Cost Effectiveness of Human Papillomavirus-16/18 Genotyping in Cervical Cancer Screening

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Abstract

Background There is limited understanding of the health economic implications of cervical screening with human papillomavirus (HPV)-16/18 genotyping.

Objective The aim of this study was to evaluate the cost effectiveness of cervical cancer primary screening with a HPV-16/18 genotyping test which simultaneously detects 12 other high-risk HPV types.

Methods A Markov cohort model compared four strategies: (1) cytology with reflex HPV testing for atypical squamous cells of undetermined significance (ASC-US); (2) co-testing with cytology and HPV testing; (3) HPV with reflex to cytology; and (4) HPV with 16/18 genotyping and reflex cytology (ASC-US threshold). Screening began at age 30 and was performed triennially over 40 years. Screening sensitivity and specificity values for cervical intraepithelial neoplasia (CIN) 3 were obtained from the Addressing THE Need for Advanced HPV Diagnostics (ATHENA) trial. Outcomes for a 1-year follow-up scenario wherein persistent disease was detected were estimated. Screening and cancer treatment costs were calculated from a US payer’s perspective in 2013. Costs and quality-adjusted life-years (QALYs) were discounted at 3% annually.

Results Applying a US$50,000/QALY threshold, strategy (4) dominated strategies (2) and (3) by reducing costs and cancer incidence and improving QALYs, and was cost effective versus strategy (1). Accounting for persistent ≥CIN 3 at 1 year, strategy (4) was cost effective versus all other strategies. Detecting HPV-16/18 resulted in earlier diagnosis of clinically relevant ≥CIN 3 at initial screening and efficient use of follow-up resources. Outcomes were most influenced by strategy performance.

Conclusions Incorporating HPV-16/18 genotyping is cost effective and may improve detection of CIN, thereby preventing cervical cancer.

Key Points for Decision Makers

Access to cervical cancer screening strategies that facilitate early detection of clinically relevant cervical intraepithelial neoplasia associated with HPV-16/18 allows women to seek treatment sooner and may thereby improve cancer protection.

Stratifying cancer risk via HPV-16/18 genotyping in women aged ≥30 years may be cost saving compared with cotesting with cytology and HPV testing by reducing the number of screening tests and overall screening costs.

HPV-16/18 genotyping with reflex cytology is cost effective in cervical screening and represents a beneficial alternative approach to cervical screening from both a health and economic perspective.
1 Introduction

Organized screening programs aimed at early detection of high-grade cervical intraepithelial neoplasia (CIN) have steadily reduced cervical cancer mortality (Alliance for Cervical Cancer Prevention, Preventing Cervical Cancer Worldwide, 2004). Nonetheless, individual screening tests can suffer from shortcomings, particularly when applied indiscriminately to the general screening population. Subjective interpretation of Pap cytology (“cytology”) renders a high degree of variability [1]. In populations with low prevalence of human papillomavirus (HPV), cytology demonstrates low sensitivity and poorer positive predictive value due to fewer high-grade lesions [1]. Also, primary HPV pooled testing exhibits lower specificity in excluding the absence of high-grade CIN [2]. Accordingly, ongoing efforts aim to identify strategies incorporating more prudent use of individual screening tests.

Since HPV-16 and HPV-18 account for more than 70% of cervical cancer cases [3, 4] and confer a higher risk for neoplasia [5], testing for these genotypes may be clinically useful for stratifying cancer risk [6]. A qualitative in vitro assay (cobas® HPV Test), which simultaneously detects 14 high-risk HPV types and provides specific genotyping information for HPV-16 and HPV-18, was validated in the Addressing THE Need for Advanced HPV Diagnostics (ATHENA) trial [6, 7]. The trial concluded that primary HPV testing and triage of HPV-positive women with HPV-16/18 genotyping and cytology may balance screening benefits and harms by maximizing sensitivity while limiting the number of diagnostic tests [7]. The US Food and Drug Administration recently approved the cobas® HPV Test for primary cervical cancer screening in women aged ≥25 years. To explore the economic implications, we compared the relative cost effectiveness of HPV-16/18 genotyping and three other primary cervical screening strategies.

