PCR detection of HIV proviral DNA (gag) in the brains of patients with AIDS: comparison between results using fresh frozen and paraffin wax embedded specimens

S F An, A Ciardi, F Scaravilli

Abstract

Aims—To adapt the polymerase chain reaction (PCR) technique of HIV detection to paraffin wax embedded brain tissue and to compare the results with those obtained using frozen tissue.

Methods—HIV antigen and HIV proviral DNA were detected in specimens of frontal lobe using immunohistochemistry and PCR, respectively. DNA was extracted from fresh tissue using standard methods whereas the technique for extracting DNA from paraffin wax embedded tissue was partly modified.

Results—Twenty cases were examined. HIV DNA was detected in 16 cases in frozen specimens. Of these, 15 were also positive when paraffin wax embedded material was analysed.

Conclusions—This study shows that HIV proviral DNA can be detected in formalin fixed, paraffin wax embedded brain tissue by PCR. The results obtained from paraffin wax embedded specimens showed a similar degree of reliability to those from fresh frozen brain. Factors such as fixative, fixation time, and delay in performing post mortem examinations did not seem to influence PCR amplification as positive results were obtained with specimens left in fixative for up to eight months, as well as in cases where post mortem examinations had been delayed for up to four days.

The polymerase chain reaction (PCR), which permits specific and rapid amplification of nucleic acid sequences from a variety of tissues, has also been used to detect HIV DNA in brain tissue of patients with AIDS and of HIV positive subjects. In previous studies carried out in this department Sinclair and Scaravilli and Sinclair et al have successfully used fresh frozen samples from brain tissue obtained at post mortem examination. However, fresh HIV positive specimens represent a safety hazard which considerably restricts their use. By contrast, formalin fixed, paraffin wax embedded tissue has several advantages: this fixative renders the tissue non-infectious and prevents further deterioration of proteins and nucleic acids.

Moreover, as embedding formalin fixed material in paraffin wax is the most widely used method in histopathology, application of PCR to this tissue would enable large retrospective studies to be undertaken and permit comparison of material without risk of contamination. To be acceptable, however, this technique must not only yield reproducible results but its rate of success must be comparable with that using fresh tissue.

The purpose of this study was, therefore, to adapt the PCR technique used to detect HIV proviral DNA in frozen tissue to paraffin wax embedded specimens and to compare the results obtained using each technique.

Methods

Twenty brains of patients with AIDS, from which both fresh frozen and paraffin wax embedded specimens were available, were chosen from the files of the Department of Neuropathology, Institute of Neurology, London. The frontal poles of brain obtained at post mortem examination were removed and frozen whilst the rest of the brain was fixed in 10% buffered formalin for a variable length of time (table). Fifteen blocks were taken from each case and routinely processed for histology. For the purpose of this study, paraffin wax embedded tissue from sites adjacent to the frozen sample was used.

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The primary antibody HIV p24 (Du Pont Diagnostics, Stevenage, UK), which detects the HIV core protein p24, was used to screen paraffin wax embedded sections of anterior frontal lobes from all cases in this study. Briefly, paraffin wax embedded sections were dewaxed and endogenous peroxidase was blocked with 0-03% hydrogen peroxide in methanol. Incubation for one hour with primary antibody (HIV p24) was followed by incubation with biotinylated rabbit anti-mouse (30-60 minutes) and peroxidase conjugated avidin (30 minutes). 3,3-Diaminobenzidine was used as a substrate to reveal the peroxidase.

DNA was extracted as previously described by Pang et al and Sinclair and Scaravilli. Briefly, blocks of semi-frozen frontal lobes were stripped of leptomeninges and diced into small pieces. Tissue was digested overnight with proteinase K (final concentration 0-25 mg/ml in 10 mM TRIS-HCl, pH 7-4, 1 mM EDTA, and 0-5% sodium dodecyl sulphate (SDS)) at 56°C. DNA was subsequently purified using standard methods.

Paraffin wax embedded sections (10 μm) were cut and placed in a 1-5 ml Eppendorf
PCR detection of HIV proviral DNA in AIDS

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<th>Fixation (months)</th>
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PML = progressive multifocal leucoencephalopathy; CMV = cytomegalovirus; FPL = focal pontine leucoencephalopathy; F = frozen tissue; P = paraffin wax embedded tissue.

Results of PCR detection with pathological and immunohistochemical findings

The number of sections varied according to the area of the tissue block; as our frontal blocks have a standard size, 10 sections were cut from each case. Paraffin was removed by washing with xylene for 15 minutes with three changes and then with 100% ethanol for five minutes, followed by drying at 95°C for about 30 minutes. Tissue was digested with proteinase K at a final concentration of 5 mg/ml in 10 mM TRIS-HCl, pH 7-4, 1 mM EDTA, and 0·5% SDS. Digestion took place at 55°C for 24 to 48 hours in tubes which were shaken periodically. The supernatant fluid was extracted once with an equal volume of phenol (equilibrated with 100 mM TRIS-HCl, pH 8-0, and protected by 0·1% hydroxyquinoline), subsequently several times with a 1/1 mixture of phenol and chloroform (a 24/1 (v/v) mixture of chloroform and isomylalcohol), and finally once with chloroform. The supernatant fluid was passed through a Microcon-30 filter (Amicon, Gloucs, UK) using an Eppendorf centrifuge at 13 000 × g for 10 minutes. The initial volume of 0·5 ml was finally reduced to about 5–20 µl. The residue was washed once with 0·5 ml distilled water and used for amplification. DNA was quantified using the ethidium bromide saran wrap method.

