Evaluation of a Commercialized In Situ Hybridization Assay for Detecting Human Papillomavirus DNA in Tissue Specimens from Patients with Cervical Intraepithelial Neoplasia and Cervical Carcinoma

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To evaluate a commercialized in situ hybridization (ISH) assay for detecting human papillomavirus (HPV) DNA, we compared the ability of a new ISH probe, Inform HPV III (Ventana Medical Systems, Tucson, AZ), to that of PCR assays to detect HPV DNA in cervical tissue specimens with normal cervix (20 cases), cervical intraepithelial neoplasia (CIN; CIN 1, 27 cases; CIN 2, 28 cases; and CIN 3, 33 cases), and cervical carcinoma (29 cases). General HPV DNA was detected using consensus primer-mediated PCR assays. HPV genotyping was performed by using EasyChip HPV blot (King Car Yuan Shan Institute, I-Lan, Taiwan). HPV16 integration status (E2/E6 ratio) was determined by using quantitative real-time PCR. Our findings showed that the ISH and PCR had fair to good agreements in detecting HPV DNA across all CIN categories without significant differences (Kappa coefficient, 0.34 to 0.63; P = 0.13 to 1.0). However, ISH detected significantly fewer HPV-positive cases in carcinoma than PCR did (Kappa coefficient, 0.2; P = 0.03). Eleven cases with ISH PCR+ results had HPV types that can be detected by Inform HPV III. Five carcinoma cases with ISH-PCR+ results showed a significantly higher level of integrated HPV16 (P = 0.008) than did the ISH--cases. As a consequence, lower copy numbers of episomal HPV16 in carcinoma might be the cause for the false-negative ISH results. Although the punctate signal pattern of HPV significantly increased with the severity of disease (P trend = 0.01), no significant difference in the HPV16 integration status was observed between the cases with a punctate signal only and the cases with mixed punctate and diffuse signals (P = 0.4). In conclusion, ISH using the Inform HPV III probe seems comparable to PCR for detecting HPV DNA in cervical tissue with CINs. False-negative ISH results appear to be associated with the lower copy numbers of the episomal HPV16 but not with the ability of the Inform HPV III probe to detect specific HPV types. In addition, signal patterns, especially a mixed punctate and diffuse pattern of HPV, cannot be reliably used to predict viral integration status.

Oncogenic human papillomavirus (HPV) infection is a well-established major risk factor for the development of more than 99% of cervical carcinomas and precancerous lesions (high-grade cervical intraepithelial neoplasia [CIN]; i.e., CIN 2/3 (6, 35, 39). In recent years, HPV DNA testing has been utilized as a molecular maker in gynecological cytology either for the triage of women with mildly abnormal Papanicolaou (Pap) test results (36) or in conjunction with Pap tests for predicting CIN 2/3 in women who are 30 years or older (37). In histopathology, HPV DNA testing in formalin-fixed, paraffin-embedded (FFPE) cervical tissue can be used for the following purposes: (i) to help resolve diagnostic discrepancy in patients with CINs and distinguish endocervical immature squamous metaplasia from high-grade dysplasia or endocervical glandular reactive changes from glandular dysplasia, (ii) to provide valuable information for assessing the risk of CIN progression or of disease recurrence in women who were treated for CIN 2/3 or carcinoma (1, 26, 27, 34), and (iii) to provide valuable information in cervical cancer research.

Different HPV DNA testing assays in FFPE cervical tissue specimens have been used, such as PCR and in situ hybridization (ISH). Each has its strengths and drawbacks. Although the PCR assay is highly sensitive for detecting HPV DNA, there are disadvantages to using PCR for HPV DNA detection in FFPE tissue specimens (5, 34). In particular, PCR inhibitors in DNA extracts from FFPE cervical tissue can interfere with the ability of PCR to detect HPV (10). In addition, although PCR assays for HPV DNA testing are commercially available, highly trained laboratory personnel are required to perform the assays, and strict laboratory conditions are needed to avoid contamination. More importantly, morphological context, which is essential for histopathological specimen interpretation, is lost when a PCR assay is used.

In contrast, ISH, a direct signal detection assay, has the advantage of preserving the morphological context with HPV DNA signals. The ISH assay can be automated along the same lines as immunohistochemistry staining to minimize intra- or interassay inconsistency. The interpretation of ISH is also similar to that of immunohistochemistry staining in tissue sections. Furthermore, the ISH signal patterns of HPV DNA have been

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reported to be associated with the physical status of HPV in infected cells, i.e., epimodal or integrated forms (9, 11). Because the integration of oncogenic HPV into the human genome is a critical step for cervical cancer carcinogenesis, the signal pattern of HPV, which may suggest viral integration status, could be a useful marker for predicting precancerous lesion progression. Despite these advantages, the ISH assay has a low sensitivity, which is a major concern for pathologists who use ISH to detect HPV in tissue (10).

