Low Expression of Human Histocompatibility Leukocyte Antigen-DR Is Associated with Hypermethylation of Human Histocompatibility Leukocyte Antigen-DRα Gene Regions in B Cells from Patients with Systemic Lupus Erythematosus

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Abstract

The relationship between the expression of HLA-DR antigens and the HLA-DRα gene methylation was examined in systemic lupus erythematosus (SLE). Using permanent B cell lines, we found reduced DR expression in SLE. The low DR expression was correlated with high anti-DNA antibody titers in patients’ sera. The amounts of DRα message were lower in SLE cells than in normal controls, suggesting that the low expression of DR antigens is associated with gene functions.

The extent of DNA methylation was examined at five CCGG sites in the HLA-DRα locus. DNA from both SLE and normal cells showed variable methylation patterns. Since the DRα gene is a single-copy gene, such a variability is the result of assaying a mixture of transformed clones containing methylated DRα gene, with other clones containing unmethylated DRα gene. A distinctive feature of normal cells was a consistent methylation pattern: 12 normal cell lines showed exactly the same pattern. In contrast, 28 SLE cell lines showed a cell-line-specific methylation, and hypermethylation at the DRα locus. The hypermethylation is often associated with transcriptionally inactive genes. Thus, our results suggest that (a) B cells with hypermethylated DR genes might express no or few DR antigens; (b) the ratio of cells with differently methylated DR genes is consistent in normal individuals, while, in SLE patients, cells with hypermethylated DR genes predominate, resulting in apparently reduced DR antigen expression; and (c) the aberrant DR expression could be associated directly with immunoregulatory dysfunctions in SLE disease.

Introduction

Systemic lupus erythematosus (SLE) is a human autoimmune disease. Patients with SLE develop a variety of autoantibodies that react with autologous antigens including cellular, nuclear, and cytoplasmic components (1-4). A genetic predisposition to the illness and numerous immune defects found in patients with SLE have suggested that the disease may result from abnormal gene expression in components of the immune system (1, 2, 4).

The immune recognition and response are considered to be controlled by the major histocompatibility complex (MHC) or HLA in humans. The HLA complex contains two classes of cell surface antigens: class I and class II. The class II products are considered to play an important role in self-recognition, especially in the immune response by regulating the interaction between T cells and B cells (5). The class II antigens, or Ia antigens, contain at least three subsets; HLA-DR, DQ, and DP (5-7). Among them, HLA-DR molecules are dominant, composing up to 60% of total class II antigens at the membrane. The HLA-DR antigen consists of two different polypeptides: α-chain and β-chain. The genes encoding these polypeptides have been isolated and sequenced (8-13). The genomic hybridization analysis indicated that the DRα chain is encoded by a single-copy gene (12, 13).

There is strong evidence for the involvement of the HLA in diseases (14). Many of the HLA-DR-associated disorders are of the autoimmune type (15). Insulin-dependent juvenile diabetes is associated with DR3, myasthenia gravis with DR3, and rheumatoid arthritis with DR4. An increased frequency of DR3 has been reported in SLE patients (6, 15). It has been suggested that a particular HLA haplotype makes the individual either susceptible or resistant to certain diseases (15). However, few surveys have studied the relationship between disease activity and the cellular expression of HLA molecules.

Vertebrate DNA contains 5-methylcytosine (mC) as a minor component, which occurs at a frequency of ~1% of the total bases (16). The mC locates primarily in the CpG dinucleotide in mammalian DNA (17). The methylation of cytosines is a post-replicative enzymatic reaction catalyzed by DNA methyltransferase (18). The physiological functions of DNA methylation have not necessarily been clear. Surveys on the relationship between gene expression and gene methylation have revealed that transcriptionally active genes are often hypomethylated (19-21). It was found that the gene function is not correlated proportionally to the overall methylation level in a given genome. The methylation at particular sites in that gene is critical for its expression (22). In the β-globin gene, only a few CCGG sites at the 5′ extragenic region were demethylated when the gene was activated (23). In some cases, the methylation of only one CpG site was enough to silence the gene (24, 25). The methylation of hypoxanthine phosphoribosyltransferase locus was studied on both active and inactive states (26). The active HPRT allele was associated with a consensus pattern of DNA methylation: hypomethylation at the 5′ region, and hypermethylation at the

1. Abbreviations used in this paper: C, cytosine; cDNA, complementary DNA; EBV, Epstein-Barr virus; GC, glucocorticoids; kb, kilobases; mC, 5-methylcytosine; mRNA, messenger RNA; SLE, systemic lupus erythematosus.