2 Materials and Methods

2.1 Study Population

The model evaluated the outcomes of a hypothetical cohort of 1,000 non-hysterectomized women who were asymptomatic for cervical cancer and had participated in cervical screening in a US healthcare setting over a 40-year period. The starting age of the cohort was 30 years. Because the risk of HPV persistence increases with age [8], HPV testing in women aged ≥30 years may identify more clinically significant infections and, therefore, the performance of HPV-16/18 genotyping in this population is of particular interest.

2.2 Comparator Screening Strategies

We assessed the cost effectiveness of four primary cervical screening strategies: (1) cytology with reflex HPV testing for triage of atypical squamous cells of undetermined significance (ASC-US) (“Cytology”); (2) co-testing with cytology and HPV testing (“Co-testing”); (3) HPV with reflex cytology for positive HPV (“HPV HR [human papillomavirus, high-risk] Only”); and (4) HPV with 16/18 genotyping and reflex cytology (ASC-US threshold) (“HPV with Genotyping”). (See Fig. 1 for detailed algorithms). Cytology and Co-testing were selected based on the American Cancer Society (ACS)–American Society for Colposcopy and Cervical Pathology (ASCCP)–American Society of Clinical Pathologists (ASCP) guidelines. These strategies represented practices most likely to be implemented in community settings [9, 10].

Routine screening was performed triennially with some women returning for 1-year retesting based on primary screening results. We assumed that all women complied with scheduled screening visits as well as recommendations for colposcopy and biopsy. For all strategies, women with false positive results returned for follow-up screening as recommended by the ASCCP [9]. Women exited screening at age 70 years in the model based on clinical guidelines advising cessation of screening after the age of 65 years, but extending beyond age 65 for women with a history of CIN 2, CIN 3, or cervical cancer [9].

2.3 Markov Model Structure and Health States

The Excel-based Markov cohort model (Microsoft Corporation, Redmond, Washington) incorporated a ‘no-screening’ model which simulated the natural history of oncogenic HPV infection in an unscreened population. This natural history model served as the framework within which the effects of each screening strategy were applied and the outcomes compared. Since the analysis reflected clinical practice in the US, ‘no screening’ was not considered a comparator option.

Applying literature-based transition probabilities [11–19], women transitioned annually across seven possible states: (1) No high-risk HPV infection; (2) High-risk HPV infection (no CIN); (3) CIN 1; (4) CIN 2; (5) CIN 3 or worse; (6) Cervical Cancer, and (7) Dead (Fig. 2). At baseline, the cohort was apportioned among the Markov states based on the age-adjusted prevalence of high-risk HPV infection, the probability of CIN, and the age-adjusted cervical cancer incidence (Table 1). Age-specific probabilities for high-risk HPV infection and CIN clearance, progression, and regression were obtained from published epidemiologic, cost-effectiveness, and mathematical models simulating HPV infection and cervical
carcinogenesis (Table 1) [17–23]. Annual transition probabilities for progression to CIN 2 and ≥CIN 3 from no CIN were computed from the 36-month cumulative hazard rate for progression from incident HPV-16 infection to CIN 2 and ≥CIN 3 (16.5 %) reported for the placebo arm (n = 42) in a randomized controlled trial of a HPV-16 vaccine [21]. Progression to invasive cervical cancer occurred within a single cycle from ≥CIN 3 only, whereas
regression from Cervical Cancer to the CIN states did not occur. Given that we primarily intended to address the screening effect of HPV with Genotyping, we did not stratify Cervical Cancer by stage; rather, we modeled health states for pre-cancer lesions arising from persistent HPV infection. Other models utilizing a single health state to depict invasive cervical cancer have also been published in the literature [24, 25].

The no-screening model was manually calibrated to approximate published estimates of cervical cancer incidence and mortality in an unscreened population. In calibrating a natural history model for HPV infection and cervical cancer, Taylor et al. [26] noted that manual calibration was one of two methods achieving the best fit, yielding a 10 % mean deviation from optimal fit. Our final model generated an annual cervical cancer incidence of approximately 19.47 cases per 100,000 women-years. This projection was 10.1 % higher than the age-standardized incidence rate of 17.70 per 100,0000 women-years for an unscreened US population cited by Gustafsson et al. [27], but 1.1 % lower than the incidence (20.64 per 100,000) predicted by a cost-effectiveness screening model by Vijayaraghavan et al. [11].

