RESEARCH ARTICLE

Undiagnosed HIV infections among gay and bisexual men increasingly contribute to new infections in Australia

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Abstract
Introduction: We determined the contribution of undiagnosed HIV to new infections among gay and bisexual men (GBM) over a 12-year period in Australia where there has been increasing focus on improving testing and HIV treatment coverage.

Methods: We generated annual estimates for each step of the HIV cascade and the number of new HIV infections for GBM in Australia over 2004 to 2015 using relevant national data. Using Bayesian melding we then fitted a quantitative model to the cascade and incidence estimates to infer relative transmission coefficients associated with being undiagnosed, diagnosed and not on ART, on ART with unsuppressed virus, or on ART with suppressed virus.

Results: Between 2004 and 2015, we estimated the percentage of GBM with HIV in Australia who were unaware of their status to have decreased from 14.5% to 7.5%. During the same period, there was a substantial increase in the number and proportion of GBM living with HIV on treatment and with suppressed virus, with the number of virally suppressed GBM increasing from around 3900 (30.2% of all GBM living with HIV) in 2004 to around 14,000 (73.7% of all GBM living with HIV) in 2015. Despite the increase in viral suppression, the annual number of new infections rose from around 660 to around 760 over this period. Our results have a wide range due to the uncertainty in the cascade estimates and transmission coefficients. Nevertheless, undiagnosed GBM increasingly appear to contribute to new infections. The proportion of new infections attributable to undiagnosed GBM almost doubled from 33% in 2004 to 59% in 2015. Only a small proportion (<7%) originated from GBM with suppressed virus.

Discussion: Our study suggests that an increase in HIV treatment coverage in Australia has reduced the overall risk of HIV transmission from people living with HIV. However, the proportion of infections and the rate of transmission from undiagnosed GBM has increased substantially. These findings highlight the importance of HIV testing and intensified prevention for Australian GBM at high risk of HIV.

Keywords: Australia; gay and bisexual men; mathematical model; transmission dynamics; undiagnosed HIV; HIV cascade

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1 | INTRODUCTION

People with undiagnosed HIV contribute disproportionately to HIV transmission because the early stages of infection are the most infectious, and without treatment an individual remains infectious [1-7]. Being unaware of their infection undiagnosed individuals are also less likely to take precautions to prevent transmission [8]. In high-income countries, with concentrated epidemics among gay, bisexual and other men who have sex with men (GBM), the proportion of people living with HIV with undiagnosed infection ranges from 10% to 50% [2-5,9-12] with modelling suggesting the majority of new HIV infections originate from GBM who are undiagnosed or diagnosed but not in care [1,6,10,13].

Reducing undiagnosed HIV and increasing treatment uptake through more frequent testing and timely diagnosis have become critical to global efforts to reducing HIV transmission [14]. Many countries monitor their progress towards global targets using a HIV cascade [15,16]. The cascade starts with the number of people living with HIV and shows the number or proportion of people who are diagnosed, retained in care, on treatment and virally suppressed.

In Australian jurisdictions, the proportion of undiagnosed GBM was reported to be 20 to 31% in 2007 to 2008
[17,18] compared to 9% nationally in 2014 [19]. While there were differences in study methodology and wide confidence intervals for earlier estimates, these studies suggest that the prevalence of undiagnosed HIV has fallen over the last decade. In parallel, there have been increases in HIV testing, substantial increases in HIV treatment coverage and reductions in population viral load [11]. Yet during the same period, annual HIV notifications increased in Australia, stabilizing at around 1000 to 1100 infections per year, with 65% to 70% occurring in GBM [11]. Similar increases in HIV incidence have been seen in other locations with concentrated epidemics among GBM despite an increase in treatment coverage (like British Columbia as of 2015 [20,21]). Whereas in San Francisco, a decline in undiagnosed infection following increases in HIV testing and treatment coverage coincided with a decline in HIV incidence [22-24].

