PCR in situ hybridisation detection of HPV 16 in fixed CaSki and fixed SiHa cell lines

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Abstract

Aims—To investigate the feasibility of using fixed cells with the polymerase chain reaction (PCR) in situ hybridisation and to investigate possible reasons for reaction failure.

Methods—Fixed SiHa and CaSki cells were used in an experimental model of PCR in situ hybridisation for the detection of low and intermediate copy number viral infection in fixed cells.

Results—PCR in situ hybridisation was able to detect one to two copies of human papillomavirus (HPV) 16 in SiHa cells, using small fragment amplicons (120 base pairs), confirming the high detection sensitivity and flexibility of the technique. Problems were encountered with localisation of PCR amplified product in CaSki cells (200-300 copies of HPV 16 per cell) owing to diffusion of product post amplification. Overall, 40% of reactions were successful, which confirms the current unreliability of the technique. Within cell preparations, about 50% of cells contained amplified product.

Conclusion—PCR in situ hybridisation represents the marriage of two revolutionary molecular pathological techniques. However, it is currently unreliable, with reaction failure common. Standardised, dedicated equipment is urgently required if the technique is to achieve universal acceptance. In the future, the technique may be used to detect chromosomal translocations in human tumours and to study cellular gene expression.

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The polymerase chain reaction (PCR) permits the selective in vitro amplification of a particular region of nucleic acid by mimicking the phenomena of in vivo DNA replication. PCR has been extensively used to amplify specific DNA sequences for use in the molecular analysis of many diseases, for which formalin fixed, paraffin wax embedded tissue has been used. However, one limitation for the histopathologist was that it was impossible to localise amplified DNA sequences in cells or tissue sections. Recently, several studies have described a new technique which combines PCR with in situ hybridisation, permitting the localisation of specific amplified DNA signals within isolated cells and tissue sections. This technique has been variously known as in situ PCR or PCR in situ hybridisation.

In situ PCR refers to use of a biotin or digoxigenin labelled nucleotide directly in the PCR mix and subsequent detection of the labelled amplified product using standard in situ detection protocols. PCR in situ hybridisation occurs when PCR amplification of a tissue section or cell suspension is carried out, followed by application of a biotin or digoxigenin labelled probe to the amplified product and detection of the hybrid by standard in situ detection protocols. The technique was first described in 1990 by Haase et al., who amplified lentiviral DNA in infected cells and subsequently detected the amplified DNA using in situ hybridisation. Nuovo et al further modified and developed this technique for the identification of different types of human papillomavirus (HPV) in formalin fixed, paraffin wax embedded samples.

Why should this technique work? Many people question the scientific basis of the in situ PCR process, with many centres reporting failure of the technique. Theoretically, fixed cells should act like sponges or at least semipermeable dialysis bags. It is reasonable, therefore, to assume that PCR reagents (Taq, primers, etc.) can diffuse through the cell membrane and into the cytosolic and nuclear components of the cell.

Preliminary results from Nuovo et al using the technique, suggested that most amplified product appeared to localise, although “leakage did occur into the surrounding medium.” The technique can be conveniently performed on a normal conventional thermal cycling block, on specially designed thermal cyclers for slide PCR, or in a microprocessor controlled oven.

In this paper we examine PCR in situ hybridisation in relation to the detection of low and intermediate copy HPV infection in fixed SiHa and CaSki cell lines and address some of the fundamental technical problems encountered when using the technique for the identification of viral pathogens in histopathological material.