Age-adjusted annual probabilities of death for women without cervical cancer were derived from the general population estimates reported in the US Social Security Actuarial Publications Period Life Table. To estimate cervical cancer mortality in the unscreened population, the age-specific 5-year relative cervical cancer survival rates reported by the National Cancer Institute Surveillance Epidemiology and End Results (SEER) Cancer Statistics Review (1975–2007) were converted to annual probabilities and multiplied by the respective annual numbers of expected cervical cancer cases [28]. Our model estimated an annual cervical cancer mortality rate of 15.5 per 100,000 unscreened women. Published cervical cancer mortality rates in unscreened populations vary by geographic locality, population age, and study year, but have exceeded 20 deaths per 100,000 women in some studies [29]. For the screening cohorts, women who were diagnosed with cervical cancer were assigned a 5-year relative survival rate of 90 % per the ACS based on the assumption that cancers in asymptomatic women receiving regular screening with 100 % compliance would tend to be early-stage, curable cancers (ACS, Cancer Facts and Figures, 2010) [30].

In addition to face validation of the no-screening model, analytical methods (including programming, coding, and equations), configuration of screening algorithms, screening results, and costs were reviewed by individuals with expertise in the area of cervical cancer screening (including two clinicians) and were found to be reasonable within the structural limitations of a Markov cohort model.

2.4 Model Inputs

2.4.1 Screening Performance and Key Assumptions

Baseline performance values were obtained from the ATHENA trial, the largest US-based screening accuracy
cohort study for HPV-16/18 genotyping which included more than 47,000 women ages ≥21 years from clinical sites across 23 states [7, 12]. A subanalysis of 34,254 participants aged ≥30 years (mean 44.7 years) compared ten different cervical screening strategies for detecting high-grade cervical disease [7]. In addition to HPV-16/18 genotyping, the analyses evaluated strategies without HPV-16/18 genotyping that were considered of interest in clinical practice (several of which were included in the present model). Given the large cohort, expected frequency of CIN [12], and pertinent strategies evaluated in a single population, the ATHENA trial was considered an appropriate source for screening performance data.

Screening performance was defined as the sensitivity and specificity for ≥CIN 3 and referred to overall strategy performance as opposed to that of component tests. Since approximately 90 % of CIN 1 and 88 % of CIN 2 cases regress within 1 year [31, 32], the model accounted only for sensitivity in detecting incident and persistent cases of ≥CIN 3. At publication, the ATHENA analysis reported only the baseline performance of the strategies as representing usual clinical practice. Thus, comparing the ‘true’ sensitivities (defined as performance achieved by completing a full screening cycle) was not feasible since outcomes could be only partially ascertained from the initial screening. Therefore, we examined a scenario wherein all cases deferred at baseline returned for 1-year follow-up. Screening sensitivities were re-calculated by assuming that deferred ≥CIN 3 cases would be captured at the 1-year visit. Detecting additional ≥CIN 3 effectively increased the overall sensitivities of the strategies, particularly those deferring more patients at baseline; this resulted in similar performance between the HPV with Genotyping and HPV HR Only strategies and improved sensitivity for co-testing (Table 2).

### 2.4.2 Costs

Costs were modeled from the perspective of the US healthcare payer and reported in 2013 US dollars (Table 2). Unit costs represented the median national Medicare payments for the respective Common Procedural Terminology codes (Centers for Medicare and Medicaid Services Physician Fee Schedule, 2013). For each strategy, the total annual cost included the costs for screening (including routine examinations, triage, and retesting) and treatment for CIN and cervical cancer. Screening costs included the costs for individual tests and office visits. Screening and

### Table 1 Model assumptions for the prevalence of high-risk HPV, HPV-16/18, and health state transition probabilities by age group