Many models have been used to estimate the role of undiagnosed infection in sustaining epidemics and assessing the impact of interventions along the treatment cascade. Many of these models are complex static HIV transmission models (providing point estimates) or dynamic HIV transmission models (to provide estimates over time) incorporating risk group, sexual behaviour and disease progression data [1,3,6,13,25,26]. Such models require a large amount of demographic, behavioural, and clinical data and considerable effort to calibrate and update. An alternative approach, previously developed by Kelly and Wilson [27], uses a simple calculation to estimate the average infectivity of people living with HIV over time. We applied this approach to estimate the proportion of new HIV infections in Australia ascribed to each step of the HIV cascade over time.

2 | METHODS

Using the Australian HIV cascade, we obtained annual estimates for the number of Australian GBM living with HIV who have undiagnosed infection $N_u$, have been diagnosed but are not on treatment $N_{u\text{a}}$, are taking treatment but have an unsuppressed viral load $N_{u\text{b}}$, and are on treatment and have a suppressed viral load $N_s$ (Data S1). We then linked estimates for the annual number of new GBM infections $I$ to each stage of the cascade, inferring a transmission coefficient, equal to the “average” number of transmissions from people living with HIV in each step of the cascade each year (denoted by $\beta_u$, $\beta_{ua}$, $\beta_{ub}$, and $\beta_s$). Mathematically, this is described by the following equation

\[ I(t) = \beta_u(t)N_u(t) + \beta_{ua}(t)N_{u\text{a}}(t) + \beta_{ub}(t)N_{u\text{b}}(t) + \beta_s(t)N_s(t) \]  

(1)

for each point in time $t$. If estimates for the number of new infections and the number of GBM in each step of the HIV cascade are available over time, it is possible to fit the transmission coefficients, which can then be used to estimate the proportion of new infections attributable to each step of the cascade; for example, for undiagnosed this is $\beta_u N_u/I$. This model does not explicitly capture changes in behaviour or clinical factors that cause changes in the transmission coefficients over time. The coefficients simply describe the overall transmission likelihood from people at each step of the cascade.

We now summarise the methods we used to estimate the number of GBM in each step of the cascade in Australia, the number of new infections, and the transmission coefficient fitting methodology (details provided in the Data S1). We conducted this analysis using R version 3.2.2 [28]. Reproducible code, cleaned input data and summary results are available online [29]. No ethical approval or consent was sought as this was a mathematical modelling study using publicly available data.

2.1 | HIV cascade for Australian gay bisexual men

We estimated four steps of the HIV cascade for Australian GBM using a variation in the methodology described in national surveillance reporting [11]. To estimate the proportion undiagnosed, we used the European Centre for Disease Control (ECDC) HIV Modelling Tool [30,31]. The ECDC tool is a multi-state back-calculation model that fits diagnoses rates over time using data on new HIV and AIDS diagnoses, estimates for the number of annual deaths and emigrations within people living with diagnosed HIV, and estimates for the rate of CD4 decline. We applied the ECDC model to GBM using HIV surveillance data from the Australian National HIV Registry [11]. The calculations for the other steps of the cascade are described in Data S1.

For each step of the cascade, we provide an annual best estimate and range, rounded to the nearest 10. We defined GBM living with HIV who have never been diagnosed in Australia as “undiagnosed”, those previously diagnosed but not on ART as “diagnosed”, those on ART but with viral load ≥200 copies/mL as “unsuppressed”, and those on ART and with viral load <200 copies/mL as “suppressed.”

2.2 | New infections

The ECDC tool also produces estimates for new infections. As arrival date for people previously diagnosed overseas is incomplete in the HIV registry, we ran the ECDC tool under two scenarios, once with people previously diagnosed overseas included in the notifications (as used for the undiagnosed estimate) and once with them excluded. We used the midpoint as the best estimate for new infections with the lower bound equalling the lower bound of the scenario with people previously diagnosed excluded and the upper bound equalling the upper bound of the scenario with people previously diagnosed included (Figure 1B). We provide the final ECDC HIV Modelling Tool output spreadsheets in the code repository [29].

2.3 | Model fitting procedure

We fitted the transmission coefficients to the HIV cascade and new infections estimates using a Bayesian melding methodology [32-34]. This allowed us to incorporate the uncertainty in the cascade and new infection estimates as well as the impact of ART on preventing HIV transmission.

