A review of Human Immunodeficiency Virus (HIV) rapid testing

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Abstract

Background: In Canada, it is estimated that 71,300 persons were living with HIV at the end of 2011. Approximately 25% (14,500 to 21,500) of prevalent cases were unaware of their HIV infection. Expanded use of HIV rapid tests may increase the detection of undiagnosed infections, enable earlier treatment and support services and prevent the onward transmission of HIV.

Objective: To examine patient acceptability, impact (defined as receipt of test results and linkage to care) and cost-effectiveness of HIV rapid tests.

Methods: A search was conducted for systematic reviews on HIV rapid testing, with studies from both developed and developing countries, published in English and between 2000 and 2013. The Assessment of Multiple Systematic Review (AMSTAR) tool was used to assess the included systematic reviews for methodological quality. Results were summarized narratively for each of the outcomes.

Results: Eight systematic reviews were included. Acceptability of HIV rapid tests was generally high in medical settings (69% to 98%) especially among pregnant women and youth attending emergency rooms but was lower in non-medical settings (14% to 46%). The percentage of people who obtained their test results was variable. It was high (83% to 93%) in emergency rooms but was low in a rapid care setting with regular business hours (27%). Impact on linkage to care was limited. Only one systematic review examined cost-effectiveness of rapid testing and concluded that HIV rapid tests were cost-effective in comparison to traditional methods; however, results were all based on static models.

Conclusion: Overall, HIV rapid tests demonstrated generally high acceptability, variability in receiving test results and limited impact on linkage to care. While these findings suggest that HIV rapid tests may be useful, further research is needed to confirm in whom, when and where they are best used and how to ensure better linkage to care.

Introduction

At the end of 2011, an estimated 71,300 persons were living with Human Immunodeficiency Virus (HIV) and Acquired Immunodeficiency Syndrome (AIDS) in Canada and an estimated 25% were unaware of their HIV status (1). Those unaware of their status are unable to take advantage of available support services and care, are at increased risk of transmitting HIV and are at increased risk of acquiring other sexually transmitted and blood-borne infections. Effective screening strategies that lead to earlier diagnosis and treatment can contribute to improved individual and population health outcomes (2).

With the emergence of new diagnostic technologies, there are increasing options for HIV testing. Rapid tests for HIV are available worldwide including oral fluid tests and finger prick tests using whole blood or plasma. HIV rapid tests can be either self-administered or administered by trained staff. In Canada, HIV rapid tests can only be carried out by trained staff in point-of-care (POC) settings (e.g., doctors’ offices, clinics, emergency departments) (3-5). In addition, the Public Health Agency of Canada recommends that HIV rapid tests be administered in conjunction with pre- and post-test counselling (5).
Only one HIV rapid test is licensed for use in Canada (6). In October 2005, Health Canada approved the INSTI™ HIV-1 Antibody Test (a single use rapid test for HIV) for use in POC settings. In 2008, the license was amended to include the INSTI™ HIV-1/HIV-2 Antibody Test (6). This test is a preliminary antibody screening test that can be performed on site where the patient can receive their results immediately (< 1hr) (7-10). If the patient receives a preliminary reactive result, a confirmatory test using traditional laboratory-based testing is required. If the test result is negative (non-reactive), no further testing is necessary (3,5).

Previous studies suggest POC testing has the potential to improve the management of infectious diseases by identifying new infections, reducing the numbers of those who are unaware and facilitating linkage to care (11,12). To ensure HIV rapid tests are feasible, they should also be cost-effective. The objective of this rapid review was to examine the most current evidence on patient acceptability, impact (defined as receipt of test results and linkage to care) and cost-effectiveness of HIV rapid tests.

Methods

We followed the Ottawa Hospital Research Institute’s methods for conducting rapid reviews (13). This method is designed to provide decision-makers with a synthesis of an extensive literature in a timely manner (13). A protocol was developed for the rapid review a priori that included: question development and refinement; a systematic literature search; screening and selection of systematic reviews; assessing the quality of the evidence; and a narrative synthesis of included studies. (13).