J. Clin. Invest. 0021-9738/85/10/1314/09 $1.00
Volume 76, October 1985, 1314-1322
3' region. The inactive allele showed nonuniformity and less extensive hypomethylation at the 5' cluster. It was suggested that both the methylated and unmethylated sites were important determinants of gene activity (26).

In an attempt to understand SLE at the molecular level, we asked whether HLA-DR antigen is expressed normally in SLE and if the genes encoding HLA molecules are expressed normally in SLE. To answer these questions, we have performed the following experiments. First, we have established permanent B cell lines from SLE patients by Epstein-Barr virus (EBV) transformation. Second, using these materials, we have assayed the amounts of HLA-DR antigens on the cell surface. Third, we analyzed the methylation patterns of the HLA-DRα gene by using methylation-sensitive restriction endonucleases and a DRα complementary DNA (cDNA) probe. In this communication, we report that, in SLE, HLA-DR expression is reduced and that such reduction is correlated with reduced amounts of DRα mRNA and with hypermethylation of DRα gene regions.

Methods

Cells. Blood samples were from SLE patients followed at the Arthritis Branch, National Institute of Arthritis, Diabetes, Digestive and Kidney Diseases, National Institutes of Health, Bethesda, MD. All patients fulfilled the American Rheumatism Association criteria for SLE. Normal blood samples were obtained either from National Institutes of Health or from the Blood Component Laboratory, Dana-Farber Cancer Institute, Boston, MA. Permanent B cell lines were established by transforming peripheral blood lymphocytes by EBV as described (27). Briefly, peripheral blood mononuclear cells were isolated by Ficoll-Hypaque (Pharmacia Fine Chemicals, Piscataway, NJ) density gradient centrifugation. Collect lymphocytes were cultured at a density of 2 × 10⁶ cells/ml. To 9 ml cell suspension, 1 ml culture supplement containing EBV (B95-8) was added. After 4–8 wk, cells were subcultured and used for further assay. Established transformed cells were designated as LC lines with appropriate numbers. Normal cell lines were marked with capital N after the number.

Determination of immunoglobulin phenotypes. Surface immunoglobulins (Ig) were determined by the conventional immunofluorescence method using fluorescein isothiocyanate-labeled anti-human γ, μ, or δ (Dakopatts Lab, Uppsala, Sweden and Behringwerke, Marburg, West Germany), and by the indirect antilgG rosetting reaction (28). Quantitative radioimmunoassay for the total IgG and IgM secreted from cells was performed in microplates as follows. A 50-µl sample was incubated with 0.5 µl ³²P-labeled anti-human γ rabbit Ig F(ab')₂ or with ³²P-labeled anti-human μ rabbit Ig F(ab')₂ in the well that had been coated with IgG (0.5 µg/ml) or with IgM (0.5 µg/ml) at 37°C for 90 min. After washing with phosphate-buffered saline (PBS) three times, the radioactivity in each well was counted by a counter. For each radioimmunoassay, a fifteen-point standard calibration curve was prepared for IgG and IgM (0.2 ng–50 µg).

Flow cytometry analysis. An EPICS V flow cytometer (Coulter Electronics, Inc., Hialeah, FL) was used to measure cell surface antigens as described previously (29). The antibody binding reaction was performed as described previously (29). The murine monoclonal antibodies used were L243, P3, and 4F2. L243 reacts with nonpolymorphic regions of HLA-DR antigens (30); P3 is the myeloma antibody produced by the P3X63 cell line; 4F2 recognizes a 40,000–80,000 kD glycoprotein-activation antigen that is present on all tumor and EBV-transformed cell lines. P3 and 4F2 were used for negative and positive controls, respectively, together with L243 at each run. Depending upon setting and conditions of the machine, the distribution spectrum differed slightly at each run. To normalize the difference, one reference sample of the normal cells was included at each run.

