The cascade of HIV care in British Columbia, Canada, 1996–2011: a population-based retrospective cohort study

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Summary

Background The cascade of HIV care has become a focal point for implementation efforts to maximise the individual and public health benefits of antiretroviral therapy. We aimed to characterise longitudinal changes in engagement with the cascade of HIV care in British Columbia, Canada, from 1996 to 2011.

Methods We used estimates of provincial HIV prevalence from the Public Health Agency of Canada and linked provincial population-level data to define, longitudinally, the numbers of individuals in each of the eight stages of the cascade of HIV care (HIV infected, diagnosed, linked to HIV care, retained in HIV care, highly active antiretroviral therapy (HAART) indicated, on HAART, adherent to HAART, and virologically suppressed) in British Columbia from 1996 to 2011. We used sensitivity analyses to determine the sensitivity of cascade-stage counts to variations in their definitions.

Findings 13140 people were classified as diagnosed with HIV/AIDS in British Columbia during the study period. We noted substantial improvements over time in the proportions of individuals at each stage of the cascade of care. Based on prevalence estimates, the proportion of unidentified HIV-positive individuals decreased from 49.0% (estimated range 36.2–57.5%) in 1996 to 29.0% (11.6–40.7%) in 2011, and the proportion of HIV-positive people with viral suppression reached 34.6% (29.0–43.1%) in 2011.

Interpretation Careful mapping of the cascade of care is crucial to understanding what further efforts are needed to maximise the beneficial effects of available interventions and so inform efforts to contain the spread of HIV/AIDS.

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HIV prevalence for British Columbia from the Public Health Agency of Canada, derived from a multiple-method approach based on back-calculation from HIV/AIDS surveillance data and other sources. Estimates from the Public Health Agency of Canada preceded the construction of the linked database used in this study.

We used a series of linked provincial datasets (comprising the linked database of the STOP HIV/AIDS initiative) to estimate the number of identified HIV-positive individuals in the various stages of the cascade of HIV care (HIV infected, diagnosed, linked to HIV care, retained in HIV care, HAART indicated, on HAART, adherent to HAART, and virologically suppressed). The BC Centre for Disease Control is the provincial agency that centralises all HIV testing data and receives reports of new HIV diagnoses from the British Columbia Public Health Microbiology and Reference Laboratory, which does all confirmatory testing in the province. Furthermore, mandatory HIV reporting legislation has been in place in British Columbia since 2003; individuals can choose to have their identifiable information suppressed in HIV case reports to the Public Health Microbiology and Reference Laboratory (non-nominal vs nominal reporting). For individuals aged 18 months or older, the BC Centre for Disease Control uses a screening test (ELISA) to detect HIV antibodies, with HIV diagnosis confirmed on the basis of a reactive western blot or nucleic acid amplification test.

We used data for plasma viral load, CD4 cell count testing, and HAART use from the BC Centre for Excellence in HIV/AIDS population-based registries. The BC Centre for Excellence is the agency that centrally distributes all antiretroviral drugs in the province. It maintains comprehensive clinical guidelines for the management of HIV/AIDS, which have remained consistent with those published by the International AIDS Society (IAS)-USA every 2 years since 1996. All measurements of plasma viral load in British Columbia are done under the auspices of the BC Centre for Excellence at the virology laboratory of St Paul’s Hospital (Vancouver, BC), thus 100% of data for plasma viral load are captured. Additionally, an estimated 80% of all

### Panel 1: Operational definitions for the eight stages of the cascade of HIV care

**HIV infected**
Based on HIV prevalence estimates reported by the Public Health Agency of Canada

**HIV diagnosed**
Defined as the first instance of any one of:
- a confirmed HIV-positive test
- detectable plasma viral load
- an HIV-related MSP billing or hospital admission
- a reported AIDS-defining illness
- dispensation of antiretroviral therapy

**Linked to HIV care**
Among HIV-diagnosed individuals, defined as:
- the first instance of an HIV-related service after HIV diagnosis (for individuals with confirmed HIV test)
- or the first instance of an HIV-related service ≥ 30 days after derived HIV diagnosis date (for individuals with no confirmed HIV test)