Search strategy

The following databases were searched: Medline, Embase, Scopus, Social Policy and Practice, Proquest Public Health and Google Scholar. Articles were included if they were published between January 2000 and September 2013; included studies from developed or developing countries; and/or published in English. The search strategy included the following key words: (“human immunodeficiency virus” OR “HIV”) AND (“Point of care” OR “point-of-care”, “rapid test” OR “home-based test” OR “screen*”) AND (“linkage to care” OR “follow-up” OR “barrier*”, “intervention*” OR “access*” OR “diagnos*”) OR (“acceptab*”, “willing*”, “satisf*”, “preference*”) OR (“feasib*”, “economic*”, “financ*”, “cost*”). Articles that reported results on HIV prevalence or studies with no mention of HIV rapid testing were excluded from the review.

Quality assessment of the studies

Each systematic review was evaluated using the Assessment of Multiple Systematic Review (AMSTAR) tool for methodological quality (14). The AMSTAR tool consists of an 11-item questionnaire that assesses the following criteria: use of an a priori design; duplicate study selection and data extraction process; comprehensive literature search; use of publication status as an inclusion criterion; characteristics of included studies; list of included/excluded studies; assessment of the quality of studies; appropriate use of scientific quality in forming conclusions; appropriate methods used to combine study findings; assessment for publication bias; and acknowledgement of conflict of interest. To ensure reliability of the assessment, two of the authors (SH, SF) evaluated the systematic reviews using the AMSTAR tool. Where there was discrepancy, a third person (DP) was invited to assess the criterion in question.

Data extraction

For each of the included systematic reviews, two authors (SH, SF) extracted data on population; search years; number of included studies; locations of included studies; study objective; type of intervention; and outcomes. Outcomes of interest included: acceptability, receipt of HIV test results, linkage to care and cost-effectiveness. After data extraction, both authors compared their findings to ensure consistency.

Results

The initial search yielded a total of 892 articles on rapid testing for HIV. After limiting to systematic reviews (n=12), eight review articles met the inclusion criteria (Figure 1).
Figure 1: Algorithm of literature search and study selection of systematic reviews on rapid HIV testing

A description of the included reviews and the respective AMSTAR score out of 11, are presented in Table 1. Three had perfect AMSTAR scores and another was of high quality (with a score of 8). Reasons for a systematic review having a score less than eight included: it did not specify a duplicate study selection and data extraction process; assessment and documentation of the quality of the studies; or assessment of publication bias.
Table 1: Description of included systematic reviews with the AMSTAR¹ scores

<table>
<thead>
<tr>
<th>Reference</th>
<th>Objective(s)</th>
<th>Population and location</th>
<th>Search period, intervention and number of included studies</th>
<th>AMSTAR score (out of 11)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bateganya (2007) (17)</td>
<td>To identify and critically appraise studies addressing the implementation of home-based HIV voluntary counselling and testing; to assess the effect of this intervention compared to facility-based HIV counselling and testing.</td>
<td>Population: Adults (&gt;15 years) Location(s): Uganda and Zambia</td>
<td>Search period: 1980 - 2007 Intervention: Voluntary counselling and testing for HIV No. included studies: 2</td>
<td>11</td>
</tr>
<tr>
<td>Pant Pai (2007) (15)</td>
<td>To summarize the overall diagnostic accuracy of rapid HIV tests in pregnancy; evaluate outcomes and impact of testing; and identify practical challenges related to the implementation of voluntary HIV testing and counselling in pregnant women.</td>
<td>Population: Pregnant women (18 to 44 years) Location(s): South Africa, United States, Latin America, South-East Asia, Jamaica</td>
<td>Search period: 1991 - July 2005 Intervention: HIV POC testing in pregnancy No. included studies: 17</td>
<td>8</td>
</tr>
</tbody>
</table>
Roberts (2007) (16)  
To review the outcomes of blood and oral fluid rapid HIV testing.  
**Population:** Various  
**Location(s):** United States, Kenya, Brazil, Zimbabwe, Burkina Faso, Mexico  
**Search period:** January 2000 - June 2006  
**Intervention:** HIV rapid testing  
**No. included studies:** 26  

Turner (2013) (19)  
To review preferences and acceptability of rapid POC testing in youth, to document notification rates and to identify sociodemographic factors associated with youth choosing rapid HIV POC testing over traditional testing.  
**Population:** Youth (<25 years)  
**Location(s):** United States  
**Search period:** January 1990 - March 2013  
**Intervention:** HIV POC testing  
**No. included studies:** 14

Acceptability

Almost all of the reviews (7/8) examined acceptability. Acceptability was defined in these reviews as the population’s uptake of a rapid test (8, 9, 15-17) or as the patient’s preference for a rapid test when offered the choice of a rapid test or traditional laboratory-based test (18, 19).