Anti-DNA antibody assay. Total plasma (~20 ml) was used as the source of antibody. The proteins were precipitated repeatedly with 40% saturated ammonium sulfate as described (31). Immunoglobulin (Ig) was purified by affinity chromatography of Sepharose-bound anti-human IgA, IgG, and IgM (Miles Laboratories, Inc., Elkhart, IN) as described (32). Anti-DNA antibody activity was measured by the binding of ³²P-labeled DNA. Calf thymus DNA was sheared by sonication and subjected to nick-translation with [α-³²P]dNTP (Amersham Corp., Arlington Heights, IL) to give a specific activity of ~10⁶ cpm/µg. A 100-µl reaction mixture contained 10 µg Ig and 1 µg DNA (10⁶ cpm) in PBS, and incubated at 37°C for 30 min. The reaction mixture was diluted twofold with cold PBS and applied onto a nitrocellulose paper that was transferred previously with 0.5 M KOH as described (32). After being washed with cold PBS, the nitrocellulose paper was dried and counted for radioactivity. The antibody activity was scored on a 0–10 scale, where 0 is no DNA binding and 10 is 100% binding of input DNA under the experimental conditions. At least three different tests, each with duplicated tubes, were done and average values were calculated.

DNA and RNA hybridization. DNA was extracted from B cell lines as described (33). DNA samples were digested with appropriate restriction endonucleases, subjected to agarose gel electrophoresis, and transferred to nitrocellulose paper as described (32). RNA was extracted from B cells and transferred to nitrocellulose paper after agarose gel electrophoresis as described (34). The CDNA probes were p34RFl-3 (generous gift from Professor S. Weissman, Yale University) for the HLA-DRα gene, and pHDS4 (generous gift from Professor G. Attardi, California Institute of Technology) for the dihydrofolate reductase gene. The probe was nick-translated and hybridized to sample DNAs as described (35).

Nearest neighbor analysis. The sample DNA was subjected to nick-translation in the presence of [α-³²P]dGTP (Amersham Corp.; specific activity, 800 Ci/mmol) as described (35). The radioactively labeled DNA was purified with repeated precipitation with ethanol and Sephadex G-50 filtration, and hydrolyzed to 3'dNMP with micrococcal nuclease and spleen phosphodiesterase as described (35). The 3'dNMP was separated with two-dimensional thin layer chromatography, and the ratio between mc and cytosine (C) was estimated from the radioactivity in each nucleotide as described (35).

Results

Establishment and characterization of EBV-transformed cell lines. B cell lines were established by EBV-transformation of peripheral blood lymphocytes from both SLE patients and apparently healthy donors. The immunoglobulin phenotypes were examined on the cell lines derived from 13 SLE patients, 10 control donors, and 2 patients with type I diabetes (as the disease control). Results are shown in Table I. All cell lines showed heterogeneous immunoglobulin production, indicating that EBV-transformation led to polyclonal cell activation. In all cell lines examined, surface α and δ were dominant. Some cells expressed γ and α. The immunoglobulin phenotype did not differ between SLE and control B cells. This indicates that B cell populations that were transformed by EBV are the same in SLE patients and controls. Thus, EBV-transformed cell lines provide a unique circumstance in which the biochemical aspects of SLE B cells can be compared with those of healthy controls.

Low expression of DR antigens in SLE cell lines. The expression of DR antigens was determined by monoclonal anti-DR antibody binding to B cells derived from SLE patients and control donors. Fig. 1 shows the representative patterns of DR expression analyzed by flow cytometry. First, we analyzed 12 control cell lines, which showed similar distribution spectra with a single, symmetrical peak as represented by LC63N (Fig. 1). Then we analyzed 46 SLE cell lines. 25 lines showed no difference from the control lines (Fig. 1 A). We refer to these cell lines as type II. 21 cell lines showed reduced DR expression. The representative patterns are shown in Fig. 1 B and C. The spectrum
**Table I. Immunoglobulin Phenotypes of Cell Lines**

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Amounts of secreted Ig</th>
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<tbody>
<tr>
<td></td>
<td>IgG (µg/ml)</td>
</tr>
<tr>
<td></td>
<td>%</td>
</tr>
<tr>
<td>LC65</td>
<td>8.8</td>
</tr>
<tr>
<td>LC66</td>
<td>2.1</td>
</tr>
<tr>
<td>LC67</td>
<td>3.3</td>
</tr>
<tr>
<td>LC68</td>
<td>54.1</td>
</tr>
<tr>
<td>LC69</td>
<td>52.4</td>
</tr>
<tr>
<td>LC70</td>
<td>40.8</td>
</tr>
<tr>
<td>LC71</td>
<td>28.4</td>
</tr>
<tr>
<td>LC72</td>
<td>32.8</td>
</tr>
<tr>
<td>LC73</td>
<td>27.4</td>
</tr>
<tr>
<td>LC74</td>
<td>33.9</td>
</tr>
<tr>
<td>LC75</td>
<td>47.2</td>
</tr>
<tr>
<td>LC76</td>
<td>44.7</td>
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<td>LC77</td>
<td>54.1</td>
</tr>
<tr>
<td>LC78</td>
<td>53.6</td>
</tr>
<tr>
<td>LC79</td>
<td>45.7</td>
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</table>

Cell lines 1 through 13 were from SLE patients, 14 through 23 were from healthy donors, and 24 and 25 were from diabetes patients.