**Retained in HIV care**
Among individuals linked to HIV care, defined as:
- HIV-related physician visits or diagnostic tests (CD4 cell count or plasma viral load test) ≥ 3 months apart within the calendar year
- or at least two antiretroviral drug dispensations ≥ 3 months apart, within the calendar year

**HAART indicated**
Among individuals retained in HIV care but not currently on HAART, defined as meeting the primary or secondary IAS-USA initiation criteria within the calendar year:
- 1996: CD4 count < 500 cells per μL, plasma viral load ≥ 30 000 copies per mL, or AIDS-defining illness
- 1997–99: plasma viral load > 5000 copies per mL or AIDS-defining illness
- 2000–01: CD4 count < 500 cells per mL, plasma viral load ≥ 30 000 copies per mL, or AIDS-defining illness
- 2002–07: CD4 count ≥ 200 cells per μL, or AIDS-defining illness
- 2008–09: CD4 count ≥ 350 cells per μL, or AIDS-defining illness
- 2010–11: CD4 count ≥ 500 cells per μL, or AIDS-defining illness

**On HAART**
Among individuals with HAART indicated, defined as receiving at least two antiretroviral drug dispersions ≥ 3 months apart, within the calendar year

**Adherent to HAART**
Among individuals on HAART, defined as having at least 80% adherence in the calendar year, or from the point of antiretroviral initiation for those who began treatment within the calendar year

**Virologically suppressed**
Among individuals adherent to HAART, defined as having at least one episode (≥ 3 months) with an undetectable plasma viral load within the calendar year

MSP=medical services plan. HAART=highly active antiretroviral therapy.
IAS=International AIDS Society. *Unpublished data (Archibald C, Public Health Agency of Canada, personal communication). ‡Based on plasma viral load testing technology available at the time of measurement. Virological suppression thresholds: < 500 copies per mL for 1996, < 400 copies per mL for 1997–98, and < 50 copies per mL for 1999–2011. §Refers to the number of days of drugs dispensed, divided by the total number of days in care.
CD4 cell count measurements done in the province were captured in the BC Centre for Excellence data.\textsuperscript{20}

We supplemented these data with the medical services plan physician billing database, which captures all fee-for-service care in the province, including HIV-related physician visits and other services; the provincial discharge abstract database, which records inpatient care; the British Columbia PharmaNet database, which captures all non-antiretroviral drug dispensations (used to assess administrative loss to follow-up); and the British Columbia Vital Statistics database.

Linkage and preparation of the de-identified individual-level database was facilitated by the British Columbia Ministry of Health, and fully described in a previous report.\textsuperscript{21} We excluded individuals from the cascade of care after death or administrative loss to follow-up, defined as having no record of death and no health administrative records from any of the linked databases for a period of at least 18 months before the end of study follow-up (March 31, 2012). Intermittent losses to follow-up were therefore not excluded by this definition; any return to care within 18 months of the conclusion of follow-up would entail continued inclusion in the cohort.

The BC Centre for Excellence received ethical approval to do this study from the University of British Columbia ethics review committee at St Paul’s Hospital, Providence Health Care site (P05-123). The programme also conforms to British Columbia’s Freedom of Information and Protection of Privacy Act.

Definitions

Whenever possible, our definitions for the eight stages of the cascade of HIV care (panel 1) followed or were adapted from evidence-based standards.\textsuperscript{22–24} HIV infection, diagnosis, and linkage to care were all fixed classifications. Once infected, diagnosed, or linked to care, an individual is counted as such for each subsequent calendar year until death or administrative loss to follow-up. Individual classifications in each of the subsequent stages of care varied over time: from one calendar year to another, individuals can be lost to care after a period of retention, become ineligible for HAART, drop out of HAART, or become non-adherent and not virologically suppressed after periods of stable, suppressive treatment.

The denominator in each step of the cascade is the sum of the preceding stage—ie, the number of individuals with suppressed plasma viral load is calculated as a proportion of the number who are adherent to HAART, the number of individuals adherent to HAART is calculated as a proportion of the number on HAART, and so on.

