Genotype-specific methylation of HPV in cervical intraepithelial neoplasia

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ABSTRACT

Objective: Hypermethylation of human papillomavirus (HPV) and host genes has been reported in cervical cancer. However, the degree of methylation of different HPV types relative to the severity of the cervical lesions remains controversial. Studies of the degree of methylation associated with the host gene and the HPV genome to the severity of cervical lesions are rare. We examined the association of methylation status between host genes and late gene 1 (L1) regions of HPV16, 18, 52, and 58 in cervical brushings.

Methods: Cervical brushings from 147 HPV-infected patients were obtained. The samples comprised normal (n=28), cervical intraepithelial neoplasia (CIN) 1 (n=45), CIN2 (n=13), and CIN3/carcinoma in situ (n=61). The methylation status of HPV and host genes was measured using bisulfite pyrosequencing and quantitative methylation-specific polymerase chain reaction (PCR).

Results: The degree of methylation of L1 in HPV16, 18, and 52 was associated with the severity of the cervical lesion. In HPV52, C-phosphate-G (CpG) sites 6368, 6405, and 6443 showed significantly higher methylation in lesions ≥CIN3 (p=0.005, 0.003, and 0.026, respectively). Methylation of most HPV types except HPV52 (r<−0.1) was positively correlated with the degree of methylation of host genes including PAX1 and SOX1 (0.4≤r≤0.7). Combining HPV methylation with PAX1 methylation improved the clustering for ≥CIN2.

Conclusion: Our study showed that the degree of L1 methylation of HPV16, 18, and 52 but not 58 is associated with the severity of cervical lesions. The association between HPV methylation and host gene methylation suggests different responses of host cellular epigenetic machinery to different HPV genotypes.

Keywords: DNA Methylation; Viruses; Methylation; PAX1 Transcription Factor; Cervical Intraepithelial Neoplasia; Human Papillomavirus 16; Human Papilloma Virus
INTRODUCTION

In 2012, there were approximately 528,000 new cases of cervical cancer and 266,000 deaths caused by cervical cancer worldwide [1]. Human papillomavirus (HPV) infection is an important risk factor for cervical cancer: 99.7% of cervical cancers are associated with HPV infection. More than 100 HPV types have been identified and can be categorized into high-risk and low-risk based on their carcinogenic ability. The high-risk HPV (hrHPV) types include 16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, 59, 68, 73, and 82 [2]. About 71% of cervical cancer is associated with infection with HPV16 and HPV18. HPV prevalence rates and types differ by ethnic group, and infection with HPV52 and HPV58 is more common in Asia than in other regions [3]. Infection with hrHPV is considered the major essential risk factor for cervical cancer and its precursor lesions [4]. HPV E6 and E7 are the 2 most important oncogenic proteins required for cervical carcinogenesis, and are known to activate oncogenic pathways and repress tumor suppressor pathways. Degradation of p53 by E6 and of retinoblastoma (Rb) by E7 results in dysregulation of the host cell cycle and cell growth [5]. Most HPV infection is transient, but persistent infection with hrHPV lasting more than 2 years can develop into high-grade cervical intraepithelial neoplasia (CIN) [6]. However, infection with hrHPV is necessary but not sufficient to cause cervical cancer [7]: other factors such as epigenetic alterations are also critical in cervical cancer development.

Epigenetic alterations such as DNA methylation and histone modifications can result in heritable gene regulation without changes to the genetic sequence and are recognized as important causes of cancer [8]. DNA methylation is an important mechanism of gene regulation, and the accumulation of alterations in DNA methylation has been shown to be related to disease severity. Alteration in DNA methylation is an early and frequent event during carcinogenesis. Such changes have been found in precancerous and malignant cervical tissue that involved global hypomethylation and the local hypermethylation of tumor suppressor gene (TSG) promoters [9]. The hypermethylation of promoters inactivates their associated genes, resulting in abnormalities in many biological functions. Genes affected include RASSFIA and MGMT, involved in DNA repair, CDKN2A, involved in cell-cycle control, PCDH13, involved in apoptosis, APC and SFRP1, involved in Wnt signaling [10]. This abnormal methylation of genes could serve as a marker for early diagnosis and cancer screening [11].

