive participants

|                          | Women        | Men          | Total        |
|--------------------------|--------------|--------------|--------------|
|                          | (N = 1085)   | (N = 338)    | (N = 1423)   |
| Age, years               |              |              |              |
| 15–19                    | 58 (5.4)     | 12 (3.6)     | 70 (4.9)     |
| 20–34                    | 537 (49.5)   | 140 (41.4)   | 677 (52.5)   |
| 35–44                    | 281 (25.9)   | 105 (31.1)   | 386 (27.1)   |
| 45–59                    | 209 (19.3)   | 81 (24.0)    | 290 (20.4)   |
| Marital Status           |              |              |              |
| Never Married            | 753 (69.4)   | 236 (70.0)   | 989 (69.6)   |
| Married/Living Together  | 265 (24.4)   | 80 (23.7)    | 345 (24.3)   |
| Divorced/Separated/Widowed | 67 (6.2)    | 21 (6.2)     | 88 (6.2)     |
| Education                |              |              |              |
| No schooling             | 96 (8.9)     | 31 (9.2)     | 127 (8.9)    |
| Primary                  | 448 (41.3)   | 161 (47.8)   | 609 (42.8)   |
| Secondary                | 513 (47.3)   | 137 (40.7)   | 650 (45.7)   |
| Tertiary                 | 28 (2.6)     | 8 (2.4)      | 36 (2.5)     |
| Place residence          |              |              |              |
| Urban                    | 202 (18.6)   | 79 (23.4)    | 281 (19.8)   |
| Rural                    | 883 (81.4)   | 79 (23.4)    | 1142 (80.3)  |
| Mobility                 |              |              |              |
| Residents                | 876 (80.7)   | 265 (78.4)   | 1141 (80.2)  |
| Migrants                 | 170 (15.7)   | 55 (16.3)    | 225 (15.8)   |
| Visitors                 | 39 (3.6)     | 18 (5.3)     | 57 (4.0)     |
| Occupation               |              |              |              |
| Employed                 | 520 (47.9)   | 192 (56.8)   | 712 (50.0)   |
| Not employed             | 565 (52.1)   | 146 (43.2)   | 711 (50.0)   |
Viral suppression was lower in individuals on ART compared to those not on ART. Among those on ART, 89.8% (95% CI: 87.3–91.9) had viral suppression below 1000 copies/mL, compared to 66.6% (95% CI: 61.5–71.2) among those who reported being on ART but did not report ART or ARVs detected. Thus, in individuals aged 15–19 years, viral suppression was lower among those who reported being on ART than among those in whom ARV was detected: 56.5% (95% CI: 35.6–75.4) vs 87.0% (95% CI: 63.9–95.2).

Resistance tests were performed in the 54 patients who reported being on ART for more than 6 months and had viral load greater than 1000 copies/mL. Among the participants who reported ART, 61.1% (33/54) had at least one resistance mutation while this proportion detected, 25.0% (13/52) compared to participants who did not report ART but had ARV detected, 70.2% (40/57). Among the 30 individuals aged 15–19 years with ART reported or ARVs detected, 8 reported being on ART but ARV were not detected and in 7 individuals ARVs were detected but not reported. Thus, in individuals aged 15–19 years, viral suppression was lower among those who reported being on ART than among those in whom ARV was detected: 56.5% (95% CI: 35.6–75.4) vs 87.0% (95% CI: 63.9–95.2).
increased to 72.4% (21/29) in individuals who had ARVs detected. There were NNRTI resistance mutations in 31/54 (57.4%), NRTI mutations in 25/54 (46.3%) and PI mutations in 5/62 (9.3%) (Additional file 1: Table S1). Among the participants with reported ART and no ARVs detected, 48.0% (12/25) had at least one resistance mutation: 7 to NRTIs, 10 to NNRTIs and 4 to PIs. Among participants taking NNRTIs as per blood test, 74.1% (20/27) had a positive resistance test to any of the drugs in the regimen. The 4 participants taking LPV as per DBS test did not have any resistance mutation detected to PI.

