Competing biomedical HIV prevention strategies: potential cost-effectiveness of HIV vaccines and PrEP in Seattle, WA

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Abstract

Introduction: Promising HIV vaccine candidates are steadily progressing through the clinical trial pipeline. Once available, HIV vaccines will be an important complement but also potential competitor to other biomedical prevention tools such as pre-exposure prophylaxis (PrEP). Accordingly, the value of HIV vaccines and the policies for rollout may depend on that interplay and tradeoffs with utilization of existing products. In this economic modelling analysis, we estimate the cost-effectiveness of HIV vaccines considering their potential interaction with PrEP and condom use.

Methods: We developed a dynamic model of HIV transmission among the men who have sex with men population (MSM), aged 15-64 years, in Seattle, WA offered PrEP and HIV vaccine over a time horizon of 2025-2045. A healthcare sector perspective with annual discount rate of 3% for costs (2017 USD) and quality-adjusted life years (QALYs) was used. The primary economic endpoint is the incremental cost-effectiveness ratio (ICER) when compared to no HIV vaccine availability.

Results: HIV vaccines improved population health and increased healthcare costs. Vaccination campaigns achieving 90% coverage of high-risk men and 60% coverage of other men within five years of introduction are projected to avoid 40% of new HIV infections between 2025 and 2045. This increased total healthcare costs by $30 million, with some PrEP costs shifted to HIV vaccine spending. HIV vaccines are estimated to have an ICER of $42,473/QALY, considered cost-effective using a threshold of $150,000/QALY. Results were most sensitive to HIV vaccine efficacy and future changes in the cost of PrEP drugs. Sensitivity analysis found ranges of 30-70% HIV vaccine efficacy remained cost-effective. Results were also sensitive to reductions in condom use among PrEP and vaccine users.

Conclusions: Access to an HIV vaccine is desirable as it could increase the overall effectiveness of combination HIV prevention efforts and improve population health. Planning for the rollout and scale-up of HIV vaccines should carefully consider the design of policies that guide interactions between vaccine and PrEP utilization and potential competition.

Keywords: HIV vaccines; pre-exposure prophylaxis; cost-benefit analysis; models; economic; costs and cost analysis; economic competition; disease transmission; infectious; preventive medicine

Additional information may be found under the Supporting Information tab for this article.

1 INTRODUCTION

Experts say an HIV vaccine is necessary, but not sufficient to end HIV [1,2]. A 50% effective vaccine may be good enough, but not enough. There are many exciting biomedical HIV prevention candidates in the research and development pipeline. Combinations of evidence-based HIV treatment and prevention interventions will be necessary for eradication. Investment and policy decisions consider not only effectiveness, but also aspects of access, acceptability, behaviour change and costs. This is an important question because current prevention interventions are imperfect. Decision-makers weigh population-level tradeoffs for opportunities that offer small benefits to a large number of individuals or large benefits to a small number of individuals. For an affordable public health programme, substantially reduced drug prices will likely be needed [3,4].

In the United States, Seattle, Washington is a national leader and early adopter of novel evidence-based HIV strategies. Seattle-King County Public Health surveillance rigorously monitors epidemic indicators and care cascade milestones, and it was the first US urban city to reach the “90-90-90” goal set by WHO. In King County, health officials estimate 6980 residents lived with diagnosed HIV infection in 2014, totaling...
more than half of all HIV cases in the state [5]. Approximately 50,000 men who have sex with men (MSM) live in King County. In 2014, 281 people were newly diagnosed with HIV, with local data suggesting rectal gonorrhoea or early syphilis as one of the strongest risk factors [6].

Seattle has combated new infections with pre-exposure prophylaxis (PrEP). As a complement to national guidelines for prescribing and monitoring PrEP [7], the local Public Health Seattle & King County with the Washington State Department of Health guide medical providers to recommend and discuss PrEP with target populations [8]. PrEP users are recommended to get tested every three months for HIV and other sexually transmitted infections (STIs); adherence and retention can be a challenge. One US study of patients prescribed PrEP at least six months beforehand (n = 171), 72% were retained in care at three months and 57% were retained in PrEP care at six months [9]. Long-acting injectable cabotegravir for PrEP is also under investigation. Future evaluation of novel PrEP products may deem placebo-controlled trials unethical.

Recent progress in HIV vaccine development means another biomedical product for prevention is approaching the horizon of availability [10]. A breakthrough 2009 Phase 3 trial in Thailand found significant HIV vaccine efficacy averaging 31% fewer infections over three years [11]. Confirmatory trials are ongoing in South Africa, with modifications to improve the Thai regimen – powered to detect HIV vaccine efficacy of 50% [12]. Based on prospectively defined immunogenicity thresholds, criteria were met for the Pox-Protein Public Partnership (P5) to support the launch of a Phase 2b/3 study in Africa: HIV Vaccine Trials Network (HVTN) 702 Study [13].