In recent years, ISH assays that use improved signal-detecting methods, such as the enzyme–categorized signal detecting system, have shown a higher sensitivity (8, 31–33). Inform HPV (Ventana Medical Systems), a commercially available ISH assay for HPV DNA testing, can be used in both cytological and histological specimens. Recently, Inform HPV III, a new generation of ISH probe, became available for HPV DNA testing in tissue specimens. Inform HPV III utilizes a stacked antibody approach (the iView Blue Plus kit) to enhance its sensitivity. Briefly, the primary antibody is directed against the DNP hapten. Signal amplifications are generated through antibody stacks consisting of a secondary antibody and a biotinylated tertiary antibody. Finally, a streptavidin–alkaline phosphatase conjugate is added to generate color precipitates. Published studies of systemic analyses of the sensitivity of ISH using Inform HPV III to detect HPV DNA in cervical tissue are limited (38). Knowing the efficacy of the ISH assay using the Inform HPV III probe in cervical tissue not only would allow us to better use ISH-based HPV DNA testing in tissue specimens but also would provide valuable information for the cytological application of ISH in Pap specimens. Therefore, in the present study, we compared the ISH assay using the Inform HPV III probe to PCR assays using consensus primer sets for detecting HPV DNA in FFPE cervical tissue specimens with CINs and carcinoma. Using Inform HPV III, the nuclear signal patterns of HPV were also compared to HPV16 integration status by using quantitative real-time PCR (qRT-PCR) assay to determine whether the nuclear signal pattern of HPV might be reliable for estimation of HPV integration status and therefore a potential useful marker to predict precancerous lesion progression.

MATERIALS AND METHODS

The Institutional Review Board of The University of Texas M. D. Anderson Cancer Center approved this study.

Cervical tissue specimen selection. Archived, FFPE cervical tissue specimens obtained from 2004 to 2005 were retrieved from the Department of Pathology at M. D. Anderson Cancer Center. In consecutive order, 200 cervical tissue specimens from punch biopsies, loop electrosurgical excision procedures, cone biopsies, and hysterectomies were selected. Three pathologists independently reviewed the hematoxylin and eosin (H&E)-stained slides. Specimens were eliminated from the study for any of the following reasons: (i) no consensus in diagnosis; (ii) no lesion in the last section of the specimen; (iii) insufficient DNA for analysis by spectrophotometry; and (iv) negative β-globin amplification by PCR. After exclusion of specimens for these various reasons, there were 137 specimens with the following pathological diagnoses: normal cervix (20 cases), CIN 1 (27 cases), CIN 2 (26 cases), CIN 3 (33 cases), and cervical squamous cell carcinoma (29 cases).

Tissue sectioning for in situ hybridization and PCR. Serial sections were cut to a thickness of 4 μm and mounted on positively charged glass slides for ISH. The paraffin rolls were cut from each block (10 sections, 5 μm in thickness) for DNA extraction. The extra sections cut before and after each tissue sections were stained with H&E and used to determine specimen quality for testing. To avoid cross-contamination, the microtome was cleaned with a new blade for each case.

Inform HPV III ISH. Inform HPV III probe sets able to detect 13 types of oncogenic HPV (i.e., types 16, 18, 31, 33, 35, 45, 51, 52, 56, 58, 59, 68, and 70) were provided by Ventana Medical Systems. The ISH assay was performed according to the manufacturer’s guidelines using the BenchMark automated slide staining system (Ventana Medical System). HPV control slides consisted of formalin-fixed, paraffin-embedded sections containing three separate collections of cells on a singl Slides (Ventana Medical Systems). These cells consisted of the CaSkii cervical cancer cell line (containing 200 to 400 copies of HPV16 per cell); the HeLa cervical cancer cell line (containing 10 to 50 copies of HPV18 per cell); and the C-33A cell line, which served as a negative control. Reagent negative control was set using negative control probes provided by Ventana Medical Systems.

Three pathologists independently reviewed the Inform HPV III slides. The pathologists agreed on the results of ISH in 95% of the cases (130 of 137). Consensus was obtained in the remaining seven cases after review. Nuclear staining was considered a positive result for HPV DNA. The signal patterns of HPV in nucleus using Inform HPV III were classified as follows: diffuse, signals that are condensed and uniformly packed in the nucleus (Fig. 1A); and punctate, signals that are dot-like and sparsely distributed in the nucleus (Fig. 1A).

DNA extraction. DNA was extracted from tissue specimens using the DNeasy kit (catalog no. 69506; Qiagen, Valencia, CA) according to the manufacturer’s instructions. The extracted DNA was quantified by using a Nanodrop spectrophotometer (Nanodrop Technologies, Wilmington, DE).

