Health Impact of Delayed Implementation of Cervical Cancer Screening Programs in India: A Modelling Analysis

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Abbreviations used: CIN: cervical intraepithelial neoplasia; GDP: gross domestic product; HPV: human papillomavirus; VIA: visual inspection with acetic acid; WHO: World Health Organization.
**Novelty and Impact:** India has the largest burden of cervical cancer in the world. To quantify the consequences of delaying implementation of organized cervical cancer screening in India, we used mathematical models to estimate the cumulative number of cervical cancer cases and deaths averted under different program implementation scenarios. Delays in implementation of screening programs in high-burden settings will result in substantial morbidity and mortality among women beyond the age for adolescent HPV vaccination.
ABSTRACT
India has the highest burden of cervical cancer in the world. To estimate the consequences of delaying implementation of organized cervical cancer screening, we projected the avertable burden of disease under different implementation scenarios of a screening program. We used an individual-based microsimulation model of human papillomavirus (HPV) infection and cervical cancer calibrated to epidemiologic data from India to project age-specific cancer incidence and mortality reductions associated with screening (once-in-a-lifetime among women aged 30 to 34 years) with one-visit visual inspection with acetic acid (VIA) and one- and two-visit HPV DNA testing. We then applied these reductions to a population model to project the lifetime cervical cancer cases and deaths averted under different implementation scenarios taking place from 2017 to 2026: 1) immediate implementation of screening with currently available screening tests (one-visit VIA, two-visit HPV testing); 2) immediate implementation of screening with currently available screening tests, with a switch to point-of-care one-visit HPV testing in five years; and 3) five-year delayed implementation of screening with current screening tests or point-of-care HPV testing. Immediate implementation of two-visit HPV testing with a switch to one-visit HPV testing averted 574,100 cases and 382,500 deaths over the lifetimes of 81.4 million 30 to 34-year-old women screened once between 2017 and 2026. Delayed implementation with a one-visit HPV test averted 209,300 cases and 139,100 deaths. Delays in implementation of screening programs in high-burden settings will result in substantial morbidity and mortality among women beyond the age for adolescent HPV vaccination.
Background

India has the largest burden of cervical cancer in the world, with an estimated 123,000 incident cases occurring annually.\(^1\) This accounts for approximately 23% of cases worldwide, and 32% of cases in less developed countries.\(^1\) Yet the disease is preventable through either prophylactic vaccination against human papillomavirus (HPV)—the sexually transmitted virus that causes cervical cancer—or screening and treatment of precancerous lesions caused by persistent HPV infection. While the introduction and scale-up of adolescent HPV vaccination programs would substantially reduce the number of cervical cancer cases in years to come, the full benefits of HPV vaccination will not be realized for more than 30 years; to date, there are several HPV vaccination demonstration projects in India, but the vaccine is not available through the national immunization program.\(^2\) In the interim, screening is the only form of prevention for the two to three generations of women beyond the target age of adolescent vaccination. Only an estimated 3.1% of women in India reported receiving a Pap smear in the last 3 years.\(^3\)

The World Health Organization (WHO) recommends screening with HPV DNA testing where resources are available; visual inspection with acetic acid (VIA) represents an acceptable alternative in low-resource settings.\(^4\) Recent guidelines from the American Society for Clinical Oncology recommend HPV testing for all resource levels, and if resources are not available, VIA should be offered with the goal of developing screening infrastructure and switching to HPV testing as soon as possible.\(^5\) Promising developments to improve the delivery of preventive care in low-resource settings include the introduction of low-cost HPV tests with samples collected by either a health provider or the woman herself;\(^6\) advances in the use of mobile phones for reaching patients;\(^7\) and introduction of preventive treatment technologies that do not rely on compressed gas, which may be difficult and costly to obtain.\(^8\) Additionally, new clinically validated technology exists to facilitate point-of-care (i.e., one-visit screen-and-treat) HPV testing, but has not yet been implemented outside of small demonstration projects.\(^9\) But like other lower-middle income settings, India faces shortages of health workers, insufficient referral
processes, and financial constraints that have impeded the success of organized screening that systematically targets eligible women. While operational guidelines developed by the Ministry of Health recommend screening with VIA every 5 years for women over age 30 years, with initial implementation in 100 districts providing data, states will need to fund and implement broader efforts. Without substantial political will and the injection of government and donor funds, the implementation of screening programs will likely continue to stall.