To examine the possibility that EBV-transformed B cells produced anti-DR antibody in vitro that blocked the assay system, we did the following experiments. Before staining with L243 monoclonal anti-DR antibody, we mixed the control cells with culture supernatant of SLE cells with low DR antigens (LC65). If the culture supernatant contained anti-DR antibody, the binding with L243 should be blocked. Results were negative (data not shown). It was concluded that the transformed B cells did not produce detectable anti-DR antibody in vitro. Another possible experimental barrier was considered: i.e., SLE cells produce normal amounts of DR antigens, but abnormally shed it. Shed DR antigens could compete with cells for L243 binding. Since before L243 binding, cells were washed twice with plain PBS, shed, or secreted DR antigens in the culture medium should be absent in L243 binding system.

**Correlation of DR expression with anti-DNA antibody quantity.** To see if the reduced DR expression of B cells from SLE patients was associated with disease activity, we compared the DR-expression types with the quantity of anti-DNA antibody. After lymphocytes were isolated from patients' blood samples, immunoglobulin fractions were obtained from the remaining plasma fraction, and were assayed for DNA binding activity. Fig. 2 shows the results. The relative activity of anti-DNA antibody was plotted in three groups; normal controls, type II (apparently normal DR expression), and type I (reduced DR expression). The distribution of antibody activity was heterogeneous, but the type I group appeared to contain the highest amounts of anti-DNA antibodies. The rank sum analysis indicated the P value to be <0.05 between type I and type II (P < 0.05).

**Reduced amounts of HLA-DRα mRNA.** The level of DRα mRNA was analyzed by Northern blotting. Representative results are shown in Fig. 3. The amounts of mRNA in three SLE cell lines were markedly reduced compared with that of control normal cells (LC63N). These cell lines showed reduced DR antigens as analyzed by flow cytometry (type I). Particularly, LC66 showed very low levels of the message, being consistent with results shown in Fig. 1. The data indicate that the low expression of DR antigens on SLE B cell surface is the result of reduced transcriptional activity of the DR genes.

**Characterization of the DRα locus and probe.** It has been found that the gene expression is often inversely correlated with DNA methylation. The transcriptionally active genes are less

![Figure 1. Comparison of the fluorescence profiles of B cells stained with L243 monoclonal anti-DR antibody. (A) Comparison of LC63N (normal cell line) with LC66 (SLE cell line). This type was defined as type II. (B) Comparison of LC63N with LC65 (SLE cell line). (C) Comparison of LC63N with LC66 (SLE cell line). These cell lines expressing reduced antigens were defined as type I.](image1)

![Figure 2. Relationship between DR expression and amounts of anti-DNA antibody. The DR expression was estimated by flow cytometry. The anti-DNA antibody titer was estimated on a 0–10 scale, where 0 is no binding to DNA, and 10 is 100% binding of the input 32P-labeled DNA. SLE-I represents cells that express reduced amounts of DR antigens (type I), and SLE-II represents cells that express apparently normal amounts of DR antigens (type II).](image2)
methylated than inactive genes. To see if the reduced DR expression in SLE patients is associated with DR gene methylation, we did Southern hybridization experiments using HLA-DRα cDNA as the probe. The methylation was analyzed using two isochizomers of restriction endonucleases, MspI and HpaII. The HpaII does not cleave the CCGG tetramer when the internal C is methylated, while MspI does. Fig. 4 summarizes the restriction map of the CCGG sequence in the vicinity of the probe of 3.4 kb. The probe was constructed from an EcoRI-digested 3.4-kb fragment, which contained exons 2, 3, 4, and 5 of the HLA-DRα gene (8). The MspI site was close to the EcoRI site, giving a 3.1-kb fragment. Consequently, MspI digestion generated almost exclusively a 3.1-kb fragment by hybridization, since the flanking fragments of 3.0 kb and 1.1 kb do not hybridize well with the probe.

