Detection of human papillomavirus genes in human oral tissue biopsies and cultures by polymerase chain reaction

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Summary We have used the polymerase chain reaction to detect DNA sequences related to human papillomavirus type 16, by simultaneous screening with oligonucleotides from the E6 and L1/L2 open reading frames of the HPV16 genome. The HPV16-related sequence is present at low levels in normal oral tissue, in addition to biopsies and cell cultures from patients with benign and malignant disease. Ultimate analysis of the amplified sequences from the E6(120bp) and L1/L2(173bp) regions of HPV16 was achieved by gel electrophoresis and comparative nucleotide sequencing. The oral carcinoma biopsies and tissue cultures contained DNA sequences which were identical to the E6 region of HPV16, but only rarely contained sequences closely related to the L1/L2 region. The PCR technology should permit the detection, identification and cloning of latent viruses from extremely small tissue biopsies.

Materials and methods

Source of tissues and cell cultures

Cubes of freshly frozen (in liquid N2) tissue biopsies were obtained from the Bristol Dental Hospital (oral samples), Birmingham Maternity Hospital (cervical cancers) and Bristol Royal Infirmary (cervical smears). Duplicated independent histological analyses were performed for all the oral and cervical cancer samples. The oral tumours reported here were all classified as well differentiated squamous cell carcinomas. In all cases the tissue extracted for DNA was adjacent to that from which frozen sections were cut for histology to minimise the effects of tissue heterogeneity on the extracted material.

Keratinocyte cultures were established from human oral squamous cell carcinoma tissue, and passaged without cloning as described by Crane et al. (1986) for rat oral squamous cell carcinoma tissue. DNA was extracted from healthy subconfluent cultures at various passages after initial establishment as a primary culture (see below). Cell cultures of CaSki were obtained from the ATCC.

Extraction and purification of DNA

Nucleic acids (both DNA and RNA) were extracted according to a variation of the method of Chirgwin et al. (1979) as previously described (Maitland et al., 1987b) from all of the tissue cultures and biopsies. Briefly, this involved homogenisation of the tissue in a detergent/guanidinium thiocyanate mixture, followed by ultracentrifugation for 16 h through a double cushion of caesium trifluoroacetate (Pharmacia) containing ethidium bromide. DNA formed a u.v. fluorescent band between the two cushions, whereas the RNA formed a solid pellet at the bottom of the centrifuge tube. This modification allows the maximum information to be obtained from a small (1 mm3) tissue biopsy.

Polymerase chain reaction

PCR was carried out essentially according to the manufacturer's instructions for 30 cycles of denaturation (30 s at 91°C), annealing (30 s at 25°C) and synthesis (5 min at 55°C) in a buffer containing 16.6 mM ammonium sulphate, 67 mM Tris.HCl pH 8.8, 6.7 mM MgCl2, 10 mM mercaptoethanol, 6.7 µM ethylene diamine tetraacetic acid (EDTA), 33 µM each of dATP, dCTP, dGTP and dTTP, 10% dimethylsulphoxide (DMSO), 0.5–1.0 µg of human DNA target and 300 ng of each primer DNA (the reaction will work with considerably lower primer concentrations) in a final volume of 50 µl. After

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The reaction mixture is prepared, and the first denaturation (7 min at 91°C) completed, 1 unit of Taq DNA polymerase (Anglian Biotec) was added and the annealing, synthesis and denaturation were carried out for 10 cycles, when a second 1 unit of Taq polymerase was added. After 20 cycles a further 1 unit of the polymerase was added and the cycles continued up to 30. After this time half of the reaction products were removed, 1/10 volume of gel loading dye (0.25% bromophenol blue, 0.25% xylene cyanole, 25% Ficoll 400) was added to each and the samples were analysed on a polyacryl- amide gel (12%) and electrophoresis at 45 mA for 60 min. The final amplified oligonucleotide products were visualised by ethidium bromide stain (1 ug ml⁻¹ in Tris/borate buffer) and ultraviolet illumination.