HPV DNA testing using consensus primer-mediated PCR. HPV DNA positive cases were determined by using consensus primer-mediated PCR assays independently performed in the genomic labs at M. D. Anderson Cancer Center, Houston, TX, and Tri-Service Hospital, National Defense Medical Center, Taipei, Taiwan. Each PCR of the HPV DNA testing battery was carried out random, without knowledge of the pathological diagnoses. At M. D. Anderson Cancer Center, PCR assays for HPV DNA detection were performed as described previously (16). Briefly, the amplification of β-globin (268 bp) was performed to determine the quality of the extracted DNA using primer pC01Igh20 according to the method of Bauer et al. (4) and using AmpliTaq Gold polymerase (Applied Biosystems, Foster City, CA). A 25-ng aliquot of genomic DNA was added to a PCR master mixture containing 1× PCR buffer (100 mM Tris-HCl [pH 8.3]), 200 μM concentrations of each deoxynucleoside triphosphate, 200 nM primer, 1.5 mM MgCl2, and 2.5 U of AmpliTaq Gold DNA polymerase. PCR was performed under the following permissive cycling conditions: 10 min at 94°C, followed by 1 min at 94°C, 1 min at 40°C, 90 s at 72°C, and 5 min at 72°C for 39 cycles.

Specificity positive for β-globin were screened for HPV DNA using GPS+/GP6+ consensus primer-mediated PCR, which generates 150-bp ampiclons and can detect all of the oncogenic HPV genotypes that Inform HPV III detects. PCR was performed according to the method of Jacobs et al. (21). A 25-ng aliquot of genomic DNA was added to the PCR master mixture, which contained 1× PCR buffer, 200 μM concentrations of each deoxynucleoside triphosphate, 200 nM primer, 3.5 mM MgCl2, and 2.5 U of AmpliTaq Gold DNA polymerase. The cycling conditions were the same as those used for β-globin amplification. Specifically, if there were false positive for β-globin but negative for GPS+GP6+ were rescreened for HPV DNA with the second consensus primer set (PGM90/11), which generates 450-bp ampiclons and also detects all of the oncogenic HPV genotypes that Inform HPV III detects. A 25-ng aliquot of genomic DNA was used for amplification in a PCR master mixture containing 1× PCR buffer, 200 μM concentrations of each deoxynucleoside triphosphate, 200 nM primer, 4.0 mM MgCl2, 7.5 U of AmpliTaq Gold DNA polymerase. The PCR was performed under the following cycling conditions: 9 min at 94°C, followed by 1 min at 94°C, 1 min at 55°C, and 1 min at 72°C for 39 cycles. Specimens with known HPV were used as positive controls. Sterile water was used as a negative control for each set of amplifications. The PCR products were visualized with etidium bromide staining on a 4% low-melting-point agarose gel.

At the Tri-Service Hospital, National Defense Medical Center, Taiwan, specimens were tested for HPV DNA using MY11/GP6+ primer sets to amplify a fragment of 192 bp as described previously (19, 24). GAPDH/FADHPR primer sets were used to amplify a 136-bp amplicon of GAPDH (glyceraldehyde-3-phosphate dehydrogenase) for validation. Briefly, PCR for HPV DNA was performed in a final reaction volume of 25 μl with 20-ng aliquot of genomic DNA in a PCR master mixture containing 15 mM Tris-HCl (pH 8.0), 2.0 mM MgCl2, 50 mM KCl, 0.25 mM concentrations of each deoxynucleoside triphosphate, 0.6 μM primer, and 0.5 U of DNA polymerase (Ventana Medical Systems). A PCR assay was performed as follows: 10 min at 95°C, followed by 40 cycles of 30 s at 95°C, 30 s at 45°C, and 30 s at 72°C, with a final extension of 5 min at 72°C. PCR for GAPDH was performed in a final reaction volume of 25 μl with 10-ng aliquot of genomic DNA.

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in a PCR master mixture containing 15 mM Tris-HCl (pH 8.0), 2.5 mM MgCl₂, 50 mM KCl, 0.2 mM concentrations of each deoxynucleoside triphosphate, 0.2 µM primer, and 0.5 U of HotStart Tag DNA polymerase. A PCR assay was performed as follows: 10 min at 95°C, followed by 40 cycles of 15 s at 95°C, 1 min at 57°C, and 30 s at 72°C, with a final extension of 5 min at 72°C. An aliquot of 5 µl of PCR product was analyzed by electrophoresis on a 2% agarose gel and stained with ethidium bromide.

Specimens that tested positive for HPV DNA in one of the consensus HPV PCRs were subjected to genotyping.

