Identification of Hodgkin and Reed-Sternberg cell-specific genes by gene expression profiling

Ralf Küppers,1,2 Ulf Klein,3 Ines Schwering,1,2 Verena Distler,1,4 Andreas Bräuning,4 Giorgio Cattoretti,3 Yuhai Tu,5 Gustavo A. Stolovitzky,5 Andrea Califano,6 Martin-Leo Hansmann,4 and Riccardo Dalla-Favera3

1Institute for Genetics, and
2Department of Internal Medicine I, University of Cologne, Cologne, Germany
3Institute for Cancer Genetics, Columbia University, New York, New York, USA
4Department of Pathology, University of Frankfurt, Frankfurt/Main, Germany
5IBM T.J. Watson Research Center, Yorktown Heights, New York, New York, USA
6First Genetic Trust Inc., Lyndhurst, New Jersey, USA

Hodgkin lymphoma (HL) is a malignancy of unknown pathogenesis. The malignant Hodgkin and Reed/Sternberg (HRS) cells derive from germinal center B cells (or rarely, T cells) but have a heterogeneous and largely uncharacterized phenotype. Using microarrays, we compared the gene expression profile of four HL cell lines with profiles of the main B cell subsets and B cell non-HLs to find out whether HRS cells, despite their described heterogeneity, show a distinct gene expression, to study their relationship to other normal and malignant B cells, and to identify genes aberrantly or overexpressed by HRS cells. The HL lines indeed clustered as a distinct entity, irrespective of their B or T cell derivation, and their gene expression was most similar to that of EBV-transformed B cells and cell lines derived from diffuse large cell lymphomas showing features of in vitro-activated B cells. Twenty-seven genes, most of which were previously unknown to be expressed by HRS cells, showed aberrant expression specifically in these cells, e.g., the transcription factors GATA-3, ABF1, EAR3, and Nrf3. For five genes, expression in primary HRS cells was confirmed. The newly identified HL-specific genes may play important roles in the pathogenesis of HL, potentially represent novel diagnostic markers, and can be considered for therapeutic targeting.


Received for publication August 9, 2002, and accepted in revised form December 17, 2002.

Address correspondence to: Ralf Küppers, University of Cologne, Department of Internal Medicine I, LFI E4 R706, Joseph-Stelzmannstrasse 9, D-50931 Cologne, Germany. Phone: 49 221 478 4490; Fax: 49 221 478 6383; E-mail: ralf.kuppers@uni-koeln.de.

Ralf Küppers, Ulf Klein, and Ines Schwering contributed equally to this work.

Conflict of interest: The authors have declared that no conflict of interest exists.

Nonstandard abbreviations used: Hodgkin lymphoma (HL); Hodgkin and Reed/Sternberg (HRS); germinal center (GC); lymphocyte predominant (LP); B cell non Hodgkin lymphoma (B-NHL); diffuse large cell lymphoma (DLCL); Burkitt lymphoma (BL); follicular lymphoma (FL); B cell chronic lymphocytic leukemia (B-CLL); lymphoblastoid cell lines (LCL).
Based on the GC B cell origin of HRS cells in most cases, it is reasonable to perform the analysis of differential gene expression between these cells and normal mature B cells as well as other types of B cell non-Hodgkin lymphomas (B-NHLs). Since it is not yet technically feasible to create large-scale gene expression profiles from the rare primary HRS cells, we generated the gene expression profiles (approximately 9,000 genes) of a number of HL-derived cell lines and compared them with the main subsets of normal mature B cells and various types of B-NHLs. Expression of several markers identified in the analysis was confirmed in HL cell lines at the protein level and in primary HRS cells by RT-PCR. With the analysis, we wanted to address three main questions: (a) Given the reported phenotypic heterogeneity of HRS cells, do the gene expression profiles define HRS cells as a distinct entity? (b) To which other normal or malignant B cells are HRS cells most closely related in terms of gene expression? (c) Can we identify HRS cell–specific genes? With this last aspect, we aimed to identify genes involved in the pathogenesis of HL and genes that may represent potential diagnostic or therapeutic markers.

Methods

Cell lines and culture conditions. The HL cell lines used originate from patients with HL of the nodular sclerosis (L428 and HDL2M) or mixed cellularity subtype (KMH2 and L1236). HDL2M is of T cell origin, and the three other HL cell lines are of B cell origin, as shown by Southern blot analysis of antigen-receptor gene rearrangements (13–15). All cell lines were grown in RPMI-1640 with GlutaMAX-I (Invitrogen, Karlsruhe, Germany), supplemented with 10% FCS and 100 U/ml penicillin/streptomycin at 37°C in an atmosphere containing 5% CO2.