Normal cell lines have a consistent methylation pattern at DRα locus. Nuclear DNA was isolated from established cell lines and the methylation at DRα locus was examined by HpaII/ MspI cleavage patterns. First, we examined the control B cells. As shown in Fig. 5, MspI generated primarily a 3.1-kb fragment. HpaII generated 3.1, 4.2, 4.5, 6.1, 7.2, and 11-kb fragments, indicating that the methylation at these sites were variable within a cell line. Since the intensity of each band is almost equal, it seems that the ratio of DNA having mC at a given CCGG site to the total DNA is similar. Although it is difficult to estimate the exact ratio from the present data, ~20–30% of the total DNA seems to contain mC at an arbitrary CCGG site. In other words, a large portion of DNA contains unmethylated cytosines at any CCGG site examined.

A characteristic feature of the DRα locus was the uniformity of methylation in all cell lines derived from normal donors. We examined 12 control cell lines (7 samples are shown in Figs. 4 and 5), and found that all of them showed similar HpaII cleavage patterns, giving 3.1, 4.2, 4.5, 6.1, 7.2, and 11-kb fragments. Thus, normal cell lines contain the consensus methylation pattern at the DRα locus.

SLE cell lines have variable methylation patterns at DRα locus. Figs. 6 and 7 show some representative cleavage patterns of DNAs from SLE B cells by HpaII and MspI. MspI generated a 3.1-kb fragment in all cell lines (LC38, LC39, LC65, and LC66).

![Figure 3](image)

**Figure 3.** Expression of DRα mRNA in B cell lines. A 10-μg total cellular RNA was applied to a slot of 1.1% agarose gel and electrophoresed at 90 V for 6 h. RNA was transferred to nitrocellulose paper and hybridized with the DRα probe, pp34RI-3. Samples were LC63N (normal cell line), LC68 (SLE), LC66 (SLE), and LC38 (SLE).

![Figure 5](image)

**Figure 5.** Southern hybridization analysis of DNAs from normal cell lines with HLA-DRα cDNA probe. Samples (10 μg each from LC48N, LC59N, LC62N, and LC63N) were digested with either MspI (30 μ, New England Biolabs, Inc., Beverly, MA) or HpaII (30 μ, New England Biolabs) for 16 h at 37°C, and subjected to an electrophoresis on 0.8% agarose gel. The marker was constructed from various restriction fragments of pBR322 and pBR322-containing plasmids of known size.

indicating that the gene itself does not differ between control and SLE DNAs. On HpaII digestion, DNAs from four SLE cell lines were methylated to different degrees. LC38 lacked 3.1- and 4.2-kb fragments, showing that these sites are heavily methylated. LC39 lacked 4.2, 6.1, and 7.2-kb fragments. LC65 and LC66 showed a similar pattern with only 4.2 kb plus high molecular weight fragments. These cell lines were all of the type I group (see Fig. 1). Other examples are shown in Fig. 8. Each cell line
showed a unique methylation pattern. It is clear from these results that, in contrast to the normal cell lines, the methylation at the DRα locus was cell line-specific in SLE B cells.

Hypermethylation at DRα locus in SLE B cells. Table II summarizes the extent of CCGG methylation in the DRα locus of SLE and control cells. Cell lines are classified according to the DR expressing types. The appearance of HpaII fragments in each cell line was examined and its frequency was estimated. In DNA from type I cells, the 3.1-kb fragment was observed in six cell lines out of 14 cell lines examined (43%). In DNA from type II cells, the 3.1-kb fragment was seen in 10 lines out of 14 lines (71%). In DNA from the controls, all cell lines showed the 3.1-kb fragment (100%). The frequencies of other fragments were similar to that of the 3.1-kb fragment. Thus, DNA from type I cells was methylated at elevated levels compared with that from control cells. The methylation extent of DNA from type II cells was between that of type I and that of normal cell DNAs. These experiments show that certain regions in the HLA-DRα gene are hypermethylated in B cells from SLE patients.

Methylation of CpG dinucleotide and of the dihydrofolate reductase gene locus. To see if the DNA hypermethylation is general in SLE cells, we examined the methylation of regions other than HLA-DRα. Since most of DNA methylation in mammalian cells occurs at the CpG dinucleotide, we analyzed the methylation frequency of CpG in the total DNA from control and SLE cells. Results are shown in Table III. The ratio of mCpG to total CpG was ~58% in all DNA examined regardless of the DR expressing types, indicating that the overall methylation in SLE cells does not differ from that in normal cells.