Real tradeoffs have to be made when offering imperfect prevention products. This is an important question because all available biomedical products for HIV prevention are imperfect, and the evidence of PrEP cost-effectiveness is mixed [4,14-16]. One framework for optimal resource allocation for investments is a static optimization model to evaluate potential combination HIV prevention strategies [17]. Previous modelling studies have separately examined the cost-effectiveness of PrEP [4,15,18] and the potential cost-effectiveness of HIV vaccines [19-29]. Two models to date have evaluated the expected combined impact of vaccine and PrEP, assuming independent coverage targets are achieved with each tool, and both found lower costs and improved health outcomes compared with PrEP alone [27,30]. This is the first modelling study to examine the economic impact of potential competition and interaction between HIV vaccine and PrEP considering the potential interaction between them.

2 | METHODS

We conducted a health economic modelling analysis to estimate the impact and cost-effectiveness of an HIV vaccine offered alongside PrEP in Seattle, WA.

2.1 | Study population

The study population includes MSM ages 15-64 in Seattle, WA, using Public Health Seattle-King County and the Washington State Department of Health reports as the primary source for population data [5,31,32]. Costs and benefits are evaluated over the time horizon 2025-2045. The Department of Health estimates MSM account for 5.4% of the population in this age range [5]. Within this population, 80% self-identify as gay and more than 31% had six or more male sex partners in the last 12 months [31].

2.2 | Model overview

We developed a deterministic dynamic compartmental mathematical model (Figure 1) to simulate the HIV epidemic among MSM in Seattle beginning in 2004. The model was already used to study the effectiveness of rapid antiretroviral therapy initiation among MSM in Peru [33]. It consists of a system of differential equations describing HIV transmission and disease progression through a series of health states. Over time, MSM enter the population at age of sexual debut and exit the population at age 64. The population is stratified into groups by HIV infection status (susceptible and infected), age (<25, 25-40, >40 years), risk of infection (low and high) and prevention modality (PrEP use and/or vaccination status). Infected MSM progress through a series of health states based on CD4-count, treatment status and viral suppression (Appendix S1).

The model is used to simulate the HIV epidemic without an HIV vaccine to provide a reference scenario for the evaluation of the vaccine impact. PrEP is introduced in 2015 followed by HIV vaccine availability in 2025. The effectiveness of the intervention over the 2025-2045 period is evaluated by comparison of the intervention to the reference scenario assuming no changes in the current Centers for Disease Control (CDC) guidelines for HIV prevention and treatment [7]. Reported metrics represent the mean outcome of 100 simulations using the preselected sets of epidemic parameters identified in the calibration procedure described below.

We followed recommendations from the ISPOR-SMDM Dynamic Transmission Modeling Task Force and the Second Panel on Cost-Effectiveness in Health and Medicine [34-36]. The model was developed in C++ and R version 3.4.2 [37].

2.3 | Model parameterization and calibration

The model is parameterized with epidemiological data representative of the HIV epidemic among MSM in Seattle. Calibration occurs from 2004 to 2014. Demographic and sexual behaviour characteristics including average number of partners per year, frequency of sex acts, proportion of acts protected by condoms, and lifetime duration of sexual activity were collected from published data (Tables 1, S2 and S3). Parameterization of age and risk group sexual mixing patterns were imputed from a meta-analyses of MSM that included a sample from Seattle [38]. King County HIV prevalence was calibrated to the 2015 HIV/AIDS Epidemiology Report from the Public Health Seattle-King County and Washington State Department of Health [5] and the CDC-sponsored 2014 Seattle area National HIV Behavioral Survey of Men Who Have Sex with Men Sexual (NHBS-MSM4) informed values for sexual risk behaviours, HIV testing and PrEP use [39]. “High-risk” was defined as MSM having five or more partners in the past 12 months, as a surrogate for the many risk factors identified in Seattle’s clinical guidelines for PrEP use [7].
2.4 | Interventions

2.4.1 | Pre-exposure prophylaxis

Daily oral Truvada® (Gilead Sciences, Inc., Foster City, CA, USA), a combination product of 200 mg emtricitabine (FTC) and 300 mg tenofovir disoproxil fumarate (TDF), for PrEP is introduced in the model beginning in 2015. After 2014, the model assumes that 25% of high-risk MSM and 6% of low-risk MSM start using PrEP annually with 20% discontinuation rate. This assumption closely reproduces the expansion of PrEP usage in Seattle up to 2017 [32] and is expected to result 25% overall PrEP coverage by 2025, with close to 50% of the high-risk MSM using PrEP (see Figure 2A). Figure 2A visualizes the utilization rates of each product at a population level when all of the model inputs and assumptions are combined into the dynamic model. Evidence from completed efficacy trials shows that PrEP efficacy depends strongly on adherence [40-48]. Results from most recent clinical studies [41,42], conducted after Truvada has already proven efficacy, suggested that PrEP reduces the HIV risk by more than 80% which motivated the efficacy assumption in our model. Conservatively, we assumed that PrEP does not reduce infectivity once infected. Condom replacement, also known as risk compensation or behavioural disinhibition, is a decrease in condom use that may occur among people using biomedical HIV prevention modalities [46]. In Seattle, on average, 63% of MSM sex acts are protected by a condom; while using PrEP, only 12.5% of MSM sex acts are protected by condoms [32]. We do not include a disutility to account for PrEP adverse events, assuming that individuals with intolerable side effects would discontinue its use.

