Hodgkin’s disease: immunoglobulin heavy and light chain gene rearrangements revealed in single Hodgkin/Reed-Sternberg cells

F Deng, G Lü, G Li, G Yang

Abstract

Aim—To corroborate and investigate the nature of Hodgkin/Reed-Sternberg cells (H/R-S) of various subtypes of Hodgkin’s disease.

Method—Single H/R-S cells were micro-picked from frozen sections of tissues affected by Hodgkin’s disease. The DNA from these cells was amplified by the polymerase chain reaction (PCR) with immunoglobulin heavy chain (IgH) gene FRH/JH primers and light chain gene family specific primers.

Results—Fifty two of 135 isolated cells gave specific reaction products (36%). IgH and V<sub>λ</sub> gene rearrangements were found repeatedly in many H/R-S cells from one case of lymphocyte predominant Hodgkin’s disease. Repeated V<sub>μ</sub> and individual IgH/V<sub>λ</sub> rearrangements were seen in one case, and individual IgH and V<sub>λ</sub>/V<sub>κ</sub>, rearrangements were seen in another case of nodular sclerosis-type Hodgkin’s disease. Repeated IgH/V<sub>λ</sub>, and individual V<sub>λ</sub>/V<sub>κ</sub>, rearrangements, repeated V<sub>μ</sub> and individual IgH/V<sub>λ</sub>, rearrangements, and repeated IgH and individual V<sub>λ</sub>/V<sub>κ</sub> rearrangement were detected, respectively, in three cases of mixed cellularity-type Hodgkin’s disease. Repeated and individual IgH rearrangements were found in another two cases of mixed cellularity-type Hodgkin’s disease.

Conclusion—The H/R-S cells isolated from lymphocyte predominant Hodgkin’s disease had IgH and V<sub>λ</sub> gene rearrangements, which supports the conclusion that this disease results from a proliferation of neoplastic B cells. The IgH and κ and/or λ light chain gene rearrangements seen in H/R-S cells isolated from classic Hodgkin’s disease (mixed cellularity-type and nodular sclerosis-type) support the theory that these cells derive from B lineage cells at various stages of differentiation. To our knowledge, this is first time that λ light chain gene rearrangements have been detected in H/R-S cells.

Keywords: Hodgkin’s disease; Hodgkin and Reed-Sternberg cells; immunoglobulin gene rearrangements

Although Hodgkin’s disease has been known for a long time, the lineage of its neoplastic elements—H/R-S cells and their variants—is still a matter of controversy. B cell, T cell, and monocyte markers have been detected by immunohistochemistry on H/R-S cells. How-ever, the detection of B and T cell associated antigens does not necessarily imply that these proteins, which might be absorbed from the microenvironment, are synthesised by the H/R-S cells. In situ hybridisation detected κ light chain mRNA in lymphocytic and/or histiocytic cells of lymphocyte predominant Hodgkin’s disease<sup>1–4</sup>; however, a similar positive signal has not been detected in H/R-S cells of classic Hodgkin’s disease (R Von Wasielewski, et al. Presented at the third international symposium on Hodgkin’s lymphoma, Cologne Germany, 1995). Southern blot hybridisation and routine polymerase chain reaction (PCR) for immunoglobulin or T cell receptor gene rearrangement does not allow an accurate explanation of the origin of clonal or polyclonal gene rearrangements,<sup>1</sup> and it is most likely that these gene rearrangements result from the large background population of cells in the tissue of Hodgkin’s disease. Therefore, it is very difficult to determine the cell lineage of H/R-S cells because Hodgkin’s disease has special histological characteristics distinct from other tumours: a small number of H/R-S cells (usually less than 1%) among a large number of reactive background cells. A major obstacle in defining the nature of H/R-S cells has been obtaining a pure population for study. Molecular analysis of single H/R-S cells is the best way to confirm the nature of these cells.

