Cost-effectiveness of dual maternal HIV and syphilis testing strategies in high and low HIV prevalence countries: a modelling study

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Summary

Background Dual HIV and syphilis testing might help to prevent mother-to-child transmission (MTCT) of HIV and syphilis through increased case detection and treatment. We aimed to model and assess the cost-effectiveness of dual testing during antenatal care in four countries with varying HIV and syphilis prevalence.

Methods In this modelling study, we developed Markov models of HIV and syphilis in pregnant women to estimate costs and infant health outcomes of maternal testing at the first antenatal care visit with individual HIV and syphilis tests (base case) and at the first antenatal care visit with a dual rapid diagnostic test (scenario one). We additionally evaluated retesting during late antenatal care and at delivery with either individual tests (scenario two) or a dual rapid diagnosis test (scenario three). We modelled four countries: South Africa, Kenya, Colombia, and Ukraine. Strategies with an incremental cost-effectiveness ratio (ICER) less than the country-specific cost-effectiveness threshold (US$500 in Kenya, $750 in South Africa, $3000 in Colombia, and $1000 in Ukraine) per disability-adjusted life-year averted were considered cost-effective.

Findings Routinely offering testing at the first antenatal care visit with a dual rapid diagnosis test was cost-saving compared with the base case in all four countries (ICER: –$26 in Kenya, –$559 in South Africa, –$844 in Colombia, and –$454 in Ukraine). Retesting during late antenatal care with a dual rapid diagnostic test (scenario three) was cost-effective compared with scenario one in all four countries (ICER: $270 in Kenya, $260 in South Africa, $2207 in Colombia, and $205 in Ukraine).

Interpretation Incorporating dual rapid diagnostic tests in antenatal care can be cost-saving across countries with varying HIV prevalence. Countries should consider incorporating dual HIV and syphilis rapid diagnostic tests as the first test in antenatal care to support efforts to eliminate MTCT of HIV and syphilis.

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Introduction

Dual elimination of mother-to-child transmission (MTCT) of HIV and syphilis is a public health priority. Worldwide, 1.4 million maternal HIV infections and 988,000 maternal syphilis infections occur annually. Although maternal treatment is highly effective at preventing MTCT of both HIV and syphilis, gaps in maternal testing and treatment coverage lead to 180,000 infant HIV infections, 355,000 adverse congenital syphilis births, and 306,000 non-clinical congenital syphilis cases every year.6,7 WHO has set goals to reach elimination of MTCT of HIV and syphilis, including at least 95% of pregnant women receiving antenatal care, 95% tested for HIV and syphilis, and 95% treated for their infection(s).8 Yet, by 2020, only 14 countries had received validation by WHO for achieving the elimination of paediatric HIV or congenital syphilis.9

Globally, as HIV testing coverage has increased, more pregnant women with HIV are aware of their status, of whom 85% have accessed treatment; whereas, only 66% of pregnant women are tested for syphilis, of whom 78% receive treatment.10 Global efforts for prevention of MTCT (PMTCT) of HIV have led to substantial reductions in new paediatric HIV infections, but PMTCT of syphilis has received considerably less attention and success.11

Integrating syphilis testing and treatment into existing HIV PMTCT programmes might avert additional syphilis morbidity and mortality. 57% of congenital syphilis cases resulting in adverse birth outcomes were attributed to an absence of syphilis screening for women attending antenatal care.12 Testing coverage for HIV is often several times higher than for syphilis, suggesting that integrated testing could improve syphilis test coverage.13 Although
rapid diagnostic tests are standard for HIV, many countries still rely on laboratory-based testing for syphilis. Three dual HIV and syphilis rapid diagnostic tests, which allow concurrent antenatal care testing that circumvents laboratory testing obstacles for syphilis, are WHO prequalified.9

With restricted resources (including for programmes to support funding for staff training, time to offer testing, and test kits) and a need to prioritise highly effective interventions in resource-limited settings, policy makers need health economic evidence to decide where and how to implement dual HIV and syphilis testing. Dual HIV and syphilis testing in antenatal care has been shown to be a cost-effective approach in Malawi,1 but benefits and costs of this approach in diverse geographical regions with different health systems and HIV and syphilis burden have not been examined.

We aimed to compare the use of a dual rapid diagnostic test with individual tests at the first antenatal care visit and assess the effect of retesting for HIV and syphilis during pregnancy using a dual rapid diagnostic test. Many countries already retest for HIV during late antenatal care and delivery,10 and retesting for HIV and syphilis testing in antenatal care has been shown to be a cost-effective approach within antenatal care to improve maternal and infant health outcomes.