2.4.2 | HIV vaccine

The model simulates vaccination with a five-dose regimen of a canarypox-based vaccine ALVAC-HIV vCP2438 DNA prime (Sanofi Pasteur, Paris, France) and bivalent gp120 protein sub-unit boost with MF59® adjuvant (GSK, Brentford, UK) beginning in 2025 (Table S1) [13]. The durability of vaccine protection is expected to wane over time [49,50]. We assume an average efficacy of 50% reduction in risk of infection lasting five years in duration [51,52]. We simulate vaccination campaigns every five years with coverage of 90% for those on PrEP and 60% for those who are not on PrEP (Figure 2A). We incorporate condom replacement among vaccinated MSM who continue to use PrEP but not to vaccinated non-users. We assume the vaccine has no disutility and that future technologies in HIV testing will overcome any previously reported social risks from vaccine-induced sero-positivity [53,54].

2.4.3 | Interactions between PrEP and vaccine

The model explores potential interactions among HIV vaccines, PrEP and condom use that risk mitigation of clinical and economic impact. We assume the protection from dual-use of PrEP and vaccine is multiplicative. Condom use may decrease as PrEP use increases. The demand for PrEP may decrease when another biomedical HIV prevention choice is on the market and HIV vaccine utilization increases. We explore the following utilizations of PrEP and HIV vaccine delivery: (i) vaccine licensure in 2025; (ii) PrEP is targeted to high-risk MSM while the vaccine is targeted to all MSM; (iii) HIV vaccination campaign cover 60% of low-risk MSM every five years beginning in 2025 (red line in Figure 2A); (iv) PrEP users being three times more likely to receive an HIV vaccine; (v) after vaccination PrEP users continue on PrEP for an additional year. This hypothetical interaction was specified by soliciting a collection of expert opinions about plausible changes expected in utilization rate.

2.5 | Approach to health outcomes

The following metrics of effectiveness are evaluated for each scenario over 20 years of intervention: cumulative number and fraction of new HIV infections prevented, reduction in HIV prevalence, and quality-adjusted life years (QALYs) gained.
Table 1. Dynamic transmission model inputs

| Parameter                                                                 | Value          | Source                                      |
|---------------------------------------------------------------------------|----------------|---------------------------------------------|
| Population size, men who have sex with men, ages 15-64 years, King County, 2004 | 45,000         | US Census [67]                              |
| Fraction young, 15-24 years                                              | 0.168          | US Census Reporter [68]                     |
| Fraction middle-aged, 25-44 years                                        | 0.463          | US Census Reporter [68]                     |
| Male maturation rate, rate of ageing into the population                 | 0.03           | Estimated                                   |
| Fraction of high risk MSM of HIV infection (>6 partners in the last 12 months) |               |                                             |
| Among young, 15-24 years                                                 | 0.310          | Seattle HIV/AIDS Epi Report [31]            |
| Among middle-aged, 25-44 years                                           | 0.099          | Seattle HIV/AIDS Epi Report [31]            |
| Among old, 45+ years                                                     | 0.065          | Seattle HIV/AIDS Epi Report [31]            |
| Insertive anal sex role, fraction of population with the role group "insertive" |               |                                             |
| Versatile anal sex role, fraction of population with the role group "versatile" |               |                                             |
| Number of sexual partners in the past 12 months                          |                |                                             |
| High-risk with young adults                                              | 10.5           | Table 11, CDC 2016 [69]                     |
| High-risk with middle-aged                                               | 10.5           | Table 11, CDC 2016 [69]                     |
| High-risk with older adults                                              | 6.0            | Table 11, CDC 2016 [69]                     |
| Low-risk with young adults                                               | 1.5            | Wall 2015 [70]                              |
| Low-risk with middle-aged                                                | 1.5            | Wall 2015 [70]                              |
| Low-risk with older adults                                               | 1.0            | Wall 2015 [70]                              |
| Death rate, non-AIDS, probability of dying between age x (midpoint of age category) and x + 1 |       |                                             |
| Ages 15-24 years                                                         | 0.001319       | Life tables, Arias 2016 [71]                |
| Ages 25-44 years                                                         | 0.001574       | Life tables, Arias 2016 [71]                |
| Ages 45-64 years                                                         | 0.008438       | Life tables, Arias 2016 [71]                |
| HIV vaccine efficacy                                                     | 0.50           | Expert opinion                              |
| HIV vaccine durability, average, years                                   | 5              | Expert opinion                              |
| HIV prevention effectiveness                                              |                |                                             |
| Condom efficacy, reduction in susceptibility per act                      | 0.7-0.9        | Smith 2015 [72]                             |
| Fraction of acts protected by a condom for unvaccinated and not using PrEP | 0.63           | Seattle HIV/AIDS Epi Report [5]             |
| PrEP users                                                               | 0.125          | Montano 2017 [32]                           |
| Vaccinated, low-risk                                                     | 0.125          | Assumed similar to PrEP users               |
| Vaccinated, high-risk                                                   | 0.63           | Assumed similar to susceptible low-risk men |
| PrEP efficacy, reduction in susceptibility per act                       | 0.80           | Molina 2015, McCormack 2015 [40,41]        |
| Calibration targets                                                      |                |                                             |
| HIV prevalence among MSM in King County                                  | 0.13-0.17      | Seattle HIV/AIDS Epi Report [31]            |
| Fraction of population who are diagnosed                                 | 0.72-0.93      | Seattle HIV/AIDS Epi Report [31]            |
| Fraction of diagnosed MSM who are engaged in care                        | 0.88-0.94      | Seattle HIV/AIDS Epi Report [31]            |
| Fraction of infected MSM on ART who are virally suppressed               | 0.8-0.86       | Seattle HIV/AIDS Epi Report [31]            |
| Utilities                                                                |                |                                             |
| Acute infection                                                          | 0.69           | Whitham 2016 [73]                           |
| CD4 count > 500 or viral suppression                                      | 0.73           | Whitham 2016 [73]                           |
| CD4 count 350-500                                                        | 0.71           | Whitham 2016 [73]                           |
| CD4 count 200-349                                                        | 0.69           | Whitham 2016 [73]                           |
| CD4 count < 200                                                          | 0.69           | Whitham 2016 [73]                           |
due to the vaccine programme. Person-time in each health state is multiplied by the corresponding preference-based utility weight, discounted 3% annually, and summed over the time horizon to calculate total QALYs [55-59].

