Projecting 10-yr, 20-yr and Lifetime Risks of Cardiovascular Disease in Persons Living with HIV in the US

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Running head: Projected lifetime risk of CVD in HIV

Summary:
Cardiovascular disease (CVD) is an increasing cause of morbidity among persons living with HIV (PLWH).

CVD prevention strategies could offer important health benefits for PLWH and should be evaluated.
ABSTRACT

Background

Cardiovascular disease (CVD) is an increasing cause of morbidity among persons living with HIV (PLWH). We projected cumulative CVD risk in PLWH in care compared to the US general population and persons HIV-uninfected, but at high risk for HIV.

Methods

We used a mathematical model to project cumulative CVD incidence. We simulated a male and female cohort for each of three populations: 1) US general population; 2) HIV-uninfected, at high risk for HIV; and 3) PLWH. We incorporated the higher smoking prevalence and increased CVD risk due to smoking to the HIV-infected and HIV-uninfected, at high risk for HIV populations. We incorporated HIV-attributable CVD risk, independent of smoking.

Results

For men, life expectancy ranged from 70.2-77.5 years and for women from 67.0-81.1 years (PLWH-US general population). Without ART, lifetime CVD risk for HIV-infected males and females was 12.9% and 9.0%. For males, by age 60, cumulative CVD incidence was estimated at 20.5% in PLWH in care compared to 14.6% in HIV-uninfected high risk persons, and 12.8% in the US general population. For females, cumulative CVD incidence was projected to be 13.8% in PLWH in care compared to 9.7% for high risk HIV-uninfected persons, and 9.4% in the US general population. Lifetime CVD risk was 64.8% for HIV-infected males compared to 54.8% in the US general population males, but similar among females.

Conclusions
CVD risks should be a part of treatment evaluation among PLWH. CVD prevention strategies could offer important health benefits for PLWH and should be evaluated.

**Key words:** HIV/AIDS, cardiovascular disease, lifetime risk
INTRODUCTION

Due to major advances in the treatment of HIV disease, the life expectancy of treatment-adherent people living with HIV (PLWH) is approaching that of the general population (1). Aging PLWH and their health care providers now face new challenges related to prevention and treatment of common chronic conditions, including cardiovascular disease (CVD) (2).

While numerous studies have focused on the increased risk of CVD in PLWH (3-7) as evidenced by abnormal biomarkers of chronic inflammation and abnormal lipid metabolism, these studies have not estimated lifetime CVD risk at the population level (8). Additionally, many studies compare CVD risk between PLWH and the US general population, but these studies may overestimate CVD risk attributable to HIV by not accounting for the higher prevalence of traditional CVD risk factors, such as smoking, among PLWH (9, 10).

Through simulation modeling, we estimated the lifetime CVD risk in treatment-adherent PLWH, considering competing mortality due to HIV. We compared this lifetime CVD risk to that in the general population, as well as to that in a “high risk” population without HIV infection but with an increased prevalence of behavioral risk factors associated with HIV and CVD and their increased risk of competing mortality.
METHODS

Analytic Overview:

We utilized the Cost-Effectiveness of Preventing AIDS Complications (CEPAC) model, a validated computer simulation of HIV disease that simulates both PLWH and HIV-uninfected persons (11-14). We expanded CEPAC to incorporate age- and sex-specific CVD prevalence, incidence, and attributable mortality. We simulated male and female cohorts for each of three populations: 1) the US general population; 2) HIV-uninfected, at high risk for HIV (due to behavioral risk factors); and 3) PLWH in care. The model followed each cohort from age 36 until death (15). We projected cumulative incidence of CVD in each cohort by the ages 40, 50, and 60, and over a lifetime, accounting for competing mortality from HIV and other causes.