Purification of primary cells. The isolation of the primary cells used in this analysis has been described in detail elsewhere (16). Briefly, normal B cell subsets were isolated by magnetic cell separation using the MACS system (Miltenyi Biotech, Bergisch Gladbach, Germany) according to the following marker expression: naive B cells (CD27–, CD10–, CD3–, CD38low, CD14–, IgD–), centroblasts (CD77–, CD38high, CD39–, CD3–, CD10+), and memory B cells (CD10–, CD3–, CD38low, CD14–, CD27+). Because gene expression profiles for CD77+ GC B cells and CD77– GC B cells showed surprisingly few differences (17), these cell populations were collectively designated GC B cells. An explanation for the few differences may be that CD77 staining does not completely distinguish between these two cell populations, although it is commonly used as a centroblast marker.

Generation of cRNA and microarray hybridization. Microarray hybridization was performed as described (16), starting from 5 µg of purified total RNA and hybridizing each 15-µg biotin-labeled cRNA to U95A microarrays (Affymetrix Inc., High Wycombe, United Kingdom). After scanning, the expression values for the genes were determined using Affymetrix Microarray Suite 4.0 software and its Global Scaling option. The expression data (average differences) were processed as follows: small and negative expression levels were clipped-off to be equal to a cutoff value arbitrarily chosen as 20. The logarithm of this clipped-off data was subsequently used throughout the analyses.

Biostatistical analysis. The hierarchical clustering algorithm used to generate the dendrogram is based on the average-linkage method (18). To construct the dendrogram, a subset of genes on the microarray was used, whose expression levels vary the most among the 23 samples and thus are most informative (see ref. 16). The expression values of each selected gene is normalized to have zero mean and unit SD. The distance between two individual samples is calculated by Pearson distance with the normalized expression values.

For supervised hierarchical clustering, we used the Genes@Work software platform, which is a gene expression analysis tool based on the pattern discovery algorithm SPLASH (structural pattern localization analysis by sequential histograms) (19). For details, see ref. 16. The primary data are available through http://ICG.cpmc.columbia.edu/faculty.htm.

Western blotting. Immunoblotting was performed using standard methods. The amount of protein was quantified using the DC Protein Assay (Bio-Rad Laboratories, Munich, Germany). The following Ab’s were used: anti-Neogenin, sc-6536; anti-Gas1, sc-9585; anti-GATA-3, sc-268 (all from Santa Cruz Biotechnology Inc., Santa Cruz, California, USA); anti-ABF1 antigen (a kind gift of C. Murre).

Flow cytometry. Cells were incubated with anti-IL-1R2 Ab (HyCult Biotechnology BV, Uden, the Netherlands), followed by staining with FITC-coupled anti-mouse Ab’s and analyzed on a FACSCalibur (Becton Dickinson Biosources, Franklin Lakes, New Jersey, USA).

Characteristics of cases used for RT-PCR analysis of microdissected HRS cells. Lymph nodes of seven classical and two LP HL patients were analyzed. The biopsy from patient 4 was obtained at relapse; the lymph nodes of the other classical HL patients were from first presentation (no information is available for the two other patients).

Staining of frozen sections and microdissection. Five-micrometer-thick frozen sections were mounted on membrane-covered slides and incubated with hematoxylin containing RNase inhibitor (200 U/ml; Roche, Mannheim, Germany) for 4 min. Sections were washed in diethylpyrocarbonate-treated water for 2 min, incubated in 2% eosin for 15 s, washed again, and then dried at 37°C for 3 h. HRS and non-HRS cells were catapulted into 20 µl of Purescript lysis buffer (Biozym, Hamburg, Germany) using an UV-laser beam (PALM, Bernried, Germany) (20) and pooled in groups of 25 or 50 cells.

RNA isolation, cDNA synthesis, and amplification. The Purescript RNA Isolation Kit (Biozym) was applied using glycogen as a carrier and reducing all reagents to one-tenth of the amounts given in the standard protocol. In the analysis of rab13 expression, a DNase I digestion (Stratagene, La Jolla, California, USA) was performed.
performed because of the existence of processed pseudogenes. Isolated RNA and 0.6 \( \mu \)M of gene-specific reverse primer (PRAMErev, 5'-AGG GCA AGG AGC TGA TCA TCC G-3'; IPLUTRev, 5'-CTA GCC TTCG TCC GAC TCG TCC-3'; rab13rev, 5'-CTC ATC CGT GAT GTC GTA TAC TAG G-3'; EAR3rev, 5'-TGG ATT GGG CTG GGT TGG AGG C-3'; FERrev, 5'-CGG ACA AAC CCC TAA GCT GAA GG-3') were heat denatured at 70°C for 10 min.