Over 20 genes, including PAX1, SOX1, ZNF582, PCDHA4, and PCDHA13 have been confirmed to be hypermethylated in cervical brushings diagnosed as CIN2 and more severe lesions. Studies have shown that their sensitivity for CIN2+ or CIN3+ is in the range of 69%–74% for clinical specimens, with a specificity in the range of 66%–82% [3,12,13]. Meta-analyses have shown the utility of PAX1 methylation status as a promising biomarker for cervical cancer screening. In addition, combining analysis of host gene methylation with hrHPV DNA testing appears to be a new strategy in cervical cancer screening. Better specificity for host gene methylation status could improve the accuracy of detection of advanced cervical abnormalities in HPV-positive women [14]. ZNF582 and a methylation panel tested in parallel with testing for hrHPV DNA has a greater accuracy for detection of CIN2 and more severe lesions than gene methylation alone [15,16]. Genetic or environmental factors may also contribute to the progression of cervical lesions. Epigenetic components such as DNA methylation constitute an interface of the genome with the environment, including viral infections. In addition to changes in methylation of the host genome, the HPV genome also shows methylation in cervical cancer [17].
Infection with viruses such as adenovirus, Epstein-Barr virus (EBV), hepatitis B virus (HBV), and retroviruses causes DNA methylation of the host genome. DNA methylation is a common cellular defense mechanism to silence invading foreign DNA and viral genomes. For adenovirus, methylation of viral promoters leads to long-term viral gene silencing [18]. For EBV and HBV, viral replication is inhibited by methylation of promoters in viral DNA and is reversed by demethylating agents [19,20]. For retroviruses, de novo methylation of retrovirus genomes will block viral gene expression [21]. The HPV genome itself is also a target of DNA methylation.

Although the correlation between the methylation status of the HPV genome (long control region [LCR], early genes, and late genes) and disease severity has been studied, the results are controversial. Some reports showed a positive correlation while others revealed a negative correlation [22]. Different HPV genotypes may be differently methylated. For HPV16 and HPV18, some studies have shown that LCR and late gene 1 (L1) methylation is associated with the severity of cervical lesions, while methylation of some C-phosphate-G (CpG) sites may be weakly associated with cervical disease severity [23,24]. However, there have also been reports demonstrating no correlation [22,25]. For HPV31 and 45, studies showed that L1 methylation status weakly distinguished cervical lesions of different severity, but that LCR methylation showed no correlation with disease severity [24,25]. It appears that the breadth and magnitude of methylation of CpG sites differ widely between HPV genotypes. The role of HPV methylation in cervical cancer remains controversial. Most studies have focused on HPV16 and HPV18, with other types such as 31, 45, 52, and 58 being rarely studied. Therefore, the role in cervical cancer of HPV methylation in other hrHPV types and in other ethnic groups needs to be validated. The correlation between HPV and host gene methylation is also an interesting but less studied issue.

In this study, we aimed to validate the methylation status of L1 CpG sites in HPV16, 18, 52, and 58 in cervical lesions from Asian patients. We further investigated the correlation of the methylation status of each CpG site in HPV with disease severity and the correlation between the degree of methylation of virus and host genes to highlight the epigenetic mechanisms involved in HPV infection and their clinical relevance.

**MATERIALS AND METHODS**

1. **Clinical samples**

We collected 147 cervical brushings from patients from the Gynecologic Committee of the Taiwan Cooperative Oncology Group (TCOG) cohorts [12] who were infected with HPV and who were diagnosed as normal (n=28), with CIN1 (n=45), CIN2 (n=13), and CIN3/carcinoma in situ (CIS, n=61). Cytological, histological, and clinical data for all patients were reviewed by a panel of colposcopists, cytologists, and pathologists. The patients’ final diagnosis was reported as the most severe grade of abnormality identified in punch or cone biopsy specimens. Including criteria were patients aged ≥20 years and enrolled from December 2009 to November 2010. Exclusion criteria were pregnancy, chronic or acute systemic viral infections, a history of cervical neoplasia, skin or genital warts, an immune-compromised state, the presence of other cancers, or a history of surgery to the uterine cervix. The infecting HPV types included 16, 18, 52, and 58. The HPV genotype was determined by reverse line blot hybridization (the prototype Roche linear array HPV test; Roche Molecular Systems Inc., Pleasanton, CA, USA) and consensus polymerase chain reaction (PCR) of the L1 sequences. The demographic details of patients are listed in Supplementary Table 1. Informed consent
was obtained from all patients, and this study was approved by the Institutional Review Boards of all participating medical centers.

