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**Title:** Behavioural not biological factors drive the HCV epidemic among HIV-positive MSM: HCV and HIV modelling analysis including HCV treatment-as-prevention impact

**Running Head:** Modelling HCV in HIV-infected MSM

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**Tables:** 2  **Figures:** 5

**Abbreviations:** HCV, hepatitis C virus; HIV, human immunodeficiency virus; MSM, men who have sex with men; DAA, direct acting antiviral;
Declaration of interests: NKM and PV has received research grants from Gilead, and NKM has received honoraria from Merck, AbbVie, and Gilead. LM has nothing to report.

Author contributions: PV and NKM designed the study. CM undertook preliminary model analyses. LM undertook the statistical analyses, model development, simulations and analyses. PV, NKM, and LM wrote the first draft of the article. LM, CM, FH, PW, MH, NKM, and PV interpreted the data, edited the article, and approved of the final version.

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**Abstract - 202**

**Background:** Uncertainty surrounds why hepatitis C virus (HCV) is concentrated amongst HIV-positive men who have sex with men (MSM). We used mathematical modelling to explore reasons for these infection patterns, and implications for HCV treatment-as-prevention.

**Methods:** Using a joint MSM HIV/HCV transmission model parameterised with UK behavioural data, we considered how biological (heightened HCV infectivity and reduced spontaneous clearance among HIV-positive MSM) and/or behavioural factors (preferential sexual mixing by HIV-status and risk heterogeneity) could concentrate HCV infection in HIV-positive MSM similar to commonly observed (5-20 times the HCV prevalence in HIV-negative MSM; defined as the HCV-ratio). We explored HCV treatment-as-prevention impact under differing HCV-ratios.

**Results:** Biological factors produced low HCV-ratios (<3), not explaining the skewed epidemic. However, combining preferential mixing by HIV-status with sexual risk behaviour heterogeneity produced high HCV-ratios (>10) that were highly sensitive to both factors. Irrespective of the HCV-ratio or behavioural/biological factors, HCV treatment of HIV-diagnosed MSM markedly reduced HCV prevalence among HIV-positive MSM, but less impact was achieved among all MSM for lower HCV-ratios.

**Conclusion:** Sexual behaviour patterns likely drive observed HCV infection patterns among HIV-positive MSM. Changes in these patterns could disseminate HCV amongst HIV-negative MSM, limiting the impact of targeting HCV treatment to HIV-diagnosed MSM.

**Key words:** hepatitis C virus, HIV, men who have sex with men, antiviral treatment, prevention
Key messages

Biological factors alone do not explain why the HCV epidemic has been strongly concentrated among HIV-positive MSM;

Sexual behavioural risk heterogeneity and HIV preferential mixing amongst sexual partners is likely to explain this observation. Changes in sexual mixing patterns could reshape the epidemic.

Targeted HCV treatment-as-prevention amongst HIV-diagnosed MSM could be an important tool for combating the current HCV epidemic, but will have less impact in settings which have, or develop, a substantial burden of HCV among HIV-negative MSM, underscoring the importance of monitoring in this population.
Introduction

An epidemic of hepatitis C virus (HCV) continues unfolding amongst HIV positive men-who-have-sex-with-men (MSM) in the UK, Europe, the US and Australia[1]. HCV is a leading non-AIDS cause of death among MSM with HIV[2]. The incidence of HCV among HIV-positive MSM is generally 5-20 times higher than HIV-negative MSM[1, 3]. In the UK, HCV seroprevalence among HIV-negative MSM was estimated to be 1.2% (0.6-2.1%) in 2009[4], but was 9.9% amongst HIV-positive MSM in 2012[5].

Behavioural and biological factors have been proposed to account for the large discrepancy in HCV burden between HIV-positive and HIV-negative MSM.[6, 7] Biological factors include reduced chances of spontaneously clearing the HCV virus[8, 9] and higher HCV viral loads, potentially leading to greater infectivity in HIV-positive MSM[10, 11].

Behavioural factors include sexual partner selection based on HIV status and heterogeneity in sexual risk[6, 12]. Heterogeneities in risk and precautionary behaviours could incorporate differences in numbers of sexual partners and use of condoms, different preferences for ano-brachial insertion (fisting) and injecting illicit drugs[6, 13-15]. Additionally, partner selection and risk behaviours may interact, such as reduced condom use occurring between couples that assume they are sero-concordant[12]. These heterogeneities in sexual risk and mixing could lead to groups with higher-risk, and thus greater sexually transmitted infection (STI) and HIV prevalences.