Evidence before this study

Globally, HIV testing coverage is high during pregnancy; by contrast, syphilis testing coverage is frequently lower, but it could be increased if combined with HIV testing. As countries strive to achieve elimination of mother-to-child transmission of both infections, studies that identify strategies to efficiently optimise implementation of maternal HIV and syphilis testing are crucial to maximise the number of infant infections prevented. We searched PubMed for studies published from the inception of the database to July 28, 2020, with no language restrictions. Using the search terms “HIV infections” OR “HIV infections/transmission” OR “HIV” OR “syphilis, congenital” AND “mass screening” OR “screening” (tiab) OR “testing”(tiab) AND “repeat”(tiab) OR “retest”(tiab) AND “maternal”(tiab) OR “pregnancy” AND “cost-effectiveness” OR “cost-benefit analysis” OR “mass screening/economics” eight studies were found with this specific combination. Previous cost-effectiveness analyses of dual HIV and syphilis testing or syphilis retesting during pregnancy have focused on a single country. Dual HIV and syphilis testing was shown to be cost-effective in Malawi and retesting for syphilis has shown mixed cost-effectiveness results in the USA. None of the previously published modelling analyses provided results relevant to a variety of contexts and settings.

Added value of this study

To our knowledge, this is the first study to model the health and economic effects of using a single rapid diagnostic test for both HIV and syphilis in the context of antenatal care in multiple countries that represent diverse geographical and HIV and syphilis prevalence settings. To our knowledge, this exploratory analysis of retesting for syphilis and HIV using the dual rapid diagnostic test is the first analysis of the cost-effectiveness of maternal retesting for syphilis outside the USA, and the first cost-effectiveness analysis of retesting using a dual rapid diagnostic test.

Implications of the available evidence

We find that using a dual rapid diagnostic test for HIV and syphilis at the first antenatal care visit is cost-saving compared with using a rapid HIV test and laboratory-based syphilis test in all four countries evaluated, regardless of HIV prevalence. Dual rapid diagnostic test for HIV and syphilis at the first antenatal care visit is projected to avert many congenital syphilis cases and disability-adjusted life-years, but similar numbers of infant HIV infections as independent HIV and syphilis tests. Retesting in late pregnancy with a dual rapid diagnostic test was more effective and less costly than independent HIV and syphilis tests during late pregnancy. This analysis highlights potential health and economic benefits of integrating maternal HIV and syphilis testing. Programmes might consider incorporating dual rapid diagnostic tests for HIV and syphilis as a cost-effective approach within antenatal care to improve maternal and infant health outcomes.

Methods

Model overview and testing scenarios

Using a Markov decision analytical model, we did a cost-effectiveness analysis of maternal HIV and syphilis testing using individual tests and dual rapid diagnostic tests in Kenya, South Africa, Colombia, and Ukraine—four countries that represent a range of HIV prevalence, syphilis prevalence, and geographical settings. Countries were classified as having low (<5%; Colombia and Ukraine), intermediate (5–15%; Kenya), and high (≥15%; South Africa) HIV prevalence.9 We compared four HIV and syphilis testing scenarios, varying the assay type and timing of testing (table I). The primary analysis assessed the cost-effectiveness of dual HIV and syphilis testing compared with individual tests at the first antenatal care visit, and the secondary analysis evaluated the effect of retesting for HIV and syphilis during late antenatal care. We modelled the same test types across all four countries...
because of heterogeneity within countries and to enable comparison between countries. We modelled the use of dual rapid diagnostic test and individual HIV and syphilis tests followed by confirmatory testing. We assumed HIV diagnoses followed two consecutive reactive rapid diagnostic tests. Individual syphilis testing was modelled as a laboratory-based rapid plasma reagin test with confirmatory testing for reactive results with the Treponema pallidum haemagglutination assay. No confirmatory test for syphilis was modelled for dual rapid diagnostic test scenarios.

In the base case scenario, we assumed that HIV and syphilis testing with individual tests was done at the first antenatal care visit, defined by country-specific average gestational age at the first visit. In scenario one, we modelled testing at the first antenatal care visit using a dual rapid diagnostic test. Additionally, because maternal HIV retesting is already recommended in many countries, we modelled two scenarios that incorporated maternal HIV retesting to address operational considerations. To estimate the relative health and economic effect attributed to HIV versus syphilis, we compared retesting with individual tests (scenario two) with retesting during late antenatal care and delivery with a dual rapid diagnostic test (scenario three). Retesting scenarios included women who missed HIV or syphilis testing at first antenatal care visit (ie, because they did not attend antenatal care before the gestational age when retesting would occur or because of gaps in test coverage). Retesting scenarios also allowed for HIV retesting at delivery for women not tested during late antenatal care (dual and individual tests), but not for syphilis retesting at delivery. For scenarios in which dual rapid diagnostic tests were modelled (scenarios one and three), we assumed that all women tested received the test regardless of previous HIV diagnosis or syphilis infection.