To provide information on the consequences of further delays in the implementation of cervical cancer screening, we quantified the potential health impact of an organized national screening program in India by estimating the cumulative number of cervical cancer cases and deaths averted over the lifetimes of women aged 30 to 34 years under different implementation scenarios taking place from 2017 to 2026: 1) immediate implementation of screening with currently available screening tests (i.e., VIA, HPV DNA testing); 2) immediate implementation of screening with currently available screening tests, with a switch to point-of-care HPV testing in five years; and 3) five-year delayed implementation of screening with current screening tests or point-of-care HPV testing.

**METHODS**

**Study design and data sources**

We used an individual-based microsimulation model of the natural history of HPV infection and cervical cancer—calibrated to epidemiologic data on cervical disease burden in India—to estimate the reductions in cervical cancer incidence and mortality attributable to screening with either VIA or HPV DNA testing. Assumptions and data inputs pertaining to the effectiveness of each test, including coverage of the target population, test performance, compliance with recommended follow-up, eligibility for cryotherapy, and treatment efficacy, are presented in Table 1. We then applied the percent reductions in cervical cancer incidence and mortality in each five-year age group from ages 30
to 79 years to population and disease burden data using the Excel-based CERVIVAC model, a tool developed for the Pan American Health Organization’s (PAHO) ProVac Initiative that contains separate modules for evaluating the costs and health impact associated with either HPV vaccination or cervical cancer screening and treatment of precancerous lesions. Female population data by birth cohort were based on estimates for India through 2100 from the United Nations Population Division (the 2012 revision). Cervical cancer incidence and mortality rates in the absence of screening were based on GLOBOCAN 2012, produced by the International Agency for Research on Cancer (IARC). The CERVIVAC model was used to estimate the numbers of cervical cancer cases and deaths averted for each implementation scenario based on screening test, age at screening, screening coverage of the target population, and year of implementation.

Statistical analysis

The individual-based microsimulation model of the natural history of HPV infection and cervical cancer, as well as the model calibration process, have been previously described. In brief, the model tracks individual girls who enter the model at age 9 years with a healthy cervix and then transition between mutually exclusive health states that include HPV infection, cervical intraepithelial neoplasia (CIN) (grade 2 or 3), and invasive cervical cancer. As individuals age, they can acquire HPV infections, which may either clear or progress to CIN2 or CIN3. Women with CIN2 or CIN3 may regress or progress to invasive cancer, which can be detected at the local, regional, or distant stage. Transition probabilities may vary by age, HPV type (stratified by HPV 16, 18, 31, 33, 45, 52, 58, other high-risk types, and low-risk types), duration of infection or CIN status, and naturally acquired immunity from prior HPV infection. Death from non-cervical causes can occur in any health state, depending on a woman’s age, and stage-specific excess mortality due to cervical cancer can occur after its onset.
To calibrate the model to epidemiologic data on age-specific high-risk HPV prevalence and cervical cancer incidence from India, we estimated baseline “prior” input parameter values for natural history transitions using longitudinal data. To reflect heterogeneity in age- and type-specific HPV incidence between India and input source study locations, as well as uncertainty surrounding natural immunity following initial infection and progression and regression of CIN2 and CIN3, we set plausible ranges around these input parameter values. Repeated model simulations (in the absence of any intervention) selected a single random value from the plausible range for each uncertain parameter, creating a unique natural history input parameter set. For each unique parameter set, we computed a goodness-of-fit score by summing the log-likelihood of model-projected outcomes to represent the quality of fit to epidemiologic data from India (i.e., calibration targets). We selected the top 50 good-fitting input parameter sets to use in analyses as a form of probabilistic sensitivity analysis. Natural history model inputs for the India microsimulation model, as well as model fit to epidemiologic data, are presented in the Appendix.

We used the calibrated microsimulation model to estimate the average percent reduction in cervical cancer incidence and mortality in each five-year age group (from ages 30 to 79 years; reductions in women aged 80 to 100 years were assumed to be the same as in women aged 75 to 79 years) associated with the following screening tests, administered once in a lifetime, relative to no intervention: 1) one-visit VIA; 2) two-visit HPV DNA testing; and 3) one-visit point-of-care HPV DNA testing (Table 2) at screening coverage levels of 10% and 20% of women aged 30 to 34 years. For VIA, we assumed that women who were screen-positive and eligible for cryotherapy were treated at the same clinical visit. For two-visit HPV DNA testing, we assumed women self-collected a vaginal HPV specimen during an initial clinical encounter, and subsequently received results and treatment (if HPV-positive and eligible for cryotherapy) in a separate visit. For one-visit HPV DNA testing, we assumed that self-collection and treatment of HPV-positive eligible women took place in a single visit. In all strategies,
women who were not eligible for cryotherapy were referred to further diagnostic testing with
colposcopy, and subsequent treatment if they received a histologic diagnosis of CIN1 or higher. Loss to
follow-up rates were assumed to be 20% per clinical encounter.

