## BRIEF COMMUNICATION

## Clonal Loss of Heterozygosity in Microdissected Hodgkin and Reed-Sternberg Cells

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The typical Hodgkin and Reed-Sternberg (HRS) cells are thought to represent the malignant cellular elements of Hodgkin's disease (HD). The detection of immunoglobulin gene rearrangements in HRS cells indicates that they are clonally derived from B cells (1–8), but immunogenotyping, as such, does not provide any information on specific gene alterations possibly involved in the molecular pathology of HD. In many tumors, highly informative polymorphic DNA markers may identify loci harboring clonal loss of heterozygosity (LOH) and thus help to trace tumor suppressor genes (9,10). Although in HD, cytogenetic data suggest that nonrandom chromosomal deletions may occur at several loci, including 1q42, 4q26, and others, very little (if anything) is known about clonal LOH in HRS cells at the molecular level (11-15). We, therefore, set out to study microdissected HRS cells from classical types of HD at candidate loci for LOH with a highly sensitive microsatellite polymerase chain reaction (PCR) assay.

In seven patients with classical HD, HRS cells and surrounding cells including bystander lymphocytes were laser microdissected from formalin-fixed, paraffin-embedded tissue sections (Table 1). From a patient with nodular-sclerosing HD (patient 6), samples were taken at presentation (6a) and at relapse (6b). In two patients (patient 6 and patient 7, a patient with mixed cellularity-type HD), frozen sections of lymphoma tissue yielded high-molecular-weight control DNA from microdissected cell populations. Buccal smears provided

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constitutional high-molecular-weight DNA in patients 5 and 6 (16,17). Patients gave their informed consent to include their material in this study. For molecular analyses, tetranucleotide repeat microsatellites were selected through the Genome Data Base at loci with a high frequency of chromosomal deletions in HD, which were predicted by the cytogenetic literature (11–15), and located at 1q42 (D1S517), 4q26 (D4S2301), 9p23 (D9S254), and 11q23 (D11S1294). Their PCR primer sequences are available in the Genome Data Base (http://www.gdb.org). A highly sensitive seminested PCR assay included a first round that used the primers indicated above. In a second PCR. one of the primers in each pair was replaced by an internal forward primer (Fint) or an internal reverse primer (Rint): 1q42 (D1S517Rint) 5'-CATGTGTCCATCAATGGTAG-3': 4q26 (D4S2301Fint) 5'-GATGAGT-GCTTAGACCATAGTA-3'; 9p23 (D9S254Rint) 5'-GTCTCCAATGCAT-GANCTT-3'; and 11q23 (D11S1294Rint) 5'-CTGGTTTGCTTTCCCTTTCTT-3' (software = "Primers! For the WWW"; http://www.williamstone.com) (Fig. 1, A). Allelic dropout during amplification of polymorphic microsatellite fragments may randomly affect either allele mimicking LOH. To exclude this pitfall, pooled samples of 10 purified HRS cells from a given patient were amplified (18,19). Ten picograms of DNA (corresponding to the DNA content of a single diploid cell) was consistently detected by our PCR assay. We also used a seminested PCR (1,3,20-22) to examine HRS cells from patients 3, 4, 6, and 7 for clonally rearranged immunoglobulin heavy-chain genes created through joining of variable, diversity, and joining immunoglobulin gene regions (V-D-J joining). The primers were a framework III immunoglobulin gene VH primer 5'-ACACGGC(C/T)(G/C)T(G/A)TAT-TACTGT-3' and a consensus JH primer (21) 5'-ACCTGAGGAGACGGT-GACC-3'. The seminested primer was from VLJH sequences (5'-GTGAC-CAGGT(N)CCTTGGCCCCA-3') (22) (Fig. 1, B).