The first step specifies prior probability distributions (priors) for each transmission coefficient. Rather than specifying priors for the number of people in each step of the HIV cascade and each transmission coefficient, we used the point
estimates for each cascade step and merged the uncertainty in the cascade values into the transmission coefficients. We did this as follows, letting \( N_x \) represent the point estimate for each cascade step then the range of values for each step can be written as \( N_x E_x \), where \( E_x \) represents a multiplicative factor essentially representing the uncertainty in the population estimate for step \( x \). Each term in Equation 1 can then be written as \( \beta \times E \times N \). As the transmission coefficients \( \beta \) and uncertainty terms \( E \) are multiplied, they can be represented by one overall transmission coefficient \( \beta' = \beta \times E \), which implicitly includes the uncertainty in both the cascade values and the transmission coefficient. Using the point estimates for each cascade step in this way means we only require priors for each \( \beta' \) term. For the remainder of this study we implicitly assume the transmission coefficients \( \beta \) in Eq. 1. multiply the corresponding point estimate for each cascade step.

Figure 1. Estimates for new infections and the Australian GMb HIV cascade. (A) Estimated percentage of people undiagnosed over 2004 to 2015 compared to the 2014 COUNT estimate. The red line is the average of the two estimates produced by the two ECDC HIV Modelling Tool scenarios. The dark red band is the range in the percentage undiagnosed from the ECDC tool and the lighter band is the overall range in the percentage undiagnosed once the uncertainty in the number diagnosed is included. The black point and error bar show the point estimate and 95% confidence interval for percentage undiagnosed nationally from the COUNT study. (B) Estimated number of new infections from the ECDC HIV modelling tool with people previously diagnosed overseas (blue) and with those people excluded (red). The blue and red points correspond to the respective number of notifications attributed to male-to-male sex. The black line is the best estimate used for the analysis. The HIV cascade for Australian GMb during 2004 to 2015 with the number (C) and proportion (D) of GMb living with HIV who are undiagnosed, diagnosed but not on ART, on ART but with detectable (unsuppressed) VL, and on ART with undetectable (suppressed) VL. (Time trends for each population and uncertainty in the estimates are shown in Data S1).
We assumed uniform priors for the transmission coefficients for the non-suppressed steps with a range large enough to capture the plausible values for these parameters (capturing the uncertainty in transmission and the cascade estimates; see Table 1). For the suppressed population we used an exponential distribution with a 95% confidence interval 0 to 0.0084 based on the results from the PARTNER Study [35]. Using the specified priors, we applied a sampling-importance-resampling algorithm to determine the posterior probability distributions (posterior) for each transmission coefficient (described in Data S1). We took 5 million samples of the prior distributions and then re-sampled the resulting set of parameters 100,000 times with probability proportional to the calculated sampling weights to approximate the posterior distribution for each transmission coefficient.

We initially applied this procedure assuming constant transmission coefficients. However, we obtained a better fit to new infections by assuming a linear change from a start value in 2004 to an end value in 2015. This is the simplest time varying assumption but is reasonable as relevant behavioural indicators (such as condomless anal intercourse with casual partners, partner numbers and testing rates) have changed gradually and linearly over time [36-39]. For each transmission coefficient, we sampled two priors from the same distribution, one for the start value in 2004, and one for the end value in 2015. We assumed the transmission coefficient related to suppressed virus was constant over time as the current evidence suggests the transmission probability from an infected person with suppressed virus is close to zero [35,40]. This means changes in behaviour (such as a reduction in condom use) will have a minimal effect on transmission from the suppressed population. Data S1 of the Supplementary Material show the priors and the resulting posteriors.

We used the mean or median of the full posterior set to produce all the results with the 2.5th and 97.5th percentiles used to produce 95% credible intervals (95% CrIs). We rounded the final cascade estimates to the nearest whole number.