T4 gene 32 protein (Ambion Inc., Austin, Texas, USA) was added at a concentration of 1.5 \( \mu \)g per sample during the initial denaturation. Reverse transcription was performed using the OneStep RT-PCR Kit components (QIAGEN Inc., Hilden, Germany), including 10 U of RNase inhibitor. The reaction mixture was preheated to 50°C for 2 min, after which 2 \( \mu \)l of QIAGEN OneStep RT-PCR enzyme mix was added. The cDNA synthesis was performed at 50°C for 30 min.

**PCR amplification of cDNA products.** One round of PCR was performed applying the reverse primers used for cDNA synthesis and the corresponding forward primers (PRAMEfor, 5'-GTA TAT CGC CCA GTT CAT C-3'; IPLfor, 5'-GAG CCC TCG GAG CCC TCC AGG-3'; rab13for, 5'-GCT TGA TCC TTC GTT CAG AGG-3'; EAR3for, 5'-CCA ATT CAC CTG CGA GGG CTG C-3'; FERfor, 5'-GCC AGA AAC TGC CTG GTG-3'). All primer pairs were intron-spanning. For amplification of a 352-bp product of the PRAME gene, samples were adjusted to 50-\( \mu \)l reaction mixtures supplemented with 2 \( \mu \)M MgCl\(_2\), 200 \( \mu \)M dNTP, and 0.125 \( \mu \)M of each primer, and the HotStarTag DNA polymerase system was used. Cycling conditions were 15 min at 95°C, 60 s at 65°C, 60 s at 72°C, and 39 cycles of 30 s at 95°C, 30 s at 65°C, and 45 s at 72°C, followed by a final 5-min incubation at 72°C. For analysis of PRAME in patients 4, 6, 8, and 9, a second round of PCR (additional 30 cycles with the same PCR conditions) was needed. A seminested PCR was performed using an internal reverse primer (PRAMErev, 5'-GTT TTC CAA GGG GTT CAT CAC G-3') and the forward primer. For amplification of a 98-bp fragment of IPI, PCR conditions were as described above (except the annealing temperature was 67°C and MgCl\(_2\) concentration was 1.5 mM). Two rounds of PCR were performed to analyze the expression of rab13, EAR3, and FER. The first round comprised 20 cycles; the second round of PCR, applying the corresponding internal forward primers (rab13for, 5'-CTA GAA GCT TGG GAC ACG GCT GG-3'; EAR3for, 5'-CTT ACA CAT GCC GTG CCA ACA AGG-3'; FERfor, 5'-CAT TAA ATG GAC AGC ACC GGA AGC-3'), comprised 30 cycles. For amplification of a 211-bp fragment of rab13 in the first round, PCR conditions were as described above, except the annealing temperature was 61°C, and the MgCl\(_2\) concentration was 1.5 mM. In the second round a 104-bp fragment was amplified (annealing temperature 61°C, MgCl\(_2\) concentration 2 mM). For amplification of a 210-bp fragment of EAR3 or a 227-bp fragment of FER, in the first-round PCR conditions were as described above. In the second round a 151-bp fragment of EAR3 or a 114-bp fragment of FER was amplified. PCR conditions were as described above (MgCl\(_2\) concentration was 2.5 mM for the analysis of FER). Samples of HRS cells and control cells were analyzed in parallel, together with controls containing only Purescript lysis buffer. Some PCR products were gel purified and directly sequenced. No processed pseudogenes for PRAME and IPL were detectable in the patients' genome.

**Results**

Gene expression profiling of HL lines defines them as a distinct entity that is related to EBV-transformed B cell lines and a subset of diffuse large cell lymphoma cell lines. Four classical HL cell lines were used for the present study. Except for HDLM2, which is of T cell origin, the other three lines are B cell derived. To generate gene expression profiles, Affymetrix U95A oligonucleotide microarrays comprising approximately 12,000 human mRNAs were used. The data sets were compared with the profiles of a large collection of normal B cell subsets (GC B cells, naive and memory B cells), B cell malignancies (diffuse large cell lymphoma; DLCL), Burkitt lymphoma (BL), follicular lymphoma (FL), and B cell chronic lymphocytic leukemia (B-CLL), as well as EBV-transformed peripheral blood B cell lines (LCL) and BL and DLCL cell lines. Unsupervised hierarchical clustering revealed that the gene expression profiles of the four HL cell lines were more similar to each other than to any other normal or malignant B cell (data not shown), implying that, with regard to their gene expression, the HL cell lines represent a distinct group among the various B cell types analyzed.