Methods

We evaluated the use of PCR in situ hybridisation with paraffin wax embedded cell lines for the detection of low and intermediate copy number HPV infection in virally infected SiHa and CaSki cells. HPV 16 was amplified using specific primers which amplify a 120 base pair fragment of HPV 16. Primers for
**Primer sequences used**

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
<th>Genome position</th>
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<tbody>
<tr>
<td>HPV 6B</td>
<td>Primer 1</td>
<td>CCTGTTCAGGGCCTCAGCTTTTGAAGCAACCGGCGCTTGTT</td>
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<tr>
<td>Primer 2</td>
<td>GTGAAATTATCGTGGTTCGTT</td>
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<td>Primer 1</td>
<td>TCCAAAAGGCACTGTTCCTG</td>
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<tr>
<td></td>
<td>Primer 2</td>
<td>GCTGTTCCTGTATGCTGCAA</td>
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<td>Primer 1</td>
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<td></td>
<td>Primer 2</td>
<td>ATGGCAGTCGCTTATATG</td>
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<td>HPV 33</td>
<td>Primer 1</td>
<td>AAGATTTAAAAACCTTTAAA</td>
</tr>
<tr>
<td></td>
<td>Primer 2</td>
<td>AGTTTTCTACTGCTGGAGCTC</td>
</tr>
</tbody>
</table>

Sequences obtained from EMBL database.

HPV 6, 18, and 33 were also tested on the cell lines to check the specificity of the reaction. Primer sequences are given in the table.

Parallel solution phase PCR with (i) β globin gene (to check amplifiability of the cells) and (ii) "No Taq DNA polymerase" was carried out in addition to PCR runs with unrelated primers to HPV 6, 18, and 33, to check the specificity of the reaction.

Cells were obtained from the European Collection of Animal Cell Cultures (ECACC) or American Type Culture Collection (ATCC) (USA) and grown in RPMI 1640 medium supplemented with 10% fetal calf serum and L-glutamine. The cell lines were grown to complete confluence and then harvested.

The cells were removed from the flask by digestion with trypsin (0.2% trypsin in Hank’s balanced salt solution), and collected in a fal- con tube. The cells were then pelleted by centrifugation for five minutes at 3000 rpm. Washing in phosphate buffered saline (PBS) was then performed and the cell pellet was resuspended in neutral buffered formalin, mixed well, and stored for the required fixation time. (Fixation time of 24–48 hours was usually found to be adequate and provided good penetration of the cell pellet.) After fixation the cells were centrifuged at 3000 rpm for five minutes and then resuspended in "deionized water" (50% neutral buffered formaldehyde and 50% sterile distilled water) and transferred to a 10–15 ml glass tube. The cells were washed with water and graded alcohols to 100% alcohol. The cell pellet was suspended in xylene, mixed well (end over rotation), and centrifuged at 3000 rpm for five minutes.

Xylene was discarded carefully. Paraffin wax at 65°C was carefully added and the cells resuspended overnight at 65°C. The glass tube containing the cells was then placed at –20°C for three to four hours, removed, and the glass broken with a hammer and the cell pellet retrieved. The cell pellet was re-embedded in a tissue cassette, applied to a microtome support, and 5 μm sections cut.

Glass single well and multi-well slides (PH106, PH005, Hendley Essex) were coated with 2% aminopropyltriethoxysilane, as described before for hybridisation.4

Sections (5 μm) were cut on the “welled” slides and placed on a hot plate for 24 to 48 hours to achieve maximum section adhesion. Section dewaxing was carried out by immersing the slides in xylene at 37°C, for 30 minutes, followed by xylene at room temperature for 10 minutes, and absolute alcohol at room temperature for a further 10 minutes. The slides were transferred to fresh absolute alcohol and rehydrated through a graded alcohol series to water over a 10 minute period. The prepared sections were immersed in 0.02M HCl for 10 minutes, washed twice in PBS for five minutes each, and treated with 0.01% Triton X-100 in PBS for three minutes. After this, the sections were washed twice in PBS for five minutes each and then incubated in prewarmed proteinase K buffer (50 mM TRIS-HCl, pH 7.6, 5 mM EDTA) at 37°C for 10 minutes and then treated with proteinase K (0.1–0.5 mg/ml) at 37°C for 10 to 20 minutes. The slides were washed in two changes of PBS containing 2 mg/ml of glycine for five minutes each, followed by immersion in aqueous 20% acetic acid at 4°C for 15 seconds. The sections were then washed in two changes of PBS for 10 minutes each and post-fixed in 4% parafomaldehyde in PBS for five minutes. Washing in PBS for five minutes was then performed, and the sections were dehydrated through graded alcohols.