HIV incidence estimates
HIV incidence was estimated at 1.4 new cases per 100PY (95%CI: 0.8–2.0) when employing no measure of ARV use, 1.1/100PY (95%CI: 0.6–1.7) using reported ART, and 1.2/100PY (95%CI: 0.7–1.7) using ARV detection (Table 4). Incidence estimates were similar using the 2 methods, with differences in point estimates constituting a small fraction of the span of confidence intervals. This analysis did not provide strong evidence for preferring one measure over the other.

We estimated the maximum improvement in the precision of the incidence estimate that the inclusion of any ARV use measure in the RITA may achieve. Assuming an FRR of 0.5% for the RITA without an ARV measure (substantially higher than our best estimate of 0.2%), the complete elimination of residual false recency, and no impact on MDRI, the relative standard error of the overall incidence estimate would decline from 20.8% to 17.9%. The potential benefit is therefore modest at best.

Discordance between self-report and ARV detection
Of the 1358 individuals with both self-report and ARV detection information, 655 (48.2%) had ARVs detected and reported being on ART; 58 (4.3%) had ARVs detected but did not report being on ART (non-disclosure) and 52 (3.8%) reported being on ART but did not have ARVs detected (non-adherence) (Additional file 2: Table S2).

There was non-disclosure of ART use in 8.1% (95%CI: 6.3–10.4) of the participants. Non-disclosure was higher at ages younger: 31.8% (95%CI: 15.5–54.3), 11.5% (95%CI: 7.9–16.3), 7.2% (95%CI: 4.6–11.1), 2.1% (95%CI: 0.8–5.4), in individuals aged 15–19, 20–34, 35–44 and 45–59 years respectively (Chi² test for trend \( p < 0.001 \)).

Conversely, 7.4% (95%CI: 5.7–9.5) of the participants reporting ART did not have ARVs detected, suggesting poor recent adherence to ART. The proportion with discordance was higher in individuals 15–19 years compared to individuals in the other age groups: 34.8% (95%CI: 18.0–56.5) vs 7.4% (95%CI: 4.7–11.6), 8.3% (95%CI: 5.2–12.9) and 2.7% (95%CI: 1.1–6.1) in individuals aged 20–34, 35–44 and 45–59 years respectively (Chi² test for trend \( p < 0.001 \)).

Among individuals who reported being on ART, median time since ART initiation was longer for individuals who had ARVs detected compared to those for whom ARVs were not detected: 33.8 vs 13.9 months.

Agreement between ART self-reported and ARV detection was 91.9% (kappa = 0.84, 95%CI: 0.81–0.87). There was no difference in the agreement between self-report and ARV detection among male and female participants, 92.0% (kappa = 0.84, 95%CI: 0.78–0.90) and 91.9% (kappa = 0.84, 95%CI: 0.80–0.87) respectively. Agreement between the 2 methods of determining ART exposure was lower in individuals aged 15–19 years: 77.6% (kappa = 0.50, 95%CI: 0.28–0.72) vs 92.6% (kappa = 0.85, 95%CI: 0.80–0.89), 89.5% (kappa = 0.76, 95%CI: 0.69–0.83) and 96.8% (kappa = 0.93, 95%CI: 0.88–0.97) in individuals aged 20–34, 35–44 and 45–59 years respectively.

In multivariate logistic regressions, younger individuals were at higher risk of discordant results (Table 5). Individuals younger than 45 years and particularly those aged 15–19 years had the highest risk of all types of discordance (Table 5) while migrants were less likely to have discordant results. Individuals with an education of lower than secondary school who had reported ART were more likely to test negative for ARVs than those with higher levels of education.

Discussion
We found that estimates of antiretroviral coverage were generally similar when using self-report and ARV detection to identify individuals on ART. Overall, there was good concordance between self-report of ART use and detection of ARVs in this study population. However, discordance was higher in individuals aged 15–19 years suggesting higher proportions of poor recent adherence and non-disclosure in young people. As a result, estimates of ART coverage in this group were sensitive to the measure of ART exposure used. Individuals who had moved residence in the last ten years were less likely to have discordance than permanent residents. The reason for this is unclear and qualitative research is needed to explore
differences regarding barriers to both disclosure and adherence in these populations.