**Model structure, parameterisation, and assumptions**

Maternal HIV and syphilis were modelled using separate Markov models (appendix pp 1–2). The HIV model was adapted from a previous model of maternal HIV retesting, and reflects HIV prevalence, incidence, disease progression, and treatment during pregnancy and post partum using 54 Markov states (six maternal HIV states by nine antenatal and post partum periods; Meisner J and Roberts DA, unpublished). The congenital syphilis model reflects syphilis prevalence, incidence, testing, and treatment using eight maternal syphilis states with time-varying transitions. We assumed risk of MTCT of HIV and syphilis and associated infant health outcomes were independent. Risk of infant HIV was modelled for 1 year after birth. The time step in both models was 1 week, and the time horizon was 20 years after birth—a common time horizon that allows our results to be directly compared with other cost-effectiveness analyses. The model was built in Microsoft Excel (2016).

Models were parametrised using values from the literature, in-country contacts, and expert opinion; country-specific estimates were used when available (table 2). Although HIV and syphilis models were constructed separately, they share common model parameter inputs. Both models incorporate disease prevalence, incidence, antenatal care attendance, test coverage, test sensitivity and specificity, maternal and infant mortality rates, and probability of infant infection. For each country, we modelled a population of pregnant women equal to the estimated number of annual pregnancies, or the number of livebirths if the number of pregnancies was unavailable. All pregnancies were assumed to be singleton. Fetal loss because of syphilis was modelled, but fetal loss because of HIV was not.

**Health effect**

We modelled infant HIV infection, congenital syphilis, and infant death. Infant HIV infections and deaths were estimated on the basis of total exposure to HIV in utero and post partum, with probabilities dependent on new versus established maternal HIV infection (on the basis of Fiebag stages), maternal viral suppression, maternal antiretroviral therapy (ART), infant ART, and breastfeeding practices. Congenital syphilis outcomes (stillbirth, neonatal death, preterm birth and low birthweight, clinical congenital syphilis, and non-clinical syphilis) were estimated at birth and modified by receipt and timing of adequate maternal syphilis treatment during pregnancy. We followed WHO guidelines on the definition of adequate syphilis treatment as at least 30 days before birth (modelled as ≤32 weeks) to prevent fetal infection, and set probabilities of adverse infant outcomes for later treatment equal to those for no treatment. We considered all infants born to untreated and late treated mothers to have congenital syphilis, regardless of symptoms. We accounted for duplication of outcomes between the HIV and syphilis models by calculating infant co-infection as the joint probability of syphilis and HIV infection in infants, assuming independence. Syphilis-related stillbirths and neonatal deaths were subtracted from the population at delivery in the HIV model.

### Table 1: Dual HIV and syphilis testing models scenarios

|               | First ANC visit | Late ANC visit |
|---------------|-----------------|----------------|
|               | Syphilis        | HIV            | Syphilis        | HIV            |
| Base case     | RPR and TPHA    | Rapid          | No test         | No test        |
| Scenario one  | Dual            | Dual           | No test         | No test        |
| Scenario two  | RPR and TPHA    | Rapid          | RPR and TPHA    | Rapid          |
| Scenario three| Dual            | Dual           | Dual            | Dual           |

Base case used independent tests at country-specific timepoints. Scenario one used dual HIV and syphilis testing with no retesting during late ANC. Scenario two used independent HIV and syphilis tests with retesting during late ANC. Scenario three used dual HIV and syphilis testing with retesting during late ANC. Dual test is a single, point-of-care rapid test for HIV and syphilis. Independent tests include rapid HIV tests and laboratory-based syphilis tests (RPR and TPHA to confirm reactive results). Late ANC occurred from 36 to 39 weeks’ gestation. ANC=antenatal care. RPR=rapid plasma reagin. TPHA=Treponema pallidum haemagglutination assay.