For amplification of paraffin wax embedded cell lines, the slide containing the fixed cell suspension was placed in an aluminium foil boat, trimmed to slightly larger proportions than the slide. PCR solution (20 μl) consisting of PCR buffer (50 mM KCl, 10 mM TRIS-HCl (pH 8.3) at room temperature, 4.5 mM MgCl2, 0.01% gelatin) and 200 mM each of dNTP and 5 μM primers were placed on top of the cell suspension. The well was covered with a pre-cut piece of Gel Bond (FM Bioproducts), hydrophobic side down. The slide (in the aluminium foil boat) was placed on the heating block of the thermal cycler and the temperature allowed to increase to 80°C. Once the temperature had reached 80°C, a corner of the gel bond was lifted and 2.5 μl Taq DNA polymerase (2.5 units per 12.5 μl final concentration) were added to the PCR mix contained on the slide. The coverslip (gel bond) was then replaced and the margins sealed with nail polish. Preheated mineral oil (1–2 ml) (80°C) was placed on top of the slide to ensure optimal thermal kinetics. The following PCR protocol was applied: 94°C for six minutes, 40 cycles of 55°C for two minutes, and 94°C for one minute.

The temperatures given refer to the temperatures of the glass slide. This was checked using a thermocouple device. It is important that the initial denaturation temperature and cycling temperatures are achieved on the glass slide. After amplification the slide was dipped in chloroform to remove mineral oil and the gel bond coverslip carefully removed. Gel Bond, hydrophobic side towards the cell sus- pension, allows the PCR reagent mix to remain on the tissue in the well. After amplification the slides were dipped carefully in 100% alcohol and dehydrated. Post fixation in 2% paraformaldehyde to maintain localisation of PCR product was also carried out.

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Biotinylated and digoxigenin labelled genomic HPV 16 probe contained in pBR322 (a gift from Dr Zur Hausen) was prepared using the nick translation labelling technique. The HPV 16 probe (75 μl) (200 ng/ml) in hybridisation buffer (50% dextran sulphate, 2 × SSC, 0.2% dried milk powder, 50% formamide) was applied to the centre of the well of the PH106, C A Hendley glass slide (SSC = sodium chloride and sodium citrate). For multiwell slides (PH005, C A Hendley), 8–10 μl of probe was applied to the centre of the well. Gel bond films were cut to coverslip size and placed hydrophobic side down, over each section. The gel bond was sealed in place with nail varnish.

Slides were placed onto a preheated baking tray and incubated at 90–95°C for 10 minutes. Finally, the slides were transferred to a humidified box and incubated at 37–42°C overnight or at 42°C for two hours.

For posthybridisation washes, the gel bond coverslips were removed with a scalpel blade and the slides washed in SSC with agitation according to the following protocol:

(a) 2 × SSC at room temperature for 10 minutes;
(b) 2 × SSC at 60°C for 20 min;
(c) 0.2 × SSC at room temperature for 10 min;
(d) 0.2 × SSC at 42°C for two minutes;
(e) 0.1 × SSC at room temperature for 10 min;
(f) 2 × SSC at room temperature, for one to two minutes.

The following solutions were required for hybridisation detection protocols:
- Buffer 1 (0.1M TRIS-HCl (pH 7.5), 0.1M NaCl, 2 mM MgCl₂, and 0.05% Triton-X-100).
- Buffer 2 (0.1M TRIS-HCl (pH 9.5), 0.1M NaCl, 50 mM MgCl₂).