We developed a dynamic joint HIV/HCV transmission model among MSM to explore the contribution of behavioural and biological factors to why HCV is concentrated among HIV-positive MSM. We assessed how variations in these biological and behavioural factors’ may
affect the HCV distribution, and evaluated the resulting implications for HCV treatment-as-prevention[16].

**Methods**

*Model Derivation*

A dynamic, deterministic model of HIV and HCV transmission among MSM was developed. We divided MSM into compartments (model schematic in supplementary figure S1) defined by HIV status (susceptible, undiagnosed acute infection, undiagnosed chronic infection and diagnosed HIV infection), HCV status (susceptible, chronic HCV, chronic HCV treatment failure) and low and high sexual risk groups, based on annual numbers of partners that MSM have anal sex with. A simplified model was designed, that was not intended to rigorously simulate the historical HIV epidemic as done in other studies[17].

MSM enter the model when they reach sexual maturity and exit though death or ageing out the model at 65 years of age. The model is dynamic, such that the risk of an individual acquiring HIV or HCV is related to the background prevalence of that infection, which can change over time. We assume both diseases are transmitted through sexual episodes, which may involve unprotected anal intercourse, injecting drug use and fisting[18-21]. Because individuals with more anal intercourse partners (whom we deem ‘high-risk’) also have a higher frequency of injecting drugs, fisting, and other high-risk behaviours, high-risk MSM were assumed to have higher chances of HIV and HCV exposure[6, 13-15].

Once HIV infected, individuals enter undiagnosed acute HIV infection, with heightened HIV infectivity[22], subsequently transitioning to undiagnosed chronic HIV infection. Individuals
are assumed to not become HIV diagnosed during the short (2.6 months) acute phase of HIV infection[22]. On diagnosis, MSM transition to diagnosed chronic HIV, where a proportion receive HIV antiretroviral treatment (ART) which we assume reduces HIV infectivity[23, 24] and increases survival[25, 26].

Newly HCV-infected MSM that don’t spontaneously clear HCV transition to chronic HCV. The HCV acute phase was not included due to its likely small contribution to HCV transmission[27]. Those clearing HCV remain susceptible. When included in our analysis, HCV treatment is assumed to cure infection for a proportion of individuals but not confer immunity. Successfully treated MSM remain at risk of reinfection. Unsuccessfully treated MSM move into the treatment failure class and remain infected with HCV.

To explore the implications of HIV infection on HCV infection/transmission patterns, we consider scenarios where HIV has no effect on HCV and the alternative; where HIV infection increases HCV liver-related progression and mortality[28], reduces spontaneous clearance[8, 9] and increases HCV infectivity[10, 11]. We assume HCV does not impact on HIV disease progression or ART response[25].

Model Parameterization

We modelled HIV and HCV transmission among sexually active MSM aged 15-65, parameterizing sexual behaviour with the UK component of the European MSM Internet Survey (EMIS-UK)[29]. EMIS was an online survey undertaken during June-August 2010, recruiting online and promoted offline through print media. Over 18,000 MSM living in the UK participated. From EMIS-UK data, we calculated the proportion of HIV-diagnosed MSM’s
sexual partners they assumed were HIV-positive (36.2%) and condom use in these pairings (13.0% in last sex act) compared to (68.0% in last sex act) other MSM partnerships. EMIS-UK data also determined the heterogeneity in frequencies of casual sexual partnerships. When risk heterogeneity was explored, we divided the MSM population into categories of low and high-risk by the annual number of sexual partnerships, 14 or less and 15 or greater, respectively, with 82.2% and 17.8% in the low and high-risk groups, respectively. In some scenarios, an additional risk was also associated with MSM in the high-risk group due to EMIS-UK data suggesting a greater proportion of these MSM either inject drugs (3.6% versus 1.0% amongst low-risk MSM), or undertake receptive (21.1% versus 8.6%) or insertive fisting (38.7% versus 14.0%) in the last year[6, 13-15]. Biological parameters were obtained from the literature. HCV treatment was not included in the baseline model because we were not aiming to closely model the precise HCV epidemic in the UK, and it was not considered to be an important determinant of observed epidemic patterns at existing treatment rates[5].

All model parameters are outlined in Tables 1 and 2.