The unsupervised clustering divided the 71 samples into two main groups, one comprising the normal B cell subsets and the tumor biopsies, the other the cell lines (not shown). To more specifically determine the relatedness of the HL lines to the other cell lines, we examined the corresponding gene expression profiles of the four HL cell lines, five LCLs, six BL and eight DLCL cell lines by unsupervised analysis (Figure 1). The analysis revealed that the HL lines, the LCLs, and two DLCL lines that have been proposed to be derived from a subset of DLCL that shows features of in vitro–activated B cells (ABC-type DLCL) (21, 22) together form a separate branch. In the second branch, the BL lines appear intermingled with the remaining DLCL cell lines, four of which have been assigned to a subset of DLCL that resembles GC B cells (GC-type DLCL) (21, 22). It should be noted that the EBV-status of the cell lines has no influence on the clustering: the EBV- BL lines cluster separate from the (EBV+) LCL, and the four HL cell lines are EBV-

We next examined which set of genes distinguishes the LCLs, HL and ABC-type DLCL cell lines from the group comprising the BL and the remaining DLCL cases. To this end, we employed supervised clustering using the Genes@Work software platform (18; and see ref. 16). Figure 2 shows that 37 genes are differentially expressed in the 11 LCLs, HL and ABC-type DLCL lines versus the 12 BL and DLCL cell lines. Consistently overexpressed in the first group were, for example, eight genes being components of an in vitro–activated B cell signature, namely cyclin D2, CD44, IRF-4/MUM-1, CCR7,
The gene expression profile of HL lines is related to that of LCL and ABC-type DLCL. Dendrogram showing the hierarchical clustering of gene expression data (see Methods for algorithm and criteria) generated from 23 transformed B cell lines derived from HL, DLCL, BL, and EBV-transformed peripheral blood B cells (LCL). Cell lines are color-coded according to their cellular origin: DLCL, red; BL, blue; HL, green; LCL, violet. If known, the EBV status is indicated in brackets. ABC- or GC-type DLCL-subtypes are indicated. The corresponding matrix is shown in supplementary Figure 1 (http://www.jci.org/cgi/content/full/111/4/529/DC1).

InBct, CLARP (c FLIP), TNF, and activated RNA polymerase II transcriptional cofactor 4 (21, 22). On the other hand, mRNA encoding the GC B cell marker CD10 was upregulated in the BL and GC-type DLCL, consistent with their proposed derivation from GC B cells. Differential expression of the BCL-2 and BCL-6 genes between the two groups could be detected by relaxing the criteria of the supervised analysis (see bottom of Figure 2). Overall, BCL-2 and BCL-6 showed a reciprocal expression pattern between the two groups of cell lines, with low BCL-2 and high BCL-6 expression in the cell lines with a GC phenotype, as expected.

The results of the clustering analyses show that HL lines represent a distinct entity independent from a B or T cell derivation and imply that the HL, LCL, and ABC-type DLCL lines express a common set of genes that distinguish them from cell lines that are derived from tumor types likely corresponding to transformed GC B cells.

Identification of genes specifically expressed in HL cell lines. To identify genes specifically up- or downregulated in HRS cells, supervised clustering was used to compare the gene expression profile of the four HL cell lines with those of the normal B cell subsets (naive, memory, centroblasts, centrocytes), the LCL, and the various B cell NHL (FL, BL, DLCL, and B-CLL). As shown in Figure 3, 27 genes are specifically and consistently upregulated in the HL cell lines, and 45 genes are downregulated. Three of the genes upregulated in HRS cells are still unnamed and/or only described as cDNA sequences. Among the 24 distinct named genes upregulated in HRS cells, two genes were found that are well known for their expression in HRS cells, namely the actin-bundling protein Fascin and the chemokine TARC (9, 10). Also, expression of the TNF receptor family member RANK and the metalloproteinase TIMP1 by HRS cells has been described recently (23, 24). The HRS cell marker CD30, another TNF receptor family member, was not identified as an upregulated gene with the stringent criteria used here, due to its expression in LCL and low-level transcription in some B-NHL.