The CERVIVAC model was then used to estimate the number of cervical cancer cases and deaths
averted over the lifetimes of women aged 30 to 34 years for the following implementation scenarios
between 2017 and 2026, relative to no screening: 1) immediate implementation of screening with one-
visit VIA beginning in 2017 (“Immediate VIA”); 2) immediate implementation of screening with two-visit
HPV DNA testing beginning in 2017 (“Immediate two-visit HPV”); 3) immediate implementation of
screening with VIA beginning in 2017 and switching to point-of-care HPV testing (i.e., one-visit HPV
testing) in 2022 (“Immediate VIA, switch to one-visit HPV”); 4) immediate implementation of screening
with two-visit HPV DNA testing beginning in 2017 and switching to one-visit HPV testing in 2022
(“Immediate two-visit HPV, switch to one-visit HPV”); 5) delayed implementation of screening with one-
visit VIA, beginning in 2022 (“Delayed VIA”); 6) delayed implementation of screening with two-visit HPV
DNA testing, beginning in 2022 (“Delayed two-visit HPV”); 7) delayed implementation of screening with
one-visit point-of-care HPV testing in 2022 (“Delayed one-visit HPV”). Screening modalities were
selected based on India’s operational guidelines, which recommend VIA,11 and HPV DNA testing, which
has been demonstrated as cost-effective in previous modeling work.13 For strategies involving
immediate implementation, we assumed initial coverage was 10% each year of each birth cohort of
women aged 30 to 34 years in that year for the first five years of the screening program (2017-2021),
and then increased to 20% each year of each birth cohort aged 30 to 34 years in the next 5 years of the
program (2022-2026). For strategies involving delayed implementation, we assumed no screening from
2017-2021 and 10% coverage each year of each birth cohort of women aged 30 to 34 years from 2022-
2026. Thus, depending on the year, cohort, and coverage level, effective coverage for a particular birth
cohort considered could be zero (e.g., for 34 year old women in 2017 under delayed implementation
scenarios), or nearly 100% (e.g., for 30 year old women in 2022 after immediate implementation with a switch to 1-visit HPV).

RESULTS

Without screening, the CERVIVAC model projected 3,824,700 cervical cancer cases and 2,878,300 deaths over the lifetimes of women aged 30 to 34 years between 2017 and 2026 (equivalent to lifetime risks of 2.5% and 1.9%, respectively, of the 151 million women alive in these cohorts when each first became eligible for screening). The cumulative number of cervical cancer cases and deaths that could potentially be averted over the lifetimes of women in India aged 30 to 34 years during the intervention period (2017-2026) are presented in Table 3; the timing of cases and deaths averted, by calendar year, is displayed in Figure 1. Scenarios involving immediate implementation of screening in 2017 were predicted to avert substantially more cervical cancer cases and deaths than scenarios that delayed implementation until 2022. Immediate implementation of two-visit HPV testing with a switch to one-visit HPV testing was the most effective scenario, averting an estimated 574,100 cases and 382,500 deaths. Immediate implementation of VIA with a switch to one-visit HPV testing was the next most effective scenario, and was projected to avert 521,900 cases and 346,600 deaths.

Among immediate implementation strategies involving only currently available screening tests, immediate two-visit HPV testing was more effective than immediate VIA, averting an estimated additional 149,800 cases and 110,100 deaths.

Despite comparable single-visit delivery and more favorable test sensitivity to detect CIN2 and higher, delayed implementation of one-visit HPV testing averted fewer cases and deaths than immediate implementation of VIA. Delayed implementation of one-visit HPV testing averted 364,800 fewer cases and 243,400 fewer deaths than immediate two-visit HPV with a switch to one-visit HPV.
Delayed implementation of two-visit HPV testing and one-visit VIA were projected to avert the fewest number of cases and deaths.

