Detection of high risk human papillomavirus in routine cervical smears: Strategy for screening

C S Herrington, M de Angelis, M F Evans, G Troncone, J O’D McGee

Abstract

**Aim:** To develop a methodology for direct detection of high risk human papillomavirus (HPV) infection in routine cervical smears by non-isotopic in situ hybridisation (NISH) which can be compared with cytopathological assessment of the same cells.

**Methods:** The methodology was established using cultured cells and routine cervical smears hybridised with digoxigenin labelled probes for HPV, 16, 18, 31, and 33. The technique was applied to the analysis of 53 patients from a sexually transmitted disease clinic.

**Results:** The optimal sensitivity achieved for single HPV detection in cultured cells was 1–2 copies of HPV 16 per cell and that for detection of a cocktail of HPV types in routine cervical smears was 2–5–12 copies per cell. Of parallel smears taken from patients with a normal Papinacolau-stained smear 33–3% (24) contained a HPV 16, 18, 31, and 33 signal indicating an occult HPV infection. The prevalence of these HPV types was similar in women in whom a cytopathological diagnosis of wart virus infection was made (64–77%, 17) and in patients with mild dyskaryosis (75%, 12).

**Conclusions:** The methodology evolved to localise HPV sequences directly to epithelial cell nuclei, which can be morphologically assessed by haematoxylin counterstaining. Sample contamination with exogenous viral sequences can be distinguished from true infection. In this study, a HPV signal was not found in morphologically normal epithelial cells. The methods described will permit the detection of HPV sequences in routinely collected cervical smears and the evaluation of the natural history and potential clinical relevance of HPV infection without changes in clinical practice.

The experimental evidence linking human papillomaviruses (HPV) with cervical neoplasia is strong. Clinically, certain HPV types, particularly HPV6 and 11, have been associated with low grade clinical lesions and a negligible risk of progression to invasive malignancy. Conversely, high grade clinical lesions are associated with a different spectrum of HPV types, particularly HPV16, 18, and related viral types: these types are an established risk factor for clinical progression. Recently the association between HPV and cervical neoplasia has been questioned because HPV sequences are demonstrable using the polymerase chain reaction (PCR) in cervical samples from a high proportion of normal women. However, interpretation of the data has been problematic due to inconsistencies in methodology and results from different laboratories. Estimates of the prevalence of HPV infection in both normal and abnormal cervical epithelial cells have varied widely. The possibility of contamination with exogenous HPV sequences is difficult to exclude when methods are used which require destruction of cellular morphology. In situ hybridisation has the advantage that cellular localisation of sequences can be performed microscopically and it excludes the possibility of contamination. Isotopic detection of HPV in cervical smears has been previously described and non-isotopic in situ hybridisation (NISH) has been performed on cell samples collected by cervicovaginal lavage. These methods are not, however, applicable to cervical smears collected in the routine way.

**Methods**

**PROBES**

Genomic probes for HPV 6, 11, 16, 18, 31, and 33 were labelled with biotin and digoxigenin by nick translation as previously described. Total human DNA was labelled with digoxigenin and used as a positive control.

**COLLECTION OF SPECIMENS**

Samples were obtained from the sexually transmitted diseases (STD) clinic in Oxford. Two standard smears were taken: one for cytopathological diagnosis and the other for NISH analysis. Smears were fixed in 70% ethanol/Carbowax in the conventional way. Cytopathological assessment was performed independently of NISH on a Papinacolau-stained smear. Conventional criteria for the diagnosis of wart virus infection and cervical intraepithelial neoplasia were employed.

** Pretreatment of cervical smears**

Routine smears were incubated in methanol/acetic acid (3:1, v/v) for 10 minutes at 22°C and then fixed in fresh 4% paraformaldehyde (w/v) in phosphate buffered saline (PBS: 10 mM phosphate, 150 mM NaCl, pH 7-4) for 15 minutes at 22°C. After washing in PBS containing 0-2% (w/v) glycine, smears were rinsed in PBS and endogenous peroxidase
activity blocked with 0.1% (w/v) sodium azide containing 0.3% hydrogen peroxide (v/v) for 10 minutes. After washing in PBS, smears were incubated in 1 μg/ml proteinase K (Boehringer, Germany) in PBS for 15 minutes at 37°C. Washed in PBS, postfixed in 4% paraformaldehyde for five minutes, washed in PBS/glycine, PBS, and air dried at 37°C. Smears prepared in this way were processed the same day.