**HPV genotyping by EasyChip.** Specimens with positive results of consensus primer-mediated PCR were genotyped by using EasyChip HPV Blot (King Car, I-Lan, Taiwan) (24). EasyChip array can genotype 39 HPV types (HPV types 6, 11, 16, 18, 26, 31, 33, 35, 39, 44, 45, 51, 52, 53, 54, 55, 56, 58, 59, 61, 62, 66, 67, 68, 69, 70, 72, 74, 82, CPN061, CP8304, L1AE5, MM4, MM7, and MM8, as well as three intrinsic controls). The HPV type-specific probes are immobilized on a 14.4-mm-by-9.6-mm nylon membrane, which is used for reverse blot hybridization and detects HPV DNA in a single assay. The hybridization was performed according to the manufacturer's guide. Briefly, the blot membrane was equilibrated with 2× saline-sodium citrate (SSC; 1× SSC is 0.15 M NaCl plus 0.015 M sodium citrate) at room temperature for 10 min. The blot was preincubated in hybridization buffer (2× SSC, 0.5% blocking reagent, 5% deionized formamide) at 45°C for 15 min. The membrane was hybridized with 500 µl of hybridization buffer containing 20 µl of the denatured amplicons (15 µl of HPV and 5 µl of GAPDH PCR products) by shaking at 35°C for at least 3 h. The blot was washed twice in washing buffer 1 (2× SSC, 0.1% SDS) for 5 min at 25°C and then washed twice in washing buffer 2 (0.2× SSC, 0.1% SDS) for 5 min at 35°C. The blot was equilibrated with buffer 1 (1× phosphate-buffered saline [pH 7.4], 0.05% Tween 20, 0.1% SDS) by shaking at 25°C for 5 min and then buffer 2 (1× phosphate-buffered saline [pH 7.4], 0.05% Tween 20, 0.1% SDS, 0.5% blocking reagent) at 25°C for 1 h. The blot was incubated in 500 µl of buffer 2 containing streptavidin-AP (Calbiochem; alkaline phosphatase conjugates and biotinylated antibodies, 1:1,000 dilution) at 25°C for 30 min. The membrane was hybridized with 500 µl of hybridization buffer containing 20 µl of the denatured amplicons (15 µl of HPV and 5 µl of GAPDH PCR products) by shaking at 35°C for at least 3 h. The blot was washed twice in washing buffer 1 (2× SSC, 0.1% SDS) for 5 min at 25°C and then washed twice in washing buffer 2 (0.2× SSC, 0.1% SDS) for 5 min at 35°C. The blot was equilibrated with buffer 1 (1× phosphate-buffered saline [pH 7.4], 0.05% Tween 20, 0.1% SDS) by shaking at 25°C for 5 min and then buffer 2 (1× phosphate-buffered saline [pH 7.4], 0.05% Tween 20, 0.1% SDS, 0.5% blocking reagent) at 25°C for 1 h. The blot was incubated in 500 µl of buffer 2 containing streptavidin-AP (Calbiochem; alkaline phosphatase conjugates and biotinylated antibodies, 1:1,000 dilution) at 25°C for 30 min. The blot was then washed in buffer 1 and rinsed with buffer 3 (0.1 M Tris-HCl [pH 9.5], 0.1 M NaCl) for 5 min. Then, 80 µl of nitroblue tetrazolium-BCIP (5-bromo-4-chloro-3-indolylphosphate) was added, followed by incubation for 30 min at 25°C. The reaction was stopped by adding distilled water. The HPV types were determined by a visual assessment protocol provided by King Car.

**qRT-PCR for HPV16 viral integration status.** HPV16 DNA was amplified with primers and probe targeting the E6/E7 oncoproteins of HPV16 using qRT-PCR according to the method of Gravitt et al. (15). The TaqMan minor groove binder probes were used for qRT-PCR. Minor groove binder probes were labeled with a carboxyfluorescein reporter dye at the 5′ end and a nonfluorescent quencher at the 3′ end. Primers described by Flores-Munguia et al. (12) for the E6/E7 regions were used for HPV16. For the HPV16 integration assay, primers for HPV16 E2, as described by Peitsaro et al. (28), were used. Minor groove binder probe for the assay was designed by using Primer Express software; both probe and primers were purchased from Applied Biosystems. We used the plasmid HPV16 clone for positive controls (American Type Culture Collection, Manassas, VA).

All qRT-PCR assays for HPV genotyping were performed by using an ABI Prism 7900 HT with a 96-well plate (Applied Biosystems). Briefly, 2.5 ng of genomic DNA of each specimen and control, including water as a no-template control, was added to a 25-µl reaction mixture containing 1× TaqMan Universal PCR Master Mix without AmpErase uracil-N-glycosylase, 1× gene expression assay (Applied Biosystems), 250 nM fluorescent probe, and 900 nM primer under the following cycling conditions: 10 min at 95°C, followed by 15 s at 95°C and 1 min at 60°C for 50 cycles. HPV16 was assayed on a single 96-well plate, with 40 specimens in duplicate. A standard curve to determine HPV quantity was established, also in duplicate, with a 10-fold dilution series ranging from 10 to 10⁵ copies per well using plasmid-cloned HPV, including water as a no-template control. A linear relationship was obtained between the log value of the viral copy numbers and the threshold cycle (data not shown).