See Online for appendix
We converted infant infections and adverse birth outcomes into disability-adjusted life-years (DALYs; appendix pp 16–17). Total DALYs were the sum of years lived with disability and years of life lost up to the age of 20 years. We assumed DALYs for each stillbirth and neonatal death were equal. We used disability weights from the Global Burden of Disease study and standard formulas for calculating co-infection disability weights. We converted infant infections and adverse birth outcomes into disability-adjusted life-years (DALYs; appendix pp 16–17). Total DALYs were the sum of years lived with disability and years of life lost up to the age of 20 years. We assumed DALYs for each stillbirth and neonatal death were equal. We used disability weights from the Global Burden of Disease study and standard formulas for calculating co-infection disability weights. For HIV-infected infants on ART at 1 year, the disability weight for HIV while on ART was applied over the 20-year time horizon. For HIV-infected infants not on ART at 1 year, we assumed 90% of time was spent with symptomatic HIV and 10% with AIDS over the 20-year time horizon. For infants with congenital syphilis, the disability weight was applied only to the first 3 years of life. DALYs were discounted at 3% annually.

### Costs and cost-effectiveness

We used a health-care system perspective for the model. Costs of supplies, labour, and treatment were obtained from the literature, in-country contacts, and expert

| Population of pregnant women | Kenya | South Africa | Colombia | Ukraine |
|-----------------------------|-------|--------------|----------|----------|
| HIV risk                    |       |              |          |          |
| HIV prevalence              | 6.1%  | 3.1%         | 0.4%     | 0.7%     |
| Maternal HIV incidence rate (per person-week) |       |              |          |          |
| Before first ANC visit during pregnancy | 0.000331 | 0.000227    | 0.00001  | 0.000002 |
| Between first ANC visit and delivery | 0.000331 | 0.000739    | 0.00002  | 0.00004  |
| Delivery to 6 weeks post partum | 0     | 0            | 0        | 0        |
| 6 weeks to 12 months post partum | 0.000269 | 0.0009      | 0.00003  | 0.00003  |
| Duration of incident maternal HIV infection (weeks) | 9     | 9            | 9        | 9        |
| HIV testing and prevention |       |              |          |          |
| Test kit unavailable        | 5.0%  | 5.0%         | 0.0%     | 0.0%     |
| Test acceptance             | 84.0% | 98.0%        | 89.0%    | 97.2%    |
| Received test results       | 97.8% | 98.0%        | 100.0%   | 100.0%   |
| HIV rapid test sensitivity in early infection | 66.7% | 66.7%        | 66.7%    | 66.7%    |
| HIV rapid test sensitivity in chronic infection | 100.0% | 100.0%      | 100.0%   | 100.0%   |
| HIV rapid test specificity  | 98.9% | 98.9%        | 98.9%    | 98.9%    |
| Dual test sensitivity in early infection | 66.7% | 66.7%        | 66.7%    | 66.7%    |
| Dual test sensitivity in chronic infection | 100.0% | 100.0%      | 100.0%   | 100.0%   |
| Dual test specificity       | 98.9% | 98.9%        | 98.9%    | 98.9%    |
| MTCT rate per week, acute maternal infection | 0.005-0.029 | 0.005-0.029 | 0.005-0.029 | 0.005-0.029 |
| MTCT rate per week, chronic maternal infection | 0.0005-0.023 | 0.0005-0.023 | 0.0005-0.023 | 0.0005-0.023 |
| Maternal PrEP use           | 0.0%  | 0.0%         | 0.0%     | 0.0%     |
| Health-care visits          |       |              |          |          |
| Attended first ANC visit    | 96.0% | 94.0%        | 97.4%    | 99.8%    |
| Attended late ANC visit     | 97.7% | 78.0%        | 88.1%    | 90.0%    |
| Facility delivery           | 61.8% | 96.0%        | 98.9%    | 99.0%    |
| Gestational age at first ANC visit (weeks) | 22    | 18           | 15       | 10       |
| Gestational age at late ANC visit (weeks) | 33    | 36           | 24       | 28       |
| Gestational age at delivery (weeks) | 39    | 39           | 39       | 39       |
| ARV coverage                |       |              |          |          |
| Maternal ART use            | 91.0% | 87.0%        | 87.8%    | 95.0%    |
| Virally suppressed women with HIV | 88.1% | 72.0%        | 88.4%    | 88.1%    |
| Weekly ART dropout (%)      | 0.3%  | 0.3%         | 0.3%     | 0.3%     |
| HIV-exposed infants receiving ARVs | 93.6% | 98.7%        | 96.0%    | 98.4%    |
| HIV-infected infants receiving ART* | 61.0% | 61.0%        | 57.9%    | 95.0%    |
| Breastfeeding               |       |              |          |          |
| Not breastfeeding in early post partum (0–6 weeks) | 2.5%  | 34.0%        | 97.9%    | 95.0%    |
| Not breastfeeding in mid post partum (6 weeks to 6 months) | 21.2% | 45.0%        | 97.9%    | 99.0%    |
| Not breastfeeding in late post partum (6–12 months) | 33.4% | 61.0%        | 97.9%    | 99.0%    |