DISCUSSION

This study supports the position that even short delays implementing organized screening programs in low-resource settings with a high cervical cancer burden will result in substantial morbidity and loss of life. We used a model-based approach—relying on both an individual-based microsimulation model that captures the reduction in disease burden associated with nuanced screening algorithms and a population model—to project the number of cervical cancer cases and deaths that could be averted with immediate implementation of organized cervical cancer screening in India versus a five-year delay in implementation. Our roll-out assumptions were conservative, assuming only 10% of each birth cohort of women aged 30 to 34 years received screening in each of the first five years of implementation. For immediately implemented strategies, we assumed scale-up to 20% coverage after five years (either continuing with currently available screening tests—VIA or two-visit HPV—or introducing a new one-visit HPV test). Among the strategies considered for delayed implementation was one-visit HPV testing, to determine whether waiting for technological improvements (i.e., point-of-care HPV testing) can overcome the potential loss of life due to postponing roll-out. Even under assumptions of low population coverage over the next ten years, we projected that immediate implementation of two-visit HPV testing with a switch to one-visit HPV testing can avert 574,100 cases and 382,500 deaths in India. If implementation is delayed by five years to wait for an improved HPV test that facilitates same-day screening and treatment of precancer, 364,800 fewer cases and 243,400 fewer deaths will be averted. Our model projections also indicate that even if a point-of-care test does not become widely available in five years, immediate implementation of two-visit HPV testing with moderate scale-up may still avert 480,600 cases and 328,000 deaths relative to no screening.
Three large randomized trials have demonstrated that screening can reduce cervical cancer incidence and mortality in India. In the Osmanabad district, a single round of HPV testing reduced advanced cervical cancer incidence and mortality by approximately 50%, while VIA did not. However, a single round of VIA in Dindigul district reduced incidence and mortality by 25% and 35%, respectively. Multiple rounds of VIA in Mumbai reduced cancer mortality by approximately 30%, but this was likely due to earlier detection of cancer rather than reduced incidence, and required extensive training of health workers. To successfully scale VIA, substantial training and quality control efforts will be needed to maintain test performance.

To date, HPV testing and VIA have generally been limited to clinical studies and demonstration projects in India. A notable exception is the Tamil Nadu Health Systems Project, which piloted a large-scale government-led VIA program in 2007. By 2015, the program achieved 71% coverage of women aged 30 to 60 years. While the program demonstrated the feasibility and acceptability of a cervical cancer screening program within the public health sector in one state, it also identified implementation challenges. Many of the women referred to treatment did not receive it, as women were referred to colposcopy at higher level facilities rather than receiving immediate treatment following a positive screening test. Furthermore, the project was supported by the World Bank (US$19 million for the non-communicable disease program targeting cardiovascular disease, breast cancer, and cervical cancer), highlighting the importance of multilateral institutions and the international donor community in strengthening health systems and building capacity to expand the reach of screening programs.

There are several limitations to our analysis. Our modelled screening algorithms assumed that currently available technologies could facilitate screening and treatment of precancer in one or two visits with VIA or HPV testing, respectively. While this may be the case at higher level facilities with cryotherapy available on-site, the equipment and compressed gas needed for cryotherapy are not available at most primary health facilities in India. Thus, women may be required to bear the time and
travel costs to attend a referral facility, reducing the likelihood of receiving treatment. However, the
coverage rates we assumed are lower, covering a much narrower age range (30 to 34 years) than the
71% coverage achieved in the Tamil Nadu Health Systems Project for women aged 30 to 60 years, and
might resemble program coverage rates if initial efforts focused on delivery of screen-and-treat
algorithms in referral centers and/or mobile clinics to ensure on-site availability of ablative treatment.
While it is possible that delayed implementation could yield higher population coverage than immediate
implementation, this would require very rapid deployment of resources and capacity building, as well as
social mobilization and outreach. In India, where 67% of the population lives in rural areas,\textsuperscript{20} our
assumptions of high compliance with treatment (100% with VIA; 80% with two-visit HPV testing) among
screen-positive women eligible for cryotherapy are more likely to be achievable when the number of
required visits for a screening episode are low and treatment is proximal. Furthermore, we did not
consider the impact of catch-up HPV vaccination in older women over age 30 years, for which the cost-
effectiveness has not been demonstrated in low-resource settings.

The accuracy of microsimulation model projections depends upon the validity of our calibration
approach. While we calibrated the model to fit epidemiologic data from India—using the expected
value outcomes from 50 good-fitting sets of natural history inputs to reflect uncertainty—the high-risk
HPV prevalence from a large demonstration project in Hyderabad\textsuperscript{6} and the Nagpur cancer registry\textsuperscript{18} do
not fully capture the geographic variation in disease burden across India. Furthermore, our data on age-
specific prevalence of high-risk HPV was restricted to women aged 30 to 49 years. However, exercises to
validate model projections suggest internal consistency between our calibrated model, test
performance characteristics for VIA and HPV testing drawn from a demonstration project in a public
sector clinic in Hyderabad,\textsuperscript{6} and two large randomized trials of screening impact in India that were not
used to derive model inputs (Appendix).\textsuperscript{24, 25} The calibrated and validated microsimulation model
allowed us to estimate age-specific cervical cancer incidence and mortality reductions resulting from
nuanced screening and management algorithms, given uncertainty in the natural history parameters and epidemiologic data.