We relied on multi-site US data to inform input parameters regarding CVD risk. To approximate the increased CVD risk in those uninfected, at high risk for HIV we assumed that they have the same traditional risk factors as PLWH. We used smoking as a proxy for traditional risk factors and incorporated data on smoking prevalence in PLWH and the increased CVD risk from smoking (16, 17). We accounted for increased CVD risk due to HIV (18). To account for higher all-cause mortality in those HIV uninfected, at high risk for HIV, we used standardized mortality ratios (SMRs) (13, 19). We separated non-CVD- and CVD-attributable mortality to approximate the time at risk for CVD acquisition more accurately. We used CVD age/sex stratified prevalence to distinguish between age-related (non-CVD) and CVD-attributable mortality. We assumed that CVD-attributable mortality is the same for all persons with CVD.
Model Structure

The Cost-Effectiveness of Preventing AIDS Complications Model

The CEPAC model is a validated computer simulation of HIV disease (11-14). In the model, each simulated HIV-infected individual transitions between health states defined by CD4 count, HIV RNA, history of opportunistic infection (OI), and antiretroviral therapy (ART) use.

Modeled PLWH initiate ART according to US guidelines upon diagnosis and linkage to care (20). ART decreases HIV RNA, increases CD4 count, and decreases OI incidence and HIV-associated mortality. Virologic failure leads to switching ART regimens. PLWH have additional HIV-attributable mortality that depends on CD4 count, history of OI, and ART complications.

The CEPAC model tracks survival in those without HIV whose survival is determined by age- and sex-stratified mortality derived from the US life tables for the general population and SMR-adjusted mortality for those at high risk for HIV acquisition. Details on the SMR adjustment and other model details are in the Technical Appendix (TA).

Incorporating CVD into the CEPAC Model:

To extend the CEPAC model to include cardiovascular morbidity and mortality, each simulated person is assigned a CVD status at model initiation, based on age- and sex-specific prevalence. Prevalent cases,
and those who develop CVD based on age- and sex-stratified CVD incidence, are subject to additional, CVD-attributable mortality over their remaining lifespan. Modeled CVD endpoints include any of the following: myocardial infarction (MI), stroke, angina, or coronary heart disease.

We derived CVD-attributable mortality from National Center for Health Statistics data, which reports deaths due to diseases of the heart and cerebrovascular diseases, stratified by age and sex (21). We divided the number of deaths due to “diseases of the heart” and “cerebrovascular diseases” by the number of persons with CVD in each age and sex group. The number of persons with CVD was estimated based on CVD prevalence and the US population size in 2008 (Table 1) (21, 22).

Cohorts Modeled

We simulated male and female cohorts from 3 distinct populations: 1) US general population, 2) HIV-uninfected, at high risk for HIV, and 3) PLWH. Each cohort differed in terms of CVD prevalence, CVD incidence over time, and age-related (non CVD-related) mortality (Table 1). We included the HIV-uninfected, at high risk for HIV cohort (assuming they never acquire HIV throughout their lifetime) to allow for comparison of their lifetime CVD risk to that of PLWH, thereby explicitly demonstrating the effect of HIV on CVD lifetime risk alone in the face of competing HIV-attributable mortality.

1) General population
**CVD prevalence (Table 1):** We derived age- and sex-stratified US general population CVD prevalence from the 2010 National Health Interview Survey (NHIS), adjusting for over-reporting in self-report data (23, 24). Details on CVD prevalence derivation are in the TA.

**CVD incidence:** We estimated age- and sex-stratified CVD incidence using the basic epidemiologic principal that \( \text{Prevalence} = \text{Incidence} \times \text{Duration of disease} \), estimating incidence as the ratio of prevalence to disease duration. To estimate CVD duration, we used CEPAC to ensure consistency in life expectancy estimates across all cohorts. We simulated HIV-uninfected individuals, by ‘turning off’ all HIV-related parameters. We assumed that, once diagnosed, CVD persists for life. The life expectancy of individuals in each 10-year age and sex stratum therefore approximates CVD disease duration for that decade of life. Additional details and validation are in the TA.