2. Collection of specimens and preparation of genomic DNA
The cervical specimens were collected by cervical brushing (PAP BRUSH; Young Ou Co., Ltd., Yongin, Korea). After cells were brushed off, the brush was immediately placed into 4°C sterile phosphate-buffered saline (PBS) for subsequent DNA extraction. Genomic DNA was extracted using the QiAamp DNA Mini Kit (Qiagen GmbH, Hilden, Germany).

3. Bisulfite modification and pyrosequencing
We used bisulfite pyrosequencing assay for the analysis of HPV gene methylation and used qMSP for the analysis of host gene methylation [12]. Bisulfite conversion was performed by EZ DNA Methylation™ Kit (Zymo Research, Irvine, CA, USA). The positive control is CpG Methylated Human Genomic DNA (Thermo Fisher, Waltham, MA, USA). The negative control is genomic DNA from healthy white blood cells. According to the manufacturer’s protocol, the methylation level of target CpG sites ranges from 0% to 100%. We used PyroMark Assay Design 2.0 software (Qiagen GmbH) to design the primers, and PyroMark PCR kits to amplify the DNA (20 ng of bisulfite-converted DNA, 500 nmol/L of each primer, 1×PCR Master Mix in 20 μL total volume). The PCR conditions were as follows: denaturation at 95°C for 15 minutes, 49 polymerization cycles (95°C for 30 seconds; the appropriate annealing temperature for 40 seconds and 72°C for 45 seconds), and a final extension at 72°C for 5 minutes. The bisulfite pyrosequencing (BPS) was performed according to the manufacturer’s recommendations for operation on a PyroMark Q24 System.

4. Statistical analysis
The Mann-Whitney nonparametric U test was used to identify differences in methylation level between 2 categories and the Kruskal-Wallis test for differences between 3 categories. All significant differences were assessed using a 2-tailed p<0.05. The correlation coefficient analysis of methylation profiling at 2 CpG sites used the Pearson correlation. Hierarchical clustering was used log values of methylation level to calculate the matrix of Euclidean distance. For the clustering analysis of the combining host gene and HPV methylation, the methylation values were transformed by quantile normalization. A receiver operating characteristic (ROC) curve was used to select the optimal threshold for distinguishing the control and comparison groups. The optimum threshold was calculated using Youden’s method. All analyses and graphs were generated using the statistical package in R (version 3.1.2; R Foundation, Vienna, Austria; https://www.R-project.org).

RESULTS

1. Correlation between HPV methylation status and severity of cervical lesions
To evaluate the degree of HPV methylation related to the spectrum of cervical lesions, we examined HPV methylation in cervical brushings using BPS. Three CpG sites of the HPV16 and 6 CpG sites of the HPV18 L1 gene were examined according to previous studies [23,26]. We selected 3 CpG sites of HPV52 and 4 CpG sites of HPV58 from L1 gene corresponding to HPV16 L1 regions. The locations of these CpG sites are indicated in Supplementary Fig. 1.

In HPV16, 1 of the 3 CpG sites, 6650, showed significantly increased methylation with more severe disease (Fig. 1A). In HPV18, the methylation level of 3 CpG sites in region 2, 7110, 7116, "https://ejgo.org" "https://doi.org/10.3802/jgo.2017.28.e56" 4/13
and 7122<sup>m</sup> showed significant differences, but not those in region 1. It is interesting that the methylation level in CIN1 was significantly lower than that in both normal controls and ≥CIN2 (Fig. 1B). In HPV52, the methylation levels of the 2 CpG sites were significantly different (Fig. 1C). There was no difference in the methylation level of any CpG site in HPV58 (Fig. 1D).