Because cascade-stage definitions are bound to differ across settings on the basis of data availability and differences in data-generating processes, we did sensitivity analyses on selected stage classifications to assess the effect of our definitions on the results. We did sensitivity analyses on the specific definitions of linkage to HIV care, retention in HIV care, on HAART, and viral suppression. The sensitivity analysis for definitions of viral suppression also assessed an alternative definition of the denominator for this stage that included all individuals with at least one measurement of plasma viral load within the calendar year.

Finally, although viral suppression is the most relevant endpoint for individual health benefits derived from treatment, a dose-response relation between plasma viral load and the risk of HIV transmission suggests that public health benefits from treatment may also extend to decreasing the likelihood of HIV transmission.
health benefits extend beyond full viral suppression.\textsuperscript{9,25} We therefore present data for aggregate plasma viral load for the population of individuals who received at least one plasma viral load test in each calendar year throughout the study period. We used the highest annual measurement of plasma viral load for each individual in this analysis to provide a conservative estimate of aggregate viral load for those with a recorded measurement, thus probably overestimating the aggregate amount of virus at any point in time during the calendar year.

**Role of the funding source**
The sponsor of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the report. BN had full access to all the data in the study and JSGM had final responsibility for the decision to submit for publication. The British Columbia Ministry of Health facilitated access to components of the linked database.

**Results**
Estimates of annual HIV prevalence, along with empirically derived counts of diagnosed cases are presented in figure 1. We estimated the remaining stages of the cascade of HIV care by use of linked individual data for the population in British Columbia (figure 2). \textit{13140} people were classified as diagnosed with HIV/AIDS in British Columbia during the study period. The proportion of HIV-positive individuals diagnosed with either a positive HIV test or otherwise identified as HIV-positive increased from \textit{51·0%} in 1996 to \textit{71·0%} in 2011. Although the proportion of HIV-positive individuals linked to HIV care was \textit{4·1–9·6%} less than the proportion diagnosed throughout the study period, retention in HIV care lagged far behind linkage, reaching \textit{80·5%} of those diagnosed in 2011. The numbers of individuals indicated for and accessing HAART were close to those for retention in HIV care, apart from during 2000–06, when guidelines for initiation of HAART were changed such

![Figure 3: Changes in leakage from the cascade of HIV care](image-url)

Shaded region in (A) represents HIV prevalence range estimates from the Public Health Agency of Canada (Archibald C, Public Health Agency of Canada, personal communication). HAART=highly active antiretroviral therapy.
that treatment was only indicated for patients with CD4 counts of less than 200 cells per μL. Finally, the proportion of HIV-positive individuals with viral suppression increased from 0·7% in 1996 to 34·6% (estimated range 29·0–43·1%) in 2011, with steep increases between 1996 and 2000 (from 0·7% to 13·7%) and between 2003 and 2011 (from 16·5% to 34·6%). The appendix provides absolute numbers for each stage of the cascade.

The proportion of people infected but undiagnosed fell from 49·0% (estimated range 36·2–57·5%) to 29·0% (11·6–40·7) during the study period (figure 3); during the same period the proportion of individuals linked to but not retained in care remained fairly constant at 20·0%. Among people on HAART, the proportion not adherent decreased from 24·0% in 2003 to 13·4% in 2011. The greatest gains were realised in viral suppression—the proportion of people adherent but not virologically suppressed decreased from 95·2% in 1996 to 21·6% in 2011.

Sensitivity analyses revealed some differences dependent on cascade definitions (figures 4, 5). Excluding CD4 cell counts and plasma viral load testing from definitions of linkage and retention in HIV care resulted in figures up to 18 percentage points lower than the baseline definitions. Separating HIV-related physician visits from other fee-for-service billings captured in the medical services plan dataset made little difference in proportions for linkage and retention. Our more conservative on-HAART classification, which required at least two dispensations at least 3 months apart, resulted in a difference of nearly 20 percentage points in positive classification compared with the most liberal definition of any antiretroviral dispensation within the calendar year; however, this difference decreased to less than five percentage points in 2011.