Our modeling approach assumed that the burden of HPV and cervical cancer were consistent over the lifetime of women aged 30 to 34 years between 2017 and 2026. A recent modeling analysis used age-period-cohort models to project the future number of new cervical cancer cases in six Baltic, central, and eastern European countries through 2040. Based on recorded trends in the most recent generations, these projections assumed cervical cancer incidence continued to increase across the study period. It is unclear whether cervical cancer burden is also increasing in India, although HPV transmission models (which integrate sexual behavior data) might inform estimates of future cancer risk among young women at or approaching screening age. We note that if GLOBOCAN 2012 underestimates the future cancer risk of women who will become eligible for screening over the intervention period we consider, we may underestimate the number of cases averted due to organized screening.

The present analysis does not address the cost-effectiveness or affordability of implementing a screening program, although both the value and budget impact need to be favorable in order for a program to be sustainable. Elsewhere, we have estimated the cost-effectiveness of various screening strategies in India, and found that a two-visit approach with HPV DNA testing three times in a woman’s lifetime would be very cost-effective, with an incremental cost-effectiveness ratio below India’s per capita GDP. We have found that the value of new technologies such as point-of-care HPV testing in India is potentially high if linkage to treatment can be assured. Furthermore, we have also estimated the financial cost of screening women aged 35 years in India with HPV testing or VIA once in a lifetime to be US$830 million over 10 years (or approximately US$83 million per year); however, this scenario represents an upper bound, as it assumed the program would reach full coverage within five years. To place this figure in context, US$1.08 billion of development assistance for health was disbursed to India in 2013. Decision makers will need to weigh the value and financial cost of implementing organized
cervical cancer screening in relation to other disease priority areas. Development assistance will be required for screening implementation. We present these findings—which suggest that immediate implementation of HPV testing has the potential to save the lives of hundreds of thousands of women in India who are beyond the target age of HPV vaccination—to catalyze the policy dialogue. This analysis quantifies the human cost of waiting to act, despite the availability of screening technologies with demonstrated effectiveness and cost-effectiveness; this toll only increases over time.

Implementation of organized screening programs is difficult work, requiring political will and coordination across levels of the health care system. A one-size-fits-all approach is unlikely to work across all states or districts within a state. Without injection of funds from the international donor community to strengthen health systems and support cancer prevention efforts, as well as governmental budgetary allocations and commitment to sustainability, the costs of cervical cancer screening will fall predominantly on women and their families; achieving high coverage and management of screen-positive women will remain a barrier to successful programs. Despite these challenges, our study provides quantitative evidence that the failure to begin prompt implementation of cervical cancer screening in India with currently available technologies will result in substantial morbidity and mortality among women who are critical to social and economic stability.
ADDITIONAL INFORMATION

Ethics approval and consent to participate: Not applicable.


Conflict of interest: Dr. Campos, Dr. Tsu, Ms. Regan, Dr. Resch, Dr. Clark, and Dr. Kim declare no conflict of interest. Dr. Jeronimo is a consultant for Merck, a manufacturer of HPV vaccines, and was a consultant for Qiagen Inc, manufacturer of HPV diagnostic tests, in 2017. He was co-owner and Deputy Manager of Onco Prev International, a Peruvian company, from 2012 through March 2017. Onco Prev offers cervical cancer screening services and in 2016 also began positioning for distribution of medical devices including colposcopes and the Liger thermocoagulator. Onco Prev International did not commercialize any medical instrument during the time Dr. Jeronimo was part of the company.

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Author contributions: NGC contributed to study conceptualization, study design, data collection, data analysis, data interpretation, and drafted the manuscript. VT contributed to study conceptualization and revision of the manuscript. JJ contributed to study conceptualization and revision of the manuscript. CR
contributed to data analysis, presentation of figures, and revision of the manuscript. SR contributed to study conceptualization, study design, data interpretation, and revision of the manuscript. AC contributed to programming of the CERVIVAC model, data collection, and revision of the manuscript. SS contributed to programming of the microsimulation model, data interpretation, and revision of the manuscript. JJK contributed to study conceptualization, study design, data interpretation, and revision of the manuscript.