(Table 2 continues on next page)
opinion (appendix pp 18–30). Empirical time-motion data collected in Kenya were used to calculate HIV testing costs in the country and estimates of time to testing were applied to other regions with country-specific labour and supply costs. Only incremental costs associated with testing and treatment were included. We excluded

| Maternal mortality rate (per person-week) | Kenya | South Africa | Colombia | Ukraine |
|-------------------------------------------|-------|--------------|----------|---------|
| During pregnancy                          | 0.0001| 0.0001       | 0.00002  | 0.00002 |
| Delivery to 6 weeks post partum           | 0.0006| 0.0002       | 0.0001   | 0.00003 |
| 6 weeks to 12 months post partum          | 0.0001| 0.0001       | 0.00002  | 0.00002 |

| Neonatal and infant mortality (per person-week) and survival | Kenya | South Africa | Colombia | Ukraine |
|---------------------------------------------------------------|-------|--------------|----------|---------|
| Stillbirth, syphilis positive (mother not treated)            | 21.0% | 21.0%        | 21.0%    | 21.0%   |
| Stillbirth, syphilis positive (mother treated)                | 3.8%  | 3.8%         | 3.8%     | 3.8%    |
| Neonatal mortality, birth to 6 weeks (mother had syphilis and not treated) | 9.0% | 9.0%         | 9.0%     | 9.0%    |
| Neonatal mortality, birth to 6 weeks (mother had syphilis and treated) | 1.8% | 1.8%         | 1.8%     | 1.8%    |
| Neonatal mortality, birth to 6 weeks (syphilis negative and HIV negative or HIV negative and on ART) | 0.5% | 0.3%         | 0.2%     | 0.1%    |
| Infant mortality, >6 weeks to 12 months                      | 0.03% | 0.03%        | 0.01%    | 0.01%   |
| Survival to 1 year, HIV negative or HIV positive and on ART  | 96.4% | 96.7%        | 98.7%    | 99.2%   |
| Survival to 1 year, HIV positive and not on ART              | 64.8% | 64.8%        | 64.8%    | 64.8%   |

| Syphilis |
|----------|
| Syphilis prevalence | 1.2% | 2.0% | 0.41% | 2.5% |
| Maternal syphilis incidence rate (per person-week) | 0.00008 | 0.00008 | 0.00004 | 0.000015 |
| Laboratory-based test sensitivity for syphilis | 80.0% | 80.0% | 80.0% | 80.0% |
| Laboratory-based test specificity for syphilis | 100.0% | 100.0% | 100.0% | 100.0% |
| Dual RDT sensitivity for syphilis | 87.0% | 87.0% | 87.0% | 87.0% |
| Dual RDT specificity for syphilis | 99.5% | 99.5% | 99.5% | 99.5% |
| Test correction factor† | 52.9% | 52.9% | 52.9% | 52.9% |
| Test coverage (laboratory-based testing)‡ | 73.0% | 83.0% | 62.8% | 98.0% |
| Received treatment (laboratory-based testing) | 50.0% | 90.0% | 90.8% | 99.0% |
| Received treatment (dual testing) | 62.5% | 92.5% | 93.1% | 99.3% |

| HIV costs (US$) |
|-----------------|
| Third generation rapid screening per woman | $2.64 | $7.72 | $6.68 | $3.99 |
| True-positive screening tests per woman* | $3.68 | $11.39 | $8.53 | $4.18 |
| False-positive screening tests per woman* | $26.39 | $34.17 | $74.81 | $19.80 |
| Maternal ART, per week per woman | $4.86 | $4.79 | $18.89 | $3.18 |
| Infant ARV prophylaxis (total cost) per infant | $2.32 | $3.82 | $52.10 | $4.00 |
| Maternal PrEP, per week per woman | $6.19 | $6.19 | $18.89 | $19.38 |
| Infant ART, per week per infant | $6.73 | $5.46 | $18.89 | $32.84 |

| Syphilis costs |
|----------------|
| RPR test screening per woman | $3.09 | $9.32 | $6.92 | $0.63 |
| TPHA test screening per woman | $9.59 | $13.04 | $3.43 | $1.14 |
| Benzathine benzylpenicillin injection, maternal treatment per woman | $5.79 | $8.69 | $8.21 | $1.99 |
| Intravenous benzathine benzylpenicillin, infant treatment per infant | $1.42 | $1.42 | $1.42 | $1.42 |
| Paediatric inpatient, per day per infant | $18.41 | $27.81 | $62.87 | $28.84 |

Details and sources provided in appendix (pp 18–30). All costs reported in 2017 US$. ANC=antenatal care. ART=antiretroviral therapy. ARV=antiretrovirals. MCH=maternal and child health. MTCT=mother-to-child transmission. PrEP=pre-exposure prophylaxis. RDT=rapid diagnostic test. RPR=rapid plasma reagin. TPHA=Treponema pallidum hemagglutination assay. *Based on the percent of infants with early infant diagnosis. †Test correction factor is the probability that a woman who tested positive for syphilis has an active syphilis infection. ‡Inclusive of stock-outs.