Classifications of viral suppression were sensitive to the definitions used. Our most conservative definition, which

**Figure 4:** Sensitivity analyses for cascade-stage definitions

Viral suppression thresholds: <500 copies per ml for 1996, <400 copies per ml for 1997–98, and <50 copies per ml for 1999–2011. HAART=highly active antiretroviral therapy. MSP=medical services plan. *Takes into account all diagnosed individuals in the numerator of the estimated proportion, including those who did not meet the baseline thresholds for adherence, being on HAART, HAART indicated, or retained in or linked to care.

See Online for appendix
required suppressed plasma viral load measurements at least 3 months apart, positively classified up to 49% of diagnosed individuals in 2011. An alternative definition that required two consecutive findings of suppressed plasma viral load resulted in an additional four percentage points of diagnosed individuals being classified as suppressed; use of a constant threshold of plasma viral load of less than 500 copies per mL increased the proportion by an additional nine percentage points. Finally, the most liberal definition, which required only one measurement in which plasma viral load was undetectable in a calendar year, classified roughly five percentage points of additional cases as suppressed from 1998 onwards. Use of the same threshold of a single measurement of less than 500 copies per mL, but with the inclusion of individuals not adherent to treatment (and thus not included in other classifications), resulted in 15–25 percentage points of additional viral load suppression from 1998 onward.

Aggregate plasma viral load measurements, based on the highest available measurement for each individual who received a test in each calendar year, showed decreasing proportions of people in high viral load strata and gains in viral suppression (figure 6).

**Discussion**

In British Columbia, we noted substantial improvements in the proportions of people diagnosed, on HAART, and virologically suppressed, largely as a result of increased testing intensity, changes in IAS-USA treatment initiation guidelines, improvements in compliance with HIV care guidelines, and clinical response to treatment. With the linked administrative data system established as part of the STOP HIV/AIDS initiative, the cascade of care provides an easily interpretable framework to analyse data for the numbers and demographic characteristics of people lost to care at various points on the HIV care continuum; to track HIV-related disparities and health inequities; to provide a basis to inform potential redistribution of resources to improve the efficiency and quality of care and reduce health disparities; to provide a basis for continued assessment of the effect of the various provincial governing bodies responsible for HIV/AIDS care on the coverage, use, and quality of health care for people with HIV, allowing identification of any difficulties encountered and informing future planning; and to provide a window into provincial health policy that can be used as a template for national efforts (panel 2).

Future efforts in the province should focus on the engagement of individuals linked to or retained in HIV care, but not accessing HAART. Further expansion of HIV testing is also a priority; however, uncertainty exists with respect to the number, distribution, and characteristics of undiagnosed HIV-positive individuals in British Columbia. Well designed epidemiological studies to better define this population are needed to inform future HIV testing campaigns. Furthermore, the definition of cascades of care stratified by key demographic characteristics (mode of transmission, age, ethnic origin, etc), characteristics of those lost to care at various stages, and within specific geographical regions would allow for better targeted surveillance systems that can be tracked over time to monitor progress. Such improvements are the key focal points for HIV surveillance in British Columbia in the future, and can inform similar efforts nationally and internationally.

Each stage of the cascade can be affected by several individual and systemic barriers; however, financial constraints consistently play a prominent part. Even within a universal health-care system, most jurisdictions in Canada charge some form of copayment or deductible against prescription drugs, including antiretroviral therapy. For example, as of 2006, Ontario’s annual prescription drug plan for non-elderly people with a net

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**Figure 5:** Cascade of HIV care, including estimates of HIV prevalence and ranges from sensitivity analyses
Prevalence estimates are based on unpublished data from the Public Health Agency of Canada (Archibald C, Public Health Agency of Canada, personal communication). Error bars represent plausible intervals from sensitivity analyses of cascade-stage definitions. HAART=highly active antiretroviral therapy.
annual household income of less than CAN$100 000 entailed a deductible of $150–4089, and a user copayment of $2 per prescription, with no maximum annual contribution.29 The effect of prescription drug cost-sharing on access to antiretroviral therapy in Canada is unclear; higher copayments have been associated with reduced adherence and increased treatment interruption in US settings.30,31 British Columbia is unique in Canada, because HAART and laboratory and medical monitoring of HIV-infected individuals is universally covered and fully subsidised. Our results are thus likely to represent a best-case scenario, in which individuals are not subject to financial disincentives and state-of-the-art antiretroviral management is consistently recommended and available.