The other 20 known upregulated genes have not been previously reported to be expressed in HRS cells. These 20 genes include four transcription factors, namely GATA-3, ABF1, Nrf3, and EAR3, the imprinted gene IPl, the tumor antigen PRAME, the tyrosine kinase FER, two genes implicated in growth arrest (Hep27 and Gas1), the decoy receptor IL-1R2, sarcolectin, which stimulates DNA synthesis, and the GTPase Rab13 (Figure 3).

Somewhat surprisingly, the majority of the 27 genes identified here were not found in a previous study comparing HRS and GC B cells (11). Limited sensitivity could at least partly explain this finding, because the cDNA library of HRS cells was restricted to 6,000 transcripts in that study.

The genes downregulated in HRS cells include many B cell lineage-specific genes (e.g., CD19, CD20, CD52, BCMA), two phosphatases (SBF1 and PTPN7), and several genes involved in the regulation of the cytoskeleton and/or cell migration (HSRHO2, MSF, Vanin2) (Figure 2). A detailed analysis of the downregulation of B lineage markers in HL cell lines is presented elsewhere (25).

Protein expression of HRS cell–specific genes in HL cell lines. To investigate whether the upregulated mRNA levels of HRS cell–specific genes correspond to elevated protein levels, we investigated HL cell lines for protein expression of these genes. ABF1, neogenin, Gas-1, and GATA-3 were analyzed by immunoblotting (see supplementary Figure 2, http://www.jci.org/cgi/content/full/111/4/529/DC1). The expression of IL-1R2 was evaluated by FACS analysis.

ABF1 protein is present in three out of four HL cell lines (L1236, L428, KM12), HDLM2 being negative or below detection limit for ABF1 expression. Two BL cell lines and Jurkat as a T cell line do not show ABF1 expression. The N-cam family member neogenin is strongly expressed in the L1236, L428, and KM12 cell lines. Only very low protein levels were observed in the HDLM2 cell line, in two BL cell lines, and in the Jurkat cell line. No significant difference in Gas-1 protein level was detected comparing the four HL cell lines with BL2, BL41, and Jurkat, suggesting that the increased
mRNA levels for this gene in HL lines does not result in increased protein levels. The T cell–specific transcription factor GATA-3 is present in Jurkat cells, as expected, but also in L1236, L428, and KM2. The HDLM2 and the two BL lines seem to be negative. Thus, for three of the four genes analyzed, protein expression could be demonstrated in at least three of the four HL lines. The lack of GATA-3 expression in the HDLM2 cell line might be surprising, but is in line with previous data showing that T cell–derived HL cases may lack expression of individual T cell markers (4, 26).

FACS analysis revealed that IL-1R2 protein expression can be detected only in the L1236 cell line, the three other HL cell lines being negative or below detection level (not shown), although high mRNA levels were detected in all cell lines in the array analysis.

**Analysis of primary HL cases by RT-PCR of microdissected HRS cells.** Initially, we tried to confirm protein expression of several upregulated genes by HRS cells in tumor tissues by immunohistochemistry on frozen and paraffin-embedded tissues. Unfortunately, none of the available Ab’s tested (against GATA-3, ABF1, EAR3, neogenin, GAS1, and IL-1R2) stained reliably on tissue sections. For five of the upregulated genes, namely PRAME, IPL, FER, Rab13, and EAR3, expression in HRS cells in the tissue was analyzed by RT-PCR of microdissected HRS cells, using a laser-based microdissection system. Single HRS cells from seven cases of classical

---

**Figure 2**

Identification of genes differentially expressed between HL lines, LCL, and ABC-type DLCL lines versus BL and GC-type DLCL. The gene expression profiles of four HL, five LCL, and two ABC-type DLCL lines that clustered together in the unsupervised analysis (see Figure 1) were compared with those generated from six BL and six DLCL (non–ABC-type) cell lines by supervised hierarchical clustering using Genes@Work (see Methods). Columns represent individual cell lines, and rows correspond to genes. Color changes within a row indicate expression relative to the average of the sample population. Values are quantified by the scale bar that visualizes the difference in the ζ-score (expression difference/SD) relative to the mean. Genes are ranked based on the ζ-score (mean expression difference of the respective gene between phenotype and control group/SD). The support value for supervised analysis was chosen as $n = n_0 - 1$, where $n_0$ is the number of cells in the given phenotype set, allowing for one unclustered sample per pattern in the phenotype set. The corresponding expression data for BCL-2 and BCL-6 were obtained by relaxing the criteria in the supervised clustering and are therefore shown separately at the bottom. Gene names and cell lines are indicated.
HL were pooled in groups of 25 or 50 cells, and three such samples per case were analyzed in parallel with groups of non-HRS cells (mainly lymphocytes) from the same sections. For each of the five genes, all or at least most samples of HRS cells per case were positive in the PCR (except PRAME in cases 4 and 6, and EAR3 in cases 1 and 7), including EBV-positive and -negative cases, while nearly all control samples were negative (Table 1). We also analyzed two cases of LP HL, revealing that four of the five genes are also consistently transcribed by HRS cells of this type of HL (Table 1). Thus, we confirmed expression of five genes in primary HRS
cells, further supporting that the results obtained with the HL cell lines reliably reflect the gene expression profile of HRS cells in HL tissues.