PCR ANALYSIS

Human single copy gene β-globin was used as a DNA positive control and only clearly positive samples were used for HIV PCR analysis. Ten molecules of non-infectious plasmid containing a completely rearranged HIV sequence (provided by UK Medical Research Council (MRC), AIDS Directed Programme (ADP), PCR Reference Centre) were used as the HIV positive control. DNA from brain tissue of patients not in any HIV risk group was used as a negative control.

All the primer sequences are written from 5’ to 3’. Two human β-globin primers (MRC, ADP, National Institute for Biological Standards and Control (NIBSC)) were used: ADP894-1, ACA CAA CTG TGT TCA CTA GC; ADP894-2, CAA CT T CAT CCA CGT TCA CC. A 110 base pair (position 14–123) fragment is flanked by primers ADP894-1 and ADP894-2. Two HIV primers 3855/1 and 3855/2 (Perkin Cetus Elmer UK) were used: 3855/1 (SK145), AGT GGG GGG ACA TCA AGC AGC CAT GCA AAT; 3855/2 (SK431), TGC TAT GTC AGT TCC CCT TGG TGC TCT. A 142 base pair (position 1366–1507) fragment is flanked by primers 3855/1 and 3855/2.

Of the reaction mixture, 50 µl contained 0·5 µM of each primer, 0·2 mM of dATP, dCTP, dGTP, and dTTP, and 2·5 units of AmpliTag DNA polymerase in PCR buffer (10 mM TRIS-HCl, pH 8·3, 50 mM KCl, 1·5 mM MgCl2, and 0·001% gelatin). Forty cycles of PCR were performed with 0·5–1 µg extracted template DNA, each cycle consisting of thermal denaturation at 94°C for one minute, primer annealing at 55°C for two minutes, and extension at 72°C for three minutes.

Human β-globin gene amplification was similar to HIV amplification except that the annealing temperature for β-globin gene amplification was 60°C and 0·1 µM of primers were used in the PCR reaction. Amplified products were visualised by ethidium bromide staining of agarose gel electrophoresis. HIV products

Figure 1 Results of PCR amplification to detect the HIV gag gene in a variety of samples of the PCR reference kits. (A) Electrophoresis. (B) Southern blot hybridisation. Lane A shows a negative control consisting of carrier DNA alone; lane B to F show the results using known copy numbers of HIV plasmid DNA (lane B, 1000 molecules; lane C, 0·1 molecules; lane D, 1 molecule; lane E, 10 molecules; lane F, 100 molecules; lane O in (A), DNA molecular weight marker).
were further detected by Southern blot hybridisation to ascertain their specificity.

**SOUTHERN BLOT HYBRIDISATION**

To detect the 142 base pair PCR product, modified Saluz and Jost\(^ {10} \) "filling in" methods were applied as described before.\(^ {2} \) Oligonucleotides were provided by the MRC, ADP, and NIBSC. Their sequences are: \( \texttt{gag} \) (30-mer), \( \texttt{5'-CAT CAA TGA GGA AGC TGC AGA ATG GGA TAG} \); (10-mer), \( \texttt{5'-TCC CAT TCT G} \). Briefly, an annealing mix containing 10 \( \mu \)l of the 30-mer, 10 \( \mu \)l of the 10-mer, and 2-5 \( \mu \)l of the oligonucleotide labelling buffer was prepared and incubated at 75°C for two minutes, at room temperature for 20 minutes and on ice for 10 minutes. This was followed by incorporation of digoxigenin-11-dUTP and precipitation of the probe by ethanol.

Nylon membranes (Boehringer Mannheim, UK) were prehybridised for more than one hour at 42°C in hybridisation buffer (5 \( \times \) saline-sodium citrate (SSC), 50% formamide, 0.1% N-lauroylsarcosine, 0.02% SDS, and 2% blocking reagent) and then hybridised overnight at 42°C with 200 ng/ml labelled probe. They were then washed in 0.2 \( \times \) SSC and 0.1% SDS at room temperature for one hour with four changes.

Colorimetric detection using nitroblue tetrasodium and X-phosphate was performed according to the Boehringer Mannheim protocol.\(^ {11} \) Briefly, after incubation with antidigoxigenin-alkaline phosphatase at room temperature for 30 minutes and a postincubation wash, the filter was incubated in substrate to permit colour development.