Nonetheless, we noted substantial leakage in the cascade of HIV care, particularly at the stages of retention in care, which is an independent predictor of survival.32 Late initiation of antiretroviral therapy (CD4 count <200 cells per μL or an AIDS-defining illness) was favoured by guidelines between 2002 and 2007.33–35 As of 2010, therapeutic guidelines have increasingly recommended that antiretroviral therapy be offered to most infected individuals immediately on diagnosis and, increasingly, irrespective of CD4 cell count.36 These new guidelines negate the necessity of a HAART–indicated classification in future cascades of care, and should decrease loss from the retention stage in the future. Further efforts, such as the refinement of HIV primary care guidelines, intensive case management, outreach, and quality improvement initiatives are urgently needed to ensure sustained engagement in appropriate care and to allow re-engagement by individuals lost to HIV care.

Our sensitivity analyses support the use of the selected cascade stage definitions and provide a basis of comparison for similar efforts in other jurisdictions. Some reports3,22,36 have defined retention in HIV care as having an HIV-related physician visit on two or more occasions at least 3 months apart in a 12 month period. We extended this definition to include any HIV care (physician visit, on HAART, or having a routine CD4 cell count or plasma viral load test). Data for plasma viral load testing and CD4 cell counts in the absence of HIV-related physician visits comprised a large proportion of those defined as linked and retained in care and an important component of HIV surveillance.

The sensitivity analyses for definitions of viral suppression showed the greatest variation. National estimates in the USA suggest that between 19% and 28% of the HIV-infected population were virologically suppressed in 2010.19,37 Our threshold for viral suppression required at least two plasma viral load readings below the threshold for suppression (<50 copies per mL from 1999 onwards) at least 3 months apart, consistent with definitions of retention in HIV care and being on HAART.

Figure 6: Aggregate HIV-1-RNA concentrations in HIV-positive individuals

Data represent the highest plasma viral load measurement for each individual who received a plasma viral load test in each calendar year. *Viral suppression thresholds: <500 copies per mL for 1996, <400 copies per mL for 1997–98, and <50 copies per mL for 1999–2011.
Furthermore, our denominator included only patients adherent to HAART, consistent with the cascade-of-care model. These definitions were chosen solely for their value as indicators of individual and public health benefit. Although a comparison with previous US estimates is indirect, with a threshold of one plasma viral load measurement of less than 50 copies per mL in a calendar year, among any HIV-positive individuals who received a plasma viral load test (previous thresholds of plasma viral load were <200 or <500 copies per mL), the proportion of individuals in British Columbia classified as virologically suppressed was 70% (5792/8308) of those diagnosed, and 50% (5792/11700) of the estimated infected population in 2011. Using a plasma viral load threshold of less than 200 copies per mL, the North American AIDS cohort Collaboration on Research and Design investigators reported that 72% of US participants were virologically suppressed; however, the investigators used as a denominator the number of individuals linked to HIV care. Despite being widely used, our sensitivity analysis suggests that definitions based on single measurements of plasma viral load suppression probably misclassify as suppressed a substantial proportion of individuals who are not stably engaged in treatment in a given calendar year.

Despite the comprehensive scope of the data systems used, our study has several limitations related to measurements at each stage of the cascade of care. First, the number of prevalent cases is not known and was instead derived from a national modelling effort on the basis of common assumptions about key model parameters across provinces. In view of the significantly decreasing rates of new cases of HIV in British Columbia compared with stable or increasing rates elsewhere in Canada, these estimates might be positively biased. Second, although administrative loss to follow-up was accounted for, emigration from the province, particularly in 2010 and 2011 in view of the study cutoff point, might have resulted in us overestimating the numbers of people diagnosed and linked to care, thereby underestimating proportions of individuals receiving HAART and those virologically suppressed. Earlier annual administrative losses to follow-up (between 1996 and 2009) were in the range of 1·5% (70 of 4656 diagnosed) in 1997 to 1·98% (102 of 5160 diagnosed) in 1998; therefore, this misclassification was probably small. Third, the study cohort was defined partly on the basis of health administrative data; therefore, some cases could have been misclassified, potentially missing some undiagnosed cases. We have described the procedures used to construct and validate the cohort elsewhere—specifically, we applied case-finding algorithms to identify HIV-positive individuals identified as such only from health administrative databases.