DNA extracted from buccal smears and from bystander lymphocytes showed constitutional individualspecific microsatellite patterns in all patients (Fig. 1, A). Of 259 samples of HRS cells microdissected from formalin-fixed tissue, 59 (23%) yielded a detectable microsatellite PCR product. Mock picks from tissue sections cleared of cellular material were consistently negative by PCR analysis. All patients except patient 5 reproducibly showed clonal LOH in HRS cells at one to three different microsatellite loci (Table 1). LOH was seen at all loci, but the locus most frequently altered was D4S2301 (4g26: four of five informative patients). High-molecular-weight DNA from fresh HRS cells and corresponding degraded DNA from formalin-fixed tissue vielded identical individual-specific microsatellite band patterns (Fig. 1, A). None of the HRS cell samples showed extra microsatellite bands (23). Patients 3, 4, 6, and 7 showed clonal immunoglobulin heavy-chain gene configurations in HRS cells and polyclonal patterns in bystander lymphoid cells (Fig. 1, B). In patient 6, a common rearranged immunoglobulin heavy-chain gene band in HRS cells, observed at presentation (6a) and at relapse (6b), indicated that the initial monoclonal population of HRS cells had relapsed. Our immunogenotype data from formalin-fixed HRS cells are in keeping with published data that HRS cells are mostly monoclonal B-cell populations (1–8,20,24).

The detection of LOH has several implications in HD. The patients' buccal mucosa and bystander lymphocytes showed constitutional microsatellite patterns indicating that LOH in HRS cells was an acquired specific genetic feature of HD. To the best of our knowledge, this is the first report on microsatellite PCR detection of clonal LOH in microdissected HRS cells. In contrast to our approach, classical cytogenetics permit the study of mitotic cells only and usually do not permit the identifi-

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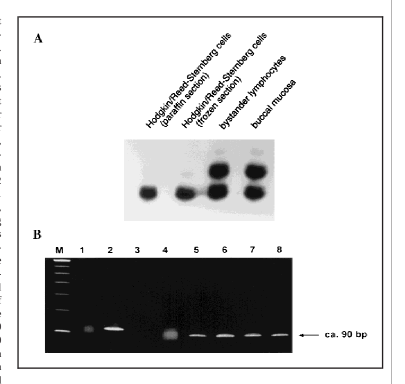
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**Table 1.** Pathology of Hodgkin's disease (HD) samples and results of microsatellite polymerase chain reaction (PCR) analysis in microdissected Hodgkin and Reed-Sternberg (HRS) cells\*

Samples							Microsatellite typing			
Histologic diagnosis (sample No.)	Sex/age, y	Stage†	CD15/30	EBER	LMP	β/gp220	D1S517 1q42	D4S2301 4q26	D9S254 9p23	D11S1294 11q22–23
Nodular sclerosing HD (1)	M/43	III	+/+	+	+	+	ni	1	nd	ni
Mixed cellularity HD (2)	F/43	III	+/+	+	+	+	2	1	ni	1
Nodular sclerosing HD (3)	F/16	III	+/+	_	nd	+	2	1	ni	1
Mixed cellularity HD (4)	F/74	III	+/+	+	+	+	1	1	1	2
Nodular sclerosing HD (5)	F/38	III	-/+	nd	nd	nd	ni	nd	ni	nd
Nodular sclerosing HD (6a)	F/69	III	-/+	nd	nd	+	2	nd	nd	nd
Nodular sclerosing HD in relapse (6b)		_	-/+	nd	_	nd	2	2	1	2
Mixed cellularity HD (7)	F/71	II	+/+	nd	-	nd	1	ni	ni	1

<sup>\*1</sup> = patient constitutionally informative and loss of one microsatellite allele (loss of heterozygosity) in HRS cells (data verified twice); 2 = patient constitutionally informative, two alleles retained in HRS cells; + = immunostaining done and positive; - = immunostaining done and negative; EBER = Epstein-Barr virus-encoded RNA; LMP = Epstein-Barr virus latent membrane protein; M = male; F = female; nd = molecular analysis not done due to lack of suitable material or failure to obtain PCR amplicons; and ni = patient not informative.