In the Roberts et al. review, the overall acceptability of rapid tests administered in both medical and community settings ranged from 14% to 98% (16). Acceptability of rapid testing was lower (14% to 46%) in alternative testing sites (e.g., bathhouses, needle exchange programs, jails and emergency departments) compared to medical settings (69% to 98%) (e.g., sexually transmitted infection clinics, labour and delivery units and hospitals) (16). The wide range of acceptance rates may have been affected by differences in the definition of acceptability and in the data collection methods.

In two reviews, acceptability for HIV rapid tests was high among pregnant women (15,16). In the Pant Pai et al. review, acceptability among pregnant women ranged from 83% to 97% (15). Similarly, in the Roberts et al. review, acceptability among pregnant women ranged from 74% to 86% in American studies and from 93% to 98% in international studies (16). Among pregnant women, the following factors were associated with high acceptability of HIV rapid testing: age (<21 years), higher education and lack of appropriate prenatal care during pregnancy (15).

Among the youth, Turner et al. found that 35% to 93% accepted HIV rapid tests when offered. The 35% acceptance rate was found in an adolescent outpatient clinic (19). However, when given the option of rapid or traditional methods, youth from the adolescent outpatient clinic selected rapid methods 70% of the time (19). The highest acceptance rates (83% to 93%) were found in emergency rooms suggesting that there is high acceptability for rapid testing among youth attending emergency departments (19).

In the Mavedzenge et al. review, acceptability was defined as the interest to self-test. Among key populations such as men who have sex with men (MSM) and emergency department attendees, the authors found that acceptability of self-testing was moderate to high (62% to 92%) (18). Reasons for preferring self-testing included privacy, autonomy, confidentiality, anonymity, convenience and speed.

Pant Pai et al. demonstrated that acceptability (choosing self-testing over the traditional laboratory-based tests) was high in supervised and unsupervised settings (9). In supervised settings, there was high acceptability (74% to 96%) among emergency department attendees, urban MSM, university students and the general urban population. Of note, an older study from 2001 reported an acceptance rate of 24% among HIV clinic attendees. In unsupervised settings, the high acceptability (74% to 84%) was only based on two studies, which focused on healthcare professionals and HIV negative MSM (9).

Acceptability of HIV rapid tests was variable across different populations, but was generally high among pregnant women, youth attending emergency rooms and in medical settings. More research is needed to explore self-testing in unsupervised settings and reasons for low acceptance rates in non-medical settings.
Receipt of HIV test results

Four of the eight (4/8) systematic reviews examined the impact of HIV rapid testing on patients’ receipt of test results. One systematic review by Roberts et al. noted that 27% to 100% of clients who attended medical and community settings for rapid testing received their HIV test results (16). The low rate of 27% was when same day results were available in an urgent care clinic with regular business hours and most participants left before the results were available (20). In the remaining studies, more than 70% of participants who underwent rapid testing at hospitals, sexually transmitted infection clinics, homeless shelters and bathhouses received their test results (16).

In a review by Bateganya et al., those who received voluntary counselling and testing (with rapid tests) at home were approximately five times more likely to receive their test results compared with those who received voluntary counselling and rapid testing at a clinic (17). The authors conducted an updated review that included one additional study, and found that 56% of individuals who had the home-based testing received their test results compared to 12% who had clinic-based testing (8). Based on these findings, receipt of rapid test results tended to be moderate to high except in urgent care clinics with regular business hours.