**Fig. 1.** The methylation level of the HPV L1 region in cervical brushings diagnosed with different levels of CIN. The methylation status detected by bisulfate-pyrosequencing in (A) HPV16, (B) HPV18, (C) HPV52, and (D) HPV58 at the indicated CpG sites, with the p-value shown above the graph. CIN, cervical intraepithelial neoplasia; CpG, C-phosphate-G; HPV, human papillomavirus; L1, late gene 1; Meth., methylation.
In Table 1, when the data were dichotomized, the methylation of HPV16-6650\textsuperscript{m} was significantly higher in ≥CIN1 than in normal controls (14.0 vs. 4.6; p=0.025). In contrast, HPV18-7122\textsuperscript{m} showed significantly lower methylation in ≥CIN1 than in normal controls (43.5 vs. 91.5; p=0.034). The methylation level of HPV52-6368\textsuperscript{m} and 6405\textsuperscript{m} showed differences between both <CIN1 and ≥CIN2. All of these HPV52 CpG sites showed significantly higher methylation in ≥CIN3 than in other stages (Table 1). Other CpG sites showed no difference and listed on Supplementary Table 2. When using ≥CIN2 as the endpoint, CpG site 6368\textsuperscript{m} had the best accuracy with the area under the ROC curve of 74%. When using ≥CIN3, CpG site 6405\textsuperscript{m} conferred the best accuracy of 81% (Table 2).

2. Boundary of HPV genome methylation
It is well known that adjacent CpG sites have similar methylation status, which suggests a methylation boundary in epigenetic modification \[27\]. Because no previous report identifies an HPV genome methylation boundary, we tested the correlation of the methylation levels at each CpG site. Our results demonstrated that all CpGs situated close together in HPV16 and HPV52 had highly correlated methylation status (r>0.9). In HPV18, the CpGs within regions 1 or 2 showed highly correlated methylation status (r>0.9; r>0.9), but the correlation of methylation between regions 1 and 2 was low (r=0.5), suggesting a possible methylation boundary in L1 at about 500 bps (Fig. 2).

3. Correlation of HPV and candidate host gene methylation
Although the HPV genome may be methylated by the host cellular defense mechanisms, HPV may also influence the host cellular methylation mechanism to alter host genome

<table>
<thead>
<tr>
<th>Table 1. Relationships between HPV methylation level and clinical lesions</th>
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<td>Variables</td>
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<tr>
<td>HPV16</td>
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<tr>
<td>6,650 bp</td>
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<tr>
<td>Case No.</td>
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<tr>
<td>Median (95% CI)</td>
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<td>HPV18</td>
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<td>7,122 bp</td>
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<td>Case No.</td>
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<tr>
<td>Median (95% CI)</td>
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<tr>
<td>HPV52</td>
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<tr>
<td>6,368 bp</td>
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<tr>
<td>Case No.</td>
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<tr>
<td>Median (95% CI)</td>
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<tr>
<td>6,405 bp</td>
</tr>
<tr>
<td>Case No.</td>
</tr>
<tr>
<td>Median (95% CI)</td>
</tr>
<tr>
<td>6,443 bp</td>
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<tr>
<td>Case No.</td>
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<td>Median (95% CI)</td>
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All p-values calculated by Mann-Whitney U test.
CI, confidence interval; CIN, cervical intraepithelial neoplasia; HPV, human papillomavirus.

<table>
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<th>Table 2. Performance of HPV52 methylation in cervical lesions</th>
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<tr>
<td>Location of CpG site (bp)</td>
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</tr>
<tr>
<td>Sensitivity</td>
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<td>6,368</td>
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<td>6,405</td>
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<td>6,443</td>
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AUC, area under the curve of receiver operating characteristic; CI, confidence interval; CIN, cervical intraepithelial neoplasia; CpG, C-phosphate-G; HPV, human papillomavirus.
methylation. The association between HPV genome methylation and host gene methylation is interesting but has not been tested previously. In an earlier study, we identified a correlation between HPV methylation and methylation of host genes [3,28]. Our results demonstrated that HPV16 methylation was positively correlated with host gene methylation, especially that of HS3ST2, PTPR, and POU4F3 (r>0.7), while HPV18 methylation was weakly correlated with methylation of PAX1 and SOX1 (r<0.4). HPV52 and HPV58 methylation status showed a weak negative correlation with host gene methylation (Fig. 3).

4. Combination of host and viral gene methylation improves cervical lesion screening
Although host gene methylation has been proposed as a biomarker for cervical cancer detection with high specificity, its moderate sensitivity requires improvement [12,16]. One appealing idea is to combine analysis of host and viral gene methylation to improve the
Methylation of $PAX1$ is one of the most promising markers for cervical cancer screening. We tested the possibility of improving the sensitivity of $PAX1$ for detecting ≥CIN2 lesions. Although the analysis was limited by the number of patients for whom both $PAX1$ and HPV type-specific methylation data were available, we used clustering analysis to test whether the inclusion of HPV methylation status could improve the sensitivity of $PAX1$ methylation status for the classification of disease. $PAX1$ alone classified a single ≥CIN2 group, while $PAX1$ and HPV methylation in combination classified 2 ≥CIN2 groups, suggesting a potential benefit of adding HPV methylation status to the analysis (Fig. 4).