Discussion

In the present study, large-scale gene expression profiling was performed to compare the gene expression profiles of HL cell lines with those of the main normal B cell subsets and B-NHL. A concern might be that the HL cell lines differ in several aspects from primary HRS cells in the tissue. Nevertheless, for several reasons we are confident that HRS cell-specific genes can be reliably identified using HL cell lines. First, a number of genes identified here as HRS cell specific are known to be expressed by HRS cells, i.e., Fascin, TARC, RANK, and TIMP1. The identification of these markers hence supports the reliability of the approach. Second, by including a number of other B cell–derived cell lines in the comparison (LCL, BL lines, DLCL lines), common putative cell culture–associated changes in gene expression will not be falsely scored as being HRS cell–specific. Third, for five of the upregulated genes, expression in primary HRS cells in the tissue was confirmed by analyzing microdissected HRS cells.

Gene expression profiling identifies HL cell lines as a distinct entity. Unsupervised hierarchical clustering revealed that the HL cell lines are more similar to each other than to any other normal and malignant B cell subset (Figure 1). Since one of the four lines (HDLM2) is of T cell origin, this finding may indicate that the development of HRS cells results in a particular gene expression profile largely independent from their derivation of either B or T cells. This raises the intriguing possibility that—opposed to most other lymphomas—the cell of origin is not the main determinant for the phenotype of the HRS cells, but that B cell–derived as well as T cell–derived HRS cells undergo (common?) transforming events resulting in an HRS cell–typical gene expression signature.

HL lines display an activated B cell signature similar to LCL and ABC-type DLCL. Hierarchical clustering of the data sets from the HL lines and the various B cell–derived lines revealed two major branches in the corresponding dendrogram, one comprising the BL as well as GC-type DLCL lines, the other containing the HL lines, LCLs, and the recently described ABC-type DLCL lines (Figure 1). BL and the GC-type DLCL are thought to represent transformed GC B cells, which is also supported by the finding that both entities show active somatic hypermutation (27, 28). The present study shows that HL cell lines, although (in most cases) also derived from GC B cells, do not show a GC B cell phenotype. This was already indicated from the lack of several typical GC B cell markers (e.g., BCL-6, CD10) (29, 30) and the lack of ongoing hypermutation in HRS cells (5, 31, 32). The lack of the GC B cell phenotype may be part of a general loss of the B lineage phenotype of HRS cells, as reported recently (25). It is an open question whether this loss of the GC B cell phenotype is somehow related to the proposed origin of HRS cells from preapoptotic GC B cells, is associated with the phenotypic changes during malignant transformation, or reflects, at least in part, a stage of GC B cell development where GC B cell markers are downregulated.

The distinction between GC- and ABC-type DLCL was based on the observation that a GC and an (in vitro–) activated B cell signature can be defined by genes specifically upregulated in these cell types and that DLCL can be clustered according to either of these signatures (21). Notably, in contrast to the GC signature, which includes several GC B cell–specific genes, the ABC-type signature is mainly composed of general activation markers (21). Most of the 37 genes that differentiate HL lines, LCL, and ABC-type DLCL lines from BL lines and GC-type DLCL lines showed upregulation in the first group of cell lines (Figure 2). Eight of these upregulated genes are part of the ABC signature. Hence, HL lines show an activated phenotype unrelated to the GC B cell phenotype. A part of this

---

### Table 1

RT-PCR analysis of microdissected HRS cells for expression of PRAME, IPL, FER, Rab13, and EAR3