In 1994, Küppers et al picked single H/R-S cells from histological sections of Hodgkin’s disease tissue samples. However, the possibility that these H/R-S cells derived from another cell lineage (T cells) cannot be excluded entirely because T cell associated antigens are found on H/R-S cells in this subtype of Hodgkin’s disease.<sup>1</sup>
In our study, we micropicked single H/R-S cells from frozen tissue sections. PCR for Ig gene rearrangements was performed with an IgH chain gene common primer and κ and λ light chain gene family specific primers, with the aim of further investigating the nature of H/R-S cells.

**Methods**

**TISSUES**

Frozen lymph nodes affected by Hodgkin’s disease (one case of lymphocyte predominant Hodgkin’s disease, two cases of nodular sclerosis-type Hodgkin’s disease, and five cases of mixed cellularity-type Hodgkin’s disease) came from the Department of Pathology, West China University of Medical Science.

**IMMUNOHISTOCHEMISTRY**

Frozen sections (10–15 µm thick) were cut and immunostained with BerH2, (Dakopatts, Glostrup, Denmark). The slides were counterstained with haematoxylin. In addition, the sections of paraffin wax embedded tissue were immunostained with anti-κ and anti-λ light chain monoclonal antibodies (Dakopatts).

**ISOLATION OF SINGLE H/R-S CELLS**

The stained, frozen sections were incubated with 5 mg/ml collagenase (Boehringer, Mannheim, Germany) for three to four hours at 56°C. Under the microscope at a certain magnification, the single H/R-S cells with CD30 positive stained cytoplasm were carefully isolated using a micropipette (20–25 µm in diameter) attached to a hydraulic micromanipulator (WR-89; Nairishiger, Japan), and then transferred to Eppendorf tubes that contained 32 µl 1× PCR buffer and stored at −20°C (fig 1).

**PCR OF SINGLE H/R-S CELLS**

The primers used for analysis of IgH chain gene rearrangement contained FR IIIa that was designed for the third framework region of the VH region, and LJH and VLJH that were designed for the JH region. The amplification of the V_{H}D_{J_{H}} fragment yields a product of ~80–120 bp in length. Family specific primers were used for detecting both light chain gene rearrangements. It is possible to differentiate monoclonal from polyclonal B cell proliferation. In a clonal population, only one or two V genes are rearranged and give rise to a PCR product in the reaction with the V primer of the respective V gene family, whereas in a polyclonal population all, or at least most, of the V gene family rearrangements are present. Both the light chain gene family specific primers were designed for the framework region I. The amplified fragment is ~350 bp in length.

A semi-nested PCR approach was chosen and carried out using a PE480 DNA hot cycle appliance. Single cells in Eppendorf tubes were incubated with 0.25 mg/ml proteinase K for two hours at 56°C. The enzyme was inactivated by heating to 95°C for 10 minutes. The first round of amplification was carried out in 50 µl volumes that contained 50 mM KCl, 10 mM Tris-HCl (pH 8.4), 0.01% gelatin, 2.5 mM MgCl{$_2$}, 800 µM dNTP{$_s$}, 2.8 nM each of the FR IIIa and LJH primers, or a mixture of all V$_{H}$ and J$_{H}$ primers, or a mixture of all V$_{L}$ and J$_{L}$ primers, and 2 U Taq DNA polymerase, hot start. The cycle programme consisted of 40 cycles of 95°C for 90 seconds, 59°C for 30 seconds, and 72°C for 80 seconds. In the first round, we added 2.8 nM each of three β-globin primers (PC40, GH2O, and PE) to determine whether it would be appropriate to perform the

![Figure 1](http://mp.bmj.com/)

**Figure 1** Procedure for isolating a single H/R-S cell. (A) A typical H/R-S cell is shown in the middle of the picture, with a double nucleus and CD30 positive cytoplasm. Haematoxylin and eosin stained; magnification, ×200. (B) Picking the cell up with the micropipette. (C) The H/R-S cell in the micropipette. (D) After isolation of the cell, there is a hole where the cell was located, with a small rim of cytoplasm and nucleus.
second round of amplification. A specific product of 268 bp, detected by polyacrylamide gel electrophoresis, implied that the DNA from a single H/R-S cell was suitable for the second round of amplification.