| Age group (years) | Reference(s) |
|-------------------|--------------|
| 25–29             |             |
| 30–34             |             |
| 35–39             |             |
| 40–44             |             |
| 45–49             |             |
| 50–54             |             |
| 55–59             |             |
| 60–64             |             |
| 65–70             |             |
| Prevalence        |             |
| HPV HR, % infected|             |
| HPV-16/18, % of HPV HR | ATHENA [12] |
| HPV HR, % per cytology | ATHENA [12] |
| Normal            |             |
| ASC-US            |             |
| LSIL/HSIL         |             |
| Annual transition probabilities used in the model |
| HPV HR (–) to HPV HR (+) | [17, 20, 23] |
| HPV HR (+) to HPV HR (–) | [17, 20–22] |
| HPV HR (+) no CIN to CIN | [18, 20] |
| HPV HR (+) no CIN to CIN 2 or CIN 3 | [18, 21] |
| CIN 1 to CIN 2 or CIN 3 | [19, 21] |
| CIN 2 or CIN 3 to Cervical Cancer | [18, 20, 21] |
| CIN1 to HPV HR (+) or HPV HR (–) | [17] |
| CIN 2 or CIN 3 to CIN1 | [17, 20] |
| CIN 2 or CIN 3 to HPV HR (–) | [17] |

ASC-US atypical squamous cells of undetermined significance, ATHENA Addressing THE Need for Advanced HPV Diagnostics, CIN cervical intraepithelial neoplasia, HPV human papillomavirus, HR high risk, HSIL high-grade squamous intraepithelial lesion, LSIL low-grade squamous intraepithelial lesion.
treatment costs were calculated by multiplying the relevant unit costs by the respective expected percentages of women utilizing each test or treatment in the strategy, and then summing the utilization-weighted costs.

2.4.3 Health Utilities

Quality-adjusted life-years (QALYs) were calculated by multiplying the expected number of life-years gained by the assigned health utility for each Markov state (Table 2). Health utility values for the CIN 1 and the CIN 2 and CIN 3 states represented the utilities for low-grade squamous intraepithelial lesions and high-grade squamous intraepithelial lesions, respectively, as cited by Sanders et al. [18] and applied to the US population in a study by the Institute of Medicine (IOM) [33]. In the latter study, Health Utilities Mark II utilities, which were derived for a general Canadian population via the standard gamble method, were assigned quality-adjusted weights to valuate various morbidity states in the US [33]. To avoid overestimating the utility for Cervical Cancer, we assigned a base value of 0.71 which represented the mean weighted health utility for cervical cancer in the treatment phase (0.79 for Stage I and 0.62 for Stages II-IV per Sanders et al.) adjusted by the SEER stage distribution for cervical cancer at diagnosis (see Table 2) [19, 28]. We assumed a 1-year disutility for detected but asymptomatic cervical cancer cases [19].

2.5 Model Outcomes

Outcomes included direct medical costs and QALYs calculated over 40 years and discounted at a yearly rate of 3.0 % [34, 35]. Cost effectiveness was expressed as the incremental cost-effectiveness ratio (ICER), which represented the additional cost per QALY gained using one strategy versus the next non-dominated strategy, and assessed against the payer’s willingness-to-pay (WTP) criterion of US$50,000/QALY. Other outcomes included the undiscounted number of life-years gained, annual cervical cancer incidence, and mortality rate (per 100,000 women) for each strategy. Additionally, we determined the expected number of colposcopies required to detect one case of ≥CIN 3, an endpoint considered by the ACS-ASC-CCP-ASCP guidelines as a primary surrogate for the potential harm of cervical screening [9].

2.6 Sensitivity Analyses

We employed one-way sensitivity analyses to assess the impact of key inputs on the relative cost effectiveness of HPV with Genotyping versus the comparators. The resulting tornado diagrams depicted the degree of influence of each parameter on the relative cost effectiveness of HPV with Genotyping when tested at high (+10 %) and low (−10 %) values; wider bars indicated greater influence. Additionally, we performed threshold analyses using an Excel what-if analysis function to examine the effects of further increasing or decreasing parameters exhibiting greater influence on visual inspection or those of particular interest (such as sensitivity and costs).

To quantify the likelihood that HPV with Genotyping is cost effective compared with the non-dominated strategies, probabilistic sensitivity analysis (PSA) was performed by simultaneously varying key inputs across values sampled 1,000 times from assigned distributions (Table 2). We applied a standard error (SE) of 1.0 % for sensitivities and specificities (based on the ATHENA data [n = 34,254]), and in the absence of other data, assumed a SE of 10 % the deterministic value for costs, health utilities, and the annual discount rate for costs and effects. The resulting cost-effectiveness acceptability curve showed the percentage likelihood that the strategy was cost effective when assessed against WTP threshold criteria. We used the net monetary benefit (NMB), a measure deemed appropriate by the International Society for Pharmacoeconomics and Outcomes Research, for generating the results of the PSA [36]. When comparing two strategies, the strategy generating a greater NMB (positive incremental NMB [INMB]) was considered more cost effective.