Outpatient care delivered in some inner-city health clinics had billing by session rather than fee-for-service, and was therefore not captured in the medical services plan database. Our definition of retention in HIV care thus incorporated antiretroviral dispensations, for which we have complete capture. Also, the number of individuals with nominal and therefore linkable HIV diagnoses was underestimated, particularly before 2003, when HIV became reportable and systematic follow-up of all new HIV diagnoses commenced, improving data quality for identifiers. As a result, our previous analysis estimated that only 52% of individuals accessing HIV care in British Columbia had a linked HIV test available, which would result in our overestimating the proportion of undiagnosed infections before 2003.

With respect to the subsequent cascade stages, incomplete capture of CD4 cell count measurements could have resulted in underestimates for retention in HIV care over time. Measurement of adherence to HAART was based on refill compliance, which might overestimate true adherence. Furthermore, an informal audit done in 2010 (Barrios R, unpublished) showed that 125 (2·4%) of 5264 individuals eligible for antiretroviral therapy in the area under the jurisdiction of the Vancouver Coastal Health Authority were on treatment and being monitored in the context of industry-sponsored clinical trials, and thereby not captured by the BC Centre for Excellence databases; our estimates for treatment uptake, adherence, and viral suppression are thus slightly lower than actual numbers, resulting in a small conservative bias. Finally, assessments of need for antiretroviral therapy were simplified from actual recommendations, focusing on standard CD4 cell counts, plasma viral load tests, and AIDS-defining illness.

Panel 2: Research in context

Systematic review
We searched PubMed for articles published in English up to April 31, 2013, that included the terms “HIV” and “cascade”, “continuum”, or “retention” in the abstract. One peer-reviewed study estimated the cascade of HIV care for the US population, and this estimate was later updated by the US Centers for Disease Control, which suggested that 40% or fewer HIV-positive individuals were retained in care, and 19–25% were virologically suppressed.

Interpretation
Although each stage in the cascade of HIV care in British Columbia has improved with time, gaps in diagnosis and retention in care remain. Our longitudinal characterisation of the cascade allows for explicit consideration of changes in cascade leakage between stages over time. The results of our sensitivity analyses showed that cascade-stage counts are sensitive to the definitions of cascade stages used, particularly with respect to how viral suppression is measured. In view of the absence of evidence-based standards for cascade-stage definitions, caution should be used when comparing our results with those from other jurisdictions. Our methods could be useful in the design of HIV surveillance systems internationally.

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thresholds for eligibility, and not taking into account secondary or individual-specific considerations.

Our results show a steady improvement in the engagement of people with HIV within the cascade of care in British Columbia during the HAART era. Careful mapping of the cascade of care is crucial to improve our understanding of how to maximise the beneficial effects of available interventions and to inform efforts to contain the spread of HIV/AIDS. A high-quality HIV surveillance system actively linked to relevant administrative health records is essential for such an endeavour.

Contributors
BN and JSGM had the initial idea for the research. BN led in the preparation of the report and contributed to the analysis. GC led the analyses. BY, KC, and VDL also contributed to the analysis. KH, MG, HS, and RSH contributed to the preparation of the report. JSGM, RB, RG, and RSH contributed to the procurement of the data on which the study is based. All authors contributed to data interpretation and approved the submitted version of the report.

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Conflicts of interest
JSGM has received grants from Abbott Laboratories, biotechnical Laboratories, Boehringer Ingelheim, Bristol-Myers Squibb, Gilead Sciences, Janssen Pharmaceuticals, Merck, and ViIV Healthcare. He has also received support from the British Columbia Ministry of Health; a Knowledge Translation Award from the Canadian Institutes of Health Research; and an Avant-Garde Award (IDPDIA026182) from the US National Institute on Drug Abuse (National Institutes of Health); the International AIDS Society; UNAIDS; WHO; the US National Institutes of Health Office of AIDS Research; the US National Institute of Allergy and Infectious Diseases; the US President’s Emergency Plan for AIDS Relief; the Bill & Melinda Gates Foundation; the French National Agency for Research on AIDS and Viral Hepatitis; and the Public Health Agency of Canada. All other authors declare that they have no conflicts of interest.