The second round of amplification was carried out in separate reactions for VLJH and each of the Vκ or Vλ family specific primers, using 2 μl of the first round reaction mixture in a 50 μl volume that contained 50 mM KC1, 10 mM Tris-HCl (pH 8.4), 0.01% gelatin, 2.5 mM MgCl2, 800 μM dNTPs, 0.2 μM of each FRIIIa and VLJH, or 0.25 μM of each Vκ and Jκ primer mixture, or each Vλ and Jλ primer mixture, and 2 U of Taq DNA polymerase, hot start. The cycle programme consisted of 40 cycles of 95°C for 90 seconds, 61°C for 30 seconds, and 72°C for 80 seconds. A 10 μl aliquot of the reaction mixture was analysed on an 8% polyacrylamide gel. The amplified products were visualised under ultraviolet light after ethidium bromide staining.

CONTROL EXPERIMENTS
One case of a follicular centre cell lymphoma was used as a positive control for the detection of IgH chain gene rearrangement. One case of lymphadenopathy was used as a positive control for amplification with both light chain gene primers. In negative controls no DNA was added.

Results
CONTROL EXPERIMENTS
In the positive control, a specific 80 bp band was seen after electrophoresis. The amplification with DNA from the tissue sample of lymphadenopathy gave rise to a product of ~350 bp in reactions with every κ and λ gene family specific primer, which indicated that a polyclonal population of B cells was present in the specimen. Therefore, we concluded that the PCR method and family specific primers that we were using were suitable for detecting Ig gene rearrangements in single H/R-S cells. No bands were seen in the negative control.

IMMUNOHISTOCHEMISTRY
Both light chain proteins and the CD30 antigen were demonstrated in H/R-S cells from each case of classic Hodgkin’s disease (mixed cellularity-type and nodular sclerosis-type). Only the κ light chain was found in H/R-S cells of lymphocyte predominant Hodgkin’s disease (fig 2).

PCR ANALYSIS OF SINGLE H/R-S CELLS
PCR specific products of the IgH, κ, and λ genes were not amplified from all single cells. There are several possible reasons for this (which might prove to be methodological limitations): (1) part of the cell nucleus may have been missing; the larger the cell, the lower the likelihood of obtaining the entire cell nucleus; (2) certain unknown V gene rearrangements might not be detected with the primers used; (3) DNA might be destroyed in the histological tissue processing; (4) some genes might have the germline configuration; and (5) the chosen primers might not cover all regions of the V gene. For example, FRIIIa is designed for the third framework region of Vκ, it hybridises to the 3’ end of most Vκ genes. However, with this primer, no information can be obtained about the particular Vκ genes rearranged, such as FRIJ/H, FRII/JH rearrangements and even bcl-2/JH translation. The sensitivity in our study was ~35.8%.

Unlike the method proposed by Küppers et al, in our method we added β-globin gene primers to the first round of amplification to determine the usefulness of performing the next round of amplification. We showed that it was not necessary to carry out the next round of amplification if the β-globin gene specific product was not detected in the first round (its presence indicates that DNA is available from the single cell), thereby reducing the number of PCR reactions that were carried out, and economising on reagents and time.

Fifty two of 135 single H/R-S cells isolated showed specific products in PCR reactions (36%). In the IgH gene, all the amplification products of single cells from a given biopsy specimen were the same length (80 bp). In the Igκ and λ light chain genes, the lengths of the products were also the same (350 bp).

In our study, many individual H/R-S cells were isolated from frozen sections. In most cases, many H/R-S cells from the same specimen produced a PCR product of the same size (80 bp for IgH, 350 bp for the light chain gene family). Therefore, there is a strong likelihood that these H/R-S cells are a clonal expansion or a monoclonal B cell population.