Model fitting and scenarios

For each different behavioural and biological risk factor scenario (detailed below), the model was calibrated to a stable HIV and HCV prevalence. The model was run with a non-least squares fitting algorithm which took point values of all parameters relevant to the scenario (shown in Table 2), except the transmission parameters for HCV and HIV which were used to fit the simulation. We calibrated the model to a 5% HIV prevalence among MSM[30] and a chronic HCV prevalence of 10% amongst HIV-infected MSM[16]. This approach gives a simplified characterisation of the HIV and HCV epidemic among MSM in
the UK. We did not fit the HCV prevalence amongst HIV-uninfected MSM. However, we explored several scenarios involving the inclusion of various biological and behavioural factors to see how they affected the HCV prevalence amongst HIV-uninfected MSM, while for each scenario assuming a 10% HCV prevalence amongst HIV-infected MSM:

1. **Baseline**: No effect of HIV infection on HCV progression, transmissibility or spontaneous clearance; no heterogeneity in sexual risk behaviour or HIV preferential mixing among MSM.

2. **Biological factors only**: Infection with HIV reduces HCV spontaneous clearance probability, increases HCV-related mortality, and increases HCV infectivity.

3. **Mixing by HIV status with biological factors**: MSM select partners preferentially based on HIV status with errors in judgement, with an additional sub-scenario assuming less condom usage among partnerships where HIV-diagnosed individuals think their partner is also HIV-positive (irrespective of whether right or not). Biological factors included as above.

4. **Heterogeneity in sexual risk behaviour with biological factors**: Heterogeneity in sexual risk behaviour based on number of casual partners. Two additional sub-scenarios further assume that a) MSM select partners preferentially based on risk group, or b) MSM select partners preferentially based on risk and assume further elevated transmission risk associated with high-risk MSM based on their higher prevalence of injecting drugs and fisting. Biological factors included as above.
5. **All factors:** Mixing by HIV status and heterogeneity in sexual risk included as described above with all associated effects from previous scenarios. Biological factors included as above.

*Model analyses and sensitivity analyses*

**Impact on the HCV ratio:** To explore the impact of these scenarios on the HCV relative burden among HIV-positive MSM, we define the “HCV ratio” as the chronic prevalence of HCV in HIV-positive MSM divided by the chronic prevalence of HCV in HIV-negative MSM. We firstly use point values for each parameter (Table 2) and assess whether each scenario produces a HCV ratio commonly observed in the UK and other settings (HCV ratio >5)[1, 3, 4, 16]. Then, to test the model’s sensitivity to parameter variation, for scenario 5 (All factors included), we undertook a univariate sensitivity analysis where we varied each parameter individually across +/-100% of their point value, and assessed the effect on the HCV ratio. These wide parameter uncertainty ranges were used to account for unknown biases and uncertainties in the data, with the same range being assumed for each parameter to see how each affected the results over the same relative range. We then performed bivariate sensitivity analyses on key parameters identified at the univariate level, quantifying their importance for three different levels of error in judgement of HIV status of sexual partners (-100%, 0% and +100% of point value).

**Impact on HCV treatment-as-prevention initiatives:** We explored the impact of HCV treatment-as-prevention for the different scenarios (Table 2). For each, we assessed the 10-year decrease in chronic HCV prevalence amongst HIV-positive MSM and all MSM achieved for an illustrative HCV treatment intervention that annually treated 10% of HIV-diagnosed
HCV co-infected MSM. We assumed a 90% sustained viral response (SVR) with interferon-free direct acting antiviral therapy (DAA)[31]. By simultaneously sampling (5000 iterations) all the parameters varied for the univariate sensitivity analysis undertaken on scenario 5, we then considered the effect of variations in the HCV ratio on the impact of the illustrative HCV treatment-as-prevention strategy. Lastly, for scenario 5 (All factors), we individually varied key parameters across +/-100% of the point value to assess their influence on the reduction in chronic HCV prevalence achieved with treatment.