Linkage to care

Six of the eight (6/8) systematic reviews assessed linkage to care although the definition of linkage to care varied among the reviews. Roberts et al. defined linkage to care as entry into medical care and found this occurred in 47% to 100% of those who were diagnosed with HIV from rapid tests (16). Mavedzenge et al. defined linkage to care as linkage to prevention, treatment and care services and concluded that data are insufficient to determine whether self-testing leads to timely linkage to care (18). Dibosa-Osadolor et al. found that rapid HIV testing resulted in a higher percentage of patients being appropriately linked to care compared to traditional HIV testing (21); however, exact percentages were not listed. Bateganya et al. did not provide a clear definition for linkage to care, but included studies that offered voluntary pre- and post-test counselling at home. Compared to those offered testing and counselling in a clinic, those tested at home were more likely to accept post-test counselling (17). In the updated review by Bateganya et al., 12% received post-test counselling from a clinic and 56% received post-test counselling at home (8). Most reviews acknowledged that information on linkage to care was sparse (9, 15, 16).

See Table 2 for a summary of the acceptability, receipt of test results and linkage to care data.

Table 2: Summary of acceptability, receipt of HIV test results and linkage to care with primary references

<table>
<thead>
<tr>
<th>Reference</th>
<th>Acceptability</th>
<th>Receipt of HIV test results</th>
<th>Linkage to care</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bateganya (2007) (17)</td>
<td>Those randomized in optional testing locations (including home-based testing) were 4.6 times more likely to accept voluntary counselling and testing than those in the facility arm (RR 4.6 95% CI 3.6-6.2) (26).</td>
<td>In the year where participants were given the option to receive their HIV test results at home, participants were 5.23 times more likely to receive their results than during the year when results were available only at the facility (OR 5.23 95%CI 4.02-6.8) (27).</td>
<td>The definition for linkage to care was unclear. It appears that those who received their results also received post-test counselling.</td>
</tr>
<tr>
<td>Bateganya (2010) (8)</td>
<td>Acceptability of pre-test counselling and HIV test was 12% vs. 57% (optional group) (26)</td>
<td>12% received post-test counselling and their test results from the local clinic; 56% received results and counselling at home (RR 4.7 95%CI 3.62-6.21) (26).</td>
<td>The definition for linkage to care was unclear. It appears that those who received their results also received post-test counselling.</td>
</tr>
<tr>
<td>Dibosa-Osadolor (2010) (21)</td>
<td>N/A</td>
<td>N/A</td>
<td>Antibody rapid testing also resulted in a higher percentage of patients being appropriately linked to care (28-31).</td>
</tr>
<tr>
<td>Napierala Mavedzenge (2013)</td>
<td>Health workers from African countries had high interest in self-testing 73% to 79% (32-34).</td>
<td>N/A</td>
<td>Insufficient data.</td>
</tr>
</tbody>
</table>
In US studies, emergency department patients and MSM\(^2\) had high acceptability ranging from 83% to 89% (35-37).