2.6 | Approach to costing

Costing of HIV prevention services follows a unit costing approach, also known as ingredients-based, while the cost of HIV treatment relies on published studies based on aggregate healthcare costs. The cost of a clinic visit for HIV prevention services with risk reduction counselling is based on Centers for Medicare and Medicaid Services reimbursement rates for the corresponding relative value units in the Physician Fee Schedule January 2018 release [60]. Medication costs reflect the Veterans Affairs National Acquisition Center Federal Supply Schedule (FSS) prices from March 2018.
Vaccine price at each dose. The launch price of an HIV vaccine is unknown. Experts suggest benchmarking on the price of a recombinant human papillomavirus (HPV) vaccine because they similarly prevent transmission of a sexually transmitted virus. The FSS price for GARDASIL-9 (Merck & Co., Inc., Kenilworth, NJ, USA), was $210 per dose in June 2019 [61]. While the HPV vaccine is delivered to adults in a two- or three-dose series, ongoing Phase IIIB clinical trials of HIV vaccines are testing a five-dose series of vaccinations. To benchmark an estimate of the launch price for an HIV vaccine series, consultation with expert opinion assumed a 30% higher cost than the FSS price for a three-dose series of the HPV vaccine, totaling $820 per series of HIV vaccine as the input for the main analysis. The cost of a clinic visit for HIV prevention counselling and laboratory tests for STIs is added to the vaccine price at each dose.

2.7 | Sensitivity analysis

Parameter uncertainty was evaluated in a sensitivity analysis [34,62-64]. One-way sensitivity analyses evaluated the effect of uncertainty from individual parameters. Scenario analysis evaluated multi-way parameter uncertainty. To understand how robust the estimates of outcomes were to the choice of calibration parameters, we performed an uncertainty analysis using 100 calibration sets of inputs within a plausible range varying with respect to one another. We conducted a two-way threshold analysis of HIV vaccine and PrEP prices to understand the maximum cost-effective price for each product in relation to the price of the other.

3 | RESULTS

Maintaining the current trends of PrEP use, rates of diagnoses, linkage to care, treatment, and viral suppression, our analysis estimates 3074 new HIV infections between 2025 and 2045. We project an HIV prevalence of 7.7% among MSM living in Seattle in 2045, a decrease from 13.9% in 2018. On the path to this decline in prevalence, the model projects almost 15,000 HIV-uninfected men using PrEP in 2045, double the number in 2018, resulting in the partial protection in one-third of MSM.

HIV vaccines are projected to decrease the number of new infections, lower HIV prevalence and gain QALYs (Table 2). Seattle’s HIV prevalence in 2045 would be 1.4 percentage points lower with 37.9% of new HIV infections avoided. Considering the imperfect protection of both PrEP and vaccine, the model projects 1164 new HIV infections would be avoided. Our simulations suggest that more than 600,000 vaccine doses will be needed to secure that at least 50% of the susceptible MSM population is protected over 20 years. This produces 63% fewer MSM using PrEP in 2045 compared to the same year with no vaccine.

