Polymorphism of major histocompatibility complex extended haplotypes bearing HLA-DR3 in patients with rheumatoid arthritis with gold induced thrombocytopenia or proteinuria

Dharam P Singal, Barbara Reid, Donna Green, Merl D'Souza, William G Bensen, W Watson Buchanan

Abstract

The distribution of DR3 and of extended haplotypes bearing DR3 was studied in three groups of subjects: 35 patients with rheumatoid arthritis (RA) with gold induced thrombocytopenia or proteinuria, 185 patients with RA without these side effects, and 300 normal healthy controls. The extended haplotypes bearing DR3 were analysed with cDNA probes for DRα, DRβ, DQα, and DQβ genes. The data showed that the prevalence of DR3 was significantly higher in patients who developed gold induced thrombocytopenia or proteinuria than in normal controls or patients with RA without these side effects. Distribution of three extended haplotypes bearing DR3 (B8,DR3; B18,DR3; non-B8,non-B18,DR3) in patients with RA with thrombocytopenia or proteinuria was significantly different from that in normal controls, but not from that in patients with RA without these toxic reactions. Southern blot analysis of DR, DQ genes with cDNA probes showed that the extended haplotype bearing B8,DR3, which carries DQA2.1 and DQB2.1 genes, was present in a significantly higher proportion of patients with RA with gold induced thrombocytopenia or proteinuria (22/24, 92%) than in patients with RA without these side effects (32/45, 71%) or normal subjects (40/61, 66%). The data suggest that the genomic region on chromosome 6 involved in susceptibility to gold induced thrombocytopenia or proteinuria should be extended to the DQA2, DQB2 gene loci.

Gold complexes are an established form of treatment for rheumatoid arthritis (RA). Numerous well controlled studies have shown the clinical efficacy of chrysotherapy in RA. Their use in the management of RA is limited by a fairly high incidence of adverse toxic reactions. For the most part, these are minor reactions; however, they may occasionally be life threatening and in rare instances have resulted in death. Of these adverse reactions, thrombocytopenia and proteinuria are the most common.

During the last few years it has been suggested that the genetic markers belonging to the human leucocyte antigen (HLA) system may be important in predicting toxic reactions to chrysotherapy. A number of investigators have described a correlation between gold induced thrombocytopenia and proteinuria and HLA-DR3, whereas three reports did not find such a correlation. We noted previously a significantly higher prevalence of DR3 in patients with RA with gold induced thrombocytopenia and proteinuria. In addition, Batchelor et al found a significantly higher incidence of DR3 in a family study of patients with RA who developed gold induced nephropathy. Despite these observations the nature of associations between development of gold induced thrombocytopenia and proteinuria and the presence of DR3 is not clear.

Extensive polymorphism in extended haplotypes bearing DR3 has been described recently by Southern blot analysis. With cDNA probes for α and β chain DR, DQ genes, two extended haplotypes bearing DR3 have been described in white populations. In addition, polymorphism in the DQA2 and DQB2 chain genes in extended haplotypes bearing DR3 can be easily identified. The objectives of this study were, therefore, to investigate the distribution of these extended haplotypes bearing DR3 by Southern blot analysis in patients with RA, with or without gold induced thrombocytopenia or proteinuria, and in normal subjects.

Patients and methods

Two hundred and twenty white patients with RA who attended the rheumatology clinics at the St Joseph's Hospital and the McMaster University Medical Center in Hamilton, Ontario, Canada were studied. All patients had moderate to severe definite classical seropositive RA. These patients had earlier been treated with non-steroidal anti-inflammatory drugs and all had had an incomplete response requiring treatment with sodium aurothiomalate. Patients were treated with sodium aurothiomalate as recommended by Gottlieb—that is, test doses of 10 mg and 25 mg intramuscularly (IM) one week apart, 50 mg IM weekly doses for 20 consecutive weeks, maintenance regimen of 50 mg IM every two weeks for three months, and thereafter 50 mg IM every three weeks indefinitely. Patients were reviewed for (a) thrombocytopenia (platelet count <100x10^9/L) and (b) significant proteinuria (>1 g/day). Eighteen patients who developed thrombocytopenia and another 17 who developed proteinuria after gold treatment were investigated for HLA-DR antigens. The remaining 185 patients with RA did not develop these side effects after gold treatment. Three hundred white, normal, healthy, unrelated subjects served as controls. Eight DR3 positive, homozygous B lymphoblastoid cell lines and members of 35 DR3 positive (20 normal controls and 15 patients with RA) families were also studied.
Peripheral blood lymphocytes were typed for HLA-A, B, C antigens by the microdroplet lymphocyte cytotoxicity test. HLA-DR, DQ typing was performed on a B cell enriched lymphocyte population separated from peripheral blood lymphocytes by the nylon wool method. HLA specificities were defined on the basis of reactivity with antisera in the 9th and 10th International Histocompatibility Workshops and with antisera in the Canadian Red Cross tissue typing trays.