The integration status of HPV16 was determined by measuring the E2/E6 ratio (28). The protocol for the E2/E6 assay was modified to include dual standard curves of E2 and E6. The E2/E6 ratio was calculated from the same reaction. An E2/E6 ratio greater than or equal to 1 was classified as a complete episomal form. No amplification for E2 was classified as a complete integration. A low E2/E6 ratio represented predominantly integrated forms. Conversely, a high E2/E6 ratio indicated high levels of episomal forms.

**Statistical analysis.** Descriptive statistics were calculated by using Kappa statistics to assess the agreement between Inform HPV III and PCR HPV DNA testing. McNemar’s tests were used to assess the homogeneity between paired categorical variables. The chi-square or Fisher exact test was used to assess the association between categorical variables. The Cochran-Armitage trend test or the Jonckheere-Terpstra trend test was used to assess the trend of the Inform HPV III results and the severity of disease. The Kruskal-Wallis test was used to assess the association between the Inform HPV III results and the HPV16 E2/E6 ratio. P values (two-sided test) of <0.05 were considered significant. All statistical analyses were carried out by using SAS 8.0 (SAS Institute, Cary, NC).
TABLE 1. Agreement between ISH using Inform HPV III and consensus primer-mediated PCR to detect oncogenic HPV DNA in CINs and cervical carcinoma.

<table>
<thead>
<tr>
<th>Diagnosis and ISH result</th>
<th>No. of cases (%) examined</th>
<th>Negative</th>
<th>Positive</th>
<th>Kappa coefficient</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>by PCR</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CIN 1</td>
<td>Negative</td>
<td>7 (77.8)</td>
<td>2 (22.2)</td>
<td>0.53</td>
<td>0.69</td>
</tr>
<tr>
<td></td>
<td>Positive</td>
<td>4 (22.2)</td>
<td>14 (77.8)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CIN 2</td>
<td>Negative</td>
<td>5 (100)</td>
<td>0</td>
<td>0.63</td>
<td>0.13</td>
</tr>
<tr>
<td></td>
<td>Positive</td>
<td>4 (17.4)</td>
<td>19 (82.6)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CIN 3</td>
<td>Negative</td>
<td>2 (40.0)</td>
<td>3 (60.0)</td>
<td>0.34</td>
<td>1.0</td>
</tr>
<tr>
<td></td>
<td>Positive</td>
<td>2 (7.1)</td>
<td>26 (92.6)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Carcinoma</td>
<td>Negative</td>
<td>1 (14.3)</td>
<td>6 (58.7)</td>
<td>0.2</td>
<td>0.03</td>
</tr>
<tr>
<td></td>
<td>Positive</td>
<td>0</td>
<td>22 (100.0)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

RESULTS

The age of women in the study ranged from 18 to 79 years, with a mean age of 40 years and a median age of 41 years. **HPV DNA detection by ISH and PCR assays.** Twenty cases of normal cervical tissue were negative for HPV DNA by ISH assay using Inform HPV III and consensus primer-mediated PCR assays using the GP5+/GP6+, PGMY09/11, and MY11/ GP6+ primer sets. In 117 cases of cervical specimens with diseases, we found that 91 cases (78%) were positive for HPV DNA by Inform HPV III, and 92 cases (79%) were determined to be positive by the PCR assays. Compared to the PCR assays, Inform HPV III showed similar HPV DNA positivity in CIN 1 (67% [18 of 27] versus 59% [16 of 27]) and CIN 2 (82% [23 of 28] versus 68% [19 of 28]) but lower HPV DNA positive rates in CIN 3 (85% [28 of 33] versus 88% [29 of 33]) and carcinoma (76% [22 of 29] versus 97% [28 of 29]). Combined HPV DNA positivities in CIN 1, CIN 2, and CIN 3 and carcinoma were 74% (20 of 27), 82% (23 of 28), 94% (31 of 33), and 97% (28 of 29), respectively. We found no association between HPV DNA positivity by Inform HPV III and the HPV genotypes (data not shown).

The agreement between the Inform HPV III and PCR assays in detecting HPV DNA is shown in Table 1. The two tests were concordant in 85% (116 of 137) of all cases. The Inform HPV III and PCR assays had moderate to good agreements (Kappa coefficients of 0.53 and 0.63, respectively) in detecting HPV DNA in CIN 1 and CIN 2 and a fair agreement (Kappa coefficient, 0.34) in CIN 3 without significant differences (P = 0.13 to 1.0). However, in cervical carcinoma, the two tests showed a significant disagreement with regard to HPV DNA positivity, with the positive rate being lower for Inform HPV III than for the PCR assay (Kappa coefficient, 0.2; P = 0.03).