**Supplementary Information:** Supplementary Information File 1 (.doc) includes model input parameters and details on model calibration and validation.
REFERENCES


Figure legend.

Figure 1. Cumulative number of expected averted cervical cancer A) cases and B) deaths are presented for the following screening strategies: 1) immediate two-visit human papillomavirus (HPV) DNA testing followed by a switch to one-visit HPV DNA testing after five years (dark blue line); 2) immediate visual inspection with acetic acid (VIA) followed by a switch to one-visit HPV DNA testing after five years (orange line); 3) immediate two-visit HPV DNA testing (red line); 4) immediate VIA (yellow line); 5) delayed one-visit HPV DNA testing (turquoise line); 6) delayed two-visit HPV DNA testing (purple line); and 7) delayed VIA (green line). Immediate implementation strategies for once in a lifetime screening assume 10% coverage each year of women aged 30 to 34 years in India between 2017 and 2021, and 20% coverage each year of women aged 30 to 34 years between 2022 and 2026. Delayed implementation strategies assume no coverage of women aged 30 to 34 years between 2017 and 2021, and 10% coverage each year of women aged 30 to 34 years between 2022 and 2026. The difference between the two-visit and one-visit HPV DNA testing strategies is that, in the two-visit strategy, 20% of women screened do not show up to receive HPV results and, if HPV-positive, treatment. In the one-visit strategy, all women who screen positive and are eligible are assumed to receive treatment with cryotherapy.
Table 1. Values and data sources for model variables.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Value</th>
<th>Data source</th>
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<tbody>
<tr>
<td>Age of target population</td>
<td>30 to 34 years</td>
<td>Assumption</td>
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<tr>
<td>Population coverage of screening program</td>
<td></td>
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<tr>
<td>Immediate implementation (2017)</td>
<td>10% (2017-2021), 20% (2022-2026)</td>
<td>Assumption</td>
</tr>
<tr>
<td>Delayed implementation (2022)</td>
<td>10% (2022-2026)</td>
<td>Assumption</td>
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<tr>
<td>Individual-based Monte Carlo simulation model</td>
<td></td>
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<td>Loss to follow-up per visit</td>
<td>20%</td>
<td>Assumption</td>
</tr>
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<td>Test sensitivity/specificity for CIN2+ VIA</td>
<td>0·55/0·92</td>
<td>Jeronimo et al. 2014</td>
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<tr>
<td>HPV DNA test (self-collection)</td>
<td>0·76/0·95</td>
<td></td>
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<tr>
<td>Eligibility for cryotherapy</td>
<td></td>
<td></td>
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<tr>
<td>No lesion or CIN1</td>
<td>100%</td>
<td>Campos et al. 2015</td>
</tr>
<tr>
<td>CIN2</td>
<td>85%</td>
<td></td>
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<tr>
<td>CIN3</td>
<td>75%</td>
<td></td>
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<tr>
<td>Cancer</td>
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<tr>
<td>Effectiveness of cryotherapy to treat CIN2/3</td>
<td>92%</td>
<td>Sauvaget et al. 2013</td>
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<tr>
<td>Test sensitivity/specificity for CIN1+, colposcopy</td>
<td>0·50/0·96</td>
<td>Jeronimo et al. 2014</td>
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<td>Effectiveness of cryotherapy/LEEP following colposcopy to treat CIN2/3</td>
<td>96%</td>
<td>Chirenje et al. 2001</td>
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<td>CERVIVAC population model</td>
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<td></td>
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<tr>
<td></td>
<td></td>
<td>Revision</td>
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<tr>
<td>Cervical cancer incidence</td>
<td>By age, in 5-year groups</td>
<td>Globocan 2012</td>
</tr>
<tr>
<td>Cervical cancer mortality</td>
<td>By age, in 5-year groups</td>
<td>Globocan 2012</td>
</tr>
<tr>
<td>Reduction in cervical cancer incidence attributable to screening</td>
<td>By age, in 5-year groups</td>
<td>Individual-based Monte Carlo microsimulation model calibrated to 1) age-specific prevalence of high-risk HPV in Hyderabad and 2) the Nagpur cervical cancer registry (1998-2002)</td>
</tr>
<tr>
<td>Reduction in cervical cancer mortality attributable to screening</td>
<td>By age, in 5-year groups</td>
<td>Individual-based Monte Carlo microsimulation model calibrated to 1) age-specific prevalence of high-risk HPV in Hyderabad and 2) the Nagpur cervical cancer registry (1998-2002)</td>
</tr>
</tbody>
</table>
a CIN: cervical intraepithelial neoplasia; HPV: human papillomavirus; LEEP: loop electrosurgical excision procedure; VIA: visual inspection with acetic acid.
b Loss to follow-up was defined as the proportion of women who do not return for each subsequent clinical encounter, relative to the previous visit. In the two-visit HPV DNA test strategy, loss to follow-up applied to the results/cryotherapy visit. In both the one-visit VIA, two-visit HPV DNA test, and one-visit HPV DNA test strategies, loss to follow-up applied to both the diagnostic confirmation and treatment visits for women who were not eligible for cryotherapy immediately following a positive screening test.
c HPV DNA test performance characteristics (for self-collection of vaginal samples) were based on a demonstration project of the careHPV test in Hyderabad. We assumed the same test performance characteristics for 1- and two-visit HPV testing strategies.
d Eligibility for cryotherapy immediately following a positive screening test was based upon a woman’s true health state in the model.
e Test performance characteristics of colposcopy in the PATH START-UP demonstration project were derived from the worst diagnosis of the local pathologist relative to the worst diagnosis by a quality control pathologist (gold standard); we applied the treatment threshold of CIN1+, although this was not the treatment threshold in the START-UP project. To derive test performance of colposcopy, we excluded histological classifications that were inadequate or with a histological classification other than negative, CIN1, CIN2, CIN3, or cancer. Because CIN1 is not a true underlying health state in the microsimulation model, performance of colposcopy in the model is based on the underlying health states of no lesion, HPV infection, CIN2, or CIN3. For a treatment threshold of CIN1, we weighted sensitivity of colposcopy for women with HPV based on the prevalence of CIN1 among women with HPV infections in the START-UP studies.
f Data from the United Nations World Population Prospects\textsuperscript{15} project female population sizes for single ages and single years from 1950 until 2100.
g Globocan 2012\textsuperscript{16} data was provided by the Institut Catala d’Oncologia for a previously published analysis\textsuperscript{17} in 5-year age groups from ages 30 to 49 years. Globocan data is collapsed into one age group for cervical cancer incidence and mortality rates over 75 years; in the CERVIVAC model, we linearly interpolated the rates in the oldest age groups, assuming that the Globocan data for 75 years and over applied to women aged 85 to 89 years and then decreased at a linear rate.
## Table 2. Screening strategies and scale-up scenarios.