Fig. 3. The correlation of methylation levels between HPV and host genes. (A) In HPV16, the correlation coefficients are classified into 3 groups: >0.7, 0.7–0.4, and <0.4. Others correlation showed in HPV18 (B), HPV52 (C), and HPV58 (D). Blue and red indicate positive and negative correlation, respectively. HPV, human papillomavirus; n, case number.

sensitivity. Methylation of $PAX1$ is one of the most promising markers for cervical cancer screening. We tested the possibility of improving the sensitivity of $PAX1$ for detecting ≥CIN2 lesions. Although the analysis was limited by the number of patients for whom both $PAX1$ and HPV type-specific methylation data were available, we used clustering analysis to test whether the inclusion of HPV methylation status could improve the sensitivity of $PAX1$ methylation status for the classification of disease. $PAX1$ alone classified a single ≥CIN2 group, while $PAX1$ and HPV methylation in combination classified 2 ≥CIN2 groups, suggesting a potential benefit of adding HPV methylation status to the analysis (Fig. 4).
DISCUSSION

The present study demonstrated that infection with different HPV genotypes leads to different HPV methylation status and that this will correlate with cervical lesion severity. Moreover, different HPV genotypes showed different correlations between HPV methylation and host gene methylation. All these data prompted us to hypothesize that different HPV genotypes may have different effects on the host epigenetic machinery, and that the host epigenetic machinery may have different responses to different HPV genotypes. HPV16 is the most common oncogenic genotype in cervical lesions. Infection with HPV16 may trigger the epigenetic defense machinery to methylate the HPV genome, but hijacking of this defense by HPV16 results in alteration of the methylation of the host genome. This could cause the observed correlation between HPV16 genome methylation and host gene methylation ranging from normal tissue to CIS. However, HPV52, although presumably an oncogenic type, may not strongly induce the host epigenetic machinery so that both viral and host genes show low levels of methylation in the late precancerous stages (Supplementary Fig. 2). This speculation warrants further investigation, which may improve our understanding of the role of epigenetics in HPV oncogenesis.

Some studies have shown that HPV infection interferes with host epigenetic regulation. HPV infection upregulated the expression of DNMT1, DNMT3A, and DNMT3B [29,30]. The mechanism of this appeared to be degradation of p53 by the E6 protein of HPV, which resulted in specificity protein 1 (Sp1) dissociating from the p53/Sp1 complex. Then, phosphorylation of Rb by E7 of HPV resulted in E2F dissociating from the Rb/E2F complex. The released Sp1 and E2F bound to the DNMT promoters and upregulated expression of DNMTs [29-31]. Previous studies showed that the E6 protein of HPV16 is capable of binding to p53, which may release more Sp1 and activate more DNMTs and alter the degree of DNA methylation. Taken together, we speculate that infection with different HPV types may result in differences in the methylation status of both the host and virus genomes. This may also affect the carcinogenic activity of different HPV types.

Increasing integration of HPV DNA into the host genome correlates with cervical cancer progression [32]. The integrated form of HPV DNA was found in 80% and 100% of HPV16- and HPV18-infected patients, respectively, but in only about 25% of HPV52-infected patients [33]. Insertion of viral DNA sequences into the host genome often triggers host defense
mechanisms such as the DNA methylation machinery, which then mediates the methylation of the virus genome [30,34]. Therefore, the low integration rate of HPV52 may induce a low level of epigenetic defenses. This is consistent with our result that the methylation of L1 in HPV52 was lower than that in HPV16 and HPV18.