Fig. 1. A) Microsatellite detection of loss of heterozygosity (LOH) at D1S517 (1q42) in Hodgkin and Reed-Sternberg (HRS) cells from patient 7, a patient with a mixed cellularity-type Hodgkin's disease (HD). DNA from bystander lymphoid cells and from buccal mucosa shows a constitutionally heterozygous microsatellite pattern (two allelic bands). HRS cells isolated from formalin-fixed, paraffin-embedded sections and fresh-frozen sections show a single lower allele, indicating LOH at 1q42. CD30<sup>+</sup> HRS cells as well as bystander lymphocytes and other infiltrating cells in HD lymphoma sections were microdissected under an inverted laser microscope (P:A:L:M; GmbH, Wolfratshausen, Germany) (17) and were subjected to a sensitive seminested microsatellite polymerase chain reaction (PCR). The first PCR round was run on a Thermal Cycler 480 (Perkin-Elmer, Rotkreuz, Switzerland) with 2 mM MgCl<sub>2</sub>, all four deoxynucleoside triphosphates (dNTPs, each at 0.1 mM), 1 U of hot-start Taq polymerase (AmpliTaq Gold; Perkin-Elmer, Rotkreuz, Switzerland), and 1 µM of each primer under the following conditions: Initial denaturation was at 95 °C for 15 minutes, 30 cycles of 95 °C for 30 seconds, 55 °C for 30 seconds, and 72 °C for 60 seconds, and the final elongation step was at 72 °C for 7 minutes. One microliter of a 1:20 dilution of first-round PCR amplicons was subjected to a second round of PCR on a Perkin-Elmer 9600 Thermal Cycler in 1.5 mM MgCl<sub>2</sub>, all four dNTPs (each at 0.1 mM), 0.5 U of Boehringer® Taq DNA polymerase, and 1 µM of each primer under the following conditions: Initial denaturation was at 94 °C for 5 minutes, 30 cycles of 94 °C for 30 seconds, 55 °C for 30 seconds, and 72 °C for 60 seconds, and the final elongation step was at 72 °C for 7 minutes. If a PCR band was identified on agarose gel electrophoresis, 1 µL of a 1:20 dilution of the respective first-round PCR sample was subjected



to a second round of PCR by use of the appropriate internal <sup>32</sup>P-labeled primer, as described above. Labeled PCR fragments were subjected to electrophoresis on denaturing 6% polyacrylamide-sequencing gels and subjected to autoradiography at -70 °C for 2-12 hours as appropriate (19). LOH seen in a given patient at a particular locus was confirmed by at least one or several repeat analyses of cells separately harvested from adjacent tissue sections. B) Immunoglobulin heavy-chain gene PCR analysis from patient 6, a patient with nodular sclerosing HD. Lane 1—peripheral blood lymphocytes from a healthy donor; lane 2—Nalm6 (monoclonal B-cell leukemia cell line with rearranged immunoglobulin heavy-chain genes); lane 3—Calu1 (lung cancer cell line with germline immunoglobulin heavy-chain genes); lane 4—bystander lymphoid cells microdissected from paraffin-embedded HD sections; lanes 5 and 6—pooled microdissected HRS cells obtained at presentation; and lanes 7 and 8—single HRS cells obtained at relapse. M = molecular weight marker (100-base-pair [bp] DNA ladder). HRS cells at presentation and at relapse show a common rearranged immunoglobulin heavy-chain gene band of about 90 bp. Lymphoid cells in lanes 1 and 4 yield multiple rearranged immunoglobulin heavy-chain gene bands indicating polyclonal B-cell populations. The first PCR round was run in 2.5 mM MgCl<sub>2</sub>, all four dNTPs (each at 0.2 mM), 1 U of hot-start *Taq* polymerase (AmpliTaq Gold; Perkin-Elmer), and 1.3 μM of each immunoglobulin gene primer under the following conditions: Initial denaturation was at 95 °C for 15 minutes; 10 cycles with a stepwise decrease of annealing temperatures from 65 °C to 56 °C were followed by 30 cycles at 95 °C for 30 seconds, 55 °C for 30 seconds, and 72 °C for 45 seconds and a final elongation step at 72 °C for 7 minutes. The second-round PCR used 1 μL of a 1:200 dilution of the first-round PCR product as template and was run under the conditions described above, except for using 2 mM MgCl<sub>2</sub> and running 40 cycles with an anneali