Results

HCV Ratio Analysis

Model projections of the HCV ratio for the different scenarios in Table 2 are shown in Figure 1. If no biological or behavioural factors are included (Scenario 1), the predicted HCV ratio is low but greater than one (1.39) due to MSM entering the model being susceptible to both diseases, so creating an increased proportion of HIV-negative MSM without HCV. Including biological factors only (Scenario 2) marginally elevates the HCV ratio (1.41) because the greater HCV transmissibility in HIV-HCV co-infected MSM increases HCV transmission in both HIV-negative and positive MSM. Similarly, including preferential mixing by HIV status (Scenario 3) cannot reproduce the high HCV ratio observed in the UK (HCV ratio of 5-20), with modelling projecting an HCV ratio of 1.7, which increases to 2.2 with inclusion of lower condom use in partnerships where HIV-diagnosed individuals assume their partner is HIV-positive.
In contrast, higher and more commonly observed HCV ratios (>5) are achieved through including heterogeneity in sexual risk behaviour (scenario 4). For instance, stratifying MSM into low and high-risk groups based on the number of casual partners, with greater injecting drug use and fisting among high-risk individuals, and preferential mixing between these groups produces a HCV ratio of 9.7.

Lastly, combining all behavioural and biological factors (scenario 5) amplifies the HCV ratio to 19.7, with different factors acting synergistically to transmit HCV amongst HIV-positive MSM but not HIV-negative MSM.

Univariate and bivariate sensitivity analyses on the HCV ratio

Univariate variations of parameters in scenario 5 around their point values (+/-100% - Table 2) identified four key parameters that have most effect on the HCV ratio (figure 2): (1) proportion of individuals preferentially mixing by HIV status (HCV ratio varies from 9.7-43.9); (2) error in HIV status judgements (HCV ratio varies 16.2-24.4); (3) ratio difference in numbers of partners between low and high-risk MSM groups (HCV ratio varies 3.4-28.6); and (4) additional relative risk for HCV transmission in high-risk MSM due to risky sexual behaviours (HCV ratio varies 8.3-33.5). Parameters that did not affect the HCV ratio as much are shown in Supplementary figure S3.

The bivariate sensitivity analysis explored the relationship between the four most influential parameters from the univariate analysis (figure 3). The two risk heterogeneity parameters were varied simultaneously, forming one combined measure. The HCV ratio is sensitive to levels of heterogeneity in sexual risk behaviour and preferential mixing by HIV status, which
amplify each other. Indeed, the figures illustrate that HCV ratios of 5-20 are possible with high levels of risk heterogeneity alone, or moderate levels of both preferential mixing by HIV status and risk heterogeneity with any level of error in HIV status judgements. Greater error in HIV status judgements dampens the HCV ratio.

**Impact of HCV treatment-as-prevention**

Annually treating 10% of HIV-diagnosed HCV co-infected MSM for HCV over 10 years reduces HCV chronic prevalence among HIV-positive MSM by a relative 40.3-50.3% across the different scenarios (Figure 4a). However, impact among the entire MSM population varies markedly, from a relative reduction in chronic HCV among MSM of 3.5% for scenario 1 to 29.3% for scenario 5 (Figure 4b). Figure 5 illustrates this effect further with the HCV ratio having a relatively small influence on the HCV treatment-as-prevention impact among HIV-positive MSM (Figure 5a), but a large influence amongst all MSM (Figure 5b). At higher HCV ratios, more of the epidemic is concentrated among HIV-positive MSM, so focusing treatment efforts on this population effectively combats the epidemic among all MSM. Univariate variations in parameters that have a large effect on the HCV ratio also affect the impact of HCV treatment amongst HIV-positive MSM on the overall HCV epidemic (supplementary figures S4 and S5).

**Discussion**

We find biological factors alone (lower spontaneous clearance rate and higher HCV infectivity and mortality amongst HIV-infected MSM) are unable to explain why the HCV epidemic is concentrated among HIV-positive MSM. Instead, we suggest that behavioural
factors (heterogeneity in sexual risk behaviour alone or combined with preferential mixing by HIV status) are highly likely to account for the higher HCV burdens among HIV-positive MSM. Thus, HCV infection and co-infection should be seen as a marker of high sexual risk behaviours, which are preferentially undertaken within partnerships with other HIV-positive MSM[12]. This is likely to have been aided by the scale-up of effective HIV treatment improving the survival of higher-risk MSM, paired with possible increases in risk behaviour due to ‘treatment optimism’[32].

Importantly, these results highlight that changes in sexual behaviour or mixing patterns could reshape the HCV epidemic. For example, decreases in preferential mixing by HIV status could occur due to reductions in perceived risk resulting from widespread ART or PrEP use reducing HIV infectivity and susceptibility, which could increase HCV transmission amongst HIV-negative MSM. Alternatively, fewer high-risk MSM acquiring HIV (due to PrEP) may also raise the likelihood of HCV transmission among HIV-negative MSM, although this may be offset by increased HCV monitoring of MSM being prescribed PrEP.