**Age-related (non-CVD) mortality:** We derived age- and sex-specific mortality from 2008 US life tables and removed CVD-attributable mortality (TA). We assumed that among persons with CVD, mortality consists of non-CVD and CVD-attributable components. Based on this assumption, we derived non-CVD mortality by subtracting CVD-attributable mortality multiplied by the CVD prevalence from total mortality for each age/sex group:

\[
\text{Non-CVD mortality} = \text{overall mortality} - \text{CVD attributable mortality} \times (\text{Prevalence of CVD}).
\]
2) **HIV-uninfected, at high risk for HIV population**

*CVD prevalence:* The increased CVD prevalence in this cohort was attributed to higher smoking prevalence (16, 17). We assumed that smoking prevalence in this cohort would be the same as among PLWH in the US (which is twice that of the general population) and stratified by age and sex (Table 1) (17). We used a relative risk of CVD compared to non-smokers of 1.9 and 1.7 for male and female smokers, from the Framingham Heart Study (16). We calculated overall age- and sex-stratified CVD prevalence in this cohort as the weighted average of CVD prevalence among smokers and non-smokers (Table 1).

*CVD incidence:* We derived CVD incidence from the age, sex, and smoking status-stratified CVD incidence in the US general population (described above), taking into consideration the higher smoking prevalence and assuming that smoking behavior does not change upon HIV diagnosis (Table 1) (TA).

*Age-related (non-CVD) mortality:* Persons at high risk for HIV may exhibit behaviors subjecting them to higher mortality, including smoking and alcohol use (2, 13, 19, 25-29). To adjust for excess mortality from these behavioral risk factors, we derived sex-specific standardized mortality ratios (SMRs) that quantified the relative change in age-related (non-CVD and non-HIV related) mortality for persons at high risk for HIV compared to the US general population; these SMRs also capture increased non-CVD and non-HIV mortality due to behavioral risk factors (e.g., mortality associated with HCV). We derived these SMRs as the average of
published risk-group-specific SMRs, weighted by the distribution of risk groups among US PLWH in care (13, 15, 19). The SMRs differed for males (1.2) and females (2.9) due to different risk group composition (TA).

3) PLWH

We derived HIV-related parameters from published literature, including demographic and clinical characteristics and treatment efficacy (Table 1). We used published data on the average age (36 years) and mean CD4 count (751 cells/µl) at seroconversion (15, 30). PLWH in the model present to care at a mean CD4 count of 351 cells/µL (31). We focused on PLWH who are treatment-adherent. (32, 33)

CVD prevalence: We estimated the prevalence of CVD among PLWH from CVD prevalence in the HIV-uninfected, at high risk for HIV cohort, but further increased to account for the additional CVD risk associated with HIV infection per se (RR=1.76) based on multi-site cohort data (18). This increased risk was independent of smoking.

CVD incidence: We derived CVD incidence in PLWH using the estimated incidence of CVD in the HIV-uninfected, at high risk for HIV population with additional adjustments for HIV-specific risk (RR=1.76) (18) (Figure 1). The HIV-specific risk factors also accounts for differences in the
prevalence of some traditional risk factors among PLWH and uninfected populations, such as glucose intolerance and dyslipidemia.

*Age-related mortality (non-CVD, non-HIV)*: Mortality in PLWH could occur due to one of three categories: HIV-attributable, CVD-attributable, or “age-related.” We derived age-related mortality in PLWH using the method described above for HIV-uninfected persons, at high risk for HIV. This age-related mortality did not include HIV-attributable mortality, which was added separately and accounts for mortality associated with chronic HIV infection (such as renal disease or anemia).