There were 21 cases with discrepancy in results between PCR and ISH tests. Of these, 10 cases showed ISH− PCR+ results, and 11 cases showed ISH+ PCR− results. In the 10 ISH+ PCR− cases, repeat DNA extractions with consensus primer-mediated PCR testing for HPV DNA were negative. Of the 11 cases with ISH− PCR+ results, 6 were carcinoma cases, with 4 cases positive for HPV16, 1 case positive for HPV16/35, and 1 case positive for HPV33/58. In the remaining five cases with ISH− PCR+ results, all showed HPV types that can be detected by Inform HPV III probe, including HPV52 (2 cases), HPV45 (1 case), HPV18 (1 case), and HPV59 (1 case) (Table 2).

**Distribution of HPV-positive cells and HPV signal patterns.** Figure 1 presents the ISH signal of HPV shown by Inform HPV III. Nonspecific background bindings were not observed, as frequently seen using Inform HPV II probes (data not shown). In the majority of cases (87%), HPV-positive cells were observed in <50% of the lesions, with more HPV-positive cells focally located in the upper epidermis in CIN 1 (Fig. 1B) or CIN 2 and more HPV-positive cells diffusely distributed in CIN 3 (Fig. 1C) or carcinoma (Fig. 1D).

A comparison of HPV signal patterns in the nucleus shown by Inform HPV III with cervical disease categories is illustrated in Table 3. Three cases with diffuse patterns were observed in CIN 1 and CIN 2 but not in CIN 3 or carcinoma. Mixed diffuse and punctate patterns were predominantly found in CIN 1 and 2. Punctate patterns were predominantly observed in CIN 3 and carcinoma cases, either alone or mixed with diffuse patterns. The punctate patterns significantly increased with the severity of disease (P = 0.01) (Table 3).

**HPV16 viral integration status and ISH signal patterns.** The signal patterns of the HPV shown by Inform HPV III were compared to the HPV16 E2/E6 ratio. The mean of the E2/E6 ratio was lower in the specimens with a punctate pattern only (E2/E6, 0.293) than in cases with mixed punctate and diffuse patterns (E2/E6, 0.532). However, no significant difference in the E2/E6 ratio was observed between these two groups (P = 0.4) (Table 4). Five carcinoma cases with ISH− PCR− results were positive for HPV16, showing a significantly lower E2/E6 ratio (E2/E6, 0.05; P = 0.008) compared to the cases shown to be positive by the PCR assays.

**Distribution of HPV genotypes in cases with ISH− PCR+ results.**

<table>
<thead>
<tr>
<th>Case no.</th>
<th>Disease</th>
<th>HPV type(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>CIN 1</td>
<td>45</td>
</tr>
<tr>
<td>2</td>
<td>CIN 1</td>
<td>18</td>
</tr>
<tr>
<td>3</td>
<td>CIN 3</td>
<td>59</td>
</tr>
<tr>
<td>4</td>
<td>CIN 3</td>
<td>52</td>
</tr>
<tr>
<td>5</td>
<td>CIN 3</td>
<td>52</td>
</tr>
<tr>
<td>6</td>
<td>Carcinoma</td>
<td>16</td>
</tr>
<tr>
<td>7</td>
<td>Carcinoma</td>
<td>16</td>
</tr>
<tr>
<td>8</td>
<td>Carcinoma</td>
<td>16</td>
</tr>
<tr>
<td>9</td>
<td>Carcinoma</td>
<td>16</td>
</tr>
<tr>
<td>10</td>
<td>Carcinoma</td>
<td>16/35</td>
</tr>
<tr>
<td>11</td>
<td>Carcinoma</td>
<td>33/58</td>
</tr>
</tbody>
</table>

**Table 2. Distribution of HPV genotypes in cases with ISH− PCR+ results.**

<table>
<thead>
<tr>
<th>HPV signal pattern</th>
<th>No. of cases (%)</th>
<th>P-trend</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diffuse</td>
<td>2 (11)</td>
<td>0</td>
</tr>
<tr>
<td>Punctate + diffuse</td>
<td>9 (50)</td>
<td>0</td>
</tr>
<tr>
<td>Punctate</td>
<td>7 (39)</td>
<td>0.01</td>
</tr>
</tbody>
</table>

**Table 3. Association between HPV signal patterns and CINs or cervical carcinoma as determined by Inform HPV III.**
be positive by Inform HPV III with either punctate or mixed signal patterns (Table 4) (Fig. 2).

