RESEARCH ARTICLE

A Model Approach for Assessing the Benefits of HPV Testing against Cytology in Screening for Cervical Cancer Precursors in Thailand

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Abstract

Objective: The aim of this study was to compare the efficacy of HPV 16/18 genotyping test, high risk HPV DNA testing, alone and in conjunction with the liquid-based cytology method in screening for cervical cancer precursors.

Methods: A Markov model was used to describe the course of the cases of CIN2+ that had been detected over a 35 year period. Screening programs started at age 30 and were performed at an interval of once every five years. The model compared three strategies of HPV 16/18 genotyping with reflex cytology triage, high-risk HPV testing alone with referral to colposcopy and cytology-based screening with referral to colposcopy. We assumed the rate of patients lost to follow-up for those referred to colposcopy would be 0%. The clinical parameters were estimated using the data from a study conducted by the Thailand National Cancer Institute. Result: Of the three screening strategies evaluated, the high risk HPV DNA testing alone was the most effective for detecting CIN2+ over the 35 year study period. It detected 143 and 510 cases per 100,000 women more than the HPV 16/18 genotyping test and cytology-based strategy, respectively. The HPV genotyping test detected 368 cases per 100,000 women more than the cytology-based approach. In addition, when viewed with five year intervals, there were missed cases totaling approximately half of the detected cases screened by the cytology strategy and 10% of cases detected with screening by the HPV genotyping test. Conclusion: This study strongly indicates that HPV/DNA testing is preferable to cytology-based screening for cervical cancer precursors. However, the balance between the benefits, burdens and cost of each screening program should be considered.

Keywords: Cervical cancer- Screening- Human papillomavirus testing- Liquid based cytology- Mathematical Models

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Introduction

Cervical cancer is the second most common cancer, globally. In 2012, more than 8,000 women per 100,000 here in Thailand were diagnosed with cervical cancer and more than half of these cases proved fatal (Bruni et al., 2015). Because of it develops over time, it is one of the preventable types of cancer. The optimal screening test and management for precancerous lesions can reduce morbidity and mortality. Papnicolaou cytology (PAP smear) has been proven to be an effective screening method. It results in the reduction of both the incidence and subsequent mortality. In 2005, the National Health Security Office and Ministry of Public Health (MoPH) of Thailand initiated a comprehensive cervical cancer screening program. Thai women at the ages of 30-60 years are encouraged to undergo a cytology based screening program once every 5 years. In 2006, the program was expanded to include a visual inspection with acetic acid (VIA) screening in certain provinces. National targets for the percentage of women aged 30-60 years who had been screened for cervical cancer once in the previous 5 years were 80% in 2013. The results of the 2010 MoPH survey have shown that 67.4% of women had been screened within the past 5 years (Joseph et al., 2015). Although it appears close to reaching stated targets, there was low screening in some subpopulations. Furthermore, almost half of the women with abnormal tests were lost to follow up. The primary reason that was given for non-attendance was lack of communication. The patient did not receive written information in the form of a letter and those who received their letters did not understand the information provided (Sriamporn et al., 2006). These are limitations of the cytology based screening program. The program requires multiple follow up visits due to its low sensitivity and the high variability of results between laboratories. This program also requires an ample amount of trained cytopathologists and efficient transport and tracking.

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The Natural history model served as the framework within which the effects of each screening strategy were applied and the outcomes compared. Women transitioned annually across 5 possible states; 1) No high risk HPV infection, 2) High Risk HPV infection, 3) Cytology abnormality, 4) CIN2 or 5) condition worsening and fatal. Regression from the CIN 2 to a no CIN state had not occurred. We have modeled health states for pre-cancerous lesions arising from HPV infections. We have not reported on the progression from CIN2 to cancer.

Age-adjusted annual probabilities of death for women without cervical cancer were derived from the general population estimates as reported in the Estimated Generation Life Tables for Thailand of Five-Year Birth Cohorts, 1900-2000 (Prasatkun and Rakchanyaban, 2002).