Rates of viral suppression were slightly higher using ARV detection to identify those on ART. This finding is expected since those with ARVs detected are likely to be currently adherent to therapy. On the other hand, individuals reporting ART with no detectable ARV are likely to be poorly adherent and viral suppression was very low in this group. Median time since ART initiation was longer for individuals who had ARVs detected which may be due to patients in this cohort struggling with adherence particularly in the period immediately after ART commencement, or because poorly adherent patients default therapy. Estimates of suppression in adolescents ranged widely depending on the method of identifying ART exposure. The prevalence of resistance among individuals on ART was also sensitive to the method used to determine the denominator (higher for ARV detection).

We found that the method of identifying ARV use had little impact on incidence estimates. Including ARV use in the RITA had only a modest impact on precision of incidence estimates.

Measures of ARV use are frequently included in recent infection testing algorithms (RITAs) on the intuition that this would reduce the false recent rate (FRR) and improve the precision of the incidence estimates. Context-specific FRR estimates for an algorithm consisting of LAg-Avidity EIA and viral load in a setting similar to the study population have been estimated as low as 0.2% [12]. Furthermore, benchmarking data are not available for quantifying the impact of including a measure of ARV use on the mean duration of recent infection (MDRI) and FRR. It is therefore questionable whether
such measures should be included in RITAs that already contain a viral load measurement. However, with early treatment becoming increasingly common and with higher treatment coverage, larger numbers of individuals on treatment but with detectable viral loads, continuing to study the impact of ARV ascertainment on RITA properties will be essential.

Based on these data, either method of ART exposure ascertainment could be used for estimation of ART coverage and viral suppression for the population as a whole. However, we found that discordant results were more common in younger individuals (9 folds higher risk). This may skew estimates of ART coverage and viral suppression, particularly in adolescent surveys. Based on our data, both non-disclosure and poor recent adherence are common in this group.

Use of a combination of methods may be useful for particular study questions: for example this would allow for exploration of adherence patterns within a survey, and for exploration of the relationship between adherence, viral suppression and viral resistance. Including ARV detection also allows for characterization of patterns of ART regimen use within the surveyed population.

This study has limitations. Adolescents formed a small proportion of individuals in the sample, and we therefore lacked power for precise estimates in this group. We did not have access to hospital and dispensing records, which would provide an objective confirmation of long-term ART.

Conclusions
In conclusion, in KZN the method of identifying ARV use had little impact on estimates of ART coverage, viral suppression rate and HIV incidence. The method for determining ART exposure should be selected based on the primary study questions. Surveys focused on young individuals, particularly adolescents, should consider using the combination of self-report and ARV detection, so that estimates are not skewed by non-disclosure or poor adherence.

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Availability of data and materials
The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

Authors’ contributions
HH conceived and designed the study. HH, FS and KC drafted the manuscript. HH and JBF performed the statistical analyses. EG performed the HIV incidence statistical analyses and interpreted the results. KC interpreted the ARV detection results. RG and GVC critically revised the manuscript. All authors reviewed, revised and approved the final paper.

Ethics approval and consent to participate
The study protocol was approved by the University of Cape Town Human Research Ethics Committee (HREC), the Health Research Committee of the Health Research and Knowledge Management Unit of KZN DOH, and the Comité de Protection de Personnes de Paris in France. Written consent for study participation was obtained from all participants prior to inclusion in the study.

Consent for publication
Not applicable

Competing interests
The authors declare that they have no competing interests

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Additional files

Additional file 1: Table S1. Resistance profile in individuals who reported being on ART for more than 6 months and with viral load $21000$ cp/mL. (DOCX 14 kb)

Additional file 2: Table S2. ART self-report and ARV detection by sex and age. (DOCX 15 kb)

Abbreviations
ART: Antiretroviral therapy (ART); ARV: Antiretroviral (ARV); ARVs: Antiretroviral drugs (ARVs); DBS: Dried blood spots (DBS); KZN: KwaZulu-Natal (KZN); LAg-Adv: Limiting antigen avidity enzyme immunoassay; NAAT: Nucleic Acid Amplification Testing (NAAT); RITAs: Recent infection testing algorithms.
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