**Sensitivity Analysis:**

We varied inputs that affected CVD risk including: CD4 count at presentation to HIV care (173-516 cells/µL); smoking prevalence (35.6/30.4 – 43.7/37.3% M/F); Framingham-based relative risk of CVD due to smoking (1.7/1.4 – 2.2/2.1 M/F); the increase in risk for CVD due to HIV (1.0-2.1); SMR adjustments (1.0/1.1 – 2.1/10.6 M/F) and age of HIV infection (26-41 years) (15-19, 31).

We also estimated the lifetime CVD risk among PLWH without ART. Combining the most influential parameters in one-way sensitivity analyses, we created ‘lowest’ and ‘highest’ scenarios for CVD cumulative risk for PLWH. The ‘lowest’ risk scenario included low CD4 at presentation (173 cells/µL), lowest relative risk of CVD due to HIV (1.0), and high SMR (2.1/10.6 M/F). The ‘highest’ risk scenario included high CD4 at presentation to care (516 cells/µL), highest relative risk due to HIV (2.1), and low
SMR (1.0/1.1 M/F). For the population of HIV-uninfected, at high risk for HIV, we varied SMRs to create similar ‘lowest’ and ‘highest’ CVD risk scenarios (1.0/1.1 – 2.1/10.6 M/F).
RESULTS

Base Case

Survival

For men, life expectancy ranged from 70.2-77.5 years and for females from 67.0-81.1 years (Table 2 and Figure 2, Panels A and B).

CVD cumulative incidence

By age 60, CVD risk was the lowest for males and females in the US general population (12.8% for males and 9.4% for females), higher in the HIV-uninfected, at high risk for HIV cohort (14.6% for males and 9.7% for females), and highest amongst PLWH (20.5% for males and 13.8% for females). For males this ranking continued through their lifetimes, with projected lifetime CVD risk at 54.8% for the US general population, 59.1% for the HIV-uninfected, at high risk for HIV cohort, and 64.8% for PLWH. For females, however, after age 70, the cumulative CVD risk of the US general population began to exceed that of the HIV-uninfected, at high risk for HIV cohort. The projected lifetime CVD risk for US general population females was 46.1%, 36.7% for the HIV-uninfected, at high risk for HIV cohort, and 43.8% for the PLWH cohort (Table 2 and Figure 2 Panels C and D).

Sensitivity analyses

The parameters that had the greatest effect on lifetime CVD risk for PLWH included SMRs, the CVD relative risk multiplier due to HIV, and CD4 count at presentation to care (Figure 3, TA). Sensitivity
analyses in which we varied age at seroconversion, rates of chronic AIDS death, CVD prevalence, and smoking prevalence across the 95% CI of reported values (35.6/30.4 – 43.7/37.3% M/F) did not have a substantial impact on cumulative CVD risk. In the absence of ART, lifetime CVD risk for PLWH was 12.9% for males and 9.0% for females.

**Standardized mortality ratios (SMRs)**

Projected lifetime CVD risks for PLWH were most sensitive to SMRs. With higher SMRs (2.1/10.6 M/F), lifetime CVD risk decreased to 21.8% for HIV-infected females and 60.9% for HIV-infected males. When SMRs were reduced (1.0/1.1 M/F), lifetime CVD risk increased to 56.9% and 66.7% for HIV-infected females and males.

**HIV-related CVD relative risk multiplier**

Decreasing the relative risk of CVD due to HIV to 1.0 resulted in lower lifetime CVD risk in PLWH: 31.3% in females and 53.0% in males. When we used the upper 95% confidence interval value (RR=2.1), lifetime CVD risk increased to 47.3% in females and to 67.1% in males (18).