Figs. 9 and 10 show the methylation of the dihydrofolate reductase gene locus. To analyze the same HpaII fragments as used for HLA-DRα probe, the nitrocellulose paper used for

Table II. Relationship between DR Antigen Expression and Frequency of HpaII Fragments

<table>
<thead>
<tr>
<th>HpaII fragment (kb)</th>
<th>Type of DR expression</th>
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<tbody>
<tr>
<td></td>
<td>SLE type I</td>
</tr>
<tr>
<td>3.1</td>
<td>6/14 (43)</td>
</tr>
<tr>
<td>4.2</td>
<td>9/14 (64)</td>
</tr>
<tr>
<td>4.5</td>
<td>6/14 (43)</td>
</tr>
<tr>
<td>6.1</td>
<td>9/14 (64)</td>
</tr>
<tr>
<td>7.2</td>
<td>3/14 (21)</td>
</tr>
</tbody>
</table>

Values are expressed as the numbers of cell lines that show the HpaII fragment in question over the total numbers of cell lines examined. In parentheses, the frequency is expressed in percent. SLE type I refers to cell lines with reduced DR antigen expression, and SLE type II refers to those with apparently normal expression of DR antigens.

Table III. Frequency of 5-methylcytosine in CpG dinucleotide

<table>
<thead>
<tr>
<th>Type of DR antigen expression</th>
<th>Average mCpG/total CpG (%)</th>
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<tbody>
<tr>
<td>SLE type I</td>
<td>57.8±7.5</td>
</tr>
<tr>
<td>SLE type II</td>
<td>58.1±7.4</td>
</tr>
<tr>
<td>Normal control</td>
<td>56.4±8.3</td>
</tr>
</tbody>
</table>

The average values were estimated from 8 samples of type I (reduced DR expression), 15 samples of type II (apparently normal DR expression), and 10 samples of normal controls.

Figure 7. Southern hybridization analysis of DNAs from normal and SLE cell lines with HLA-DRα cDNA probe. Samples were LC32N (normal cell line), LC38 (SLE cell line, DR type I), LC39 (SLE cell line, DR type I), and LC65 (SLE cell line, DR type I, see Fig. 1 B). Experimental procedures were the same as described in the legend for Fig. 5.

Figure 8. Southern hybridization analysis of DNAs from SLE cell lines with HLA-DRα cDNA probe. Samples were LC07 (DR type II), LC08 (DR type I), LC09 (DR type II), LC10 (DR type I), LC13 (DR type not measured), LC14 (DR type II), LC16 (DR type II), and LC17 (DR type II). Experimental procedures were the same as described in the legend for Fig. 5, except that samples were digested with HpaII only.
HLA-DRα cDNA hybridization was washed with NaOH, and rehybridized with the cDNA of dihydrofolate reductase. All cell lines, both SLE and normal, showed the same methylation pattern: no variation was observed. These results suggest that, in SLE cells, the aberrant methylation occurred at restricted regions in genomic DNA, including HLA-DRα locus.

**Discussion**

This paper describes the relationship between HLA-DR antigen expression and the methylation of HLA-DRα gene regions in B cells from SLE patients. Three experimental results are presented: B cell lines were established by EBV transformation, reduced DR antigens were observed in these cell lines, and reduced DR expression was correlated with reduced DRα message and with hypermethylation at the DRα locus.

**EBV-transformed cell lines as a material for biochemical studies.** We have established more than 100 B cell lines from both SLE patients and healthy donors by EBV-transformation. The established cell lines were derived from peripheral blood lymphocytes, and could be a good source for biochemical studies, since they were available in considerable amounts at any time desired. Most B cell lines so far established were derived from peripheral blood mature lymphocytes, but recent studies have shown that EBV could transform pre-B cells originated from fetal livers (36, 37) and bone marrows (37). Considering these observations, we asked two questions concerning the appropriate use of the established permanent B cell lines before further experiments: did EBV transform B cells equally between SLE and normals? And is it proper to analyze the surface antigens with EBV-transformed cell lines? The first question was answered by examining the immunoglobulin phenotypes. The expression of surface and secreted immunoglobulins was the same between SLE and controls, indicating that the B cells capable of EBV transformation do not differ between normals and SLE patients. The transformed cells were possibly derived from mature B cells in both healthy and SLE patients. This was important for experiments to be followed since DR antigens are expressed only in mature B cells, not in immature and pre-B cells (38). The second question was answered by several reports. The expression of various HLA antigens, including HLA-A, B, C, and DR, was successfully analyzed with EBV-transformed B cells (39). Therefore, we concluded that it was proper to analyze and compare the expression of DR antigens in EBV-transformed B cells between SLE patients and healthy donors.