Table 2: Key model parameters
Sensitivity analysis

Uncertainty analyses were done for scenario one. One-way sensitivity analyses were used to assess the effect of changing individual model parameters, including HIV and syphilis prevalence, test coverage, test characteristics, gestational age at first antenatal care visit, and costs. We used low and high parameter values from confidence interval bounds for test characteristics, and 20% relative changes for all other parameters. We also did scenario implementation analyses to estimate the effect of epidemic changes (declines in HIV and syphilis prevalence) or HIV test acceptance on cost-effectiveness, calculated using Microsoft Excel (2016).

Role of the funding source

Collaborators from WHO were involved in study design, interpretation of results, and manuscript development; no other funders had a role in design, analysis, or interpretation. All collaborators had full access to all the data in the study. ALD had final responsibility for the decision to submit for publication, with concurrence from all authors.

Results

We modelled populations of 163 1470 pregnant women in Kenya, 110 699 in South Africa, 346 409 in Colombia, and 363 946 in Ukraine. In our models, routinely offering a dual HIV and syphilis rapid diagnostic test at the first antenatal care visit (scenario one) was more effective...
and less costly compared with individual HIV rapid diagnostic tests and laboratory assays for syphilis in all four countries (table 3). Per DALY averted, a dual rapid diagnostic test at the first antenatal care visit saved $26 in Kenya, $559 in South Africa, $844 in Colombia, and $545 in Ukraine. Dual rapid diagnostic tests at the first antenatal care visit averted 3005 congenital syphilis cases in Kenya, 2909 in South Africa, 403 in Colombia, and 639 in Ukraine and reduced DALYs by 9770 in Kenya, 9358 in South Africa, 1258 in Colombia, and 1980 in Ukraine. The PMTCT potential for HIV using the dual rapid diagnostic test and independent tests was similar in Ukraine and Colombia but resulted in more HIV infections in Kenya (one) and South Africa (five).

Retesting in late pregnancy with a dual rapid diagnostic test (scenario three) was more effective and less costly than retesting in late pregnancy with individual tests (scenario two). Retesting with a dual rapid diagnostic test averted 2526 additional HIV infections in Kenya and 3184 in South Africa compared with testing with the rapid diagnostic test only at the first antenatal care visit (scenario one). Retesting with the dual rapid diagnostic test was cost-effective in all countries (ICER in Kenya $270, South Africa $260, Colombia $2207, and Ukraine $205). However, in countries with low HIV prevalence, fewer additional HIV cases were averted by retesting in Colombia (23) and Ukraine (four). When retesting occurred by 32 weeks, 352 additional congenital syphilis cases were averted in Colombia and 297 were averted in Ukraine. By contrast, retesting did not avert any additional congenital syphilis cases in Kenya or South Africa because testing was done after 32 weeks.

The findings from scenario one were most sensitive to gestational age at first antenatal care visit, syphilis test coverage, overall cost of testing with dual rapid diagnostic tests, and HIV test acceptance, but were robust to parameter uncertainty (figure 1). ICER outcomes in South Africa were particularly sensitive to syphilis testing coverage, which varied between 66% and 100%; by contrast, we varied coverage in Kenya between 58% and 88%. Although ICERs were below the cost-effectiveness threshold in all one-way sensitivity analyses, ICER values for Kenya moved from cost-saving to cost-effective when gestational age at the first antenatal care visit, overall testing cost for the dual rapid diagnostic test, HIV test acceptance, syphilis testing coverage, and overall testing
cost per rapid plasma regain were modified. In Colombia, ICERs were most sensitive to HIV test acceptance, syphilis testing coverage, and overall testing costs for the dual rapid diagnostic test, HIV rapid diagnostic test, and rapid plasma reagin tests. In Ukraine, ICERs were most sensitive to changes to dual rapid diagnostic test sensitivity for syphilis and syphilis testing coverage.