2.4 Sensitivity analysis

To assess the effect of changing the suppressed transmission assumption and the robustness of our methodology we ran four alternative scenarios: (1) using a prior that describes a relative reduction in transmission for the suppressed population compared to the diagnosed population which was informed by the Cohen et al. study in 2011 [40]; (2) assuming GBM living with HIV with suppressed virus are not infectious; (3) assuming no uncertainty in the HIV cascade estimates for 2004 to 2015; and (4) the methodology for the 2004 to 2014 Australian GBM HIV cascade estimates which assumed a lower overseas migration rate and defined a viral load <400 copies/mL for suppression [41]. We describe these scenarios in more detail and provide the complete results of these analyses in Data S1 respectively of the Supplementary Material.

3 RESULTS

The percentage of GBM with undiagnosed HIV infection in Australia decreased from 14.5% (range 11.2% to 18.7%) to 7.5% (range 5.0% to 11.0%) during 2004 to 2015 with the number undiagnosed declining slightly (Figure 1A, C, Table 2). The estimated percentage undiagnosed was similar to that observed in a 2014 national prevalence study with a range within the 95% confidence interval of the study (Figure 1A) [19]. This fall in the proportion with undiagnosed HIV was mainly due to the 47% increase in the number of GBM living with HIV (i.e. an increasing denominator; Figure 1C, Table 2). Figure 1C shows there was a substantial increase in the number and proportion of GBM living with HIV on treatment and with suppressed virus (see Table 2). Despite the level of viral suppression increasing, the estimated number of new infections rose 14.5% according to our ECDC model estimates (Figure 1B; Data S1) – although the number of new infections per 1000 GBM living with HIV decreased from 49.3 to 36.2 (Table 2). We provide all estimates and ranges for the HIV cascade steps and new infections in Data S1 of the Supplementary Material.

The annual posterior mean for the number of new infections closely aligns with the ECDC HIV Modelling Tool estimates during 2004 to 2013 (Figure 2A). The model posterior estimates were below the new infections estimates for 2014 and 2015 but are still well within the 95% confidence intervals and occurred when the ECDC HIV Modelling Tool had the widest uncertainty.

Despite large uncertainty in the estimates (according to the 95% credible intervals), there was a large increase in the number and proportion of infections attributable to undiagnosed infection during 2004 to 2015 (Figure 2B, Table 2), even with a small decline in the number of GBM with undiagnosed infection (see Data S1). The corresponding percentage of new infections attributable to undiagnosed infection increased from 33.2% (95% CrI: 2.4% to 80.8%) in 2004 to 59.1% (95% CrI: 20.9% to 89.0%) in 2015. The number and percentage of new infections attributed to diagnosed and unsuppressed GBM decreased during 2004 to 2015 (Table 2). New infections attributed to GBM on ART with suppressed virus was very small (<7%) but increased during 2004 to 2015 (see Table 2). This is due to the large increase in the number of virally suppressed GBM (Figure 1C) and a small but non-zero posterior probability of transmission from this group. We provide results from the four alternative assumptions for transmission while virally suppressed in Data S1 of the Supplementary Material.

Figure 3 shows the annual posterior distributions (as box-plots) for the percentage of new infections acquired from GBM in each step of the HIV cascade. The resulting posteriors show large uncertainty, reflecting the lower and upper bounds in the cascade estimates and new infections. This uncertainty also reflects our fitting methodology as we fitted seven model parameters using only 12 time points. Except for the suppressed step, this uncertainty generally reduced over time. The posterior peak for the percentage acquired from suppressed GBM remained relatively stable during 2004 to 2015, however, the variance increased resulting in the mean percentage increasing fourfold (see Table 2 and Figure 3).