<table>
<thead>
<tr>
<th>Strategy</th>
<th>Coverage level (% of women aged 30-34 years) and screening test by year</th>
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<tbody>
<tr>
<td>Immediate implementation of currently available tests with moderate scale-up</td>
<td></td>
</tr>
<tr>
<td>Immediate VIA</td>
<td>10</td>
</tr>
<tr>
<td>Immediate 2-visit HPV</td>
<td>10</td>
</tr>
<tr>
<td>Number of women screened</td>
<td>5,185,700</td>
</tr>
<tr>
<td>Immediate implementation of currently available tests followed by a switch to point-of-care HPV testing</td>
<td></td>
</tr>
<tr>
<td>Immediate VIA, switch to 1-visit HPV</td>
<td>10</td>
</tr>
<tr>
<td>Immediate 2-visit HPV, switch to 1-visit HPV</td>
<td>10</td>
</tr>
<tr>
<td>Number of women screened</td>
<td>5,185,700</td>
</tr>
<tr>
<td>Delayed implementation of currently available tests or point-of-care HPV testing</td>
<td></td>
</tr>
<tr>
<td>Delayed VIA</td>
<td>--</td>
</tr>
<tr>
<td>Delayed 2-visit HPV</td>
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</tr>
<tr>
<td>Delayed 1-visit HPV</td>
<td>--</td>
</tr>
<tr>
<td>Number of women screened</td>
<td>--</td>
</tr>
</tbody>
</table>

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1-v: 1-visit strategy; 2-v: 2-visit strategy; HPV: HPV DNA testing; VIA: visual inspection with acetic acid. VIA testing is indicated by blue squares (light blue: 10% coverage; dark blue: 20% coverage). 2-visit HPV testing is indicated by the lavender shading (light lavender: 10% coverage; dark lavender: 20% coverage). This article is protected by copyright. All rights reserved.
lavender: 20% coverage). 1-visit (point-of-care) HPV testing is indicated by the yellow shading (light yellow: 10% coverage; dark yellow: 20% coverage). Each year, the relevant coverage level was applied to each birth cohort of women between age 30 and 34 years. Thus, depending on the year, cohort, and coverage level, effective coverage for a particular birth cohort could be zero (e.g., for 34 year old women in 2017 under delayed implementation scenarios), or nearly 100% (e.g., for 30 year old women in 2022 after immediate implementation with a switch to 1-visit HPV).
Table 3. Cervical cancer cases and deaths averted over the lifetimes of women aged 30 to 34 years during the intervention period 2017-2026 in India, by cervical cancer screening implementation strategy.

