How accurately do routinely reported HIV viral load suppression proportions reflect progress towards the 90-90-90 target in the population on antiretroviral treatment in Khayelitsha, South Africa?

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Background. The Joint United Nations Programme on HIV/AIDS (UNAIDS) third 90-90-90 target requires 90% of patients on antiretroviral treatment (ART) to be virally suppressed (<1 000 copies/mL). In Khayelitsha, Cape Town, South Africa viral load (VL) suppression of <400 copies/mL was reported as 89% in 2016, but only 56% of patients had a result recorded in routine data. We conceived a VL ‘cascade’ to represent the steps required for an expected VL to be reported as complete in routine data and thus contribute to reported VL suppression: among those for whom a VL is ‘expected’, a sample must be collected and tested (‘done’), a result must be ‘filed’ in the patient folder, ‘noted’ by a clinician and electronically ‘captured’. The low reported completion suggested gaps along the VL cascade and cast doubt on the validity of reported suppression.

Objectives. To assess the validity of routinely reported VL suppression and identify barriers to VL completion.

Methods. A retrospective cohort study between 1 July 2015 and 30 June 2016, which included all Khayelitsha patients receiving ART, with a routine VL expected, was conducted. We obtained data routinely captured on site and VL data from the laboratory system. A sample of 1 035 patient folders was reviewed. VL suppression was calculated using laboratory data, including all tests done, and compared with reported suppression based on on-site captured electronic data. Successful progression through each step on the VL cascade was estimated. We used logistic regression to identify factors associated with laboratory data and reported VL testing.

Results. Of 22 991 patients for whom a routine VL test was due, 84% were done, 79% filed, 76% noted and 55% captured. Using all laboratory data, VL suppression was estimated as 82%, 87%, 89% and 91% at the 50, 200, 400 and 1 000 copies/mL thresholds, respectively, but reported suppression using captured results was 80%, 86%, 88% and 89% at those thresholds. Routine VL testing is more likely to be done in children <15 years old (adjusted odds ratio (aOR) 1.89, 95% confidence interval (CI) 1.45 - 2.48) and pregnant women (aOR 1.90, 95% CI 1.28 - 2.81) than in men, adjusted for facility.

Conclusions. Despite a low reported completion, VL testing completion was high. Reported suppression using captured data was similar to suppression calculated using all laboratory data, which provided an accurate measure of progress towards the 90-90-90 target. More work is needed to reach the 16% of patients missed by routine testing.

The Joint United Nations Programme on HIV/AIDS (UNAIDS) 90-90-90 strategy ambitiously aspires to have 90% of people living with HIV (PLWH) know their status, 90% to receive sustained antiretroviral treatment (ART) and 90% to achieve viral load (VL) suppression by 2020.[1] Many studies in different settings have assessed progress compared with these targets, with significant heterogeneity in data sources, outcome definitions, methods and results.[2]

The Khayelitsha cohort in Cape Town, South Africa (SA) is one of the oldest and largest HIV cohorts in the country. Free public sector ART was initially rolled out in SA in 2001, in partnership with Médecins Sans Frontières.[3] VL testing is the gold standard in HIV treatment monitoring.[4] Unusually for a low-resource setting, VL monitoring has been widely used in Khayelitsha since programme inception in 2001. With ~22 000 patients receiving ART and in care at provincial facilities in 2016, Khayelitsha reported almost 89% suppression in quarterly reports; however, there was only 56% completion of routine VL tests indicated by national guidelines (R Holtman, Western Cape Department of Health – personal communication, 2017). This low reported completion does not necessarily imply low actual completion, as it may be the result of incompleteness in routine data capturing.