**Reduced DR expression in SLE.** Among 46 B cell lines derived from SLE patients, about half showed reduced amounts of cell surface DR antigens. In addition, the amounts of DR antigens seemed to be correlated inversely with the amounts of anti-DNA antibody at the time of blood sampling. The decreased expression of HLA-DR antigens was also shown in peripheral blood monocytes from SLE patients (40). The involvement of DR antigens or Ia antigens in autoimmune diseases has been suggested (41–44). Self-antigens associated with self-Ia do not induce immune response through T cell reaction. When the self-Ia molecule alters, the immune response occurs (43, 44). The alteration of Ia antigens, both qualitatively and quantitatively, may occur through chronic viral infection, drugs, mutation, and gene conversion (41, 42). The theory is partly supported by experiments showing that the recognition of foreign Ia antigens by T cells led to an SLE-like disease in mouse (43, 44). Alternatively, abnormal Ia recognition may occur by T cell defect, or miseducation of helper and/or suppressor T cells (41). Whatever the mechanism is, the class II antigens play a key role in the immune regulation, and therefore, aberrant DR expression might be closely correlated with the immunoregulatory dysfunctions in SLE disease.

Two different mechanisms could result in reduced DR antigens. First, blockage of the surface antigens by autoantibodies causes an apparent reduction in DR molecules, and second, malfunctions of the DR producing system including the gene itself could account for the reduced amount of DR molecules. The first possibility is based on an observation that SLE patients produce anti-leukocyte antibodies (45, 46). Sera from SLE patients contain antibodies that block the binding of anti-Ia antibody to T cells that express Ia antigens. It has been suggested that production of anti-Ia antibody in vivo by SLE patients led to reduced suppressor T cell function (46). However, no evidence
has been presented showing that SLE B cell functions are modified by autoantibodies blocking surface Ia antigens. Also, no anti-DR antibodies were detected in vitro under our experimental conditions, and the first possibility should be excluded. The second possibility should be considered by two different mechanisms: (a) SLE cells produced normal amounts of DR antigens, but their structure is altered, and L243 anti-DR antibody did not bind to them; and (b) SLE cells produced reduced amounts of DR antigens. The first assumption is probably, but we favor it less: the structural mutation, if any, should occur at the whole DR molecule, since L243 reacts at the nonpolymorphic regions of DR antigen, most likely at the α-chain; no evidence has been presented showing the presence of “mutated DR antigens”. The second assumption is more favorable for us, considering the reduced amounts of DRα message and the methylation patterns of DRα gene regions as discussed below.

Association of hypermethylation with reduced DR expression. We examined the extent of methylation at at least five CCGG sites in the HLA-DRα locus. In both normal and SLE cell lines, the methylation at a given CCGG site was variable within a cell line. Since the DRα gene is a single-copy gene, a CCGG site from a single clone should be either methylated or unmethylated. The variable methylation patterns observed in normal and SLE cell lines could result from studying mixed populations of cells, which might be a mixture of transformed clones in which the sites are methylated, with other clones, in which the sites are not methylated. Thus, established B cells are heterogeneous in terms of the DRα gene methylation.

A striking feature of the DRα gene from healthy individuals is a consistent pattern of methylation. B cell lines from 12 healthy persons showed exactly the same methylation pattern. This indicates that the ratio of cells with differently methylated DR genes is consistent, or well balanced in healthy persons. Striking features of the DRα locus from SLE patients are (a) the absence of consensus methylation pattern, and (b) hypermethylation. B cell lines from 28 SLE patients showed a variety of methylation patterns, indicating that the ratio of cells with differently methylated DR genes is patient-specific. Since the extent of methylation in SLE was higher than that in controls, cells with hypermethylated DRα gene regions could predominate in SLE patients.