‘Lowest’ and ‘Highest’ CVD risk scenarios for the HIV-infected population

In the ‘lowest’ CVD risk scenario the cumulative CVD risk by age 60 was 16.7% for males and 8.8% for females; and 54.5% and 16.3% over a lifetime. In the ‘highest’ CVD risk scenario, the cumulative CVD risk by age 60 was 26.2% for males and 18.7% for females; and 76.3% and 66.8% over a lifetime.
DISCUSSION

Our model-based evaluation shows that treatment-adherent PLWH in the US have a greater risk of CVD at younger ages (under 60) compared to both the US general population and the HIV-uninfected, at high risk for HIV population. HIV-infected men remain at higher risk for CVD over their lifetimes, whereas HIV-infected women have lower lifetime CVD risk compared to the US general population. Because SMRs for both HIV-infected women and women at high risk for HIV were substantially higher than those for men, women had lower projected survival than men. This is in contrast to data in the general population where women’s life expectancy is generally longer than that of men. The higher SMR in women is explained by the fact that most women with or at risk for HIV have additional risk factors such as drug or alcohol abuse, which puts them at risk for lower life expectancy.

The increased relative CVD risk from HIV is similar to the increased CVD risk due to diabetes (HIV: 1.75; diabetes: 2.1/2.0, M/F) (18, 34). The projected cumulative lifetime CVD risks for PLWH (ages 50-75) that we report are also similar to the lifetime CVD risk in diabetic patients, despite the competing mortality of HIV disease (diabetes: 67.1/57.3%, M/F, HIV: 64.8/43.8%, M/F) (35). A large body of evidence supports the benefits of CVD prophylaxis for diabetes mellitus, and diabetes is explicitly incorporated into CVD prevention guidelines (36). If HIV carries similar cumulative lifetime CVD risk, it is important to investigate the benefit of CVD prevention approaches among PLWH (37, 38). While new evidence suggests that in PLWH only 50% of MIs are of the traditional type (Type 1, from plaque instability), the other 50% are Type 2 MI, or secondary to ischemia due to either increased oxygen demand or decreased supply. This type of MI is more common among persons who inject drugs, who are also less likely to receive the full benefits from ART due to suboptimal adherence (39). A large, ongoing, randomized
controlled trial will help illuminate the specific role, if any, of statins as CVD prophylaxis in PLWH given this heterogeneity in MI type (40).

Our results provide important insights into the potential impact of smoking cessation on CVD lifetime risk in PLWH. One recent study suggests that smoking cessation can dramatically improve life expectancy in PLWH (41).

This study has several limitations. We explicitly model smoking as the major CVD risk factor among persons HIV-uninfected, at high risk for HIV and among PLWH. Other risk factors are modeled implicitly and, as a result, we have limited ability to examine differences in CVD risk due other individual risk factors such as hypertension, impaired glucose tolerance, or dyslipidemia explicitly. Our estimates for lifetime CVD risk in the non-HIV population were consistent with the results of population-based studies (42). Since there is no similar independent data source for PLWH, we made every attempt to inform our estimates using data from population-based studies among PLWH. Additionally, we did not distinguish between mortality from acute and chronic CVD and used aggregated mortality rates from population-based surveys. While we recognize that different subcategories of CVD (e.g., MI, stroke, peripheral vascular disease) may have different short- and long-term mortality (43), using aggregated data allowed us to focus on overall population-based impact. We also did not explicitly investigate the influence of racial and ethnic differences, smoking duration, duration of HIV infection, or means of HIV acquisition, which could all influence CVD risk (44, 45). Lastly, we did not assess the economic impact of CVD care and potential prevention efforts in the current analysis due to the relatively low cost of primary CVD
prevention in relation to the cost of ART. A comprehensive analysis of cost related to acute CVD events among PLWH is beyond the scope of this analysis.

These results have important implications for the care of PLWH in the US. While CVD is increasingly recognized as a common cause of death in treated PLWH, additional attention and guidance should be paid to CVD screening and risk factor counseling for PLWH. These results can be used in clinical practice to facilitate discussion between PLWH and clinicians, providing quantitative evidence to guide discussion around CVD risk reduction in PLWH receiving HIV care. Further, to the extent that PLWH in Europe are also living longer, at risk for CVD mortality, and have higher smoking rates than in the US (46), the overall findings of this study may be generalized to these settings.