For this study, the concept of a VL ‘cascade’ was introduced to represent the steps required for an ‘expected’ VL to be reported as complete in routine data and thus contribute to reported VL suppression in routine quarterly reports (Fig. 1). As a first step, a blood sample was taken in patients in whom a routine VL was expected (step 1). This blood sample must then be processed by the laboratory (‘done’) (step 2), the test result printed, couriered to the facility, sorted and distributed to the appropriate registry and ‘filed’ by a clerk in the patient folder (step 3), the result ‘noted’ by a clinician in the visit summary paper stationery at the next clinical assessment (step 4) and electronically ‘captured’ by a clerk onto the primary healthcare infor...
mation system (PHCIS) (step 5). Finally, as routine quarterly ART data report on facility-based ART-naive cohorts, VL results for patients who initiated outside the facility at which they were being cared for were not included in reports (step 6).

The low reported completion for Khayelitsha in 2016 therefore raised two important questions: was the high reported suppression proportion a valid measure of progress towards the third 90, and what were the barriers to testing and recording of VL results in the context of widely available laboratory testing? This study aimed to answer these questions by describing a VL cascade from expected to reported, and estimating success and failure at each step on the cascade.

Methods

Setting

The Khayelitsha cohort has been described in detail elsewhere.[8] Briefly, Khayelitsha is the largest township in Cape Town, with a high burden of HIV. The provincial government initiated the provision of free ART services at primary healthcare facilities in 2001.[9] Eligibility criteria have evolved in response to World Health Organization (WHO) recommendations, with universal test and treat adopted nationally in September 2016.[7]

Study design

A retrospective cohort was constructed, including all patients on ART and in care at provincial healthcare facilities in Khayelitsha, with a routine VL expected during 1 July 2015 - 30 June 2016.

Population

In the ART programme in SA, routine quarterly reports follow naïve cohorts while in care at their initiating facility. Patients who die, transfer out or are lost to follow-up (defined as 90 days without medication in hand) exit the cohort. Even if they return to care elsewhere, they are no longer included in the reported suppression proportion. By contrast, the intended denominator for the ‘third 90’ includes all patients receiving ART, regardless of where they access care. This study included all patients receiving ART in Khayelitsha during the study period, irrespective of where they initiated ART, but distinguished between patients who are part of the reporting cohort (the naïve cohort at the reporting facility) and those who are not.

Outcomes

Suppression

National guidelines define VL suppression as <400 copies/mL. While the UNAIDS 90-90-90 document does not explicitly define a threshold, it is generally taken to be <1 000 copies/mL, although various thresholds have been used in different settings.[10] To assess the effect of using different thresholds and facilitate comparability across studies, we reported at multiple thresholds.

Completion

Provincial guidelines recommend routine VL testing for all ART patients at 4 months, again at 12 months and annually thereafter.[11] More frequent testing is recommended after an unsuppressed VL or regimen switch, in HIV-exposed infants, HIV-positive children <5 years of age and pregnant breastfeeding women. To calculate routine VL completion, we calculated patient-specific window periods around each expected VL due date based on the patient’s ART initiation date, using the same algorithms as those in routine reports (Fig. 2). The algorithm depicted in Fig. 2 is the same for all patients, despite the different clinical guidelines alluded to above. According to this algorithm, where >1 VL was taken within an allocation window, the VL on the date closest to the due date was used and others were excluded from reports.

Data collection

We used 3 data sources to assess completion at each step of the cascade: PHCIS, the Provincial Health Data Centre (PHDC) and the physical patient folder containing all paper records at the facility (Fig. 1). To assess completion at steps 0, 2 and 5 on
the cascade, data were requested from PHCIS and PHDC. PHCIS contains routine data digitised by clerks on site. PHDC contains all data from the National Health Laboratory Service (NHLS), which performs all VL testing for patients in care at provincial facilities in Khayelitsha. All VL testing during the study period was performed by the NHLS using an Ampliprep/COBAS TaqMan HIV-1 Test v2 (Roche, Switzerland). Step 1 could not be directly assessed, e.g. if a test was not registered by the laboratory system owing to incorrect or illegible request forms, it would not be counted as done at step 2, despite a blood sample having been taken within the window period and sent to the laboratory. Completion at steps 3 and 4 was assessed by physical folder review, for which folders were drawn on site at facilities and some data captured directly to a stand-alone encrypted electronic database. Routine patient data in all sources (physical folder, PHCIS, NHLS and PHDC) were captured and linked with the unique patient identifier used across the public health platform in Western Cape Province.191

Data analysis
The VL cascade is simplified insofar as it reports the furthest step a VL reached on the cascade and assumes that the VL successfully reached all previous steps. Only VLs done (step 2) but not captured (step 5) were sampled for physical folder review. The folder review then generated proportions of the sample, which were filed (step 3) and noted (step 4). These proportions (of the sample) could then be added to the proportion captured (of the total) to generate the proportions filed and noted on the cascade.