CaSki, HeLa 229, and SiHa cells
Cells were obtained from the ATCC (USA) and RPMI 1640 medium supplemented with 10% fetal calf serum (FCS) and L-glutamine. All three cell lines were grown to confluence and cells harvested by treatment with trypsin-EDTA solution (Gibco, UK). After washing in PBS, cells were fixed in suspension in methanol/acetic acid (3 in 1, v/v) and stored at −20°C. When required, these cells were centrifuged at 600 × g for five minutes, resuspended in a small volume of fresh cold methanol/acetic acid, spotted onto four spot multiwell slides (Hendley, Essex), dried at 37°C and treated as for routine smears. The peroxidase blocking step is not required for analysis of epithelial cell lines.

NON-ISOTOPIC IN SITU HYBRIDISATION (NISH) HPV6, 11, 16, 18, 31, and 33 probes labelled with either biotin or digoxigenin were hybridised to cervical smears pretreated as above. Cocktails of these genomic probes were used for the analysis of routine cervical smears (see Results). Cells of CaSki, SiHa (which contain integrated HPV16) and HeLa cells (which contain integrated HPV18) were hybridised with HPV16 and 18 individually. The hybridisation cocktail contained the following: 50% formamide, 5% dextran sulphate, 300 mM NaCl, 30 mM sodium citrate, 50 mM Tris HCl, pH 7.4, 0.1% (w/v) sodium pyrophosphate, 0.2% (w/v) polyvinylpyrrolidone (molecular weight 40 000), 0.2% (w/v) Ficoll (molecular weight 400 000), 5 mM EDTA, 200 ng/μl sheared human DNA and 2 ng/μl of each labelled probe. An aliquot of hybridisation cocktail (50 μl) was added to each smear and covered with a 22 × 50 mm coverslip. Cells and probes were denatured simultaneously in moist Terasaki plates at 95°C for 15 minutes and hybridised at 42°C for two hours. After hybridisation, slides were washed in 4 × (SSC) standard saline citrate twice at 22°C, then 50 mM TRIS-HCl, pH 7.2, 100 mM NaCl (TBS) containing 3% (w/v) bovine serum albumin and 0.05% (v/v) Triton X-100 (TBT) for 10 minutes.

PROBE DETECTION
All antibody/avidin incubations were carried out at room temperature for 30 minutes unless otherwise stated.

Biotinylated probes
Smears were incubated in either avidin alkaline phosphatase (Dako, UK) diluted 1 in 50 or avidin peroxidase (Dako, UK) diluted 1 in 75 in TBT containing 5% (w/v) non-fat milk (Cadbury, UK). After washing for 10 minutes in TBS, the signal was developed using either nitroblue tetrazolium (NBT)/5-bromo-4-chloro-3-indoly phosphate (BCIP) for alkaline phosphatase detection or aminoethylcarbazole (AEC; Zyomed, USA) for peroxidase detection.19

Digoxigenin labelled probes
The detection procedures consisted of three or five steps.

Three step detection Smears were incubated in mononclonal anti-digoxin (Sigma, UK) diluted 1 in 10 000 in TBT followed by biotinylated rabbit anti mouse [F(ab)2 fragment] diluted 1 in 200 in TBT and then by avidin peroxidase diluted 1 in 75 in TBT containing 5% (w/v) non-fat milk. The signal was developed using AEC as described."
Figure 1 HPV16 detection in SiHa cells. Two discrete foci of signal of similar morphology can be seen within a single nucleus.

which is known to occur in this cell line. Other nuclei in the preparations contained single signals (data not shown). No signal was obtained after hybridisation with HPV18. The three step detection system gave no signal with the HPV16 probe on SiHa cells. Both the three and five step procedures give high resolution of nuclear detail and allow precise intranuclear localisation of signal in haematoxylin counterstained cells.