<table>
<thead>
<tr>
<th>Strategy</th>
<th>Number of women screened</th>
<th>Cumulative cases averted</th>
<th>Cumulative deaths averted</th>
<th>Incremental number of cases averted relative to most effective strategy</th>
<th>Incremental number of deaths averted relative to most effective strategy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Immediate implementation (2017) followed by a switch to point-of-care HPV testing (2022)</td>
<td>81,377,000</td>
<td>574,100</td>
<td>382,500</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>Immediate two-visit HPV, switch to one-visit HPV</td>
<td>81,377,000</td>
<td>521,900</td>
<td>346,600</td>
<td>-52,200</td>
<td>-35,900</td>
</tr>
<tr>
<td>Immediate VIA, switch to one-visit HPV</td>
<td>81,377,000</td>
<td>521,900</td>
<td>346,600</td>
<td>-52,200</td>
<td>-35,900</td>
</tr>
<tr>
<td>Immediate implementation of currently available tests (2017) with moderate scale-up (2022)</td>
<td>81,377,000</td>
<td>480,600</td>
<td>328,000</td>
<td>-93,500</td>
<td>-54,500</td>
</tr>
<tr>
<td>Immediate two-visit HPV</td>
<td>81,377,000</td>
<td>330,800</td>
<td>217,900</td>
<td>-243,300</td>
<td>-164,600</td>
</tr>
<tr>
<td>Immediate VIA</td>
<td>81,377,000</td>
<td>111,200</td>
<td>73,800</td>
<td>-462,900</td>
<td>-308,700</td>
</tr>
<tr>
<td>Delayed implementation of currently available tests or point-of-care HPV testing (2022)</td>
<td>27,435,000</td>
<td>209,300</td>
<td>139,100</td>
<td>-364,800</td>
<td>-243,400</td>
</tr>
<tr>
<td>Delayed two-visit HPV</td>
<td>27,435,000</td>
<td>165,900</td>
<td>111,600</td>
<td>-408,200</td>
<td>-270,900</td>
</tr>
<tr>
<td>Delayed VIA</td>
<td>27,435,000</td>
<td>111,200</td>
<td>73,800</td>
<td>-462,900</td>
<td>-308,700</td>
</tr>
</tbody>
</table>

*Strategies are listed in order of effectiveness. The target population included cohorts of women aged 30 to 34 years during implementation between 2017 and 2026. Cumulative cases and deaths extend over the lifetime of these cohorts, through 2095. Strategies considered for immediate implementation followed by a switch to point-of-care HPV testing included 1) Immediate two-visit HPV, switch to one-visit HPV: two-visit HPV DNA testing with 10% coverage (implemented 2017-2021) scaling up to 20% coverage with one-visit HPV testing (implemented 2022-2026); and 2) Immediate VIA, switch to one-visit HPV: one-visit VIA with 10% coverage (implemented 2017-2021) scaling up to 20% coverage with one-visit HPV testing (implemented 2022-2026). Strategies considered for immediate implementation included 1) Immediate two-visit HPV: two-visit HPV DNA testing with 10% coverage (implemented 2017-2021) scaling up to 20% coverage (implemented 2022-2026); and 2)
Immediate VIA: one-visit VIA with 10% coverage (implemented 2017-2021) scaling up to 20% coverage (implemented 2022-2026). Strategies considered for delayed implementation included: 1) Delayed one-visit HPV: one-visit HPV DNA testing with 10% coverage (implemented 2022-2026); 2) Delayed two-visit HPV: two-visit HPV testing with 10% coverage (implemented 2022-2026); and 3) Delayed VIA: one-visit VIA with 10% coverage (implemented 2022-2026).
Novelty & Impact Statement:

Nearly one-quarter of cervical cancer cases worldwide occur in India. Nonetheless, while the disease can be prevented in part through screening for precancerous lesions, very few Indian women receive Pap tests. Here, the authors estimated cervical cancer burden in India assuming different screening program implementation scenarios, including immediate implementation with both two-visit and one-visit human papillomavirus (HPV) testing and delayed implementation with a one-visit HPV test. Models showed that immediate implementation of HPV testing averted more than double the number of cases and deaths from cervical cancer among 30- to 34-year-old women compared with delayed implementation with one-visit HPV testing.