Statistical analyses were performed using Stata 14 (StataCorp., USA). The sample of folders selected for review was prepared using the simple random sample function, clustered on healthcare facility.

A logistic regression model was built to investigate the effect of facility, age category and pregnancy status on VL testing completion (done) and inclusion in routine reports (reported). Healthcare facility was included to account for different operational practices at different facilities. Age category and pregnancy status were included because different models of care are provided for children and pregnant women and different clinical guidelines are followed. Adjusted odds ratios (aORs) with 95% confidence intervals (CIs) were estimated using this model.

Ethical approval
The research was approved by the Faculty of Health Sciences Human Research Ethics Committee at the University of Cape Town (ref. no. HREC: 270/2017).

Results
There were 22 991 patients receiving ART and in care at provincial healthcare facilities in Khayelitsha between 1 July 2015 and 30 June 2016 with a VL expected, of whom 18 450 (84%) had a VL performed within their window period. Of these, 11 100 (60%) were included in the reported suppression proportion in routine data: 1 790 (10%) patients were excluded because they were not part of the naive cohort at the reporting facility, and 5 560 (30%) were excluded because they were not captured (step 5). Of those not captured, 1 035 physical folders were reviewed, and completion at steps on the cascade was estimated to be 84% done (VL sample taken and tested), 79% filed, 76% noted by a clinician, and 55% captured electronically.

Actual completion (done) and reported completion (reported) varied by facility, age category and pregnancy status (Table 1). Expected VLs were less than half as likely to be done at facility C than at facility A (aOR 0.46, 95% CI 0.42 - 0.51). Expected VLs at facility C were also less likely to be included in the reported suppression proportion (aOR 0.74, 95% CI 0.69 - 0.78). Children <15 years of age were more likely than men to have a routine VL done (aOR 1.89, 95% CI 1.45 - 2.48). Despite pregnant women being almost twice as likely as men to have a VL test done within the routine window period, their results were 0.60 times as likely to be included in reported suppression proportions (aOR 1.90, 95% CI 1.28 - 2.81 and aOR 0.60, 95% CI 0.47 - 0.77, respectively). The inclusion of duration on ART in the model made no meaningful difference to the effect estimates in the model (Supplementary Table 1). *

Suppressed VLs of <400 copies/mL were slightly less likely to be included in reported suppression proportions (aOR 0.70, 95% CI 0.63 - 0.77). Actual VL suppression among all the VLs done, calculated using laboratory data, was 82%, 87%, 89% and 91% at the 50, 200, 400 and 1 000 copies/mL thresholds, respectively, but reported suppression would have been 80%, 86%, 88% and 89% at those same thresholds.

Discussion
The reported VL suppression proportion of <400 copies/mL from routine data (89%) differed by only 1% when all VL results from the laboratory were included, despite only 60% contributing to reported suppression. This confirms that Khayelitsha is very close to achieving the third 90 among those with available results. However, taking into consideration the 16% of patients not tested, the proportion of patients with confirmed suppression of <400 copies/mL declined to 75% of those receiving ART and in care. This is concerning, as it implies that even as we successfully increase ART coverage, the proportion of patients on ART who may be at risk of transmitting HIV remains between 10% and 25%.