Further, HCV treatment-as-prevention initiatives among HIV-diagnosed MSM will have greatest impact on overall levels of HCV transmission in settings where HCV is concentrated among HIV-positive MSM, as less of the epidemic is driven by HIV-negative MSM. Conversely, settings which have, or develop, a greater burden of HCV among HIV-negative MSM would also need to focus HCV treatment to the HIV-negative MSM.

Limitations

Our analysis has a number of limitations. Firstly, we utilized a simplified model of HCV and HIV transmission and ART that was calibrated approximately to the UK without recreating
historical epidemic trends, which suggest a slowly increasing HIV and HCV epidemic[5, 30]. This was because our intention was to explore qualitatively how behavioural and biological factors contribute to HCV epidemic patterns, not make detailed predictions about the epidemics’ trajectory. Importantly, this simplification should not affect the degree to which HCV propagates preferentially amongst HIV-positive MSM. A further simplification of our model involved the incorporation of injecting drug use related risk as an increased transmission risk among a subset of MSM [6, 13-15], instead of explicitly modelling injecting. We made this simplification because, although injecting drug use is a risk factor for HIV/HCV acquisition amongst MSM, it is unclear the degree to which this is due to injecting drug use itself or co-occurring high-risk sexual behaviours. Also, datasets such as EMIS only ask basic questions about undertaking injecting drug use in the last year, so preventing any explicit modelling of its role in HCV transmission amongst MSM.

Secondly, there exists uncertainty in our parameters and variation across settings, most notably amongst those related to self-reported behavioural data. We performed extensive sensitivity and scenario analyses to explore the effect of varying different behavioural factors. As such, our analyses form a platform from which to explore how variations in parameter assumptions effect observed epidemic patterns and treatment-as-prevention impact. However, care should be taken in generalising our results to non high-income settings where limited data suggests lower HCV-coinfection prevalences amongst MSM[33], and where differences in sexual behaviour and the underlying HIV and HCV epidemic are likely to heavily effect the HCV epidemic that occurs.

Thirdly, although parameterizing our model to EMIS-UK data produced realistic projections for the HCV ratio (~20), we advise caution regarding potential over-interpretation of the
quantitative accuracy of our model. For instance, the model did not incorporate all sources of HCV infection, such as amongst migrants with historic HCV infection. Conversely, compared to a national probability survey on sexual behaviours, the EMIS-UK dataset used to parameterise our model was biased towards higher-risk MSM due to their web-based convenience sampling approach[34], as well as MSM with higher education levels[34]. These MSM may have a greater interest in HIV prevention and so increased propensity to mix preferentially by HIV status and use condoms with perceived sero-discordant partners.

Finally, our estimate for increased HCV infectiousness amongst HIV-positive MSM is uncertain, although data from vertical transmission studies[11] suggests our assumption is reasonable. We assume that higher HCV viral load in blood samples amongst HIV-positive MSM translates to increased infectivity[10], but this may not be the case. However, this should not be a concern because this parameter had little effect on the resulting model projections.

Comparison with other publications

To our knowledge, this is the first modelling analysis of the joint epidemics of HIV and HCV among MSM, although many previous analyses have modelled just HIV[17] and some have also modelled other sexually transmitted infections (STI) among MSM[17, 35-38]. However, existing HIV and STI co-infection models generally considered different questions, focusing primarily on the degree to which STIs contribute to HIV transmission, and the possible impact of STI treatment on HIV epidemics. Previous analyses have also modelled the transmission of HIV and HCV amongst people who inject drugs [39-43]. Importantly, existing work by our group and others have modelled the HCV epidemic amongst HIV-diagnosed MSM, and evaluated the impact of scaling-up HCV treatment in this group[5,
These studies were limited in that they did not explicitly include HIV transmission. Our new analysis supports findings of these previous two studies by indicating that scaling-up HCV treatment among HIV-positive MSM could have substantial prevention benefits among HIV-positive MSM[16]. Additionally, it extends previous work by dynamically modelling the transmission of HCV to and from the HIV-negative population, assessing how different behavioural and biological factors could result in the observed epidemic patterns, and evaluating the implications for HCV treatment-as-prevention.

