Comparison of in situ hybridisation and polymerase chain reaction in the diagnosis of B cell lymphoma

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Abstract

Aim—To compare the sensitivity of the detection of immunoglobulin light chain messenger RNA (mRNA) restriction by in situ hybridisation (ISH) and clonal immunoglobulin heavy chain gene rearrangements by polymerase chain reaction (PCR) in the diagnosis of B cell lymphoma.

Methods—Analyses were applied to formalin fixed, paraffin wax embedded, routine diagnostic specimens from cases with a provisional diagnosis of reactive lymph node (n = 23), B cell lymphoma (n = 21), and T cell lymphoma (n = 4). Non-isotopic ISH for kappa and lambda immunoglobulin light chain mRNA was performed using both fluorescein and digoxigenin labelled oligodeoxynucleotide probe cocktails. PCR was carried out on DNA extracted from sections using primers to framework 3 (Fr3) of the V segments and to conserved sequences from the J regions of the immunoglobulin heavy chain genes.

Results—All reactive lymph node showed a polyclonal pattern of light chain mRNA by ISH, although one showed an excess of kappa positive cells. Nineteen of 21 (90%) cases of B cell lymphoma showed light chain restriction, and a further case showed a vast excess of kappa positive cells. By PCR, 20 of 23 reactive nodes (87%) showed a polyclonal pattern. In 13 of 21 B cell lymphomas (62%) a clonal band was detected.

Conclusion—In the diagnosis of B cell lymphoma in routinely processed diagnostic material ISH for light chain mRNA was more sensitive (90%) than PCR for heavy chain gene rearrangement using Fr3 and J region primers (62%).

Keywords: in situ hybridisation; polymerase chain reaction; B cell lymphoma

The demonstration of clonality is the usual method of making the final distinction between reactive and neoplastic lymphoid proliferation. In B cell lymphomas, this has most commonly been achieved by the demonstration of immunoglobulin light chain restriction by immunocytochemistry. More recently, in situ hybridisation (ISH) for kappa and lambda messenger RNA (mRNA) has been applied, which avoids the problem of high levels of background staining due to tissue immunoglobulins. Several reports have indicated a higher sensitivity for ISH than for immunocytochemistry.

Clonality may also be demonstrated by molecular genetic analysis of immunoglobulin heavy chain gene rearrangements, which initially used Southern blot analysis of fresh tissue. However, the polymerase chain reaction (PCR) and sections of fixed tissues can now be used. Although less sensitive than Southern blot analysis, it has also been reported to be more sensitive than immunohistochemical detection of kappa and lambda light chain restriction, and can be performed on paraffin wax embedded diagnostic material.

To date, there has been no reported study directly comparing ISH and PCR in the detection of clonality on the same cases. The aim of our study, therefore, was to perform such a comparison in a series of paraffin wax embedded lymph nodes submitted for diagnostic investigation to our surgical pathology service.

Methods

CARES

Lymph nodes were fixed in neutral buffered formalin and processed to paraffin wax. Based on morphological and immunohistochemical characterisation, a provisional diagnosis was made of reactive lymph node (n = 23), B cell lymphoma (n = 21), and T cell lymphoma (n = 4). Cases in each group were selected on a random retrospective basis. The B cell tumours were diagnosed in the REAL classification as follicular lymphoma (n = 7), large B cell lymphoma (n = 9), lymphoplasmacytic lymphoma (n = 1), and lymphocytic lymphoma (n = 4). Human tonsil was included as a technical control.

IN SITU HYBRIDISATION

The ISH technique was based on previous publications. Sections (4 µm thick) were cut on to silane coated slides. All solutions used before hybridisation were diethylpyrocarbonate (DEPC) treated. The sections were dehydrated to water. They were incubated in 0.2 M HCl for 20 minutes, washed in water, then immersed in 2× SSC (1× SSC = 0.15 M sodium chloride, 0.015 M sodium citrate) at 70°C for 10 minutes before digestion with proteinase K (Sigma, Poole, Dorset, UK) at 10 µg/ml for one hour at 37°C. This reaction was stopped with 0.2% glycine before washing in water and postfixing in freshly prepared 0.4% paraformaldehyde for 20 minutes at 4°C. The sections were washed again in water, then rinsed in methylated spirits, and allowed to air dry.
Two different sources of probes were used in this study: fluorescein (FITC) labelled oligonucleotide cocktails to kappa and lambda light chain mRNA (Dako, High Wycombe, Bucks, UK) were used initially. Detection was achieved using an alkaline phosphatase labelled anti-FITC antibody (Dako). When production of these probes ceased, we changed to digoxigenin labelled cocktails (R & D, Abingdon, Oxfordshire, UK), detection being attained with an alkaline phosphatase conjugated antibody to digoxigenin (Boehringer, Lewes, East Sussex, UK). Both sets of probes were subjected to the same pre- and post-hybridisation conditions.

**POST-HYBRIDISATION**

Following overnight hybridisation at 37°C, the coverslips were removed in 2× SSC, then in Tris buffered saline (TBS), 0.1% Triton X for 10 minutes. This was followed by two washes in TBS before incubation with the relevant antibody for 30 minutes. The antibody was removed with a TBS wash. The products were visualised with a standard alkaline phosphatase detection system using nitroblue tetrazolium (NBT)/5-bromo-4-chloro-3-indolyl phosphate (BCIP) as substrate, the slides being incubated in dye overnight. The slides were lightly counterstained with haematoxylin and water mounted using Glycergel (Dako).