<sup>†</sup>Cotswold staging classification.

cation of the lineage of cells being karvotyped. The detection of LOH restricted to HRS cells is consistent with the view that they are clonally derived from a common single cell of origin (16,25-27). The probability in a given patient that all HRS cells tested would have lost the same allelic microsatellite band independently or by chance is .002 (i.e.,  $0.5^{n-1}$ , where n is the number of HRS cells examined) and, therefore, is low (28). Our findings thus confirm and extend molecular immunogenotype data on the clonality of HRS cells in classical HD with a molecular marker system unrelated to the configuration of immunoglobulin genes.

The detection of LOH in tumor cells at a particular locus indicates clonality and by the same token points to a site in the genome possibly harboring inactivated tumor suppressor genes. In non-Hodgkin's lymphomas, inactivation of tumor suppressor genes has been described at the p53 gene or the  $p16^{INK4a}$ locus. Our work now shows that in classical HD, clonal LOH may be present at loci, such as 1q42, 4q26, 9p23, 11q22-23, and possibly others. The dense genomic map of microsatellite markers now available will help to narrow down such loci as a next important step for the eventual cloning of tumor suppressor genes operative in the molecular pathology of HD.

## REFERENCES

- (1) Marafioti T, Hummel M, Anagnostopoulos I, Foss HD, Falini B, Delsol G, et al. Origin of nodular lymphocyte-predominant Hodgkin's disease from a clonal expansion of highly mutated germinal-center B cells. N Engl J Med 1997;337:453–8.
- (2) Ohno T, Stribley JA, Wu G, Hinrichs SH, Weisenburger DD, Chan WC. Clonality in nodular lymphocyte-predominant Hodgkin's disease. N Engl J Med 1997;337:459–65.
- (3) Hummel M, Ziemann K, Lammert H, Pileri S, Sabattini E, Stein H. Hodgkin's disease with monoclonal and polyclonal populations of Reed-Sternberg cells. N Engl J Med 1995; 333:901–6.
- (4) Kuppers R, Kanzler H, Hansmann ML, Rajewski K. Immunoglobulin V genes in Reed-Sternberg cells [letter]. N Engl J Med 1996; 334:404–6.
- (5) Kuppers R, Rajewski K, Zhao M, Simons G, Laumann R, Fischer R, Hansmann ML.

- Hodgkin disease: Hodgkin and Reed-Sternberg cells picked from histological sections show clonal immunoglobulin gene rearrangements and appear to be derived from B cells at various stages of development. Proc Natl Acad Sci U S A 1994;91:10962–6.
- (6) Tamaru J, Hummel M, Zemlin M, Kalvelage B, Stein H. Hodgkin's disease with a B-cell phenotype often shows a VDJ rearrangement and somatic mutations in the VH genes. Blood 1994;84:708–15.
- (7) Vockerodt M, Soares M, Kanzler H, Kuppers R, Kube D, Hansmann ML, et al. Detection of clonal Hodgkin and Reed-Sternberg cells with identical somatically mutated and rearranged VH genes in different biopsies in relapsed Hodgkin's disease. Blood 1998;92: 2899–907.
- (8) Brauniger A, Kuppers R, Strickler JG, Wacker HH, Rajewski K, Hansmann ML. Hodgkin and Reed-Sternberg cells in lymphocyte predominant Hodgkin disease represent clonal populations of germinal centerderived tumor B cells [published erratum appears in Proc Natl Acad Sci U S A 1997;94: 14211]. Proc Natl Acad Sci U S A 1997;94: 9337–42.
- (9) Weissenbach J. The Human Genome Project: from mapping to sequencing. Clin Chem Lab Med 1998;36:511–4.
- (10) Pabst T, Schwaller J, Tobler A, Fey MF. Detection of microsatellite markers in leukaemia using DNA from archival bone marrow smears. Br J Haematol 1996;95:135–7.
- (11) Barrios L, Caballin MR, Miro R, Fuster C, Berrozpe G, Subias A, et al. Chromosome abnormalities in peripheral blood lymphocytes from untreated Hodgkin's patients. A possible evidence for chromosome instability. Hum Genet 1988;78:320–4.
- (12) Dohner H, Bloomfield CD, Frizzera G, Frestedt J, Arthur DC. Recurring chromosome abnormalities in Hodgkin's disease. Genes Chromosomes Cancer 1992;5:392–8.
- (13) Tilly H, Bastard C, Delastre T, Duval C, Bizet M, Lenormand B, et al. Cytogenetic studies in untreated Hodgkin's disease. Blood 1991;77:1298–304.
- (14) Kristoffersson U, Heim S, Mandahl N, Olsson H, Akerman M, Mitelman F. Cytogenetic studies in Hodgkin's disease. Acta Pathol Microbiol Immunol Scand [A] 1987;95: 289–95
- (15) Teerenhovi L, Lindholm C, Pakkala A, Franssila K, Stein H, Knuutila S. Unique display of a pathologic karyotype in Hodgkin's disease by Reed-Sternberg cells. Cancer Genet Cytogenet 1988;34:305–11.
- (16) Pabst T, Schwaller J, Bellomo MJ, Oestreicher M, Muhlematter D, Tichelli A, et al. Frequent clonal loss of heterozygosity but scarcity of microsatellite instability at chromosomal breakpoint cluster regions in adult leukemias. Blood 1996;88:1026–34.