Cost-effectiveness of various strategies were also robust to decreases in maternal HIV and syphilis prevalence, and changes in HIV testing coverage across scenarios and countries (figure 2). ICERs were most sensitive to changes in HIV and syphilis testing coverage across scenarios and countries (figure 2). ICERs were most sensitive to changes in HIV and syphilis testing coverage across scenarios and countries (figure 2). ICERs were most sensitive to changes in HIV and syphilis testing coverage across scenarios and countries (figure 2). ICERs were most sensitive to changes in HIV and syphilis testing coverage across scenarios and countries (figure 2).

Discussion
Our model-based analysis projected that incorporating a dual HIV and syphilis rapid diagnostic test at the first antenatal care visit is cost-saving in settings of high (South Africa), intermediate (Kenya), and low (Colombia and Ukraine) HIV prevalence, compared with using individual HIV and syphilis tests. Overall, our results suggest syphilis test coverage in antenatal care could increase if dual rapid diagnostic tests are used and that dual tests have the potential to overcome any limitations, such as the time to do individual HIV or syphilis tests. Cost-effectiveness of various strategies were also robust to decreases in maternal HIV and syphilis prevalence, and changes in HIV testing coverage across scenarios and countries (figure 2).

Conclusions were consistent across a range of settings with varied maternal syphilis prevalence (0-41–2.5%), and the results are consistent with a previous analysis from Malawi. Compared with individual tests for HIV and syphilis, dual HIV and syphilis testing at the first antenatal care visit averted congenital syphilis infections and DALYs, but it led to small increases in the absolute number of infant HIV infections in Kenya and South Africa. Increases in infant HIV infections are attributed to increased infant survival when syphilis-related stillbirth and neonatal deaths were averted, leading to a larger population of infants at risk for HIV.

Figure 2: Scenario implementation analyses
(A) Decreasing HIV prevalence in Kenya and South Africa. (B) Decreasing HIV prevalence in Colombia and Ukraine. (C) Decreasing syphilis prevalence in Kenya and South Africa. (D) Decreasing syphilis prevalence in Colombia and Ukraine. (E) Variation in HIV testing acceptance in Kenya and South Africa. (F) Variation in HIV testing acceptance in Colombia and Ukraine. Base case used independent tests at country-specific timepoints. Scenario one used dual HIV and syphilis testing with no retesting during late ANC. Scenario two used independent HIV and syphilis tests with retesting during late ANC. Scenario three used dual HIV and syphilis testing with retesting during late ANC. ICERs compare scenario one with the base case and scenario three with scenario one, under alternative parameter values. ANC=Antenatal care. DALY=Disability-adjusted life-year. ICER=Incremental cost-effectiveness ratio.
In all countries, scenario three, which used a dual rapid diagnostic test at the first antenatal care visit and for retesting, was less costly and more effective than scenario two, which used individual tests at the same timepoints. Cost savings were primarily driven by lower costs per woman tested with a dual rapid diagnostic test at the first antenatal care visit compared with combined overall costs of testing with individual assays. In all countries except Kenya, a dual rapid diagnostic test kit was less expensive than the combined cost of an HIV rapid diagnostic test kit and a rapid plasma reagin test. Furthermore, testing with dual rapid diagnostic tests reduced labour costs in all countries. Although the dual rapid diagnostic test remained cost-effective with 20% increases in test kit cost, it was not always cost-saving.

Because retesting did not avert additional syphilis infections in countries with intermediate and high HIV prevalence and antenatal care visits occurred after 32 weeks in these settings, retesting for HIV only (not modelled) is likely to be similarly effective and less costly than retesting with a dual rapid diagnostic test. In a previous analysis of the maternal HIV retesting model used for this analysis, retesting averted few infant HIV infections and has restricted potential for the prevention of MTCT of HIV in low HIV prevalence countries (Meisner J, unpublished data). Although retesting for maternal syphilis is not recommended by WHO, if retesting occurs early enough for maternal syphilis treatment to prevent adverse birth outcomes (≤32 weeks), it has the potential to avert stillbirths and congenital syphilis cases attributed to incident maternal syphilis infections or infections not previously detected. Our results suggest that retesting in countries with low HIV prevalence would avert few HIV infections and no congenital syphilis infections.