Figure 4 shows the change in the rate of transmission per 1000 people living with HIV in each cascade step (reflecting the fitted posterior for each transmission coefficient β). While the posteriors for each step showed large uncertainty, we estimated a relatively small change in the mean rate of transmission for diagnosed, unsuppressed and suppressed GBM during 2004 to
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Symbol</th>
<th>Prior distribution</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rate of transmission from GBM with undiagnosed HIV infection to susceptible GBM (per 1000 people)</td>
<td>$\beta_u$</td>
<td>Uniform [3.9, 2105]</td>
<td>Assumed broad range based on the number of HIV notifications per person living with HIV as described for $\beta_u$ (in the row below) multiplied by a factor reflecting potential changes in sexual behaviour post-diagnosis and a likely higher viral load during early infection pre-diagnosis. Generally GBM reduce behaviour associated with HIV transmission [8,45,46], however, we assume this factor ranges from 0.9 reflecting the potential for some GBM to increase sexual activity post-diagnosis, to 30 which reflects higher levels of behaviour contributing to the risk of acquiring HIV [8,45,46] and the relative increase in transmissibility due to early infection [40,47,48]. The resulting upper and lower bounds are then multiplied by the mean relative difference between the point estimate and lower and upper bounds of the range (0.86 to 1.17) for the number of people with undiagnosed infection over 2004 to 2015. Lower value $=$ 1000 $\times$ 0.005 $\times$ 0.9 $\times$ 0.86; upper value $=$ 1000 $\times$ 0.06 $\times$ 30 $\times$ 1.17.</td>
</tr>
<tr>
<td>Rate of transmission from GBM with diagnosed HIV infection not on ART to susceptible GBM (per 1000 people)</td>
<td>$\beta_d$</td>
<td>Uniform [3.5, 78]</td>
<td>Assumed range based on the number of HIV notifications per person living with HIV from Australia’s Annual Surveillance Report [41]. This has ranged between 6 and 4 new diagnoses per 100 people living with diagnosed HIV. Given this includes people on ART and with suppressed virus we assume an upper bound of 6 per 100 people. For the lower bound we assume a small rate (0.5 per 100 people) to reflect the potential reduction in sexual activity initially post diagnosis. These rates were then multiplied by the mean relative difference between the point estimate and lower and upper bounds of the range (0.7 to 1.3) for the number of people diagnosed but not on ART over 2004 to 2015. Lower value $=$ 1000 $\times$ 0.005 $\times$ 0.7; upper value $=$ 1000 $\times$ 0.06 $\times$ 1.3.</td>
</tr>
<tr>
<td>Rate of transmission from GBM on ART but with unsuppressed virus to susceptible GBM (per 1000 people)</td>
<td>$\beta_u$</td>
<td>Uniform [0.08, 203]</td>
<td>Assumed broad range based on the range in the number of HIV notifications per person living with diagnosed HIV as described for $\beta_u$ (in the row above) multiplied by a factor reflecting potential changes in sexual activity upon the start of ART. We assume this factor ranges from 0.05, reflecting the potential for some GBM to almost stop sexual activity, as they are aware they have not achieved viral suppression, to 2 reflecting a potential increase in sexual behaviour due to a belief that ART prevents transmission. The resulting upper and lower bounds are then multiplied by the mean relative difference between the point estimate and lower and upper range (0.33 to 1.7) for the number of people on ART with unsuppressed virus over 2004 to 2015. Lower value $=$ 1000$\times$0.005$\times$0.05$\times$0.33; upper value $=$ 1000$\times$0.06$\times$2$\times$1.7.</td>
</tr>
<tr>
<td>Rate of transmission from GBM on ART but with suppressed virus to susceptible GBM (per 1000 people)</td>
<td>$\beta_s$</td>
<td>Exponential: Mean: 1/2.8</td>
<td>Assumed distribution based on the results of the PARTNER study that reported 0 transmissions due to any sex between serodiscordant men who have sex with men (with the HIV-positive partner having suppressed virus) in 418 couple-years of follow-up [35]. Using the exact Poisson method, the study estimated the upper limit of the 95% confidence interval to be 8.4 transmissions per 1000 couple-years. We used an exponential prior with a mean equal to 2.8 transmissions per 1000 people such that the 95% quantile equals 8.4 transmissions per 1000 people. Note samples from this prior were multiplied by sampled values from a uniform distribution with a range given by the mean relative difference between the point estimate and lower and upper range (0.89 to 1.12) for the number of people on ART with suppressed virus over 2004 to 2015 to capture the uncertainty in the number with suppressed virus.</td>
</tr>
</tbody>
</table>
Table 2. Cascade estimates, transmission coefficients and new infections attributed to each step of the Australian GBM HIV cascade for 2004, 2010 and 2015