Development reagent (NBT/BCIP)—alkaline phosphate detection system. A fresh solution was made just before use and kept in the dark. This consisted of buffer 2 (3.75 ml), nitroblue tetrazolium (NBT) (16.5 μl), bromochloroindoly phosphate (BCIP) (12.5 μl), TBS (TRIS-buffered saline) (50 mM TRIS-HCl, 100 mM NaCl, pH 7.2), and TBT (blocking reagent) for the detection of digoxigenin labelled probes (50 mM TRIS-HCl, 100 mM NaCl (pH 7.2), 3% (w/v) bovine serum albumin (BSA) and 0.5% Triton X-100 (v/v)).

For the detection of biotinylated probes, the slides from the posthybridisation wash steps were transferred into buffer 1 containing 5% (w/v) BSA and incubated at room temperature for a minimum of 30 minutes. Excess buffer was wiped from the slides and the slides transferred to a slide incubation tray. A few drops of buffer 1 containing Avidin D (Vector) at a concentration of 10 μl/ml were added and the slides incubated at room temperature for 10 minutes. The slides were washed with agitation in two changes of buffer 1 for 10 minutes each. The slides were then

Figure 1  PCR in situ hybridisation of fixed SiHa cells showing two copies of HPV16 per cell nucleus using a biotin labelled probe.

Figure 2A  PCR in situ hybridisation of fixed SiHa cells showing two copies of HPV16 per cell nucleus in two adjacent SiHa cells, using a biotin labelled probe.

Figure 2B  PCR in situ hybridisation of fixed SiHa cells showing one copy of HPV16 per cell nucleus, using a digoxigenin labelled probe.
A polymerase.

Figure 4A Fixed cells subjected to DNA amplification protocol omitting Taq DNA polymerase. No signal is visible.

Figure 4B Fixed SiHa cells subjected to DNA amplification using unrelated primers (HPV 18).

Figure 5 PCR in situ hybridisation of fixed SiHa cells showing diffuse staining, with occasional cells showing dot positivity. Sometimes, interpretation of diffuse staining may be difficult and may simply reflect background staining due to detection reagents, etc.

returned to the incubation tray and a few drops of buffer 1 containing biotinylated alkaline phosphatase (Vector) at a concentration of 10 μl/ml were added to each section. Incubation at room temperature for 10 minutes was carried out. The slides were washed with agitation twice in buffer 1 for 10 minutes each. Transfer of slides into buffer 2 then took place and they were allowed to equilibrate for 30 minutes. The slides were returned to the incubation tray and the sections were covered with NBT/BCIP development reagent. Colour development was monitored after five minutes and then continuously until development looked complete. The reaction was terminated by immersion of slides in PBS or distilled water for five minutes.

For detection of digoxigenin labelled probe, the slides from the posthybridisation washes were immersed in TBT (blocking reagent) at 22°C for 10 minutes. The slides were transferred to a slide incubation tray and incubated in alkaline phosphatase conjugated antidigoxigenin (Boehringer, Germany) diluted in 1 in 600 in TBT. After this the slides were washed in TBS for five minutes, twice. Signal development using NBT/BCIP development reagent was carried out for 10–30 minutes. The colour development reaction was terminated by washing in distilled water for five minutes.

Results

Figures 1, 2A, 2B, and 3 show typical results obtained with the SiHa cell line using PCR in situ hybridisation. SiHa cells contain one to two copies of HPV 16 per nucleus and these cannot be routinely detected by non-isotopic in situ hybridisation (NISH) except for complex five-step detection procedures. The result is morphologically similar to that obtained by Herrington et al using NISH and a five-step detection protocol for digoxigenin labelled genomic probes. When Taq DNA polymerase was omitted from the reaction protocol, amplification was not achieved (fig 4A). Unrelated primers also did not yield amplification (fig 4B).