The relationship between sensitivity of detection and applied probe concentration was investigated to assess the possibility of reducing individual probe concentrations when probe cocktails were used in the analysis of cervical smears (see below). Hybridisation of CaSki cells with digoxigenin labelled HPV16 followed by detection using the three step procedure with AEC as substrate was carried out at probe concentrations of 0-1, 0-5, 1, 2, 4, and 8 ng/μl. Reduction in probe concentration below 2 ng/μl led to a reduction in sensitivity as judged by the median number of signals obtained per CaSki cells nucleus (table 1).

CERVICAL SMEARS
Methodological development
Initially, the single step detection methods were employed on routine cervical smears to evaluate the pretreatment and hybridisation methods. By using total human DNA as a probe, the effect of aldehyde fixation as assessed: as was noted previously for HPV16 detection in CaSki cells, the signal obtained in routine smears was considerably stronger in aldehyde fixed smears than in those fixed only in methanol/acetic acid (data not shown). Aldehyde fixed smears were therefore used in the remainder of the study and hybridised with a cocktail of HPV16, 18, 31, and 33 probes.

The main initial technical problem in the analysis of routine cervical smears hybridised with a cocktail of biotinylated HPV probes was intense background staining (fig 2A). By incorporating large molecular weight polymers in the hybridisation cocktail, this background staining was reduced but not abolished. For single nucleic acid detection, digoxigenin labelled probes gave lower background staining than biotinylated probes and peroxidase based systems produced cleaner results than alkaline phosphatase. However, the single most important component was the inhibition of non-specific avidin binding by the use of non-fat milk (table 2, fig 2). The cellular morphology is clarified by haematoxylin counterstaining (fig 2B). Further amplification using the five step method gave unacceptable background staining despite the incorporation of all blocking steps in table 2 (data not shown). This background was reduced by lowering the concentration of each probe in the hybridisation reaction. However, the sensitivity of probe detection is dependent on probe concentration (see above) and hence reduction of the probe concentration below 2 ng/μl negates the advantage of using the five step system.

The optimum system for HPV detection in routine cervical smears is therefore the three step peroxidase based detection of digoxigenin labelled probes with haematoxylin counterstaining (fig 2B).

Application to routine cervical smears
Having established the most appropriate method, this technique was applied to 53 cervical smears collected in the routine way from women attending a sexually transmitted disease clinic. The smears used for NISH analysis were collected by a separate pass of the spatula. The cytopathological diagnoses made on the Papinacolau stained smears were compared with the results of NISH analysis (table 3, fig 3). All epithelial cells which contained a NISH signal demonstrated at least minor morphological abnormality (see Discussion). The positive and negative controls incorporated in each batch gave the appropriate result—that is multiple nuclear HPV16 signals were obtained in CaSki cells and every nucleus in the smear was stained after hybridisation with a probe for total human DNA.

Discussion
The object of this study was to develop methods of high sensitivity and resolution for the detection of human papillomavirus DNA sequences in routine cervical smears and to apply them to the analysis of a group of women attending a sexually transmitted disease clinic. HPV6/11 infection has been associated with low grade, and HPV16/18 and related viral types with high grade, squamous cervical lesions. It is therefore of potential clinical importance to determine by molecular means the viral type present in cervical epithelial cells. Although this can be performed by a variety of methods involving the extraction of DNA from

Table 1 Sensitivity of HPV16 detection in CaSki cells

<table>
<thead>
<tr>
<th>Applied probe concentration (ng/μl)</th>
<th>Median signal</th>
<th>No</th>
<th>Range</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-1</td>
<td>0</td>
<td>1</td>
<td>(0-7)</td>
<td>123</td>
</tr>
<tr>
<td>0-5</td>
<td>1</td>
<td>6</td>
<td>(2-11)</td>
<td>111</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>11</td>
<td>(3-20)</td>
<td>107</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>10</td>
<td>(3-20)</td>
<td>106</td>
</tr>
<tr>
<td>4</td>
<td>1</td>
<td>11</td>
<td>(5-20)</td>
<td>104</td>
</tr>
</tbody>
</table>
Figure 2  HPV detection in routine cervical smears. (A) No blocking procedures were used and a biotinylated cocktail of HPV probes was detected using avidin alkaline phosphatase and NBT/BCIP. All cellular and non-cellular components of the smear are labelled, typical of biotinylated probes in the absence of blocking agents. (B) All blocking procedures were used and a digoxigenin labelled HPV probe cocktail detected using the three step peroxidase AEC method. A clear nuclear red signal is coupled with good morphological definition. An intranuclear red signal is present in several abnormal nuclei.