<table>
<thead>
<tr>
<th>Population</th>
<th>Year</th>
<th>Number of people (mean, range)</th>
<th>Percentage of all people living with HIV (mean, range)</th>
<th>Transmission coefficient (transmission per 1000 people; mean, 95% CrI)</th>
<th>Number of new infections (mean, 95% CrI)</th>
<th>Percentage of all new infections (mean, 95% CrI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Undiagnosed</td>
<td>2004</td>
<td>1880 (1590 to 2190)</td>
<td>14.5% (11.2% to 18.7%)</td>
<td>110 (8.7 to 280)</td>
<td>212 (16 to 521)</td>
<td>33.2% (2.4% to 80.8%)</td>
</tr>
<tr>
<td></td>
<td>2010</td>
<td>1690 (1420 to 2010)</td>
<td>10.6% (8% to 14%)</td>
<td>–</td>
<td>358 (147 to 559)</td>
<td>49.5% (20.1% to 77.1%)</td>
</tr>
<tr>
<td></td>
<td>2015</td>
<td>1440 (1080 to 1860)</td>
<td>7.5% (5% to 11%)</td>
<td>290 (92 to 470)</td>
<td>423 (133 to 678)</td>
<td>59.1% (20.9% to 89%)</td>
</tr>
<tr>
<td>Diagnosed not on ART</td>
<td>2004</td>
<td>4430 (2980 to 5850)</td>
<td>34.1% (21% to 49.8%)</td>
<td>31 (3 to 81)</td>
<td>136 (13 to 360)</td>
<td>21.4% (2% to 57.3%)</td>
</tr>
<tr>
<td></td>
<td>2010</td>
<td>4370 (2070 to 6660)</td>
<td>27.3% (11.7% to 46.3%)</td>
<td>–</td>
<td>165 (40 to 314)</td>
<td>22.7% (5.7% to 43.2%)</td>
</tr>
<tr>
<td></td>
<td>2015</td>
<td>2380 (0 to 5790)</td>
<td>12.5% (0% to 34.3%)</td>
<td>43 (3.7 to 93)</td>
<td>103 (9 to 222)</td>
<td>15% (1.2% to 34.3%)</td>
</tr>
<tr>
<td>On ART but unsuppressed</td>
<td>2004</td>
<td>2750 (1720 to 3740)</td>
<td>21.2% (12.2% to 31.8%)</td>
<td>100 (11 to 200)</td>
<td>282 (29 to 562)</td>
<td>43.9% (4.6% to 84.6%)</td>
</tr>
<tr>
<td></td>
<td>2010</td>
<td>1620 (0 to 3500)</td>
<td>10.1% (0% to 24.3%)</td>
<td>–</td>
<td>177 (43 to 334)</td>
<td>24.5% (5.9% to 46.7%)</td>
</tr>
<tr>
<td></td>
<td>2015</td>
<td>1200 (0 to 4410)</td>
<td>6.3% (0 to 26.1%)</td>
<td>110 (5.2 to 260)</td>
<td>138 (6 to 308)</td>
<td>19.7% (0.9 to 45.8%)</td>
</tr>
<tr>
<td>On ART and suppressed</td>
<td>2004</td>
<td>3920 (3450 to 4390)</td>
<td>30.2% (24.4% to 37.3%)</td>
<td>2.5 (0.059 to 9)</td>
<td>10 (0 to 35)</td>
<td>1.5% (0 to 5.6%)</td>
</tr>
<tr>
<td></td>
<td>2010</td>
<td>8340 (7390 to 9300)</td>
<td>52.1% (41.8 to 64.7%)</td>
<td>–</td>
<td>24 (3 to 70)</td>
<td>3.3% (0.4 to 9.7%)</td>
</tr>
<tr>
<td></td>
<td>2015</td>
<td>1,4050 (12,440 to 15,740)</td>
<td>73.7% (58.3% to 93.2%)</td>
<td>3.2 (0.086 to 12)</td>
<td>44 (1 to 167)</td>
<td>6.2% (0.2 to 22.5%)</td>
</tr>
<tr>
<td>Total</td>
<td>2004</td>
<td>12,980 (11,760 to 14,160)</td>
<td>100%</td>
<td>49.3 (39.3 to 61.6)</td>
<td>640 (556 to 724)</td>
<td>100%</td>
</tr>
<tr>
<td></td>
<td>2010</td>
<td>16,015 (14,380 to 17,680)</td>
<td>100%</td>
<td>–</td>
<td>725 (645 to 805)</td>
<td>100%</td>
</tr>
<tr>
<td></td>
<td>2015</td>
<td>19,070 (16,890 to 21,340)</td>
<td>100%</td>
<td>36.2 (25.3 to 50.8)</td>
<td>690 (539 to 858)</td>
<td>100%</td>
</tr>
</tbody>
</table>

