Monoclonality in B cell lymphoma detected in paraffin wax embedded sections using the polymerase chain reaction

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Abstract

The polymerase chain reaction (PCR) was used to develop a simple technique for detecting monoclonality at the DNA level in B lymphocyte populations in formalin-fixed, paraffin wax embedded material. Sections were dewaxed and dehydrated and the DNA was extracted by boiling in water for 45 minutes. A semi-nested PCR was performed to amplify the V-D-J region of the immunoglobulin heavy chain gene. The product was electrophoresed and viewed under ultraviolet light after ethidium bromide staining. Specimens from 26 B cell lymphomas produced a monoclonal band in 24 cases and no amplification in two cases; monoclonality was specific for this disorder. Specimens from seven T cell lymphomas produced no amplification; specimens from nine reactive nodes produced a broad smear of polyclonal material; and specimens from 12 cases of carcinoma produced either no amplification or polyclonal material.

As detection of monoclonality is strongly suggestive of neoplastic disease, this technique is likely to be of value in routine diagnosis, because of its speed, simplicity, and applicability to fixed, embedded material.

Diagnosis and classification of malignant lymphoma is still a challenge, in part due to the limitations and ambiguities of morphology and immunophenotyping. Analysis of the immunoglobulin and T cell receptor genes by Southern blotting to show monoclonality has helped in the distinction between neoplastic and reactive disorders, and to a lesser extent, in the assignment of lineage. Southern blotting, however, is of limited value for routine diagnostic work because it is slow and complex, requires the use of radioactive isotopes and a large amount (1–5 μg) of DNA, and is not well suited to the study of fixed, embedded material owing to degradation of DNA.

Trainor et al reported a technique based on the polymerase chain reaction (PCR) for rapid diagnosis of B lymphocyte neoplasms in frozen or fresh samples of blood, bone marrow, or lymph node. In this paper we extend this technique to the detection of neoplastic populations in paraffin wax embedded sections and show that it is possible to distinguish B cell lymphoma from other neoplastic and non-neoplastic conditions.

Methods

Paraffin wax embedded, formalin fixed tissue sections were obtained from the Department of Histopathology, Flinders Medical Centre, and included lymph node tissue from 26 patients with B cell non-Hodgkin's lymphoma (B-NHL), seven patients with T cell non-Hodgkin's lymphoma (T-NHL), nine patients with reactive lymph node enlargement, and other tissues from 12 patients with non-hodgkinoid neoplasms (pulmonary adenoma, carcinoma from oesophagus, stomach, colon, breast, skin, prostate).

All of the tissue samples had previously been studied and classified as B or T NHL, reactive tissue, or carcinoma, by routine histopathology, immunohistochemistry, and, for all lymphomas, by Southern blotting of the immunoglobulin and T cell receptor genes.

EXTRACTION OF DNA FROM TISSUE SECTIONS

Two slices, 10 μm thick, were cut from tissues embedded in paraffin wax blocks and placed in screw-cap Eppendorf tubes. Xylene (800 μl) was added to dissolve the paraffin. After brief vortexing samples were left for five minutes, then centrifuged in an MSE MicroCentaur at 13500 rpm for five minutes. The supernatant was removed using a fresh Pasteur pipette for each sample to avoid cross-contamination, and 800 μl ethanol were added. After vortexing for 10–20 seconds and centrifuging for another five minutes the ethanol was pipetted off. This ethanol washing procedure was repeated once more. The samples were dried under vacuum for 45 minutes. Sterile water was added (200–400 μl, according to the amount of cellular material present), and the samples were placed in a boiling water bath for 45 minutes. The supernatant was used as the DNA source for PCR amplification. Two or more DNA samples were prepared and tested from each tissue block.

PRIMERS FOR PCR

The primers used for the Joining (J) and 3' end of the Variable (V) segments of the human immunoglobulin heavy chain genes were synthesised on an Applied Biosystems 381A DNA synthesiser and their sequences were as follows.
A PCR cycle consisted of annealing for two minutes at 60°C, extension for two minutes at 72°C, and denaturation for two minutes at 94°C. A semi-nested PCR was performed: a first round of 30 cycles with primers FR3A and LJH and 10 μl of the boiled cellular extract as template; and a second round of 20 cycles with FR3A and VLJH, with 10 μl of a 1 in 1000 dilution of the first round PCR as template. Before each round the PCR reaction was heated to 94°C for four minutes, and after each round a final extension step of six minutes at 72°C was performed. Each experiment also contained a tube without DNA template (as a negative control), and a tube whose template was DNA from pre-B lymphocyte line Nalm 6, which has its immunological genes rearranged (as a positive control). PCR amplified material (5 μl) was electrophoresed in 2% agarose gels in TBE buffer (89 mM TRIS-HCl, 89 mM boric acid, 2 mM ethylene diamine N,N,N',N'-tetra acetic acid) at 100 volts for 90 minutes and the DNA was visualised under short wavelength ultraviolet light after ethidium bromide staining of the gel.