Model inputs
The probabilities for the prevalence of women who are positive for HPV 16/18, women positive for HR-HPV and women cytology positive are a requirement for the model. All of these were taken from the data of National Cancer Institute of Thailand which had enrolled 5056 Thai women, 30-65 years of age, who were undergoing cervical cancer screening. It is the largest trial done to evaluate HPV testing in Thailand. (National Cancer Institute of Thailand, unpublished data 2016)

The incidence of HPV16/18 positive women was 9.3%. The incidence of HR-HPV positive women was approximately 3.5%. The incidence of cytology positive women was 1.5%. The model outcome is the detection rate of CIN2+ for women over 35 years of age and a reduction allowance at an annual rate of 3% (Table1).

Model outcome
The model’s outcome or dependent variable is the cumulative detection rates of CIN2, CIN3 and cervical cancer cases that had been calculated over a period of 35 years. An annual reduction of 3% is assumed by the Markov model.

Sensitivity analysis
A one-way sensitivity analysis undertaken to assess the impact of the parameter uncertainty on the model’s results followed a standard Monte Carlo approach that had been based on 10,000 randomly generated simulations of parameter values.

Results
Throughout the 35 years of cervical screening, the detection rate of CIN2+ using the HPV 16/18 genotype, HR-HPV testing and Liquid base cytology were 1,389, 1,520 and 1,013 cases per 100,000 women, respectively (Table 2). The Model prediction indicates that high risk HPV-DNA testing alone was the most effective strategy. Whereas, it has been found that the least effective strategy is the cytology based screening method and this is currently the most common practice utilized in Thailand.

The graph in Figure 4 shows the comparisons of the detection rates of all three strategies over the 35 year study period. HR-HPV testing alone detected 143 and 510 more...
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cases per 100,000 women than the HPV genotyping test and the cytology-based strategy, respectively. Comparing the HPV genotyping test and cytology-based testing, the HPV genotyping test detected 368 more cases per 100,000 women than did the cytology-based test. In addition, about half of detected cases were missed in screening by the cytology strategy and 10% of the detected cases screened by the HPV genotyping test in comparison to HPV testing alone when the women were tested at the 5 year intervals.

A probabilistic sensitivity analysis (PSA) was conducted to determine how differences in the prevalence of HPV infection and women with positive cytology impact the detection rates of CIN2+. The results suggest that if the incidence of HPV infection increases at least 3 times, HPV genotyping might be the most effective strategy and have also detected approximately 1,200 cases per 100,000 more than in strategy 2. The liquid based cytology method would be more effective if the sensitivity of the cytology was increased (Table3).

Table 1. Epidemiological Parameters and Ranges Used in the Sensitivity Analysis

| Prevalence | Rate      | Range          | Ref    |
|------------|-----------|----------------|--------|
| STRATEGY 1 |           |                |        |
| HPV16/18   | 0.0093    | 0.0084 - 0.0279| NCI    |
| Other 12HR positive | 0.025    | 0.0025 - 0.075 | NCI    |
| HPV16/18+ → Colpo CIN2+ | 0.191    |                | NCI    |
| Other 12 HR +ve → LG cyto | 0.296    |                | NCI    |
| Other 12 HR +ve → LG cyto → Colpo CIN2 | 0.078    |                | NCI    |
| Other 12 HR +ve → NILM → (wait 1y) HPV+ve | 0.25   |                | †      |
| Other 12 HR +ve → NILM → (wait 1y) HPV+ve →Colpo CIN2+ | 0.2    |                | †      |
| Other 12 HR +ve → HG cyto | 0.148    |                | NCI    |
| Other 12 HR +ve → HG cyto → Colpo +ve →Conization CIN2+ | 0.06    |                | NCI    |
| Other 12 HR +ve → HG cyto → Colpo CIN2+ | 0.15    |                | NCI    |
| STRATEGY 2 |           |                |        |
| HR-HPV +ve | 0.0346    | 0.0311 - 0.1038| NCI    |
| HR-HPV +ve→ Colpo CIN2+ | 0.114    |                | NCI    |
| STRATEGY 3 |           |                |        |
| LG cyto +ve | 0.008     | 0.0072 - 0.024 | NCI    |
| LG cyto +ve → Colpo CIN2+ | 0.11    |                | NCI    |
| HG cyto +ve | 0.0067    | 0.00603 - 0.201 | NCI    |
| HG cyto → Colpo CIN2+ | 0.2      |                | NCI    |
| HG cyto → Colpo +ve → Conization CIN2+ | 0.074    |                | NCI    |