Cascade estimates rounded to nearest 10. Number of new infections rounded to nearest whole number. Transmission coefficients rounded to two significant figures. 95% CrI = 95% credible interval. Only 2004 (start) and 2015 (end) values for the transmission coefficient shown. Results for all years are provided in Data S1 in the Supplementary Material.

2015 (39% increase in the mean for diagnosed, 10% increase for unsuppressed and 28% increase for suppressed; see Table 2 and Data S1). In contrast, we estimated a large increase in the rate of transmission from undiagnosed GBM from a mean of 110 (95% CrI: 8.7% to 280) to 290 (95% CrI: 92% to 470) per 1000 undiagnosed GBM during 2004 to 2015, a 2.6-fold increase over the 12 years. While each transmission coefficient increased during 2004 to 2015 the number of infections per 1000 people living with HIV decreased from 49.3 (95% CrI: 39.3% to 61.6) to 36.2 (95% CrI: 25.3 to 50.8) due to the large increase in the suppressed population. Prior and posterior distributions for the transmission coefficients are in Data S1 of the Supplementary Material.

Our results were relatively robust to changes in the suppression prior and the HIV cascade estimates. Using a suppressed prior based on the relative reduction in transmission compared to the diagnosed step from the Cohen 2011 et al. study [33], the percentage of infections attributed to undiagnosed GBM increased from a mean of 33.3% (95% CrI: 2.2% to 81.3%) in 2004 to a mean of 57.5% (95% CrI: 13.9% to 91.1%) in 2015. This scenario attributed a similar percentage of new infections to suppressed GBM as for the main analysis. When we assumed suppressed GBM were not infectious the percentage of infections attributed to undiagnosed GBM increased more rapidly from a mean of 31.8% (95% CrI: 2.4% to 80.9%) in 2004 to a mean of 65.3% (95% CrI: 28% to 93.1%) in 2015. Assuming a zero range in the estimates for each cascade step resulted in different posteriors for the percentage of infections attributed to undiagnosed GBM with a mean of 47.7% (95% CrI: 13.4% to 83.3%) in 2004 and a mean of 73.6% (95% CrI: 51.3% to 91.7%) in 2015. However, the posterior distributions still overlap the posteriors obtained in our main analysis. Removing the uncertainty in the cascade estimates essentially reduced the variance in the posteriors, as expected, and increased the estimated contribution of undiagnosed to new infections. Finally, applying our methodology to the 2004 to 2014 HIV cascade estimates (which are slightly higher for each step due to a different methodology) only produced a marginally higher estimate for the percentage of new infections from undiagnosed GBM. We provide full details and figures for all the sensitivity results in Data S1 of the Supplementary Material.

4 | DISCUSSION

We assessed the contribution and role of GBM with undiagnosed HIV to new infections in Australia during 2004 to 2015 using a simple transmission model and Bayesian fitting procedure. Despite a 50% increase in the proportion of GBM living with HIV virally suppressed and the proportion of GBM living with HIV undiagnosed almost halving between 2004 and 2015, the estimated number of new HIV infections rose 8% (consistent with trends in HIV notifications attributed to male-to-male sex). It is likely that most new HIV infections among GBM in Australia are now attributable to those who are undiagnosed (59%, 95% CrI: 20.9% to 89.0%), having grown from around a third of infections in 2004. As the proportion
attributable to undiagnosed HIV has increased, the number of new infections has increased offsetting the infections prevented through reductions in community viral load.