Operationally, retesting using dual HIV and syphilis rapid diagnostic tests might have advantages for simplifying testing algorithms that avoid laboratory-based syphilis tests and reducing prices through increasing procurement volumes in high-HIV-burden settings. However, challenges exist for the allocation of restricted resources towards a costly retesting strategy with minimal benefits. Programmes will need to clarify unique needs for women previously diagnosed with HIV who only require syphilis testing, and for those diagnosed with syphilis during their current pregnancy who require only HIV testing. Although many PMTCT programmes use HIV rapid diagnostic tests, laboratory testing for syphilis is still common in many low-income and middle-income settings. The logistics of laboratory testing for syphilis—such as sample collection following antenatal care visits and delays in waiting for laboratory results (eg, confirmatory assays for reactive tests)—might result in women not getting tested, not receiving test results, lower syphilis treatment coverage, and more congenital syphilis cases.

Dual HIV and syphilis rapid diagnostic tests have many characteristics that appeal to individuals accessing services and health-care providers. They only require a single blood draw and provide immediate results, allowing health-care providers to make life-saving decisions regarding treatment more quickly. Training for health-care providers would be simplified because of the need for only one initial screening test for HIV and syphilis, rather than two. Additionally, dual rapid diagnostic tests have the advantage of streamlined procurement of test kits and reagents. However, countries will still need a supply of single HIV rapid diagnostic tests to avoid syphilis retesting when not indicated, to comply with national testing policies, to prevent missed opportunities to detect maternal HIV or syphilis infection, and to prevent MTCT in the event of dual rapid diagnostic test stock-outs. This could result in challenges determining which programmes should fund test kits, with siloed funding streams for HIV and syphilis. In addition, the sensitivity and specificity of the syphilis component of the dual assay has been reported to be variable under field conditions, and quality control measures should be followed to ensure accurate interpretation of results.

Our analysis should be viewed in the context of some limitations. Syphilis testing in antenatal care might be heterogeneous between and within countries, and the assumption that countries exclusively use laboratory-based syphilis testing might be inaccurate. There could be variation in the assay type within a country based on geographical differences or programmatic partners. Our results might have restricted generalisability to settings currently using rapid diagnostic tests for syphilis, and are subject to market prices for syphilis and HIV test kits and labour costs. Our retesting scenarios include women who are tested for the first time during late antenatal care, which might overstate the benefit of the retest relative to the first test. Because of the separate construction of the HIV and syphilis models, we were unable to model syphilis and HIV co-infection, and did not model risk of fetal loss due to HIV. However, whether risks of MTCT of HIV are higher in the context of maternal co-infection is unclear. Additionally, because of the small amount of data on the effectiveness of treating maternal syphilis infection after 32 weeks’ gestation, we assumed that there would not be an effect on PMTCT of syphilis after this point, which might underestimate maternal and infant benefits of retesting later in pregnancy. We also restricted syphilis-related disability to the first 3 years of life, which probably underestimates the burden of congenital syphilis. Adding additional years of disability from congenital syphilis would increase the positive health effect of dual testing. Although we used the literature and in-country experts to populate model parameters, some parameters were unavailable or based on a small amount of data, particularly related to syphilis. Future studies are needed to inform assumptions about breastfeeding practices, syphilis treatment coverage, and rates of adverse maternal syphilis outcomes. Our results were qualitatively robust to all one-way sensitivity analyses, but ICER
estimates might be imprecise. We had insufficient data to inform many parameter distributions and therefore did not do probabilistic sensitivity analysis to estimate credible intervals. Despite the limitations, our results are probably conservative estimates because we did not include maternal treatment benefits, such as improved quality of life, reduced risk of sexual transmission, or health system costs averted. Our estimate of time to complete a rapid test of 30 min per patient might be conservative in some settings. Our analysis has several strengths. We included countries that represented variations in HIV prevalence, syphilis prevalence, and geographical region. We compared HIV and syphilis testing strategies across countries using the same methods and model structure, and similar assumptions. Our analysis expanded on previous studies by including incident syphilis and HIV infections, in addition to prevalent infections.

Dual HIV and syphilis rapid diagnostic test as the first test in antenatal care can improve efforts to provide integrated services within the antenatal care setting, saving overall costs while improving health outcomes for women and their children. Breaking down siloes of vertical programmes can improve the quality of antenatal care and help to streamline service delivery, thereby accelerating progress on efforts to achieve dual elimination of MTCT of HIV and syphilis.

Contributors
PJR, DAR, JM, RB, MSj, RB, MB-D, MNO, CJ, MMT, and ALD designed the study. PJR, DAR, JM, BC, MBM, AB, AV, AS, GG, TC, CJ, MMT, and ALD collected data to populate the model. PJR, DAR, JM, MSJ, CJ, MMT, and ALD searched the literature. PJR, DAR, JM, MS, and ALD were responsible for constructing the model and analysing model results. All authors contributed to writing the manuscript and interpreting the results.

Declaration of interests
We declare no competing interests.

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