Results

Figure 2 shows typical examples of the amplified DNA. Tracks 1–3 show the discrete monoclonal band (100–120 base pairs long) of amplified DNA observed in three cases of B-NHL; tracks 4–6 show failure of amplification observed in three cases of T-NHL; tracks 7–9 show the broad smear of polyclonal amplified DNA observed in three cases of reactive lymphadenopathy; and tracks 10–12 show failure of amplification in three cases of carcinoma. The results of all experiments, summarised in the table, show that the detection of one or two discrete bands was absolutely specific for B-NHL, with a sensitivity of 92%. A diffuse polyclonal smear was seen in all cases of reactive nodes. T-NHL and carcinomas usually gave no amplification but polyclonal material was observed in several cases. The two cases of B-NHL which produced no amplified product were consistently negative on repeated testing.

Discussion

The PCR technique for detection of monoclonality, recently described by Trainor et al, is based on the events which occur during immunoglobulin heavy chain gene

Use of polymerase chain reaction to detect B-cell non-Hodgkin’s lymphoma (B-cell NHL) in paraffin wax embedded tissue sections

<table>
<thead>
<tr>
<th>Tissue</th>
<th>Monoclonal band/Total</th>
</tr>
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<tbody>
<tr>
<td>B cell NHL</td>
<td>24/26</td>
</tr>
<tr>
<td>T cell NHL</td>
<td>0/7</td>
</tr>
<tr>
<td>Reactive lymph node</td>
<td>0/9</td>
</tr>
<tr>
<td>Carcinoma</td>
<td>0/12</td>
</tr>
<tr>
<td>Total</td>
<td>54</td>
</tr>
</tbody>
</table>

Samples were scored for the presence of a discrete monoclonal band as exemplified in fig 2, tracks 1–3.

For the third framework region (at the 3' end of V):
Fr3A: 5' - 5' ACACG GC [C/T] [G/C] T GTATT ACTGTG 3'
For the J region:
LJH: 5' - 5' TGAGG AGACG GT GACC 3', or
VLJH: 5' GTGAC CAGGG T [A/G/C/T] CCT TGGCC CCAG 3'
Fr3A is based on a consensus sequence from 17 human V-segments, while LJH and VLJH are based on a consensus from the six J regions. Use of these primers would be expected to generate a fragment of about 100–120 kilo base pairs in length (fig 1). The binding site for the VLJH primer is 5' compared with that for the LJH primer.

POLYMERASE CHAIN REACTION TECHNIQUE

The PCR was performed in a Perkin-Elmer-Cetus Thermal Cycler. The reaction mix (25 μl) contained 0-125 μg of each primer, 1 mM each of deoxyadenosine, deoxyctydine, deoxyguanosine, and thymidine triphosphate, 10% v/v dimethyl sulphoxide, 170 μg/ml bovine serum albumin (Sigma), 200 μg/ml gelatin, 16-6 mM ammonium sulphate, 4-5 mM magnesium chloride, 10 mM β-mercaptoethanol and 67 mM TRIS-HCl (pH 8-8 at 25°C), 0-2 units of Taq polymerase (Amplitaq: Cetus), and a template. Each reaction mix was overlaid with 50 μl of light mineral oil (Sigma).
rearrangement. In the germline state (and therefore in all cells except B lineage lymphocytes) the V and J segments of the gene are separated by several thousand bases of DNA, and the polymerase chain reaction, which uses primers binding to these regions, cannot take place. During the differentiation of B lymphocytes, however, a V and a J segment are brought close to each other. The polymerase chain reaction can generate a DNA fragment of 100–120 base pairs which is readily detectable by electrophoresis on an agarose gel. The properties of the amplified product permit the ready distinction of a monoclonal population of neoplastic B lymphoid cells from a polyclonal population of non-neoplastic B lymphoid cells. During differentiation, varying numbers of nucleotides are added at random between the V, D, and J segments (fig 1). The two primers bind to fixed portions of the framework 3, and J regions of the immunoglobulin genes and therefore different B cell clones, which have different rearrangements of their immunoglobulin genes, will give rise to amplified segments of different sizes. A monoclonal B lymphocyte population will therefore be characterised by amplified DNA of a single size, whereas a polyclonal B-lymphocyte population will be characterised by amplified DNA of a range of sizes.

The results in this study of fixed tissue compare well with those of Trainor et al, who used fresh or frozen tissue samples. Monoclonality was detected in 92% of cases of B-NHL and was specific for this disorder. As detection of monoclonality is strongly suggestive of neoplastic disease, the present technique is likely to be of value in routine diagnosis, owing to its speed, simplicity, and applicability to fixed, embedded material. It will also permit the retrospective study of large numbers of patients. The nested primer technique used in this study is helpful for small amounts of starting material. It is more sensitive and specific than the single round approach originally described by Trainor et al.

In this study two of the 26 cases of B-NHL did not give a monoclonal band despite the detection in all of a monoclonal gene rearrangement by Southern blotting. This phenomenon has been noted before and possible explanations in these two patients are (1) atypical rearrangement, such as inversion of the immunoglobulin genes, which seems most likely, or (2) a false assignment of lineage by Southern blotting of the immunoglobulin and T cell receptor genes. The Southern blotting method detects rearrangement of the immunoglobulin of T-cell receptor gene, irrespective of the final arrangement of the gene, whereas the PCR technique detects only V-D-J rearrangements which are in the correct orientation. Thus occasional cases in which monoclonal gene rearrangement can be detected by Southern blotting but not by PCR are to be expected.

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