<table>
<thead>
<tr>
<th>Case</th>
<th>Subtype</th>
<th>EBV status</th>
<th>Samples positive/Samples analyzed PRAME</th>
<th>IPL</th>
<th>FER</th>
<th>Rab13</th>
<th>EAR3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Classical HL</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>NS –</td>
<td></td>
<td>3/3</td>
<td>3/3</td>
<td>3/3</td>
<td>3/3</td>
<td>1/3</td>
</tr>
<tr>
<td>2</td>
<td>MC +</td>
<td></td>
<td>3/3</td>
<td>3/3</td>
<td>nd</td>
<td>nd</td>
<td>nd</td>
</tr>
<tr>
<td>3</td>
<td>NS –</td>
<td></td>
<td>4/4</td>
<td>3/3</td>
<td>2/3</td>
<td>3/3</td>
<td>3/3</td>
</tr>
<tr>
<td>4</td>
<td>NS +</td>
<td></td>
<td>2/4A</td>
<td>3/3</td>
<td>3/3</td>
<td>3/3</td>
<td>3/3</td>
</tr>
<tr>
<td>5</td>
<td>NS –</td>
<td></td>
<td>3/3</td>
<td>3/3</td>
<td>2/3</td>
<td>3/3</td>
<td>3/3</td>
</tr>
<tr>
<td>6</td>
<td>MC +</td>
<td></td>
<td>0/3A</td>
<td>2/3</td>
<td>3/3</td>
<td>3/3</td>
<td>2/3</td>
</tr>
<tr>
<td>7</td>
<td>MC –</td>
<td></td>
<td>3/3</td>
<td>3/3</td>
<td>3/3</td>
<td>3/3</td>
<td>1/3</td>
</tr>
<tr>
<td>LP HL</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>LP –</td>
<td></td>
<td>1/2A</td>
<td>2/2</td>
<td>2/2</td>
<td>2/2</td>
<td>0/2</td>
</tr>
<tr>
<td>9</td>
<td>LP –</td>
<td></td>
<td>2/2A</td>
<td>2/2</td>
<td>2/2</td>
<td>2/2</td>
<td>1/2</td>
</tr>
<tr>
<td>Controls</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1–9</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Buffer controls</td>
<td>1–9</td>
<td></td>
<td>0/27</td>
<td>0/25</td>
<td>0/22</td>
<td>0/22</td>
<td>0/22</td>
</tr>
</tbody>
</table>

Groups of 25 or 50 HRS cells and non-HRS cells (small cells isolated from the same tissue sections as the HRS cells) were analyzed. Overall, 37 samples with 25 HRS cells and 81 samples with 50 HRS cells and 26 samples with 25 non-HRS cells and 92 samples with 50 non-HRS cells were analyzed. Buffer controls represent reaction tubes with buffer but without cells. The few positive control non-HRS cell samples may be due to cellular contamination of the samples by fragments from HRS cells or expression of the respective genes by some cells among the non-HRS cells. Groups of 10 FACS-sorted KMH2 cells served as positive controls. PCR for PRAME and IPL consisted of one round with 40 cycles, the three other genes were analyzed by two rounds of PCR with 20 and 30 cycles. In each of cases of 4, 8, and 9, one HRS cell sample became positive only after two rounds of amplification; control cells remained negative after two rounds of PCR. NS, nodular sclerosis; MC, mixed cellularity; nd, not done.
The Journal of Clinical Investigation | February 2003 | Volume 111 | Number 4

phenotype is based on NF-κB–regulated genes (namely cyclin D2, IRF-4/MUM-1, CCR7, JκBα, cFLIP), and HRS cells, LCL and ABC-type DLCL indeed show constitutive activity of this transcription factor (22, 33, 34).

Identification of HRS cell–specific genes. In the supervised comparison of the four HL cell lines with the collection of all other subsets, 27 distinct genes were identified as being significantly upregulated in HRS cells (Figure 2). Four of the genes (Fascin, TARC, TIMP1, and RANK) were already known from previous studies to be expressed in HRS cells, but for the other 23 genes it was not known previously that they are expressed by these cells. By immunoblotting, we showed protein expression of GATA-3, ABF1, GAS1, and neogenin. Moreover, for five genes (PRAME, IPL, Fer, Rab13, EAR3), expression by primary HRS cells was confirmed by RT-PCR on microdissected tumor cells (Table 1). As transcription of the genes was detected in EBV-positive as well as EBV-negative cases of classical HL, expression of these genes by HRS cells is independent from the EBV status of the HRS cells. Moreover, we detected transcripts of four of the five genes also in HRS cells of two cases of LP HL.