With model parameters based on US cost data over a 20-year time horizon, the total incremental cost of introducing an HIV vaccine in Seattle is estimated to be $30 million from a healthcare sector perspective. Given the assumed rate of product substitution, PrEP costs would decrease by $450 million and the cost of $532 million on HIV vaccines would be introduced (Table 2). In addition, prevention with a 50% effective vaccine is expected to reduce HIV treatment costs by $51 million.

3.1 | Cost-effectiveness

The introduction of an HIV vaccine was predicted to have an incremental cost-effectiveness ratio (ICER) of $42,473 per QALY gained. The cost per HIV infection avoided was $26,151. Scenarios in Figure 3 report the additional cost of HIV vaccines against QALYs gained, over time, highlighting each five-year increment since start of the intervention. The shaded grey area reflects the cost-effectiveness threshold range of 1-3 times gross domestic product (GDP) per capita.

3.2 | Sensitivity analysis

The greatest source of uncertainty and driver of cost-effectiveness is HIV vaccine efficacy. Vaccine efficacy of 70% had an ICER of $4136 per QALY while 30% was $108,824 per QALY compared to no vaccine (Table 3). Vaccine cost-effectiveness was also sensitive to the price of PrEP drugs. If competition from the entry of generic PrEP products reduces the medication cost to half by 2025, then the addition of an HIV vaccine would cost $336,671 per QALY gained compared to no vaccine. Assuming no condom displacement with HIV vaccines or PrEP lowered the ICER to $21,457 per QALY. The magnitude of health benefit from a vaccine was sensitive to the degree of condom replacement assumed when partially protected by PrEP and/or vaccine.

4 | DISCUSSION

This study used a dynamic transmission model to evaluate the cost-effectiveness of an HIV vaccine launch in 2025, assuming the vaccine would complement and substitute some PrEP use. The tradeoffs from competition between two imperfect biomedical HIV prevention products include (i) the opportunity to vaccinate a larger fraction of the population than with PrEP alone and (ii) the downside from potential substitution with a less effective prevention product. Assuming most high-risk and some low-risk MSM in Seattle are using PrEP in 2025, rapid uptake of HIV vaccine by 60% of men, and declines in PrEP use with increasing vaccine uptake, we found an ICER of $42,473 per QALY gained compared to PrEP with no vaccine. In this case, HIV vaccines would be cost-effective using a 1x GDP or 3x GDP per capita, cost-effectiveness threshold. Though HIV vaccines increased total healthcare costs by $240 million, some costs were offset by reduction in HIV treatment and PrEP medications.

Key uncertainties in the analysis affect the results under different scenarios. As expected, scenarios with greater vaccine efficacy were more likely to find vaccines cost-effective and led to a higher maximum threshold price where the vaccine would remain cost-effective. Given that an HIV vaccine candidate is in development, the exact frequency of vaccine administrations will be determined with more data about the durability of protection. We have previously explored the impact of HIV vaccine durability and we have been examined...
implementation approaches for three-year HIV vaccine campaigns in another study [65,66]. Results were also sensitive to assumptions about the rate of switching from PrEP to vaccines. If PrEP users who become vaccinated continued to use PrEP for the same length of time as non-vaccinated PrEP users, the vaccine has a higher ICER and is therefore less likely to be cost-effective. If all high-risk men using PrEP switched immediately to a vaccine, the vaccine has a lower ICER and would be more likely to be cost-effective, but the total population-level health benefit is slightly smaller. Assuming no change in condom use among people using PrEP or vaccines produced a lower, more likely cost-effective, ICER. Lastly, scenarios assuming generic PrEP prices in the future led to the vaccine being less likely cost-effective, while alternative scenarios simulating the launch of newer, branded, long-acting, injectable products for PrEP at higher prices affected the results by lowering the ICER for HIV vaccine introduction, meaning vaccines would be more likely to be cost-effective.

This modelling analysis yields two important lessons: (i) the cost-effectiveness of HIV vaccines will depend on the utilization and cost of PrEP at the time of launch and (ii) condom displacement with vaccines could diminish the potential benefit of vaccines for the population and lower its value. Even if, however, vaccines induce some condom displacement and decline in PrEP use, we project overall population health benefits. Policies guiding the interactions between these interventions could have substantial impact on the value of each product alone and in combination.

We compared the results of this model with two other published studies evaluating the cost-effectiveness of HIV vaccines in the United States, and our conclusions were consistent when assuming the same vaccine price. A dynamic transmission model of HIV vaccines conducted in the pre-PrEP era estimated an ICER of $91,000/QALY for universal HIV nation and net cost-savings from targeting MSM when assuming a cost of $500 per vaccine series [24]. A more recent static HIV model comparing PrEP offered with HIV vaccines to PrEP alone, assuming vaccines cost $2500 per series, also estimated a net cost-savings for MSM [30]. In our sensitivity analysis varying HIV vaccine costs, we similarly found vaccines costing $500 or $2500 per series would have a net cost-savings for MSM.