**POLYMERASE CHAIN REACTION**

A 10 µm section was cut from each block, a fresh knife being used for each new block. The sections were dewaxed using 1 ml xylene, vortexed, then centrifuged at 13 000 × g for five minutes. The supernatant was carefully dis-
carded and a fresh 1 ml aliquot of xylene was added. This procedure was repeated twice with 1 ml absolute alcohol. The pellet was dried using a vacuum desiccator. An aliquot of 100 µl of 500 µg/ml proteinase K (Sigma) was added to each sample and incubated at 37°C overnight. The enzyme was inactivated by heating for 10 minutes at 95°C and samples were stored at 4°C until needed.

The primers consisted of a consensus primer to the framework region 3 (Fr3) of the V segment and two primers directed to conserved sequences from the J region in the immunoglobulin heavy chain gene.5–8 The sequences were as follows:

Fr3: ACACGGGC(C/T)(G/C)TGTATTAC TGT
LJH: TGAGGAGACGGTGACC
VLJH: GTGACCAGGGT(N)CCTTGGCC CCAG

A semi-nested PCR method was used.7 In the first round, amplification used Fr3 and LJH primers. Two per cent of this product was used as the template for the second round of amplification, using Fr3 and VLJH primers.

A 50 µl reaction volume was used for each round. This consisted of 1× reaction buffer (10 mM Tris HCl, pH 8.3, 50 mM KCl, 0.001% gelatin), 200 µM each dNTP, 250 ng each primer, and 0.5 units of AmpliTaq (Perkin-Elmer, Warrington, Cheshire, UK). In the first round, 5 µl of DNA was used with 9 mM MgCl₂. For the second round, 1 µl of first round product and 3.5 mM MgCl₂ were used.

First round amplification was 30 cycles of 93°C for 45 seconds, 50°C for 45 seconds, 72°C for 110 seconds, then one cycle of 72°C for five minutes on a thermal cycler (Hybaid, Basingstoke, Hants, UK). The second round consisted of 20 cycles at the above conditions. The final products (15 µl) were analysed on a 10% polyacrylamide gel (Boehringer), post-stained with ethidium bromide, and viewed under UV light. A product in the 70–120 base pair (bp) range was expected.

**Results**

**IN SITU HYBRIDISATION**

The results using both sets of probes were similar, with identical results in cases selected for direct comparison. All reactive nodes showed a polyclonal pattern (fig 1), although one showed a vast excess of kappa positive cells in germinal centres. Nineteen cases (90%) of B cell lymphoma showed light chain restriction (figs 2 and 3). A further case showed a vast excess of kappa positive cells. In one case, no signal was obtained. Tumour cells from T cell lymphomas were negative, with infiltrating B lymphocytes showing a polyclonal pattern.

**POLYMERASE CHAIN REACTION**

In 18 reactive lymph nodes and in the T cell lymphomas, a ladder or a smear was seen (fig 4). Human tonsil showed a smear. In two reactive nodes, a prominent band was seen within a ladder. One of these was the case showing kappa excess on ISH. One reactive node failed to amplify. In 13 of 21 (62%) B cell lymphomas a clonal band was detected. In seven cases, a
ladder or smear was seen. One case failed to amplify (fig 4).

Discussion
Initially, ISH for kappa and lambda immunoglobulin light chain mRNAs was limited in sensitivity and a positive signal was seen only in lesions with a high copy message, such as multiple myeloma and plasma cytoma. The use of riboprobes increases sensitivity, but has the disadvantage of requiring molecular biology facilities for probe production. The development of more sensitive non-isotopic techniques based on oligonucleotide cocktails labelled with digoxigenin or fluorescein, with an extended detection step, now permits general application to diagnostic specimens. Using this technique, we have been able to demonstrate light chain restriction in 90% of 21 cases of B cell lymphoma studied. This approximates to 95% positivity in a series of extramedullary plasmacytomas. In a selected series of monocytoid B cell proliferations, 100% success was reported. We have found the technique useful for detecting clonal proliferations in two biopsies that had previously been labelled reactive but where, subsequently, a lymphoma has been diagnosed (unpublished data). While slight changes in intensity of signal could be achieved by altering the concentration of proteinase K, the concentration of 10 µg/ml used gave the appropriate pattern of staining in all cases.

PCR techniques for the demonstration of clonality are usually based on amplification of the variable region of the immunoglobulin heavy chain using primers Fr1, Fr2, or Fr3 of the variable sequence and to a lesser extent Fr4. The level of detection of 62% for clonal bands in lymphoma cases is similar to some published series, but lower than that found in others, of cases. This may be due to the high proportion of follicular and large B cell lymphomas in our series (16 of 21), because centrocytic/centroblastic tumours have been shown to amplify less efficiently than other subtypes with Fr3 primers.

The application of a second PCR with different primers will increase the yield of positive cases, but in diagnostic practice this would increase the workload and cost for each test if applied to all diagnostic cases, and the turnaround time if applied sequentially to Fr3 negative cases. The time taken for the procedure was similar for both PCR and ISH, but the other costs of ISH were approximately half those of PCR.

On the basis of our results, we would suggest that ISH is the more useful test for general application in the diagnostic histopathology laboratory to differentiate lymphomas from reactive lymphoproliferative disorders. False negative results were rare. Both probes failed to bind in a single case, suggesting a general problem with preservation of RNA.