Detection of disease-specific gene methylation is a new strategy for cancer screening, and an optimal biomarker will improve detection of cancer at an early stage. SEPT9 methylation was identified as a biomarker of colorectal cancer over 10 years ago [35] and was recently approved for clinical use by the Food and Drug Administration (FDA). In previous studies, cervical cancer-specific methylation of genes such as PAX1, SOX1, ZNF582, and POU4F3 showed their clinical usefulness in detecting ≥CIN3 precancerous samples [3,12,36]. PAX1 methylation was recently approved by Taiwan FDA as an adjunct testing for cervical cancer screening. These methylation biomarkers showed satisfactory clinical results in patients with CIN3 and more severe lesions. However, the goal of early detection is to improve the detection of precancerous lesions before the ≥CIN3 stage by, for instance, identifying new cervical cancer methylation biomarkers, possibly by including analysis of the methylation of other genes or by HPV typing. In this study, we evaluated the feasibility of combining HPV methylation with host gene methylation to improve the detection of CIN3 lesions. Our data showed that the combination of HPV methylation and specific gene methylation improved the clinical performance of classification of precancerous lesions. This result was consistent with the previous reports that HPV16 gene methylation provided some improvement in disease discrimination when combined with methylation of EPB41L3 and LMX1 [37]. A larger patient sample is required to verify the optimal combination for precancer screening of hrHPV-infected CIN patients.

The correlation coefficient for methylation of adjacent CpG sites was high in HPV16 and HPV52, but low in 2 CpG sites more than 500 bps apart in HPV18. This result might reflect the phenomenon of methylation boundaries. Sp1 and CTCCC-binding factor (CTCF) are associated with methylation boundaries in the human genome [38], and there are 2 CTCF binding motifs on the HPV18 genome that might be relevant to this phenomenon. However, HPV18 has a breakpoint at 6,775 to 6,803 bps [39] between regions 1 and 2. If HPV18 cleaves at this position and inserts into the host genome, the distance between regions 1 and 2 will change from 500 bps to more than 7,000 bps. Taken together, these results support our finding that a lower correlation between the methylation of regions 1 and 2 in HPV18 may be the result of a methylation boundary. We searched for Sp1 and CTCF binding motifs in the HPV genome, and identified 1 Sp1 binding motif in HPV52 and 3 in HPV58, but no Sp1 binding motifs in either HPV16 or HPV18. There was 1 CTCF binding motif in HPV16, 2 in HPV18, 4 in HPV52, and 5 in HPV58. Because Sp1 and CTCF protect the surrounding DNA against methylation, a greater number of binding motifs in the HPV genome implies a higher potential to form a methylation-free boundary. The abundant CTCF and Sp1 binding motifs in the HPV58 genome may explain the low methylation of HPV58.

In conclusion, our study showed that L1 methylation of HPV16, HPV18, and HPV52 is associated with cervical lesion severity. The L1 methylation of HPV16 and HPV18 had the potential to differentiate normal from ≥CIN1 cervical lesions. The L1 methylation of HPV52 had the potential to distinguish <CIN2 from ≥CIN2 and <CIN3 from ≥CIN3 cervical lesions. In addition, the combination of host and HPV genome methylation had the potential to improve the diagnosis of cervical lesions. HPV16 and HPV18 methylation was positively correlated and HPV52 methylation was negatively correlated with that of PAX1 and SOX1.
These results suggest that infection with different HPV genotypes might lead to different methylation status of the HPV genome and host genes. Increasing the number of samples and exploring wider areas of relevance are the goals of our future studies.

ACKNOWLEDGMENTS

Yaw-Wen Hsu is thankful to Graduate Institute of Life Sciences for the Ph.D. Fellowship.

SUPPLEMENTARY MATERIALS

Supplementary Table 1
Demographics related to the clinical samples

Click here to view

Supplementary Table 2
Methylation level of HPV L1 region in patients and control subjects

Click here to view

Supplementary Fig. 1
The distribution of CpG sites in 4 HPV types. The location of CpG sites and genes shown on each HPV genome by IGV. The arrows indicate regions where methylation was detected. (A) HPV16. (B) HPV18. (C) HPV52. (D) HPV58.

CpG, C-phosphate-G; HPV, human papillomavirus; IGV, Integrative Genomics Viewer.

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Supplementary Fig. 2
The epigenetic aspects of the HPV genome. We hypothesized that different HPV genotypes may have different effects on the host epigenetic machinery. In HPV16, the host methylation machinery is triggered strongly to methylate the viral and host genome. In HPV52, the viral genome methylation is accompanied by less methylation of host gene.

CIN, cervical intraepithelial neoplasia; CIS, carcinoma in situ; HPV, human papillomavirus.

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