Table 2. Model results

| Outcome                              | Current practice | HIV vaccine, 50% efficacy | Incremental difference | Relative difference (%) |
|--------------------------------------|------------------|---------------------------|------------------------|------------------------|
| HIV burden                           |                  |                           |                        |                        |
| New HIV infections, 2025-2045        | 3074             | 1910                      | –1164                  | –37.9%                 |
| New HIV diagnoses 2025-2045          | 2934             | 2121                      | –814                   | –27.7%                 |
| People living with HIV in 2045       | 4806             | 3949                      | –857                   | –17.8%                 |
| HIV prevalence (%) in 2045           | 7.7%             | 6.3%                      | –1.4%                  | –18.0%                 |
| Utilization of biomedical prevention  |                  |                           |                        |                        |
| Protected by PrEP in 2025            | 11,233           | 11,233                    | 0                      | 0.0%                   |
| Total protected by PrEP or vaccine in 2045 | 14,905          | 36,680                    | 21,775                 | 146.1%                 |
| PrEP alone (% of susceptible)        | 14,905           | 5494                      | –9412                  | –63.1%                 |
| HIV vaccine alone (% of susceptible) | 0                | 31,158                    | 31,158                 |                        |
| PrEP + HIV vaccine (% of susceptible)| 0                | 29                        | 29                     |                        |
| Health outcomes                      |                  |                           |                        |                        |
| Total LYSb                            | 1,100,665        | 1,102,750                 | 2086                   | 0.2%                   |
| Total QALYs                          | 923,770          | 924,486                   | 717                    | 0.1%                   |
| Costs                                |                  |                           |                        |                        |
| Total cost (millions $)              | $2396            | $2426                     | $30                    | 1.3%                   |
| PrEP costs (millions $)              | $675             | $224                      | –$450                  | –66.7%                 |
| HIV vaccine costs (millions $)       | $0               | $532                      | $532                   |                        |
| HIV care costs (millions $)          | $1720            | $1669                     | –$51                   | –3.0%                  |
| ICER ($ per QALY)                    |                  |                           | $42,473                |                        |

Costs are presented in a common currency of 2017 USD. Cost-effectiveness analysis uses time horizon of 2025-2045. Per capita and per capita susceptible calculations are based on the common population size of MSM projected in 2025.

ICER, incremental cost-effectiveness ratio; LYS, life years; MSM, men who have sex with men; PrEP, pre-exposure prophylaxis; QALYs, quality-adjusted life years.

The relative difference in HIV prevalence is slightly different from the relative difference in number of people living with HIV 2045 as more MSM are alive in 2045 with the vaccine – contributing to the denominator of HIV prevalence but not the number of cases living with HIV; bLY and QALYs summed among MSM ages 15-64 years between the years 2025-2045.
cost-effectiveness results, despite the differences in their structure [67]. As all available epidemiological data were used for calibration, the model has not yet been validated for time periods outside 2004-2014. The U.S. does not have a stated willingness to pay for health gains and we assumed a range of cost-effectiveness thresholds from current recommendations [36,37]. Also, the generalizability of these findings should be limited to MSM in the US. Not only is the future cost of PrEP uncertain, the products that will be used for PrEP in the future are also uncertain. With no additional approvals of for new drugs or indications, we would expect generic TDF/FTC products to become available at lower cost. If newer PrEP products offer fewer side effects, this would effectively extend the patent period and prices are unlikely to decline. Tested model interactions between vaccine and PrEP utilization are limited and rely on plausible scenarios and assumptions described by expert opinion. Insurance status and PrEP medication cost were not found to be significant barriers for obtaining PrEP in one study [9], but price elasticity is not yet understood. Adverse events from PrEP and HIV vaccines are not well defined and estimating any related disutility is difficult.

The findings from this study point to several policy considerations. First, further public investment in US HIV vaccine clinical trials is warranted to reduce the uncertainty in expected vaccine efficacy. When regulatory bodies deem an HIV vaccine as having "good enough" efficacy, commercialization may be a challenge. This economic model does not include novel incentives that may be needed to encourage industry partners to commercialize and manufacture the product for global distribution. A target product profile to guide the innovation of better, cheaper, faster and point-of-care STI diagnostics may also be needed. Second, more research on and education to prevent a decrease in condom displacement with biomedical HIV prevention products is needed to optimize the potential effectiveness and prevent further outbreak of other STIs such as syphilis and gonorrhoea. Third, value-based pricing of the vaccine at launch should consider both the risk level of the indicated population and the current cost of PrEP. Risk to public investment in immunization campaigns could be mitigated with outcomes-based risk-sharing agreements between government payers and the manufacturer with support from existing CDC surveillance systems.