Nucleotide sequence analysis of PCR products

The 298 bp amplification product from the E6 gene primers 2 and 3 was purified from a 12% polyacrylamide gel by excision of the band, overnight elution in an excess of 0.1×TE buffer at 37°C and centrifuge driven dialysis on a centriforic 30 (Amicon) as described in Higuchi et al. (1988). The 40 µl product of this was freeze dried, resuspended in up to 8 µl of dd H₂O and approximately half of this (depending on the yield) was transcribed in vitro, using T7 RNA polymerase (Davanloo et al., 1984). Sequencing of the in vitro transcripts was carried out with deoxyxynucleotide triphosphates and avian myeloblastosis virus reverse transcriptase (Pharmacia) exactly as described by Stoflet et al. (1988), using a 32P end-labelled primer 2.
Results

Standard polymerase chain reaction on oral tissue DNA using HPV16 E6 gene primers

One microgram of DNA from a number of different tissues and lesion types was subjected to the standard PCR at 55°C. We selected the E6 region of the HPV16 genome for amplification, and in particular a 120 bp fragment located close to the 3' terminus as shown in Figure 1a. The expected 120 bp amplification product was visualised by ultraviolet illumination of the ethidium bromide stained polyacrylamide gel (Figure 2), except in the fetal kidney and the carcinoma culture 145(6). The negative result with 145(6) was in contrast to the positive result with 145(8). However, reamplification of the non-visualised products from 145(6) resulted in a weak positive result. Since 145(6) is an earlier passage of 145(8), it implies that selection of HPV16 positive cells may occur in vitro.

DNA sequence analysis of PCR products from the E6 region

Preliminary comparative analysis of the PCR products from oral and cervical tissues was carried out with the restriction endonucleases shown in Figure 1a (data not shown). Since the amounts of DNA obtained from a single PCR were close to the limits required for convincing restriction endonuclease analysis, we were anxious to confirm the identity of the DNA amplified by the E6 primers as HPV16 at the ultimate level, i.e. that of the sequence of the nucleotide bases in this region. Initial experiments, in which the 120 bp E6 amplification products were directly sequenced with the sequenase system, produced disappointing results with material amplified from oral samples (whereas cervical samples gave correct sequences according to Seedorf et al. (1985) for both the E6 and L1/L2 genes). The lack of success with oral samples was attributed to the poor yields of PCR product and, to amplify these products further, we employed a third E6 primer, which was homologous to a 20 base sequence, 5' to the original primer 1. Attached to the HPV16 portion of this primer was a 28 base sequence, encoding the T7 phage RNA polymerase promoter (Davanaloo et al., 1984). After purification of the 298 bp amplification product of this reaction, it was therefore possible to amplify the products by in vitro transcription. Sequencing was accomplished with AMV reverse transcriptase from end-labelled primer 2 (Stoflet et al., 1988). The results of a sequencing of one oral sample, and an HPV16 positive cervical carcinoma biopsy control are shown in Figure 3. Over the entire legible range, the two sequences were identical (the locations of primer 1 and the restriction endonuclease sites from Figure 1 are also indicated).
Figure 3  Nucleotide sequence analysis of E6 amplification products from oral and cervical carcinoma DNA. Sequencing was carried out on the 298 bp product of a PCR with E6 primers 2 and 3 (see Figure 1c). The end-labelled sequencing primer was primer 2. The position of primer 1 in the determined sequence, and the restriction endonuclease cleavage sites from Figure 1a are also indicated.

Polymerase chain reaction on oral DNA samples using L1/L2 primers

To test whether the late regions of the HPV16 sequence were present in the oral tissues, we employed primers synthesised from the overlap region between the L1 and L2 regions (Figure 1b). The results of the PCR with these primers are shown in Figure 4. Positive controls with CaSki DNA and cloned HPV16 DNA were positive for both the early and late primers separately and on simultaneous priming with all four oligonucleotides, to give the expected 120 and 173 bp amplification products respectively. Similar results were obtained with other cervical tumour DNA samples which contained considerably lower amounts (one to two genomes per cell) of HPV 16 DNA per cell (data not shown). In contrast, DNA from the oral tissues and cultures was positive only for the E6 diagnostic fragment, even under the less stringent reaction conditions, with the possible exception of culture H157(17) and SCC biopsy 1, which were weakly positive with the L1/L2 primers. In our experience, this implies an extremely low level of late genes, perhaps in a small number of the cells in the biopsy or culture, whereas early genes are present in the majority of the cells. In some of the oral samples, a number of low molecular weight bands (<80 bp) were detected. However, these bands failed to react with a HPV16 DNA probe (data not shown). Multiple low molecular weight bands of this nature were often observed, particularly when several sets of primers were employed simultaneously, but a positive reaction was only scored when the amplification product predicted from the published DNA sequence was observed.
Discussion