Table 2  Reduction of non-specific staining in routine cervical smears

<table>
<thead>
<tr>
<th>Hybridization solution</th>
<th>Detection procedure</th>
<th>Block non-specific avidin binding using modified avidin in non-fat milk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Digoxigenin labelled probes</td>
<td>Block non-specific avidin binding using modified avidin in non-fat milk</td>
<td>Use peroxidase based system rather than alkaline phosphatase</td>
</tr>
</tbody>
</table>

Fig 2 compares the results with biotin and digoxigenin labelled probes with and without the blocking procedures.

exfoliated cervical cells (Southern blotting, 24 filter in situ hybridisation; PCR amplification), only in situ hybridisation can demonstrate HPV sequences in intact nuclei. Moreover, as cervical smears taken by general practitioners are used as the means of cervical screening in the United Kingdom, the ability to analyse these samples directly would not require changes in clinical practice.

Table 3  Comparison of the detection of HPV16, 18, 31, and 33 by NISH in routine cervical smears with cytopathological diagnosis

<table>
<thead>
<tr>
<th>Cytopathological diagnosis</th>
<th>NISH n = 24</th>
<th>Wart virus infection n = 17</th>
<th>CIN + WV1 n = 12</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative</td>
<td>16 (66.7%)</td>
<td>6 (35.3%)</td>
<td>6 (25%)</td>
</tr>
<tr>
<td>Positive</td>
<td>8 (33.3%)</td>
<td>11 (64.7%)</td>
<td>9 (75%)</td>
</tr>
</tbody>
</table>

CIN, cervical intraepithelial neoplasia.

The detection of HPV DNA in routine cervical smears poses several problems. Firstly, only one smear is taken and this is required for cytopathological diagnosis. It is thus impossible to perform controls on material from the same patient. However, the use of HPV containing cell lines and hybridisation of cervical smears with a probe to total human DNA allowed assessment of each experiment. Secondly, cervical smears contain a variable amount of mucin and may be contaminated with bacteria, fungi, and protozoa. This may lead to technical problems, particularly high background staining. As cervical smears are collected from patients randomly with respect to their menstrual cycle and by different practitioners with consequent variation in smear quality and fixation, it was necessary to develop a method with the following properties: minimal loss of material, no reliance on quality of initial fixation, low background staining, and inhibition of endogenous neutrophil peroxidase. Optimisation of fixation and unmasking conditions showed that aldehyde fixation followed by limited protease K digestion produced superior results to direct analysis of ethanol fixed smears using digoxigenin labelled total human DNA as probe. This is consistent with results obtained using CaSkI cells. 19 The combination of sodium azide and hydrogen peroxide prior to nucleic acid unmasking produced effective inhibition of endogenous peroxidase, which is particularly important in the analysis of smears containing large numbers of inflammatory cells. Postfixation of smears in aldehyde minimised the loss of cellular material, particularly as routine smears are not collected on silane-coated slides. 25 Hybridisation in the presence of excess human DNA, polyvinylpyrrolidone, and Ficoll reduced non-specific probe binding to non-cellular components of the smear. High background staining due to deposition of NBT/BCIP derived product over mucin rich areas was inhibited by blocking non-specific avidin binding by both the use of modified avidin and incubation in non-fat milk. The most effective individual step in the reduction of background staining was incubation in the avidin conjugate in the presence of non-fat milk. This suggests that the majority of the background staining observed was due to non-specific binding of avidin to non-cellular components of the smear.