**Results**

Figure 1 illustrates the results of PCR amplification with the oligonucleotide probe for the HIV \( \texttt{gag} \) gene in a variety of samples of the PCR reference kit. The result of this test shows that 10 molecules of HIV-1 DNA gave a clear signal after Southern blot hybridisation. There were no false positive results following either PCR amplification or Southern blot hybridisation.

The neuropathological changes observed in the brain tissue of the 20 patients with AIDS included in this study are summarised in the table. HIV encephalitis was present in seven cases; other neuropathological findings included cytomegalovirus encephalitis (three cases), progressive multifocal leukoencephalopathy (one case), cryptococcosis (two cases), lymphoma (one case), tuberculosis (one case), and non-specific changes (four cases); three cases did not show any abnormalities. Nine cases, including all those with neurological evidence of HIV encephalitis, were HIV p24 positive.

Specimens were regarded as positive or negative whenever PCR results were consistently positive or negative (PCR amplifications were repeated three times). Samples that were positive in only one or two of three tests (cases 12 and 17–20 in the table) were defined as indeterminate;\(^ {3} \) and underwent another five tests, on the basis of which they were reclassified as positive (four cases) or negative (one case).

Sixteen cases were positive on analysis of frozen specimens. These included all those with immunohistochemical evidence of HIV. Of these, 15 were also positive when paraffin wax embedded material was analysed. Southern blot hybridisation confirmed the positive results. None of the cases positive on analysis of paraffin wax embedded samples gave negative results when frozen tissue was examined.

Figure 2 shows the results of PCR amplification in three cases using both frozen and paraffin wax embedded samples. Case 1 (lanes A and B) was positive in both; case 2 (lanes C and D) gave a positive result with frozen material only; and case three (lanes E and F) is an example of the four cases with negative results on analysis of both frozen and paraffin wax embedded material.

**Discussion**

In this study we have shown that HIV proviral DNA can be detected by PCR in brain tissue...
of patients with AIDS using formalin fixed, paraffin wax embedded tissue. The HIV primers used were chosen so as to amplify a specific sequence of the HIV gag region. As in a previous study,1 gag primers produced more reliable results than pol or env primers, and highly conserved areas have been identified in the HIV-1 genome in long terminal repeat sequences and in the gag gene.8 Moreover, as the region spanning the gag sequence is thought to be the last to be synthesized during reverse transcription, positive results using the gag region reflect completely or almost completely synthesized viral DNA.8

In a previous investigation using PCR specimens were classified as positive, negative, or indeterminate depending on whether most of the tests were positive; all tests were negative; or one or two only were positive, respectively. In the present study each indeterminate case underwent five additional tests and samples were defined as positive when at least three of the latter gave positive results.

Of the 20 cases in this study, 16 were positive on analysis of frozen samples; of the latter, 15 were also positive when paraffin wax embedded material was studied. On the other hand, none of those which were negative on analysis of frozen samples gave positive results on analysis of paraffin wax embedded sections. Positive cases on analysis of both frozen and paraffin wax embedded material included the seven with morphological and immunohistochemical evidence of HIV encephalitis. A possible explanation for the single negative result (fig 2) could be the presence of an inhibitor of Taq polymerase.11 Although the human β-globin gene was successfully amplified in this case, the degree of inhibition may not have been uniform for the different sets of primers.14

Several investigators have drawn attention to the effects of various fixatives and fixation times on PCR results.17 In this study the specimens had been kept in 10% buffered formalin (the most widely used fixative in neuropathology) for one to eight months, and the results show that even relatively long fixation times are compatible with successful amplification of HIV. However, DNA extracted from paraffin wax blocks will probably need to be concentrated using a microconcentrator instead of ethanol precipitation (data not shown).

The PCR technique has been used to amplify specific DNA from paraffin wax embedded human tissue in a variety of disorders. Human papillomavirus was identified by Brandwein et al.18 Nawa et al.19 Ohta and Ikeda0 Salzstein et al.21 Van-Bommel et al.22 and Kellokoski et al.23 an adenovirus, cytomegalovirus, and herpes simplex virus (HSV) 1 and 2 were detected by Vesey et al.24 whilst Epstein-Barr virus (EBV) was detected by Niedermeyer et al.25 and Moshred et al.26 and hepatitis C virus by Brexsters et al.27 Central nervous system (CNS) tissue specimens were used by Woodall et al.28 for the identification of enterovirus RNA; Nicoll et al.29 detected HSV1 DNA in the brains of patients with herpes encephalitis and Lagger et al.30 detected EBV in the brains of patients with primary biliary cirrhosis. Morgello31 examined EBV and HIV in 12 AIDS related primary CNS lymphomas and detected EBV in six of the 12 tumours but could not detect HIV; Lai-Goldman et al.32 detected EBV in various organs of patients with AIDS but not in the brain. In this study we have shown not only that HIV proviral DNA can be detected by PCR in paraffin wax embedded tissue from the CNS but also that the results have the same degree of reliability as those obtained using fresh frozen material.

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