A large increase in the proportion of infections attributed to undiagnosed GBM is potentially unsurprising given the large increase in treatment and viral suppression among Australian GBM during 2004 to 2015. Such an increase might not necessarily be due to an associated increase in the transmission coefficient for undiagnosed GBM (it is possible, depending on the cascade population sizes, for the proportion of new infections attributed to undiagnosed GBM to increase while the average rate of transmission from this population decreases). The large increase in the transmission coefficient for undiagnosed GBM seen in our analysis could be due to several reasons. First, the overall population of GBM living with HIV has increased substantially since 2004 in Australia and people living with HIV are now significantly more likely to be receiving effective treatment and much less likely to be infectious. Second, condomless sex among GBM in Australia has become more common over the last decade [36,37], so men with undiagnosed HIV are more likely to engage in sexual practices that facilitate transmission. There has also been an increase in the use of risk reduction practices relying on an accurate knowledge of serostatus by HIV-negative men, such as serosorting [37,42]. If undiagnosed GBM believe they are HIV-negative and engage in serosorting or other risk reduction strategies (and do not use condoms),
then sexual transmission is more likely to occur. Third, there has been a reduction in time between HIV infection and diagnosis during 2004 to 2015 as HIV testing frequency among GBM has increased [38,39], with the ECDC model estimating a decline from a mean time of 2.8 years to 1.9 years between infection and diagnosis. At a population level this means a larger proportion of undiagnosed GBM will be in the acute phase of infection and the average viral load (and resulting infectiousness) of undiagnosed GBM will likely have increased during 2004 to 2015 [43].

An advantage of our approach is it uses annually updated surveillance data and cascade estimates and is based on a simple transmission model. Our methodology does not require detailed demographic, behavioural and clinical data and can quickly produce estimates. It is also flexible, being able to incorporate changed assumptions, priors or alternative input.

The disadvantage of this simplicity is that we cannot understand the potential casual factors affecting temporal changes across the cascade steps. Our estimates also have several limitations. The number and proportion of infections attributed to each step of the cascade have a large amount of uncertainty due to the range in cascade estimates, the use of wide uninformative priors, and because we are fitting 7 parameters with only 12 data points. If better cascade estimates for each step of the GBM cascade were available with narrower priors, then better estimates with smaller ranges would be obtained. We also assumed a linear change in transmission coefficients over time. This assumption appears to be reasonable, as changes in behaviour, such as the increase in condomless sex [36,37] and changes in HIV testing frequency among GBM have changed relatively linearly over time, and is likely robust over short periods. However, this assumption may be
stretched over the 12-year period used in this analysis as indicated by the poorer, but still acceptable, fit to new infections for 2014 to 2015. The flexibility of our methodology means we could use different assumptions for the change in the rate of transmission for each step of the cascade over time; however, the number of parameters would need to be minimal to prevent over-parameterization.

5 | CONCLUSIONS

Our study suggests most new infections in Australia are now due to transmissions from undiagnosed GBM. This concentration of new infections from undiagnosed GBM reflects the success of interventions aimed at increasing HIV treatment coverage in the growing population of GBM with diagnosed HIV. However, the rise in notifications during 2004 to 2015 highlights the need to do more to reduce new infections. Our results suggest the need for a renewed focus on interventions for HIV-negative GBM while maintaining treatment scale-up and retention in care for people already living with HIV. The increasing contribution of undiagnosed GBM to new infections highlights the importance of HIV testing and the role of novel diagnostic services (such rapid and self-testing) to reduce the time between infection and diagnosis [44]. It also highlights the need for improved prevention methods for GBM at high risk of infection.

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COMPELING INTERESTS

The authors have no competing interests to declare.

AUTHORS’ CONTRIBUTIONS

RTG developed the model and performed the modelling analyses with support from DPW. RTG wrote the modelling code and set up the online code repository. RTG took primary responsibility for drafting and redrafting the manuscript, with support from MH, DPW and RG. All authors read, commented on and approved the final manuscript.

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REFERENCES


SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article:
Data S1. Methodological details.