Our previous results (Scully et al., 1987; Maitland et al., 1987a) indicated that papillomavirus DNA, closely related to, but not identical to that from the potentially oncogenic cervical HPV type 16, could be detected in nucleic acid preparations from a number of oral lesions, including carcinoma. To further explore the nature of the oral variant of HPV16, we produced cultures from the tumour tissues, but even in these uncloned keratinocyte cultures, detection of HPV DNA was technically difficult. The weak signals could have been due to nucleotide base sequence differences between the cervical HPV16, which we used as a hybridisation probe, and the related oral virus. Equally, the positive results could have resulted from cross-hybridisation between human DNA sequences and the viral probe. By employing the PCR we were able to produce unequivocally positive results, in almost 50% of oral carcinomas tested, even with the smallest oral tissue biopsies. This value is entirely in agreement with our earlier results (Maitland et al., 1987a).

An alternative explanation for the consistently poor signals obtained in experiments to detect HPV in oral tissues may lie in the distribution of viral DNA positive cells in the cultures and the biopsies. In some of the oral keratinocyte cultures the strength of the PCR signal for HPV16 E6 varied with the passage number of the cultures, implying selection for HPV-positive cells with time in culture. It is likely that an uneven distribution of HPV positive cells does occur, both in vivo and in vitro, based on this result and also on in situ hybridisations to HPV16 positive tissue sections (unpublished observations).

The design of oligonucleotide primers for the PCR technique used in this study was based on our previous results (Maitland et al., 1987a). The E6 gene sequences are consistently retained in both cervical and oral carcinomas (Baker et al., 1987; Smotkin & Wettstein, 1986; N. Maitland, unpublished data), whereas the late genes (L1/L2) are often deleted in cervical carcinomas when the HPV16 genome is integrated into the cell genome. In addition, only a very small proportion of the oral samples in our previous study contained detectable HPV16 late genes.

Using this combination of oligonucleotide primers the results again bore out those of our previous work, that in early genes were readily detected, whereas the late genes were frequently absent in oral tissues. Confirmation of the identity of the viral DNA present in the oral tissues was achieved by direct nucleotide sequencing of a portion of the E6 gene.

Employing a third primer from this gene and in vitro transcription from the T7 phage RNA polymerase promoter, sufficient HPV template for nucleotide sequencing was produced. The sequencing indicated that the oral and cervical HPV16 E6 genes were, as predicted from the restriction endonuclease digestions, identical over the 120 bp amplified in the initial experiments, including the primer 1 sequence. It is perhaps significant in view of the essential role of E6 gene expression in tumours that the E6 genes from a total of nine oral and nine cervical HPV16 positive samples sequenced to date show no sequence variations (manuscript in preparation).

Unfortunately, we are still unable to determine, on the basis of either the Southern blotting or the PCR results, whether the frequently negative result with the late region primers is due to lack of these sequences, or to gross sequence divergence, which would make both annealing of high complexity probes in Southern blotting or low complexity oligonucleotide primers for the PCR unlikely. The detection of multiple low molecular weight bands in the simultaneous E6 and L1/L2 priming (Figure 4) experiments probably represents self-priming of the cellular DNA, but could equally be the result of viral genome rearrangements. The presence of the E6 gene in these HPV16 probe arguems in favour of the former explanation, however. If the late regions were completely absent, then the oral 'sequences' would not represent a viable virus.

The resolution of these two possibilities requires cloning of the HPV16 homologous sequences from the oral cultures, which will provide the required amounts of DNA. Using the amplified sequence for the E6 region as an homologous hybridisation probe, we are now able to clone the entire HPV16-related sequence and to determine its role in oral diseases, including cancer.

The use of the PCR has overcome some of the extreme difficulties experienced in the analysis, at the molecular level, of extremely small tissue samples and heterogeneous primary cell cultures. The technique should also assist in epidemiological studies of virus incidence, without the compromises in sensitivity which are often necessary in the handling of large sample numbers.

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