3 Results

In the primary analysis, HPV with Genotyping was more effective and less costly compared with both Co-testing and HPV HR Only, thereby dominating the two strategies (Table 3). Compared with Cytology, HPV with Genotyping was cost effective, generating an acceptable ICER of US$7,667/QALY. Further, HPV with Genotyping appeared beneficial in reducing annual cervical cancer incidence and mortality rates compared with the other strategies, while requiring a relatively moderate number of colposcopies for each ≥CIN 3 detected (Table 3).

As shown in Table 4, women were more frequently triaged with Cytology when HPV HR Only was employed for primary screening compared with HPV with Genotyping. Similarly, fewer women screened with HPV with Genotyping were deferred to 1-year retesting compared with those triaged in both HPV HR Only and Co-testing. Examining the expenditures (data not shown), the average total costs which included routine and repeat screening, triage tests (for HPV HR Only), colposcopy, and biopsy, were higher for Co-testing and HPV HR Only (US$1,737 and US$1,481, respectively) compared with Cytology and HPV with Genotyping (US$1,001 and US$1,064, respectively).
Similar trends were observed in the second scenario. Despite equivalent sensitivities, HPV with Genotyping resulted in fewer women being triaged and deferred to follow-up compared with HPV HR Only, yielding lower average screening costs (US$1,063 versus US$1,463, respectively) and an average cost-savings of US$400 per
woman screened. Co-testing generated a marginal QALY gain but was more costly than HPV with Genotyping and, by comparison, not cost effective.

In the one-way sensitivity analyses, changes in the health utilities and screening sensitivities had the largest impact on the relative cost effectiveness of HPV with Genotyping compared with the comparators (Fig. 3). HPV with Genotyping appeared more cost effective when lower utilities were assigned to the CIN and Cervical Cancer states, but remained cost effective even when the utilities were increased to 1.0. As presented in Table 5, threshold analyses showed that under the current assumptions and a WTP criterion of US$50,000, HPV with Genotyping was less cost effective than the comparators only when (i) the current cost for HPV-16/18 genotyping was more than tripled, or (ii) the sensitivity of HPV with Genotyping was reduced by ≥50% of the current estimate, or (iii) the sensitivity of Cytology approached 100%. Compared with Cytology, the only other non-dominated strategy, the PSA showed that, at a WTP threshold of US$50,000/QALY as well as with more stringent payer criteria, HPV with Genotyping was cost effective (Fig. 4).

4 Discussion

Stratifying cervical cancer risk via HPV-16/18 genotyping may constitute a cost-effective approach for primary screening in women aged ≥30 years. Under the current assumptions, this strategy conferred incremental QALY gains and reduced cervical cancer incidence and mortality, but on average, was less expensive compared with HPV HR Only and Co-testing. HR HPV Only yielded additional re-testing and diagnostic visits and incurred higher costs in the long run compared with HPV with Genotyping. Co-testing was most costly as both routine and follow-up screening required dual testing with HPV and cytology tests. Although more costly than Cytology, HPV with Genotyping improved clinical outcomes, thereby maintaining cost effectiveness.

Because baseline sensitivities were obtained from initial screening in the ATHENA trial, the values were adjusted to account for detecting new and persistent ≥CIN 3 at 1 year. It was expected that strategies deferring intermediate-risk women to 1-year follow-up would capture additional persistent ≥CIN 3. This assumption was supported by the HPV FOCAL trial which showed that ≥CIN 3 detection rates at 1 year were increased in women who had been initially high-risk HPV-positive and cytology-negative [37]. Despite identical sensitivities for ≥CIN 3 in this scenario, HPV with Genotyping was cost-saving over HPV HR Only again by reducing the frequency of triage and follow-up tests. Although both strategies deferred women who were initially negative for intraepithelial lesion and malignancy to 1-year follow-up, HPV HR Only retested all high-risk HPV-positive women whereas HPV with Genotyping retested a smaller subgroup that was positive for non-16/18 high-risk HPV only. Likewise, triage was restricted to women positive for non-16/18 high-risk HPV only in the latter, but included all women positive for high-risk HPV in the former. Stratifying high-risk patients with 16/18 genotyping at baseline initially incurred relatively higher routine screening and diagnostic costs, but resulted in fewer highest-risk patients in the long term, thereby reducing costs. Consistent with a prior lifetime model [11], triennial Co-testing increased QALYs in the second
scenario, but was more costly than HPV with Genotyping, mainly due to the greater number of initial tests.