<table>
<thead>
<tr>
<th>Study</th>
<th>Overall acceptability: %</th>
<th>N/A</th>
<th>Details of linkages to care and prevention were not reported.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pant Pai (2007) (15)</td>
<td>83% to 97% (38-42).</td>
<td>N/A</td>
<td>Overall acceptability: 83% to 97% (38-42).</td>
</tr>
<tr>
<td>Pant Pai (2013) (9)</td>
<td>74% to 96% for both supervised and unsupervised settings (7, 35, 43-49). Supervised settings: 24% to 95% (7,35,43-47) - Urban MSM: 74% (35) - Emergency department: 85% (7) - Rapid HIV testing site: 78% (45) - General urban population: 92% (46,47) - Educated students: 95% (44) - HIV Clinic attendees: 24% (43) Unsupervised settings: 78% to 84% (48,49) - Non-monogamous MSM: 84% (49) - Healthcare professionals: 78% (48)</td>
<td>N/A</td>
<td>Only one study in a US unsupervised setting was reported - 96% of those who test positive for HIV would seek post-test counselling (50).</td>
</tr>
<tr>
<td>Roberts (2007) (16)</td>
<td>14% to 98% (10, 39,51-64). - Pregnant women (US): 74% to 84% to 86% (39,52,53) - Women in labour: 95% (62) - Pregnant women in international prenatal settings: 93% to 98% (57,62,64) - Pregnant women in prenatal medical care (18% and 26%) (59,60) - Urgent care: 40% (20) - Hospitals: 60% (63) - Emergency: 29% (54) - Government Health centre (Kenya): 93% (51) - STI(^1) clinics: 65% to 87% (54-56) - County jail: 46% (54) - Bathhouse: 21% (10) - Needle exchange: 14% (10)</td>
<td>Overall receipt of HIV test results: 27% to 100% (10, 20, 51,54-56,59,63-69). - Hospital: 95% to 100% (63,65) - STI(^1) clinic: 89% to 99% (55,56,66) - Urgent care: 27% (20) - Labour / delivery unit: 68% to 94% (67) - Prenatal care: 74% and 98% (59,64) - Mobile site: 99% (68) - Community settings (e.g., homeless shelters, jail, bathhouse, needle exchange): 83% to 100% (10,54,69)</td>
<td>Overall: 47% to 100% (all US studies) (20, 54, 55, 65). Few studies examined entry rates into medical care in those who were found to be HIV+ from rapid tests.</td>
</tr>
<tr>
<td>Turner (2013) (19)</td>
<td>35% to 93% (70-80). Lowest acceptance rate was found in an adolescent outpatient clinic (35%) (73). Highest acceptance rates found in emergency departments (83% and 93%) (74, 77). When given the options of rapid and traditional testing, youth selected rapid tests 70% of the time (73).</td>
<td>Participants who chose a rapid test were more likely to receive their test results within the follow-up period, compared with those who chose traditional test (91.3% vs. 46.7%; OR 12 95%CI 3.98-36.14) (73). 100% of youth aged 13-17 years who accepted rapid testing received their results (77).</td>
<td>N/A</td>
</tr>
</tbody>
</table>

\(^{1}\)STI = sexually transmitted infections  
\(^{2}\)MSM = men who have sex with men
Cost-effectiveness

Of the 17 modelling studies reviewed by Dibosa-Osadolor et al., seven studies addressed diagnostic testing for HIV detection. Four modelling studies specifically assessed rapid testing with immediate patient notification in a clinical setting. The authors concluded that HIV rapid testing was more cost-effective than traditional laboratory-based testing with immediate patient notification (21). However, the majority of the modelling studies of rapid testing reviewed were based on static models, which do not include time dependencies. This can potentially result in an overestimation of the cost-effectiveness of infectious diseases (21, 22). In this review, there was no information on the direct and indirect costs of rapid testing or on the cost per the quality-adjusted life-year gained.

Discussion and conclusion

Our rapid review of eight systematic reviews found that HIV rapid tests demonstrated generally high acceptability, especially among pregnant women; variability in receiving test results; and limited impact on linkage to care. One review found that rapid testing was cost-effective, but the studies were based on a static versus a dynamic model; therefore, further studies are warranted to determine the impact of rapid tests on linkage to care and its cost-effectiveness.

The rapid review methodology is a fairly novel approach that has its strengths and limitations. The strength is that it is a rapid way to summarize evidence for decision-makers. In addition, the evidence is presented transparently, allowing users to assess the evidence and make informed decisions. However, there are a few limitations to consider when reviewing these results. The shortened timeframe of the rapid review process may miss studies that were not included in the reviews and therefore, may introduce bias through the absence of some relevant information. It may exclude recently published systematic reviews or those currently in press (23, 24). Moreover, data from some individual studies were cited more than once across the systematic reviews, which may inflate the confidence in the results presented in this rapid review (24, 25). Finally, the systematic reviews included studies from different countries and different types of HIV rapid tests; therefore, the results from this review may not be generalizable to other rapid tests or to the Canadian setting.

It appears that offering HIV rapid tests in settings is highly effective when test results can be readily obtained. This suggests that rapid HIV tests could decrease the proportion of individuals who are unaware of their HIV status and merits further study. Future research should compare effectiveness among different populations and settings, as well as explore ways to improve linkage to care. It would be useful to have a cost-effectiveness study based on a dynamic model.

Acknowledgements

The authors thank Margaret Gale-Rowe, John Kim, Lisa Pogany and Tom Wong for their review and input into the writing of this article. The authors would also like to thank Cindy Smalley and Elizabeth Dekens for their contributions to the literature search.

Conflict of interest

None.

Funding

This study was supported by the Public Health Agency of Canada.

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