Reported completion proportions were low, mostly owing to failure to test patients (16%) and failure to capture VL results electronically (21%). A trial is currently being performed of the inclusion of electronically imported test results from laboratory systems in routine quarterly reports in the Western Cape and would obviate the need for manual capturing of the results into PHCIS. However, the advantage of manual capturing from the folder was that it allowed for indirect monitoring of other steps in the VL cascade, as the presence of a VL result in routine data suggested successful progress through previous steps. Specifically, it implied that the result was available to the clinician at the next clinical assessment, as the clinician had to note it in the stationary for a clerk to capture electronically. Importing results will improve the completeness of data in routine reports, but it will no longer be possible to make inferences regarding intermediate steps on the VL cascade from routine data. Reassuringly, there was relatively little loss on the cascade between VL being done (step 2), filed (step 3) and noted (step 4), implying that results from VL tests that were done were usually available to clinicians at the next clinical assessment.

Our results suggest that if electronic import of test results from laboratory systems is successfully implemented, the reported VL completion proportion will increase substantially, but there will be little effect on the reported suppression proportion. Importantly, this study only assessed whether clinicians noted the VL results in the folder; it did not attempt to assess the effective use of VL results for clinical decision-making. Further research in this important area is required.

The variation in actual completion between facilities based on laboratory data suggests facility-specific challenges that require further investigation. The variation by pregnancy status and age category was expected, as clinical guidelines recommend more frequent testing for pregnant women and children.

The under-representation of pregnant women in the reported suppression proportion may be due to increased mobility during pregnancy because of movement between integrated maternal ART
Despite low reported VL completion, actual VL testing completion was high. The study confirmed the high levels of suppression that are routinely reported. More work is needed to reach the 16% of patients missed by routine testing. Most VL results were accessed and noted by clinicians, and further research is necessary to assess how effectively these results are used in clinical decision-making.

### Table 1. Patient characteristics, outcomes and adjusted odds ratios

<table>
<thead>
<tr>
<th>Facility</th>
<th>Expected, n (%)</th>
<th>Done, n (%)</th>
<th>aOR (95% CI)</th>
<th>Reported, n (%)</th>
<th>aOR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Facility A</td>
<td>7 129 (32)</td>
<td>6 313 (34)</td>
<td>1.00</td>
<td>3 815 (34)</td>
<td>1.00</td>
</tr>
<tr>
<td>Facility B</td>
<td>5 916 (27)</td>
<td>5 130 (28)</td>
<td>0.84 (0.76 - 0.94)</td>
<td>3 181 (29)</td>
<td>1.01 (0.94 - 1.08)</td>
</tr>
<tr>
<td>Facility C</td>
<td>8 946 (41)</td>
<td>7 007 (38)</td>
<td>0.46 (0.42 - 0.51)</td>
<td>4 104 (37)</td>
<td>0.74 (0.69 - 0.78)</td>
</tr>
<tr>
<td>Men</td>
<td>6 097 (28)</td>
<td>5 012 (27)</td>
<td>1.00</td>
<td>3 138 (28)</td>
<td>1.00</td>
</tr>
<tr>
<td>Non-pregnant women</td>
<td>15 011 (68)</td>
<td>12 649 (69)</td>
<td>1.17 (1.08 - 1.27)</td>
<td>7 546 (68)</td>
<td>0.96 (0.90 - 1.01)</td>
</tr>
<tr>
<td>Pregnant women</td>
<td>275 (1)</td>
<td>246 (1)</td>
<td>1.90 (1.28 - 2.81)</td>
<td>107 (1)</td>
<td>0.60 (0.47 - 0.77)</td>
</tr>
<tr>
<td>Children &lt;15 years</td>
<td>608 (3)</td>
<td>543 (3)</td>
<td>1.89 (1.45 - 2.48)</td>
<td>309 (3)</td>
<td>0.99 (0.84 - 1.17)</td>
</tr>
<tr>
<td>Years on ART, median (IQR)</td>
<td>4 (2 - 7)</td>
<td>4 (2 - 7)</td>
<td>1.09 (1.08 - 1.11)</td>
<td>4 (2 - 7)</td>
<td>1.07 (1.06 - 1.08)</td>
</tr>
</tbody>
</table>

aOR = adjusted odds ratio; CI = confidence interval; ART = antiretroviral treatment; IQR = interquartile range.


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