*, Reference form Expert’s opinion; NCI, National Cancer Institute of Thailand
Discussion

In this study, HPV based testing is superior than the cytology based strategy for detecting CIN2+ cases. The cytology method has the highest numbers of missed cases. This is in line with several studies that recommend HPV testing to replace cytology as the primary screening method (Campos et al., 2015; Huh et al., 2015; Jin et al., 2016; Kitchener et al., 2014; Wright et al., 2015). In a comparison of HPV genotyping and HPV testing alone, a study has shown that HPV testing alone is more effective than HPV genotyping. Whereas, most previous studies have revealed HPV genotyping as the most effective strategy (Wright et al., 2016; Beal et al., 2014; Huh et al., 2015). This may be explained by many reasons. First, we used the incidence and genotypic distribution of HPV in Thai women which contrasts most studies that have been

| Strategy                  | Detection rate (per 100,000 women) |
|---------------------------|-------------------------------------|
| HPV16/18                  | 1,389.98                            |
| HR-HPV testing            | 1,520.10                            |
| Cytology LBC              | 1,013.94                            |

Table 2. Outcomes of the Three Cervical Screening Strategies

| Plausible prevalence of | Strategy | Detected cases (per 100,000 women) |
|-------------------------|----------|-------------------------------------|
| HPV 16/18 positive women in strategy 1 | 0.0084 | 2 1,520 |
|                         |         | 1 1,325 |
|                         |         | 3 1,013 |
| Other 12 HR-HPV positive women in strategy 1 | 0.0225 | 2 1,520 |
|                         |         | 1 1,311 |
|                         |         | 3 1,013 |
| HR-HPV positive women in strategy 2 | 0.03114 | 2 1,375 |
|                         |         | 1 1,381 |
|                         |         | 3 1,013 |
| Low grade cytology positive women | 0.0072 | 2 1,524 |
|                         |         | 1 1,381 |
|                         |         | 3 979 |
| High grade cytology positive women | 0.00603 | 2 1,524 |
|                         |         | 3 947.7 |
|                         |         | 1 1,381 |
|                         |         | 3 2,319 |

Table 3. Probabilistic Sensitivity Analysis of the Three Strategies
done in Western countries (Wright et al., 2012; Husain et al., 2015; Kietpeerakool et al., 2015; Kantathavorn et al., 2015). In addition, each step that followed the first screening was different among several studies. Also, the population of the NCI study that has been used as clinical data was small. This may reflect on the impact of the rate of HPV infection and women who were cytology positive.

This is the first study to assess the efficacy of the HPV-based method as a primary screening cervical cancer precursor in Thailand. We used model analysis and conducted sensitivity analysis to represent the actual situations. Furthermore, base case values used in the model were based on the largest amount of and latest data from Thailand. However, there are a few limitations to this study. First, the efficacy we used for comparison is only an intermediate outcome of cervical cancer. Second, clinical input data obtained from a single institute and does not represent the entire population in Thailand. Last, excluding the non compliant cases at follow up, could have possibly over estimated the case numbers.

For a more accurate prediction of the most effective strategy, future studies should be based on the data from multi-centers in several regions of Thailand. Long term clinical outcomes such as lifetime cervical cancer risks or the impact of the women’s quality of life should be assessed. It would be helpful if either VIA and Care-HPV tests can be used as comparative algorithms, especially in low resource settings. In policy making plans, economic analysis studies should be evaluated to identify the best strategy.

In conclusion, this study strongly supports the HPV DNA testing as a preferable option to cytology-based screening for detecting cervical cancer. However, the balance between the benefits, burdens and cost of each screening program should be considered.

Statement conflict of Interest
None.

Funding Statement
None.

Presentation
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