In this study patchy amplification was seen, and reproducibility of results was not always achieved. On average, 50% of cells on most slides contained amplified product, with about 40% of PCR in situ hybridisation rea-
pathogens in fixed cells. We do not purport to have examined human structural genes. We were able successfully to amplify one to two copies of HPV 16 in fixed SiHa cells, which are not normally detectable with standard non-isotopic in situ hybridisation, except for complicated five-step sandwich techniques,

as mentioned earlier. Discrete and diffuse signals were seen within the nucleoplasm of the fixed SiHa cells (figs 1, 2A, 2B, and 3). Fixation of cells for up to 24 hours appeared to yield material suitable for use with PCR in situ hybridisation. The results confirm the sensitivity of the technique for the detection of low copy number viral infection in infected cells. Fixed CaSki cells were also amplifiable after 12 to 24 hours of fixation. Problems with localisation of signal were commonly seen when using CaSki cells (fig 6). The reason for this is that the starting copy number of HPV in CaSki cells is much higher than in SiHa cells (200–300 copies compared with one to two copies). Such diffusion of product makes cytological and histological interpretation difficult. To overcome such diffusion problems, we advocate the use of paraformaldehyde and 100% alcohol dehydration after amplification. We have also clearly shown in this study that small amplicon PCR in situ hybridisation (120 base pairs) can be carried out successfully on fixed tissues using a single primer pair and the "hot start" PCR modification. In both experimental systems not all cells contained amplifiable product. This was primarily due to leakage of amplified product from truncated cells which have been cut by the microtome blade during sectioning of cells for the reaction—very much like a leaky bucket.

The success rate of the reaction (40%) was low, with failure and patchy amplification common. This is a poor success rate when compared with conventional in situ hybridisation. The choice of reporter molecule (biotin/digoxigenin) did not seem to affect the success rate of the reaction with equally sensitive results obtained with both reporter molecules.

Many centres have experienced consistent failure using PCR in situ hybridisation or in situ PCR. The following points should therefore be considered:

1. The cytoskeleton of the cell should be made rigid before PCR in situ hybridisation, to prevent cytoskeletal collapse during amplification. Cells or tissues should be fixed with formaldehyde. However, fixation of cells and tissues influences solution phase PCR and also seems to influence the results obtained with in situ PCR or PCR in situ hybridisation.

2. The most important feature of formaldehyde fixation in relation to in situ PCR techniques is its ability to cross-link histone proteins that normally coat DNA, subsequently effecting the progression of Taq DNA polymerase along native DNA template. In solution phase PCR attempts to override cross-linking using extensive digestion steps are made. However, in PCR in situ hybridisation extensive proteolytic digestion is not possible (in most cases 10 to 15 minutes, as in this study), so full dissociation of histone pro-
some parts of the tissue section or cell suspension. This may be due to inadequate digestion during pretreatment, lingering DNA cross-linking phenomena (because of prior fixation of cells), and variations in the volume of PCR reactants over the slide during the reaction.

7 The thermal profile of the reaction can differ at several sites on the slide/cell suspension and indeed varies from top to bottom of the slide and between the slide and the thermal block of the PCR machine (fig 8). This is the overwhelming reason why it is necessary to use a thermocouple to optimise heat transfer kinetics during amplification.

Currently, PCR in situ hybridisation or in situ PCR has little to offer over conventional in situ hybridisation. In the future, the technique may be useful for the detection of low copy number viral infections, in the study of the cellular expression of cellular oncogenes, growth factors, etc., and in the diagnosis of chromosomal translocation in certain tumour types (lymphomas, primitive neuroectodermal tumours, and Ewing’s sarcoma). But the technique is still not reliable and PCR plus NISH techniques are inherently complex and expensive. Standardisation of instrumentation and simplification of the technique are required in the near future, if the technique is to receive universal acceptance.

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3 Haase AT, Rietzel EF, Staskus KA. Amplification and detection of lentiviral DNA inside cells. Proc Natl Acad Sci USA 1990;87:4971-5.