**Concluding remarks**

Overall, our work indicates that sexual risk behaviour heterogeneity and HIV preferential mixing likely explain why the HCV epidemic amongst MSM is strongly concentrated among HIV-positive MSM, with HCV co-infection possibly signifying high-risk behaviours as suggested by others[15]. Targeted HCV treatment-as-prevention amongst HIV-diagnosed MSM could be an important tool for combating the current HCV epidemic, but will have less impact in settings which have, or develop, a substantial burden of HCV among HIV-negative MSM. This could occur if sexual risk behaviours increase amongst HIV-negative MSM or if higher-risk MSM do not become HIV-infected as frequently. This underscores the importance of monitoring HCV among HIV-negative MSM, to assess any shifts in the patterns of the HCV epidemic.
References

Figures

Figure 1. Modelled HCV Ratio (ratio of HCV chronic prevalence among HIV-positive compared to HIV-negative MSM) for various scenarios incorporating different biological and/or behavioural factors as detailed in Table 2. *These scenarios also include the biological factors. IDU denotes injecting drug use.
Figure 2. Effect of univariate changes in individual parameters (A-D) on the HCV ratio for the “All effects” scenario 5. All other parameters are set to their point values in Table 2. Only those parameters that markedly affect the HCV ratio are shown. Numbers shown on x-axis are -100% of the point value, the point value and +100% of point value.
Figure 3. Contour maps showing how the HCV ratio (contour lines - produced by the “All effects” scenario 5) is affected by both the level of HIV preferential mixing (y axis) and sexual risk heterogeneity (x axis) for three levels of error in judging the HIV status of a sexual partner: (A) zero error (e=0%), (B) medium error (e=25%), (C) and high error (e=50%). All other parameters are set to their point values in Table 1. Sexual risk heterogeneity is the simultaneous variance of the ratio in the average number of partners between the low and high-risk groups and the additional relative risk for HCV transmission in high-risk MSM due to risky sexual behaviours, from -100% to +100% of their estimated point values, which vary respectively from 1 at -100% to 20.0 at +100% and 1 at -100% to 5.4 at +100%.
Figure 4. Impact of HCV treatment on the relative reduction in HCV chronic prevalence (%) among (A) HIV-positive MSM and (B) all MSM, achieved by treating 10% of HIV-diagnosed MSM with HCV per year for 10 years. Projections assume the point value of parameters for each scenario in Table 1 and assume 90% HCV treatment efficacy. *These scenarios also include the biological factors.
Figure 5. Effect of variations in the HCV ratio on the impact of HCV treatment-as-prevention (% relative reduction in chronic HCV prevalence at 10 years when treating at a rate of 10% of HIV diagnosed HCV co-infected MSM annually, y-axis) among (A) HIV-positive MSM, and (B) all MSM. We assume 90% HCV treatment efficacy, and uniformly sampled other parameters randomly between +/-100% of their point values.
<table>
<thead>
<tr>
<th>Parameter</th>
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<th>Source</th>
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<td>Excess death rate due to mono-infection with HIV untreated (annual)</td>
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<td>Decreased mortality hazard ratio for HIV mono-infection due to ART</td>
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<td>[25, 26]</td>
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<td>[28, 45]</td>
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</tr>
<tr>
<td>(When heterogeneity is turned on, low and high risk group in brackets)*</td>
<td>7.4 [2.9, 29.1]</td>
<td>EMIS</td>
<td>See supplementary material for details</td>
</tr>
<tr>
<td>Mean number of casual sex partners for anal intercourse. (When</td>
<td>2.7</td>
<td>EMIS</td>
<td>See supplementary material for details</td>
</tr>
<tr>
<td>heterogeneity is turned on, low and high risk group in brackets)*</td>
<td></td>
<td>[6, 13-15]</td>
<td></td>
</tr>
<tr>
<td>Increased overall risk ratio of HIV and HCV transmission due to</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>injecting drugs and fisting between low and high risk group*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixing parameter for choosing partners by risk behaviour category*</td>
<td>0.2</td>
<td>EMIS</td>
<td>See supplementary material for details</td>
</tr>
</tbody>
</table>

* See supplementary material for details

**Table 1: Model parameter values**
Table 2: Parameterization of the scenarios with point values shown for each model. Sub-scenarios within the main scenarios take the parameter values corresponding to the values given by * and ** in the table where relevant.