- (17) Becker I, Becker KF, Rohrl MH, Minkus G, Schutze K, Hofler H. Single-cell mutation analysis of tumors from stained histologic slides. Lab Invest 1996;75:801–7.
- (18) Garvin AM, Holzgreve W, Hahn S. Highly accurate analysis of heterozygous loci by single cell PCR. Nucleic Acids Res 1998;26: 3468–72.
- (19) Hughes AE. Optimization of microsatellite analysis for genetic mapping. Genomics 1993;15:433–4.
- (20) Delabie J, Tierens A, Gavriil T, Wu G, Weisenburger DD, Chan WC. Phenotype, genotype and clonality of Reed-Sternberg cells in nodular sclerosis Hodgkin's disease: results of a single-cell study. Br J Haematol 1996; 94:198–205.
- (21) Segal GH, Wittwer CT, Fishleder AJ, Stoler MH, Tubbs RR, Kjeldsberg CR. Identification of monoclonal B-cell populations by rapid cycle polymerase chain reaction. A practical screening method for the detection of immunoglobulin gene rearrangements. Am J Pathol 1992;141:1291–7.
- (22) Reed TJ, Reed A, Wallberg K, O'Leary TJ, Frizzera G. Determination of B-cell clonality in paraffin-embedded lymph nodes using the polymerase chain reaction. Diag Mol Pathol 1993;2:42–9.
- (23) Mark Z, Toren A, Amariglio N, Schiby G, Brok-Simoni F, Rechavi G. Instability of dinucleotide repeats in Hodgkin's disease. Am J Hematol 1998;57:148–52.
- (24) Hummel M, Marafioti T, Stein H. Clonality of Reed-Sternberg cells in Hodgkin's disease [letter]. N Engl J Med 1999;340:394–5.
- (25) Fey MF, Peter HJ, Hinds HL, Zimmermann A, Liechti-Gallati S, Gerber H, et al. Clonal analysis of human tumors with M27β, a highly informative polymorphic X chromosomal probe. J Clin Invest 1992;89:1438–44.
- (26) Wainscoat JS, Fey MF. Assessment of clonality in human tumors: a review. Cancer Res 1990;50:1355–60.
- (27) Fearon ER, Hamilton SR, Vogelstein B. Clonal analysis of human colorectal tumors. Science 1987;238:193–7.
- (28) Sidransky D, Frost P, Von Eschenbach A, Oyasu R, Preisinger AC, Vogelstein B. Clonal origin of bladder cancer. N Engl J Med 1992;326:737–40.

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