Our analysis is noteworthy as it incorporated data from over 34,000 women aged ≥ 30 years participating in the largest US diagnostic accuracy cohort study for cervical screening to date. Contrary to other models, we employed screening performance inputs from a single trial. Screening performance can vary between clinical trials, impacted by factors such as study design, disease prevalence, and statistical methods. Applying data from the ATHENA trial facilitated comparison of the relative performance of the strategies within the same cohort, thereby reducing variability.

We note several limitations. First, excluding the impact of non-compliance at follow-up may have over-estimated screening sensitivities and costs, particularly for Co-testing and HPV HR Only, which deferred more patients to re-testing. Second, we did not examine co-testing every 5 years because while this is a preferred strategy for women aged 30–65 years, the American Congress of Obstetricians and Gynecologists and other clinical experts have contended that triennial co-testing is the more likely scenario in the community as patients are unlikely to feel comfortable with extended screening intervals [10]. Thus, we believe co-testing every 3 years is relevant for a US model. Nevertheless, we recognize that screening practices may vary considerably, the effects of which may be investigated in future research.

Under the current assumptions, we projected cervical cancer incidence rates which exceeded the 2007–2011 US age-adjusted incidence of 7.8 per 100,000 reported by the SEER. However, our model included non-hysterectomized women aged ≥30 years only. Rositch et al. [38] reported hysterectomy-corrected age-standardized cervical cancer incidence rates in the US. For the age groups spanning from 30 to 74 years, incidence rates (per 100,000) ranged from 11.5 to 27.4 following correction for hysterectomy prevalence [38], which was higher than our model results. These findings indicate that cervical cancer incidence may be higher than the national estimate when correcting for hysterectomy, implying the need for appropriate adjustments when evaluating cervical screening outcomes.

Annual cervical cancer mortality rates in the model exceeded the 2006–2010 SEER age-adjusted rate of 2.4 per 100,000 women. While the effects of hysterectomy prevalence on cervical cancer mortality rates have not been extensively investigated, an early study by the Centers for Disease Control and Prevention found that correcting for hysterectomy prevalence resulted in a 39 % relative increase in cervical cancer mortality rates in the US during the period 1965–1988 [39]. We assumed that cancers were detected at an earlier stage in the screened population, which was associated with 90 % 5-year relative survival.

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Table 4  Expected annual number of cervical screening examinations and colposcopies required for detecting ≥CIN 3 in the base case and 1-year follow-up scenario

| Strategy                                      | Number of screening examinationsa,b | No. of colposcopies per ≥CIN 3 detected |
|-----------------------------------------------|-------------------------------------|----------------------------------------|
| Routine                                      | Triage                              | Retest                                 | Colposcopies                 |
| Base case                                    |                                     |                                        |                           |
| Cytology (reflex HPV test for ASC-US)         | 30,364                              | 1,282                                  | 0                          | 2,104                        | 4.76                       |
| HPV with genotyping (reflex cytology [ASC-US])| 23,065                              | 6,372                                  | 1,024                      | 2,159                        | 3.06                       |
| HPV HR only (with reflex to cytology [ASC-US])| 22,872                              | 8,405                                  | 6,557                      | 2,339                        | 3.95                       |
| Co-testing (with cytology and HPV testing)    | 22,848                              | 0                                      | 6,686                      | 2,967                        | 4.79                       |
| 1-year follow-up scenario                     |                                     |                                        |                           |
| Cytology (reflex HPV test for ASC-US)         | 30,294                              | 1,282                                  | 0                          | 2,104                        | 4.76                       |
| HPV with genotyping (reflex cytology [ASC-US])| 23,201                              | 6,265                                  | 1,021                      | 2,130                        | 2.73                       |
| HPV HR only (with reflex to cytology [ASC-US])| 23,201                              | 8,079                                  | 6,332                      | 2,249                        | 2.89                       |
| Co-testing (with cytology and HPV testing)    | 23,161                              | 0                                      | 6,479                      | 2,915                        | 3.67                       |