Given that the projected CVD risk among PLWH was similar to those with diabetes, we believe that HIV should be considered a major risk factor for CVD and that PLWH could benefit from preventive strategies similar to persons with diabetes mellitus. It is critical to test the effectiveness of CVD primary prevention therapies for PLWH.
NOTES:

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Potential conflicts of interest.

P. E. S. has served as a consultant to Abb-Vie, Janssen, Bristol-Myers Squibb, Gilead, GlaxoSmithKline/ViiV, and Merck, and has received grants from Bristol-Myers Squibb, GlaxoSmithKline/ViiV, and Gilead. M. C. W. has served as a consultant for OptumInsight on topics unrelated to human immunodeficiency virus. All other authors report no potential conflicts.
References


Legend to Figures

**Figure 1.** Visual schematic of step-by-step estimation of CVD incidence for cohort of HIV-uninfected, at high risk for HIV and PLWH.

**Figure 2.** Model projected survival (A, B) and CVD cumulative incidence curves (C, D). Males are shown in blue (A, C) and females in red (B, D). The median survival for the US general population, the HIV-uninfected, at high risk for HIV population, and the PLWH population is 77.5, 76.4, and 70.2 years for males. For females, life expectancy was 81.1, 73.4, and 67.0 years. Cumulative CVD risk for males was 54.8%, 59.1%, and 64.8% for the US general, HIV-uninfected, at high risk for HIV, and PLWH populations. Cumulative CVD risk for females was 46.1%, 36.7%, and 43.8% for the US general, HIV-uninfected, at high risk for HIV, and PLWH populations. Kinks in the cumulative incidence curves are due to limitations in age stratification from our source data. Around age 70 in females, the cumulative incidence in the US general population begins to exceed that of the HIV-uninfected, at high risk for HIV population (Panel D, arrow).

**Figure 3.** Tornado diagrams summarizing sensitivity of the CVD lifetime risk among PLWH to variation in key input parameters. Males (A) are in blue and females (B) are in red. Input base case values and ranges are in parentheses.
Table 1. Model input parameters for analysis of CVD risk in PLWH in the U.S.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Base Case Value</th>
<th>References</th>
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<tr>
<td><strong>Cohort Characteristics</strong></td>
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<tr>
<td>Age, years (SD)</td>
<td>36 (0)</td>
<td>(15)</td>
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<tr>
<td>Distribution of initial CD4 (mean cells/µl)</td>
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<tr>
<td>± SD</td>
<td>751 (267)</td>
<td>(30)</td>
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<td><strong>HIV RNA Distribution after acute infection, %</strong></td>
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<tr>
<td>&gt;100,000 copies/ml</td>
<td>25</td>
<td></td>
</tr>
<tr>
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<td>3,001-10,000 copies/ml</td>
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<td><strong>Overall first-line ART efficacy</strong></td>
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<td>HIV RNA suppressed at 6 months, %</td>
<td>91</td>
<td>(32)</td>
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Table 1, continued:

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<th>HIV-uninfected, at High Risk for HIV and PLW (M/F)</th>
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<td>&gt;50 years</td>
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<th>CVD-Attributable Mortality, % monthly M/F</th>
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Table 1, continued:

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<tr>
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<th>HIV-uninfected, at High Risk for HIV†</th>
<th>PLWH†</th>
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<td>0.6/0.5</td>
<td>0.7/0.6</td>
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<td>1.7/1.4</td>
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<td>40-49</td>
<td>3.2/2.7</td>
<td>3.5/2.9</td>
<td>6.1/4.9</td>
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<tr>
<td>Age (years)</td>
<td>General Population</td>
<td>HIV-uninfected, at High Risk for HIV</td>
<td>PLWH</td>
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<td>5.9/4.6</td>
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<td>9.8/6.2</td>
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<td>43.7/26.0</td>
<td>76.8/45.7</td>
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*Derived from National Health Interview Survey using positive predictive values (see Methods and Technical Appendix).
†Derived using CEPAC model as described in the Methods.
All values are reported as male/female.
Abbreviations: ART: antiretroviral therapy; CVD: cardiovascular disease; M/F: male/female; SD: standard deviation.
Table 2. Life expectancy and cumulative risk of CVD