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The HIV care cascade through time

HIV care and treatment can prevent morbidity, mortality, and virus transmission. Optimum care for individuals and communities of people living with HIV involves identification of infected individuals, linkage to initial HIV care, long-term retention in care, and treatment adherence—the so-called cascade of care. However, in many settings, the scope of the cascade is such that few patients actually achieve undetectable viral loads, the end goal of engagement in care. Understanding how to measure and intervene to improve engagement in HIV care is a subject of intense debate.

In The Lancet Infectious Diseases, Bohdan Nosyk and colleagues1 from the STOP HIV/AIDS Study Group chart the longitudinal changes in the cascade of HIV care in British Columbia, Canada, from 1996 to 2011. Their study is the first longitudinal examination of the HIV care cascade. The investigators assessed the numbers and proportions of individuals in eight distinct stages of the cascade: HIV infected, diagnosed, linked to HIV care, retained in care, antiretroviral treatment indicated, receiving antiretroviral treatment, adherent to antiretroviral treatment, and virologically suppressed.

The study’s strengths derive from the extensive use of comprehensive linked databases from national and provincial health programmes, and population-based registries from the BC Centre of Excellence in HIV/AIDS (Vancouver, BC, Canada)—the sole provincial agency providing HIV diagnostic testing and distribution of all antiretroviral drugs. Additional information was derived from provincial hospital, pharmacy, and vital statistics databases. The analysis shows that overall engagement in care and use of antiretroviral treatment improved between 1996 and 2011, but that substantial numbers of individuals are still lost from each step of the cascade.

In 2011, an estimated 29% of HIV-infected individuals remained undiagnosed, an additional 4–10% were not linked to HIV care, and another 20% were not retained in care. Overall, viral suppression increased from 1% to 35% of the HIV-infected population over the study period.

Nosyk and colleagues’ study shows us the value of looking longitudinally at the use of HIV care. Although changing standards for when to begin antiretroviral treatment limit the ability to analyse trends in viral suppression over time, increasing numbers of individuals are achieving this important benchmark. However, only a minority of HIV-infected individuals in British Columbia are virologically suppressed, and this finding is surprising and disappointing. As the investigators suggest, emigration from the province might account for some losses to follow-up; in a recent US study, about 15% of individuals emigrated from the state in which they were diagnosed during 3–5 years of follow-up. Other potential losses of data in British Columbia, such as receiving care through participation in clinical trials, seem to have had little effect on estimates of viral suppression.

The implications of persistent gaps in cascade steps before administration of antiretroviral treatment and viral suppression are particularly worrying. Compared with research from the USA, the investigators in British Columbia report fairly similar proportions of HIV underdiagnosis, linkage to care, and retention in care. Factors embedded in health-care systems, stigma, and discrimination probably drive these consistent deficits. Solving these issues will need thoughtful application of best practices and additional research.

Antiretroviral treatments for HIV have greatly improved with time, offering the promise of normal life expectancy for people living with HIV who successfully navigate the cascade. Yet, in North America, and around the world, losses from the various cascade steps can jeopardise the health outcomes of many people. Jurisdictions should develop metrics of care cascades and population-based assessments of HIV viral loads. Such data could provide a broad overview of the HIV treatment and prevention landscape and point to areas in need of improvement.

In this context, understanding the limiting factors in each of the cascade steps is important, with the recognition that what causes people to be lost from the cascade is likely to vary by region and individual. Although it is tempting to use cascades to compare countries, provinces, states, and other jurisdictions, differing HIV care guidelines, clinical practice, data reporting, and data availability make such comparisons difficult. Instead, the value of the cascade lies in its ability to guide local programmes and jurisdictions to address problematic cascade transitions.

By following the cascade over time, jurisdictions...
can assess the effectiveness of programmes and interventions. Nosyk and colleagues’ study is an excellent example of the usefulness of reviewing cascades of care over time—we look forward to seeing more studies of this kind in the future. Shared successes will hopefully help the global HIV community to improve care worldwide.

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