For routine nucleic acid detection, the most effective protocols were those involving peroxidase based detection of digoxigenin labelled probes. Amplification of the detection
of these probes enabled HPV16 to be detected in SiHa cells. However, application of this system to the detection of probe cocktails in routine cervical smears produced unacceptable and variable background staining, due to the high total probe concentration. However, reduction in probe concentration leads to a marked reduction in sensitivity, which is predictable as the in situ hybridisation reaction is carried out with excess probe. The three step digoxigenin detection system was the most routinely applicable, giving the most reproducible signal with high sensitivity (approximately 2.5–12 HPV copies per cell),16 high resolution, and low background staining. The combination of this protocol with haematoxylin counterstaining provides good morphological contrast and is therefore of potential routine use.

This methodology was applied to cervical smears collected from a sexually transmitted disease clinic as there was likely to be a high prevalence of HPV infection amongst these patients. This analysis showed the presence of "high risk" sequences (HPV16, 18, 31, and 33) in 33.3% of cases (24) in which no cytological abnormality was demonstrated on a parallel Papinacolau-stained smear. One possible explanation for these findings is that the cervical sampling of the two smears differs, with the second smear containing cells shed from an abnormal area not sampled in the first. This situation is known to arise with moderate frequency.26 Careful analysis of the morphological features in these cases showed that the positive cells were not morphologically normal, although the abnormalities present were not sufficient to warrant a diagnosis of wart virus infection or dyskaryosis (fig 3). These would be classified as borderline nuclear abnormalities or inflammatory changes by routine cytological criteria. In addition, many positive cells were present in large clusters of epithelial cells where morphological assessment is difficult and any abnormalities present would not necessarily have been recognised on routine Papinacolau screening. No evidence was found for the presence of HPV sequences in morphologically normal epithelial cells and, if this occurs, it is likely to be an uncommon event. In this study, the presence of HPV sequences was always associated with at least minor morphological abnormality. This does not exclude the possibility of HPV infection of morphologically normal epithelial cells, as the sensitivity of the technique is not sufficient to detect fewer than 2.5–12 copies of HPV per cell, but these observations are consistent with the fact that HPV sequences have never been shown in normal epithelial cells in cervical biopsies by in situ hybridisation.17 27 Moreover, a study in which HPV sequences were detected by PCR amplification and compared with the morphology of parallel sections showed that there was absolute concordance between morphological abnormality and the presence of HPV; no HPV sequences were shown in normal cervical epithelium, even directly adjacent to a focus of HPV infection.28 However, irrespective of the reason for the discrepancy between cytopathology and NISH, HPV infection was shown in one third of patients with no abnormality on routine cervical screening. These data concur with those derived from a separate patient cohort from the same clinic, analysed by dual in situ hybridisation, in which 25% of parallel smears (n = 128) from patients with a Papinacolau smear showing no abnormality contained high risk HPV sequences.29 The technique used in the latter study had a lower sensitivity (approximately 40–50 copies per cell) than that used here. The possibility that the high prevalence of HPV in women with parallel Papinacolau stained smears showing no abnormality is due at least in part to sampling error may explain the wide variation in estimates of prevalence of HPV sequences in the normal population.4–12 The application of this technique to a large unselected population of patients will permit assessment of the true prevalence of HPV infection of normal epithelial cells and its role in the natural history of HPV infection.

The demonstration of HPV sequences by molecular techniques in the absence of clinical or cytopathological abnormality has been defined clinically as latent HPV infection.3 Since all HPV positive cells in this study showed minor morphological abnormalities and HPV sequences were not found in morphologically normal cells, it would be wise to use the term occult rather than latent HPV infection: the term latency has a different biological implication.

The analysis of smears with morphological evidence of wart virus infection showed the presence of high risk HPV sequences in morphologically abnormal cells in 64.7% of cases (17). The high prevalence of cases infected with high risk types is at variance with the commonly held view that morphological evidence of wart virus infection is indicative of HPV6, 11—that is low risk infection. These data agree with those from a study of a separate cohort of patients from the same clinic in which 58% of cases (50) were infected with high risk


25. CSH held a Cancer Research Campaign (UK) Clinical Research Fellowship and is a Junior Research Fellow, Green College, Oxford. This work was supported by grants to J O'D McGee from the Cancer Research Campaign (UK). We are grateful to Drs G Orth and A Lorincz for the HPV31 and 33 probes.