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<th>Males, Age (years)</th>
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<th>60</th>
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<th>80</th>
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<td>77.5</td>
<td>0.7</td>
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<td>12.8</td>
<td>25.6</td>
<td>44.6</td>
<td>54.8</td>
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<td>HIV-uninfected, at high risk for HIV</td>
<td>76.4</td>
<td>0.8 (0.7-0.8)</td>
<td>4.6 (4.5-4.7)</td>
<td>14.6 (14.0-14.8)</td>
<td>29.1 (27.5-29.6)</td>
<td>49.5 (46.2-50.7)</td>
<td>59.1 (55.3-61.0)</td>
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<tr>
<td>PLWH</td>
<td>70.2</td>
<td>1.3 (1.1-1.5)</td>
<td>7.1 (5.9-8.8)</td>
<td>20.5 (16.7-26.2)</td>
<td>37.9 (30.8-47.8)</td>
<td>57.9 (47.9-70.0)</td>
<td>64.8 (54.5-76.3)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Females, Age (years)</th>
<th>LE (y)</th>
<th>40</th>
<th>50</th>
<th>60</th>
<th>70</th>
<th>80</th>
<th>Lifetime</th>
</tr>
</thead>
<tbody>
<tr>
<td>US general population</td>
<td>81.1</td>
<td>0.6</td>
<td>3.4</td>
<td>9.4</td>
<td>17.3</td>
<td>33.5</td>
<td>46.1</td>
</tr>
<tr>
<td>HIV-uninfected, at high risk for HIV</td>
<td>73.4</td>
<td>0.6 (0.6-0.6)</td>
<td>3.5 (3.4-3.6)</td>
<td>9.7 (8.0-10.3)</td>
<td>17.3 (12.3-19.2)</td>
<td>30.4 (16.3-36.8)</td>
<td>36.7 (16.7-49.9)</td>
</tr>
<tr>
<td>PLWH</td>
<td>67.0</td>
<td>1.0 (0.9-1.2)</td>
<td>5.5 (4.2-7.0)</td>
<td>13.8 (8.8-18.7)</td>
<td>23.3 (12.7-32.7)</td>
<td>38.0 (16.0-55.2)</td>
<td>43.8 (16.3-66.8)</td>
</tr>
</tbody>
</table>
Abbreviations: CVD: cardiovascular disease; LE: life expectancy.
Figure 1. Schematic for derivation of CVD incidence for HIV-uninfected, at high risk for infection and PLWH cohorts.
Figure 2: Projected survival and CVD cumulative incidence for the US general population, HIV uninfected at high risk for HIV, and PLWH populations
Figure 3: The sensitivity of CVD lifetime risk among PLWH to changes in model input parameters.

A) Males

- Mean CD4 at presentation to care (351; 173-515)
- SMR (1.2; 1.1-1.0)
- Increased risk of CVD due to HIV (1.8; 1.0-2.1)
- Age of infection (36; 28-41)
- Smoking prevalence (39.5; 35.6-43.7)
- Increased risk of CVD due to smoking (1.9; 1.7-2.2)

B) Females

- SMR (2.9; 10.6-1.1)
- Mean CD4 at presentation to care (351; 173-516)
- Increased risk of CVD due to HIV (1.8; 1.0-2.1)
- Age of infection (36; 28-41)
- Increased risk of CVD due to smoking (1.7; 1.4-2.1)
- Smoking prevalence (33.7; 30.4-37.3)