In addition, IgH and Vκ family products were found repeatedly in many H/R-S cells from one case of lymphocyte predominant Hodgkin’s disease. In two cases of nodular sclerosis-type Hodgkin’s disease, repeated Vκ and individual IgH/Vκ products and individual IgH and Vκ/Vκ family products were seen (fig 3). In five cases of mixed cellularity-type Hodgkin’s disease, repeated IgH/Vκ, and individual Vκ/Vκ, repeated Vκ, and individual IgH/Vκ, and repeated IgH and individual Vκ/Vκ family products were detected, respectively (fig 4).

In patient 3 (nodular sclerosis-type Hodgkin’s disease) and patient 7 (mixed cellularity-type Hodgkin’s disease), one H/R-S cell had two light chain gene family rearrangements (Vκ and Vλ family in patient 3, Vκ and Vλ family in patient 3, Vκ and Vλ family in patient 7).

Figure 2 Patient 1 (lymphocyte predominant Hodgkin’s disease). A typical Hodgkin cell is seen in the middle of the section, with the cytoplasm immunostained positively for the κ light chain. Avidin–biotin complex stained; magnification ×1000.
Lymphocyte predominant Hodgkin’s disease is a distinctive and rare form of the disease that is clinically indolent and has an excellent prognosis. The presence of H/R-S cells among a background of small lymphocytes and histiocytes has recently been classified as a special entity, distinct from classic Hodgkin’s disease. H/R-S cells of lymphocyte predominant Hodgkin’s disease have special immunophenotypic and genotypic characteristics, different from those of other subtypes of Hodgkin’s disease. Expression of CD19, CD20, and CD45R, but not of CD15 and CD30, is common. The detection of restricted Ig light chain mRNA supports the theory that lymphocyte predominant Hodgkin’s disease is a monoclonal proliferation of B cells. In many cases of lymphocyte predominant Hodgkin’s disease, H/R-S cells with the molecular features of a monoclonal population of B cells were found, which were negative for Epstein-Barr virus (EBV), except in a few cases. However, some investigators have reported evidence of the B cell origin of H/R-S cells and the monoclonal nature of the disease. Our experiments corroborate the view that H/R-S cells of lymphocyte predominant Hodgkin’s disease originate from B lymphocytes because of the IgH and \( \kappa \) light chain gene rearrangements revealed in individual H/R-S cells of this type of Hodgkin’s disease. These cells might represent a separate B cell clone, indicating a polyclonal population.

During B cell development, an IgH gene undergoes recombination through a joining of the V, D, and J regions, giving rise to a three-part \( V_{\mu}D_{\mu}J_{\mu} \) unit, which is called the complementary determining region III, and is a clonal signature of an individual B cell. If PCR is applied across this region, the lengths of the reaction products of different clonal B cells tend to differ. If single H/R-S cells from different areas of a specimen from a patient with Hodgkin’s disease have the same size PCR products, we can assume that these H/R-S cells originate from the same B cell clone. However, the clonality of H/R-S cells is discordant. Hansmann’s group studied 14 cases of Hodgkin’s disease, mainly of the two subtypes in mixed cellularity and nodular sclerosis types of Hodgkin’s disease have been identified variously as B or T lymphocytes, monocytes, and even as dendritic cells. The major difficulty in determining the origin of these malignant cells lies in the disturbance caused by background cells. However, the “molecular histology” technique proposed by Küppers et al enables one to pick out individual H/R-S cells with typical morphology and surface markers in a histological section of tissue affected by Hodgkin’s disease and determine whether the monoclonal or polyclonal rearrangements are derived from H/R-S cells or from other background cells in the specimen. Using this approach, ourselves and other workers have shown that Ig gene rearrangements occur in single H/R-S cells, so that it has gradually become clear that B cell malignant transformation occurs in these cells.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Picked cells</th>
<th>( \beta )-globin positive cells</th>
<th>No of cells with rearrangements of Ig gene</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (LPHD)</td>
<td>14</td>
<td>8</td>
<td>2, 2</td>
</tr>
<tr>
<td>2 (NSHD)</td>
<td>14</td>
<td>7</td>
<td>1, 1, 1</td>
</tr>
<tr>
<td>3 (NSHD)</td>
<td>14</td>
<td>8</td>
<td>1, 2, 1</td>
</tr>
<tr>
<td>4 (MCHD)</td>
<td>26</td>
<td>21</td>
<td>6, 1</td>
</tr>
<tr>
<td>5 (MCHD)</td>
<td>14</td>
<td>7</td>
<td>1, 2, 1</td>
</tr>
<tr>
<td>6 (MCHD)</td>
<td>9</td>
<td>5</td>
<td>2</td>
</tr>
<tr>
<td>7 (MCHD)</td>
<td>48</td>
<td>42</td>
<td>14, 2, 1</td>
</tr>
<tr>
<td>8 (MCHD)</td>
<td>4</td>
<td>4</td>
<td>1</td>
</tr>
</tbody>
</table>