Table 3. Sensitivity analysis

| Scenario                        | Inc Cost       | Inc QALYs | ICER ($/QALY) |
|---------------------------------|----------------|-----------|---------------|
| Main analysis                   | $30,439,907    | 717       | $42,473       |
| HIV vaccine 70% efficacy        | $3,518,673     | 851       | $4136         |
| HIV vaccine 30% efficacy        | $59,386,518    | 546       | $108,824      |
| No condom replacement            | $12,135,182    | 566       | $21,457       |
| PrEP half price                 | $241,288,044   | 717       | $336,671      |
| HIV vaccine half price           | $180,422,645   | 717       | Dominant      |
| PrEP price doubled               | $391,256,368   | 717       | Dominant      |
| Doubled price HIV vaccine       | $452,165,011   | 717       | $630,908      |

Incremental results represent Seattle population-level health gains and healthcare payer costs during the period 2025-2045. Costs in 2017 USD. Dominant scenarios gained health and had lower cost than the reference comparator of PrEP with no HIV vaccine. ICER, incremental cost-effectiveness ratio; PrEP, pre-exposure prophylaxis; QALYs, quality-adjusted life years.
5 | CONCLUSIONS

Moderately effective HIV vaccines have the potential to be a cost-effective intervention implemented alongside PrEP in Seattle. The potential population health gains from and value-based price of an HIV vaccine in this setting depends on the degree of interaction and substitution with PrEP. Dual methods could be implemented as complementary products if lower PrEP pricing could be negotiated. Access to an HIV vaccine is desirable as it could increase the overall effectiveness of combination HIV prevention efforts and improve population health. HIV vaccines may have the potential to reach subpopulations that PrEP has been unable to reach. The barriers to implementation of and access to vaccines could be lower with provision at clinic visits compared to prescription drugs that require high adherence to be effective. Planning for the rollout and scale-up of HIV vaccine in the United States.