**DISCUSSION**

In this study, we observed that the ISH assay using the Inform HPV III probe is comparable to the PCR assays in detecting oncogenic HPV DNA in FFPE cervical tissue from patients with CINs and can be used in cervical tissue specimens. Further, the punctate pattern of HPV is closely associated with CIN 2/3 and carcinoma. However, the use of signal patterns, especially the mixed HPV signal pattern to estimate the viral integration status, is limited because of the wide range of viral integration status in cases with mixed HPV signal patterns. Finally, we conclude that low copy numbers of episomal HPV16, as a result of high level of viral integration, may contribute in part to the false-negative results of ISH in carcinoma specimens. Based on our observation, since the punctate signal pattern is more frequent in CIN 3 and cervical carcinoma, it becomes more difficult to recognize or interpret the ISH signals, especially when these signals are weak and sporadic. In a few cases of CIN 3 and carcinoma, high-power objective lenses were required to identify the punctate signals. Other factors that may contribute to the false-negative ISH results include HPV types that were not covered by Inform HPV III probes. In our study, such cases were not observed. In addition, the heterogeneous distribution of HPV in low-grade CIN might cause signal absence in tissue sections, resulting in false-negative results.

**Comparison of Inform HPV III with PCR.** The ISH assay using the Inform HPV III probe to detect HPV DNA showed fair to good agreement with PCR assays in FFPE cervical tissue specimens with CINs. This represents a significant improvement in sensitivity for the Inform HPV III probe. We compared the Inform HPV III probe to the previous generation of Inform HPV probe, Inform HPV II, and observed significantly higher HPV-positive rates shown by Inform HPV III compared to Inform HPV II, especially in CIN 3 and carcinoma cases (data not shown). Recently, Kong et al. reported similar observations on Inform HPV II and Inform HPV III. (23).

With regard to the cause of false-negative ISH results in carcinoma specimens, it seems unlikely that the majority of such cases were due to undetected HPV types, which were not covered by the ISH probes. In our study group, all 11 cases with ISH− PCR+ results tested positive for one or more specific HPV genotypes that were actually covered by the Inform HPV III probe. Of these, five cases of cervical carcinoma were positive for HPV16. These five cases showed a significantly lower HPV16 E2/E6 ratio compared to those in ISH− cases, either with a punctate signal pattern only or with mixed signal patterns. This indicates a significantly higher proportion of integrated HPV16 in ISH− PCR+ carcinoma cases than in ISH+ cases (16). We therefore speculate that the low level of episomal HPV16, as a consequence of high viral integration status, might be associated with the false-negative results in these carcinoma cases. Based on our observation, since the punctate signal pattern is more frequent in CIN 3 and cervical carcinoma, it becomes more difficult to recognize or interpret the ISH signals, especially when these signals are weak and sporadic. In a few cases of CIN 3 and carcinoma, high-power objective lenses were required to identify the punctate signals. Other factors that may contribute to the false-negative ISH results include HPV types that were not covered by Inform HPV III probes. In our study, such cases were not observed. In addition, the heterogeneous distribution of HPV in low-grade CIN might cause signal absence in tissue sections, resulting in false-negative results.

Our study demonstrated the limitation of PCR assays for HPV DNA detection in FFPE tissue specimens, and optimizing the PCR assay to detect HPV DNA in FFPE tissue specimens is required. Limitations of PCR assays in detecting HPV DNA in FFPE tissue specimens have been observed. Even with the combination of type-specific and consensus primer sets, the

<table>
<thead>
<tr>
<th>HPV signal pattern</th>
<th>No. of cases</th>
<th>Mean E2/E6 ratio</th>
<th>SD</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative</td>
<td>5</td>
<td>0.050</td>
<td>0.101</td>
<td>0.008</td>
</tr>
<tr>
<td>Punctate</td>
<td>20</td>
<td>0.293</td>
<td>0.291</td>
<td>0.4</td>
</tr>
<tr>
<td>Punctate + diffuse</td>
<td>17</td>
<td>0.532</td>
<td>0.407</td>
<td></td>
</tr>
</tbody>
</table>