Illustration: for example, in patient 4, a total of 28 H/R-S cells were isolated. In the first round of amplification, \( \beta \)-globin specific product was detected in 21 cells, of which six cells revealed IgH rearrangement, one cell revealed \( V_{\gamma} \) family rearrangement, and one cell revealed \( V_{\mu} \) family rearrangement in the second round. And the like. LPHD, lymphocyte predominant Hodgkin’s disease; MCHD, mixed cellularity Hodgkin’s disease; NSHD, nodular sclerosis-type Hodgkin’s disease.
Ig gene rearrangements in single H/R-S cells

which more H/R-S cells are present. In 13 of these cases, all the H/R-S cells from a given biopsy specimen had the same, or very similar Ig gene rearrangements. These cells, therefore, were B cells derived from a single clone. The related and unrelated pattern of IgH gene rearrangements in 12 patients were observed by Hummel et al. They gave rise to a possibility of monoclonal and polyclonal H/R-S cells in the tumour tissues, although cellular contamination might account for these results. On the other hand, classic Hodgkin’s disease with cells coexpressing CD20 and CD30 accounts for only 5–30% of all cases of Hodgkin’s disease and the results of Hummel et al are therefore not relevant to most cases of Hodgkin’s disease. 

In our series, patients 1, 4, 6, and 7 had many cells expressing the same size of IgH rearrangement (80 bp). In addition to IgH, κ and λ light chain gene rearrangements were detected in all patients except patient 8. The same light chain gene rearrangements were present simultaneously in two individual H/R-S cells from the same specimen (Vκ in patients 3 and 5; Vλ in patient 7). This result was confirmed by identifying identical rearrangements in cells from different sections. However, the family of different light chain gene rearrangements was also detected in cells from the same specimen (patients 2, 4, 5, and 7). Thus, it is likely that these cells in our series represented a different clonal B cell population at various stages of development. These H/R-S cells in a given biopsy specimen might be descendants of a single B cell and therefore constitute a monoclonal population, or each H/R-S cell might be from a separate B cell clone.

Why does light chain restriction almost always involve the κ but not the λ light chain? During B cell development, functional heavy chain gene rearrangement is followed by κ gene rearrangement. If κ gene rearrangement fails, the cell may proceed with λ gene rearrangement to yield a functional λ light chain gene. In humans, ~40% of B cells express the λ light chain. For this reason, at least some H/R-S cells should have rearranged λ light chain genes. Our study showed this to be true: in addition to Vκ, (patients 2 and 5) and Vλ family rearrangement, Vκλ, two different family rearrangements, which might be on just two chromosomes or on more than two chromosomes in hyperplid H/R-S cells, were detected.

There are still many questions to be answered about Hodgkin’s disease, including: why H/R-S cells do not behave like EBV transformed B cells or the cells in any common B cell non-Hodgkin’s lymphoma; how H/R-S cells stay alive; and the clonality of H/R-S cells. It is hoped that more carefully planned studies in the future will provide new insights that will clarify this enigmatic disease.

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1 Drexler HG. Recent result on the biology of Hodgkin and Reed-Sternberg cells. I. Biopsy material. Leuk Lymphoma 1992;2:283–313.
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