REFERENCES

1. Fauci AS, Marston HD. Ending AIDS – is an HIV vaccine necessary? N Engl J Med. 2014;370:495–8.
2. Fauci AS. An HIV Vaccine is essential for ending the HIV/AIDS pandemic. JAMA. 2017;318:1535.
3. Ong KJ, Desai S, Field N, Desai M, Nardone A, van Hoek AJ, et al. Economic evaluation of HIV pre-exposure prophylaxis among men-who-have-sex-with-men in England in 2016. Euro Surveill. 2017;22:17-00192. https://doi.org/10.2807/1560-7917.ES.2017.22.42.17-00192.
4. Bernard CL, Brandeau ML, Humphreys K, Bendavid E, Holodny M, Weyant C, et al. Cost-effectiveness of HIV pre-exposure prophylaxis for people who inject drugs in the United States. Ann Intern Med. 2016;165:10–9.
5. HIV/AIDS Epidemiology Unit Public Health – Seattle & King County and the Infectious Disease Assessment Unit Washington State Department of Health. HIV/AIDS epidemiology report 2015. Vol. 84. Seattle 2015. Available from: https://www.kingcounty.gov/depts/health/communicable-diseases/hiv-std/patient–/media/depts/health/communicable-diseases/documents/hivstd/2015-hiv-epidemiology-anual-report.aspx
6. Menza TW, Hughes JP, Celum CL, Golden MR. Prediction of HIV acquisition among men who have sex with men. Sex Transm Dis. 2009;36:547–55.
7. US Public Health Service. Preexposure prophylaxis for the prevention of HIV infection in the United States – 2014: a clinical practice guideline. Atlanta (GA): US Public Health Service; 2014.
8. Public Health – Seattle & King County WSD of H. Pre-exposure prophylaxis (PrEP) implementation guidelines 2015. 2015. Available from: https://www.kingcounty.gov/depts/health/communicable-diseases/documents/PrEP-implementation-guide-lines.aspx
9. Chan PA, Mena L, Patel R, Oldenburg CE, Beauchamps L, Perez-Brumer AG, et al. Retention in care outcomes for HIV pre-exposure prophylaxis implementation programmes among men who have sex with men in three US cities. J Int AIDS Soc. 2016;19:20903.
10. Hsu DC, O’Connell RJ. Progress in HIV vaccine development. Hum Vacc Immunother. 2017;13:1018–30.
11. Rerks-Ngarm S, Pitisuttithum P, Nitayaphan S, Kaewkungwal J, Chiu J, Paris R, et al. Vaccination with ALVAC and AIDSVAX to prevent HIV-1 infection in Thailand. N Engl J Med. 2009;361:2199–209.
12. National Institute of Allergy and Infectious Diseases. NIH-sponsored HIV vaccine trial launches in South Africa: early-stage trial aims to build on RV144 results. NIH News. 2015 [cited 2015 Jul 31]. Available from: http://www.niaid.nih.gov/news/newsreleases/2015/Pages/HVTN100.aspx
13. Russell ND, Marovich MA. Pox-Protein Public–Private Partnership program and upcoming HIV vaccine efficacy trial. Curr Opin HIV AIDS. 2016;11:614–9.
14. Paltiel AD, Freedberg KA, Scott CA, Schackman BR, Losina E, Wang B, et al. HIV preexposure prophylaxis in the United States: impact on lifetime infection risk, clinical outcomes, and cost-effectiveness. Clin Infect Dis. 2009;48:806–15.
15. Gomez GB, Borquez A, Case KK, Wheelock A, Vassall A, Hankins CD. The cost and impact of scaling up pre-exposure prophylaxis for HIV prevention: a systematic review of cost-effectiveness modelling studies. PLoS Med. 2013;10: e1001401.
16. Juusola JL, Brandeau ML, Owens DK, Bendavid E. The cost-effectiveness of pre-exposure prophylaxis for HIV prevention in the United States in men who have sex with men. Ann Intern Med. 2012;156:541–50.
17. Juusola J, Brandeau M. HIV treatment and prevention: a simple model to determine optimal investment. Med Decis Making. 2016;36:391–409.
18. Ouellet E, Durand M, Guerin JR, LeLorier J, Tremblay CL. Cost-effectiveness of “on demand” HIV pre-exposure prophylaxis for non-injection drug-using men who have sex with men in Canada. Can J Infect Dis Med Microbiol. 2015;26:23.
19. Adamson B, Dimitrov D, Devine B, Barnabas R. The potential cost-effectiveness of HIV vaccines: a systematic review. Pharmacoeconomics. 2017;1:1–12.
20. Bos JM, Postma MJ. The economics of HIV vaccines: projecting the impact of HIV vaccination of infants in sub-Saharan Africa. Pharmacoeconomics. 2001;19:937–46.
21. Amirfar S, Hellenberg JP, Abdool Karim SS. Modeling the impact of a partially effective HIV vaccine on HIV infection and death among women and infants in South Africa. J Acquir Immune Defic Syndr. 2006;43:219–25.
22. Ono S, Kurotaki T, Nakasone T, Honda M, Boon-Long J, Sawanpanyalert P, et al. Cost-effectiveness analysis of antiretroviral drug treatment and HIV-1 vaccination in Thailand. Jpn J Infect Dis. 2006;59:168–73.
23. Long EF, Brandeau ML, Owens DK. Potential population health outcomes and expenditures of HIV vaccination strategies in the United States. Vaccine. 2009;27:5402–10.
24. Long EF, Owens DK. The cost-effectiveness of a modestly effective HIV vaccine in the United States. Vaccine. 2011;29:6113–24.
70. Wall K, Stephenson R, Sullivan PS. Frequency of sexual activity with most recent male partner among young, internet-using men who have sex with men in the United States. J Homosex. 2013;60:1520–38.
71. Arias E, Heron M, Xu J. United states life tables, 2012. Natl Vital Stat Rep. 2016;65:1–65.
72. Smith DK, Herbst JH, Zhang X, Rose CE. Condom effectiveness for HIV prevention by consistency of use among men who have sex with men in the United States. J Acquir Immune Defic Syndr. 2015;68:337–44.
73. Whitham H, Kuppermann M, Shrestha R, Hutchinson A, Grund B, Shouse L, et al. Health utility estimates among persons living with HIV in the U.S. – implications for cost-effectiveness modeling and future research needs. In: 38th Annual North American Meeting of the Society of Medical Decision Making, Vancouver: MDM; 2016.
74. Centers for Medicare & Medicaid Services. 2018 National physician fee schedule relative value file January release. Baltimore (MD); 2018. Available from: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/PhysicianFeeSchedules/downloads/2018_Fee_Schedule.pdf
75. National Alliance of State & Territorial AIDS Directors (NASTAD). Billing coding guide for HIV prevention. Washington (DC); 2016. Available from: https://www.nastad.org/sites/default/files/BillingCodingGuide_v4_Final_2016.pdf
76. Centers for Medicare and Medicaid Services (CMS). 2017 Clinical diagnostic laboratory fee schedule; 2017. Available from: https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/CLVlntSched/index.html
77. Gebo KA, Fleishman JA, Conviser R, Hellinger J, Hellinger FJ, Josephs JS, et al. Contemporary costs of HIV healthcare in the HAART era. AIDS. 2010;24:2705–15.
78. Farnham PG, Gopalappa C, Sansom SL, Hutchinson AB, Brooks JT, Weidle PJ, et al. Updates of lifetime costs of care and quality-of-life estimates for HIV-infected persons in the United States. J Acquir Immune Defic Syndr. 2013;64:183–9.

SUPPORTING INFORMATION

Additional information may be found under the Supporting Information tab for this article.

Appendix S1. Supplementary material.