**FIG. 2.** Comparison of HPV16 integration status (E2/E6 ratio) with HPV signal patterns determined by Inform HPV III. P, punctate pattern; D, diffuse pattern.
sensitivity is limited (3). In our study, ISH⁺ PCR⁻ results were observed mainly in cases with CINs. Unger et al. reported similar findings in FFPE cervical carcinoma using ISH and PCR (34). In both studies, optimizing the PCR assay for detecting HPV DNA was attempted. In our study, all of the specimens were selected within 2 years of the study; strict measures were followed for tissue sectioning, DNA extraction, and the PCR assays to avoid cross-contaminations among specimens; and general PCR and HPV genotyping tests were randomized without the knowledge of the pathological diagnoses and independently performed using three primer sets at two centers. Repeat DNA extraction with consensus primer-mediated PCR was conducted in the specimens with ISH⁺ PCR⁻ results. Even though the PCR assays were optimized, factors that affect the efficiency of the PCR assay in detecting HPV DNA in FFPE tissue specimens cannot be completely eliminated. The ISH⁺ PCR⁻ results in our study could be partially due to the presence of PCR inhibitors for amplification (5, 34). The heterogeneous distribution of HPV in CINs, especially when the viral load is low, can result in the target molecule missing, leading to a negative result. DNA extraction can also affect PCR efficiency for HPV DNA (6, 10). In our study, the amplification was suboptimal in 8% of all initially selected cases because of no β-globin amplification. The results are similar to those in carcinoma cases in which the PCR assay was required to be optimized in tissue specimens (34). The primer sets selected for PCR assays can also affect the PCR testing results. Recently, Chan et al. compared GP5+/GP6+ and PGMY09/11, the two most commonly used primer sets for HPV DNA testing. These researchers found that although both the GP5+/GP6+ and the PGMY09/11 primer sets broadly cover most of the oncogenic HPV types, each primer set has a certain complementary coverage, with PGMY09/11 detecting more cases with multiple viral infections (7). Using GP5+/GP6+ consensus primers, we demonstrated a high HPV positivity, especially in cervical carcinoma, a finding which is consistent with the published pooled data (25). Using the second primer set, PGMY09/11, 6% more positive results were obtained, predominantly in CIN 3 and carcinoma cases (16). Four additional positive cases were identified by using the GP6+/MY11 primer sets. Using three consensus primer sets, Baay et al. reported only 72.8% positive rate of HPV DNA in carcinoma cases and an 87.6% positive rate with adding type-specific primer sets (3). Compared to the limitation of using the PCR assay in FFPE to detect HPV DNA, the application of ISH with Inform HPV III and the interpretation of the ISH results are relatively simple. With improved sensitivity for detecting HPV DNA, the ISH assay using Inform HPV III has potential use in cervical tissue specimens with CINs.

**HPV signal pattern by ISH and HPV16 integration status.**

The punctate pattern of HPV observed by ISH has been linked to the integrated HPV forms in the host genome using a Southern blot assay (9) and has been reported to be associated with CIN 2/3 or cervical carcinoma (18, 20, 22). Since HPV integration is an important step in the carcinogenesis of cervical carcinoma (29, 30), it has the potential to be used as a marker for CIN progression. In evaluating HPV signal patterns in CINs and cervical carcinoma, we found that a punctate pattern was more frequently observed in CIN 3 and cervical carcinoma, while diffuse or mixed patterns were predominantly found in CIN 1 and 2, a finding consistent with those of previous studies (18, 20, 22).

To evaluate the association between HPV signal patterns and HPV integration status, we compared HPV signal patterns with the HPV16 E2/E6 ratio using qRT-PCR. The E2/E6 ratio has been used to estimate HPV viral integration into the host genome. HPV16 integration occurs by disruption or deletion of the viral E1 and/or E2 open reading frame. Therefore, lower E2 amplification indicates a higher integrated form of HPV16. The E2/E6 ratio has been reported to be correlated well with the severity of CINs or cervical carcinoma (2, 13, 17, 28). Using qRT-PCR to measure the HPV16 E2/E6 ratio, we demonstrated that HPV16 was integrated more significantly into cervical carcinoma cells than into CIN 2/3 cells (16). In comparing the HPV16 E2/E6 ratio to the HPV signal patterns, however, we did not observe a significant difference in the E2/E6 ratio between cases with a punctate signal only and cases with mixed punctate and diffuse signals. We speculate that this could be caused by a variation in the percentage of the two patterns of HPV signals in cases with mixed signal patterns; this makes quantitative analysis difficult and therefore affects an accurate estimation. The presence of episomal forms of HPV DNA or RNA also has been reported to mask the integrated HPV DNA when the amount of the integrated viral forms is small (18), causing an inaccurate estimate of the level of integrated HPV. Furthermore, the heterogeneous distribution of HPV-positive cells shown by ISH assay in tissue sections may cause inaccurate classification. Although our observation supports the notion that the pure punctate pattern of HPV indicates a high level of viral integration and may be a marker for CIN progression, the level of HPV integration cannot be accurately determined in the cases with mixed signal patterns that encompass a wide range in viral integration status. Recently, Fijii et al. reported similar findings in their comparison of HPV signal patterns shown by ISH with the integrated HPV DNA when the amount of the integrated viral forms is small (18), causing an inaccurate estimate of the level of integrated HPV. Furthermore, the heterogeneous distribution of HPV-positive cells shown by ISH assay in tissue sections may cause inaccurate classification.

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