ASC-US atypical squamous cells of undetermined significance, CIN cervical intraepithelial neoplasia, HPV human papillomavirus, HPV HR human papillomavirus, high-risk

a Per 100,000 women and annualized over a 40-year time horizon
b Triage tests included HPV test for Cytology and cytology for both HPV with genotyping and HPV HR only

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a Per 100,000 women and annualized over a 40-year time horizon
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Fig. 3  One-way sensitivity analysis: tornado diagrams of INMB of HPV-16/18 genotyping with reflex cytology (‘HPV with Genotyping’) versus alternative primary cervical screening strategies. The INMB of HPV with Genotyping compared with Cytology (a), HPV HR Only (b), and Co-testing (c) decreased primarily when higher values were assigned to the health utilities for the CIN states, or when screening performance (sensitivity) was improved for the comparator strategies or reduced for HPV with Genotyping. The INMB was calculated based on a WTP threshold of $50,000 per QALY and represented the difference between the net monetary benefit of the two strategies. CIN cervical intraepithelial neoplasia, DNA deoxyribonucleic acid, HPV human papillomavirus, HPV HR human papillomavirus, high-risk, INMB incremental net monetary benefit, QALY quality-adjusted life year, WTP willingness-to-pay.
Lastly, we did not consider HPV immunization effects, which would theoretically decrease the prevalence of high-risk HPV and thereby reduce the sensitivity of the HPV strategies. Analyses of data from eight US managed care organizations participating in the Vaccine Safety Datalink, however, found that over the 5-year period following approval of the quadrivalent HPV vaccine (2006–2011), 56 % of females ages 9–26 years had received at least one vaccine dose, but only 42 % had completed the three-dose series [40]. Given these modest vaccination rates and generally lower prevalence of high-risk HPV in women ≥30 years, including vaccination effects may not have substantially impacted our results.

Our findings suggest that with proper management, earlier detection of clinically relevant CIN attributable to HPV-16/18 may improve protection against cervical cancer. In the POpulation-BAse d SCreening study AMsterdam (POBASCAM), significantly more CIN 2+ cases were
detected at baseline in women randomized to HPV DNA and cytology co-testing compared with those tested with cytology only \((p = 0.015)\) [41]. Conversely, at 5-year follow-up, significantly fewer HPV-16–positive ≥CIN 3 lesions were detected with co-testing compared with cytology-based ASC-US triage \((p = 0.012)\). Detection of non-HPV-16–positive ≥CIN 3 did not differ between groups. These results support the utility of HPV testing in primary screening for women aged ≥30 years.

Despite technological advancements, the goals of cervical cancer screening have remained unchanged and aim to identify as many women at risk for cancer as possible, while protecting others against potential harms of unnecessary interventions. Strategies employing a highly sensitive HPV test in primary screening must be balanced with a more specific test (such as cytology) to ensure that the right women receive appropriate interventions. Incorporating such a strategy, however, must account for the population screened as well as economic considerations. A Dutch simulation cost-effectiveness model evaluating nine primary screening strategies concluded that HPV screening was preferred for women aged >30 years in many scenarios simulating screening situations in Europe [42]. In contrast, primary cytology was preferred only in scenarios bearing high HPV prevalence and high HPV testing costs [42]. For developing countries where screening effectiveness was suboptimal, the World Health Organization recommended screening in focused geographies and in high-risk women (aged 35–49 years or 30–50 years) once or twice in their lifetime, using a highly sensitive test with high coverage of the targeted population [43].

HPV with Genotyping may detect clinically relevant high-grade CIN earlier and facilitate efficient use of healthcare resources in the long run. Detecting disease at the initial visit may also optimize protection against cervical cancer. Further analyses of women screened in real-world settings may assist in confirming the cost effectiveness of HPV-16/18 genotyping in primary cervical screening.

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Author contributions W Huh is the guarantor for the manuscript content. W. Huh, E. Williams, and J. Huang contributed to the development and design of the economic model, selection of data sources, and analyses. All authors participated in the writing, critical review, editing, and approval of the manuscript for submission.

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