Notably, four transcription factors were identified among the upregulated genes, namely GATA-3, ABF1, Nrf3, and EAR3. GATA-3 is a T and NK cell–specific transcription factor that is essential for T cell development and involved in the regulation of T1/2 cytokine expression. It was reported that HL cell lines lack expression of GATA-3 (35); however, we detected expression of this protein in three of the four HL lines by immunoblot analysis. The ectopic expression of this T cell marker in the B cell–derived HL lines L428, L1236, and KMH2 underscores the aberrant phenotype of these cells. ABF1 is a member of the basic helix-loop-helix family of transcription factors and was found to be strongly expressed only in LCL and in vitro–activated B cells (36). ABF1 can form heterodimers with both products of the E2A gene, i.e., E12 and E47, and likely represses their activating functions (36). E2A transcription by the HL lines is detected on the array, so that ABF1 may indeed function in these cells, at least in part, by repressing E2A proteins, which are essential regulators not only during early B cell development but also in mature, activated B cells (37). However, ABF1 may also bind to DNA as a homodimer, suggesting that it may have so far not understood functions in addition to repressing E2A proteins. Nrf3 belongs to the family of transcription factors with Cap’n collar-type basic region leucine zipper domains (38). Low-level expression was detected in various tissues, including the spleen and peripheral blood leukocytes. Nrf3 heterodimerizes with small Maf proteins and binds to Maf recognition elements (MARE). The fourth transcription factor with increased expression in HL lines is EAR3, a transcriptional repressor belonging to the steroid hormone receptor superfamily (39). This repressor is mainly expressed in the central nervous system during development and counters the activation of various genes by hormones (39). The relevance of aberrant expression of these transcription factors in HRS cells remains to be determined.

The nonreceptor tyrosine kinase FER is widely expressed, but was found to be significantly upregulated in HL cell lines (Figure 3). This kinase has been shown to activate STAT-3 (40). Because constitutive activation of STAT-3 has been shown for HRS cells (41, 42), the overexpression of FER may be involved in this activation.

The list of genes showing deregulated expression in HL lines also includes the imprinted gene IPL, which shows normally high-level expression only in the placenta (43). Its function is not yet known, but a mouse gene highly similar to IPL is essential for Fas expression, pointing to a potential role of IPL in the regulation of apoptosis. Rab13 is a Ras-associated small GTPase, which functions presumably in vesicular traffic. The common tumor antigen PRAME is expressed on melanoma cells, by a fraction of myeloid and lymphoid leukemias, and by multiple myeloma cells, but only rarely by B-NHL (44–46). Because expression in normal tissues is absent or barely detectable, PRAME may represent an interesting target for immunotherapy in HL.

Implications for HL pathogenesis, treatment, and diagnosis. By comparing gene expression profiles of HL cell lines to profiles of the main mature B cell populations, LCL, and the main types of B-NHL, we identified genes that show specifically upregulated expression in HRS cells of classical HL. To our surprise, four of these markers turned out to be expressed also in HRS cells of LP HL (Table 1). This was unexpected since the tumor cells of these subtypes are well known for their phenotypic differences (2, 12). The present RT-PCR data may indicate a closer relationship of classical and LP HL than thought previously.

Of particular interest for the pathogenesis of HL may be the deregulated expression of four transcription factors. It will be interesting to study the role of these factors in HRS cells, e.g., by inactivating them in HL cell lines. The HRS cell markers identified here may become valuable tools for the differential diagnosis of HL, which is still sometimes difficult with the standard staining routinely used (1). Moreover, while nearly 90% of patients with HL have a good prognosis with current treatment regimens, the remaining patients do not respond well (47). Thus, there is a need to identify prognostic factors that allow one to identify cases with a bad prognosis, so that the treatment for these patients can be adjusted accordingly. The HRS cell–specific genes identified here may represent good candidates for biological prognostic factors. Finally, some of the aberrantly expressed genes, such as the common tumor antigen PRAME, may become targets for immunotherapy of HL.

Taken together, by comparing large-scale gene expression profiles of four HL cell lines to a large number of normal and malignant B cells, we show that (a) HL
lines represent a distinct entity, irrespective of the cellular origin of the HRS cells; (b) HL lines are most similar to LCL and ABC-like DLCL lines, suggesting an activated lymphocyte phenotype; and (c) over 20 genes are specifically upregulated in HRS cells, which may play an important role in the pathogenesis of HL.

Acknowledgments

We are grateful to V. Miljkovic, M. Fahrig, J. Jesdinsky, and Y. Blum for technical assistance. We thank C. Murre for supplying anti-ABF1 antiserum. This work was supported through the Deutsche Forschungsgemeinschaft by SFB502 and a Heisenberg Award to R. Küppers. U. Klein was recipient of a fellowship granted by the Human Frontiers Science Program. G. Cattoretti is a recipient of an Esther Aboodi Associate Professorship.