The DNA hypermethylation is often associated with transcriptionally inactive genes (19–21). Cells with unmethylated DR genes may express DR antigens, and cells with methylated DR genes may express no or few DR antigens. The apparently low expression of DR antigens in SLE could be the result of predominance of cells with hypermethylated DR genes among total B lymphocytes. Note, however, that type II cells, which express apparently the same amounts of DR antigens as normal cells, also showed hypermethylation in HLA-DRα regions. The methylation extent of type II cells was between that of normal cells and that of type I cells, which express reduced amounts of DR antigens. The question why type II cells express DR antigens normally despite the apparent hypermethylation can be explained by a theory that, for the normal gene functions, methylation/demethylation is critical at the particular sites rather than at the whole genome (22–25). In mammalian DNA, the frequency of the CCGG tetramer, of which methylation was examined in the present study, is ~5% of the total methylatable CpG dinucleotide (47). Since 60% of CpG was methylated in B cell lines, the methylation at CCGG represents only 8% of the total methylation. Thus, it is reasonable to speculate that the DRα gene contains some critical regions other than CCGG, and that their methylation might be associated with relatively high methylation levels of other sites, including CCGG. The critical sites might not be methylated in type II cells, and might be methylated in type I cells, resulting in full expression in the former and reduced expression in the latter. There is a report that might support this speculation. In DNA from T cells that do not express DRα messenger RNA (mRNA), all CCGG sites in the DRα locus examined were methylated. T cells that were infected with T cell leukemia virus (HTLV) expressed DR antigens, and the DRα locus was found to be demethylated at some CCGG sites (48).

Relation to normal and SLE B cell differentiation. Ia antigens are not expressed in stem B cells: as B cells mature, Ia molecules are gradually co-expressed with surface immunoglobulin at the membrane (38, 49). Thus, Ia genes that are silent in stem cells are activated during the course of B cell differentiation. Silent genes are usually methylated: as such genes are transcriptionally activated, demethylation occurs. Examples are the β-globin gene (50) and immunoglobulin genes (51). The same sequence may occur with HLA-DR genes. In normal cells, the genes become demethylated to express DR antigens. In many SLE cells, the demethylation probably does not occur, leaving the DR genes transcriptionally silent even after B cells mature.

Peripheral blood B cells from SLE patients were found to have reduced responsiveness to B cell mitogens compared with those from normal persons (2, 52). The low response was attributed to higher spontaneous B cell proliferation in SLE patients (2). Also, SLE patients showed the greater numbers of Ig-producing cells (2, 4). It was concluded that SLE patients have larger amounts of activated B cells in vivo than do normal persons (2, 52), i.e., SLE B cells could be more differentiated than normal cells in terms of antibody production. Nevertheless, we observed the reduced DR expression in SLE B cells, i.e., SLE B cells are less differentiated in terms of DR expression. These observations suggest that the abnormal B cell functions in SLE might occur through disturbance of a concerted regulation of genes involved in differentiation. Compared with normal cells, SLE cells might contain some genes that are suppressed and others that are activated. In lupus-prone mice, primary B cell abnormalities have been identified (53–57), and the involvement of inherent genetic factors was suggested in lupus disease (55–57). We propose that one of factors that are involved in B cell gene regulation is DNA methylation, and that aberrant methylation/demethylation during differentiation results in dysfunctions in SLE B cells.

Finally, note that our results obtained with EBV-transformed cells might reflect the molecular aspects of limited B cell clones in vivo. First, the target of EBV seems to be restricted to certain populations of B cells. Particularly, since most SLE patients are treated with glucocorticoids (GC), the susceptibility of SLE B cells to EBV might be influenced by GC therapy. It has been reported that GC treatments lowered the Ig production in human lymphocytes (58) and promoted B cell differentiation (59). When peripheral B cells were fractionated by density gradient centrifugation, only high density resting cells were infected. Activated low density cells were resistant to EBV infection (60). Since the phenotype of established cell lines from both SLE and normals did not differ (Table I), the susceptibility to EBV was the same between SLE and normal B cells. In other words, even if GC treatment affected the population ratio of B cells, as far as concerned with EBV-transformed cells, the biochemical properties...
should be comparable between SLE and normals in vitro. But we should keep in mind that GC might alter the ratio of B cell subsets in vivo. Second, GC treatment might influence the DR gene expression at the transcriptional and translational levels. Several lines of evidence have indicated that steroid treatment stabilized some mRNAs (61) and promoted and/or inhibited some mRNA productions (62). We cannot exclude a possibility that the same sequence might occur in HLA-DRα mRNA production, although no evidence has so far been presented to support this idea. Further experiments are necessary to clarify these problems.

Acknowledgments

This work was supported by grants from National Institutes of Health (AM31240) and from the Vehicle Racing Commemorative Foundation.

References