Samples of genomic DNA (7 μg), prepared from peripheral blood lymphocytes, were separately digested with 20 units of different restriction enzymes—for example, Bgl II, Bam HI, etc., for 18 hours at 37°C with buffer conditions specified by the manufacturer (Bethesda Research Laboratories) and were electrophoresed on a 0.9% agarose gel in TAE buffer (40 mmol/l TRIS-acetate, 1 mmol/l EDTA) for 18 hours at 50 mA. Molecular weight markers on each gel contained 1.5 μg of Hind III-digested lambda DNA (BRL). After electrophoresis, gels were denatured and neutralised before transfer (alkaline) to nylon membrane (Biotrace RP, Gelman) by Southern’s method. After transfer, filters were prehybridised at 42°C and hybridised with oligonucleotide labelled DRα, DRβ, DQα, and DQβ cDNA probes at 42°C for 48 hours. After hybridisation, blots were first washed for five minutes twice at room temperature with 2xSSPE (0.3 mol/l NaCl, 20 mmol/l NaH₂PO₄, H₂O, and 2 mmol/l EDTA); the filters were then washed for 15 minutes twice at 65°C, once with 2xSSPE and 0.5% sodium dodecyl sulphate and once with 0.5xSSPE. The filters were autoradiographed with intensifying screens on Kodak XAR film at -70°C for 3–10 days.

Statistical analysis of the results was performed by the χ² test. Relative risk (RR) values were calculated by the method of Woolf.

Table 1: Prevalence (%) of HLA-DR3 and DR4 in normal controls, in patients with rheumatoid arthritis (RA) with thrombocytopenia or proteinuria, and in patients with RA without these toxic reactions. Number (%) of patients is shown.

<table>
<thead>
<tr>
<th>HLA-DR antigen</th>
<th>Normal healthy controls (n=300)</th>
<th>Patients with RA with thrombocytopenia or proteinuria (n=35)</th>
<th>Patients with RA without these toxic reactions (n=185)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DR3</td>
<td>61 (20)</td>
<td>24 (69)*</td>
<td>45 (24)</td>
</tr>
<tr>
<td>DR4</td>
<td>68 (23)</td>
<td>13 (37)</td>
<td>105 (57)†</td>
</tr>
</tbody>
</table>

*Significantly higher than normal controls (p<10⁻⁴; RR=8.55), and than patients with RA without side effects (p<10⁻⁴; RR=6.79).
†Significantly higher than normal controls (p<10⁻⁴; RR=4.48), and than patients with RA with thrombocytopenia or proteinuria (p<0.04; RR=2.22).

Table 2: Distribution of extended haplotypes bearing HLA-DR3 in normal healthy controls, in patients with rheumatoid arthritis (RA) with thrombocytopenia or proteinuria, and in patients with RA without these toxic reactions after gold treatment.

<table>
<thead>
<tr>
<th>Number of DR3 haplotypes</th>
<th>B8,DR3 No (%)</th>
<th>B18,DR3 No (%)</th>
<th>Non-B8,non-B18,DR3 No (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal healthy controls</td>
<td>61</td>
<td>40 (66)</td>
<td>6 (10)</td>
</tr>
<tr>
<td>Patients with RA without toxic reactions</td>
<td>45</td>
<td>32 (71)</td>
<td>3 (7)</td>
</tr>
<tr>
<td>Patients with RA with thrombocytopenia or proteinuria*</td>
<td>24</td>
<td>22 (92)</td>
<td>1 (4)</td>
</tr>
</tbody>
</table>

*Distribution of extended haplotypes bearing DR3 significantly different from that in normal healthy controls (p<0.05).
the normal healthy controls, but not from that in patients with RA without adverse reactions.

We examined the RFLP in the α and β chain genes of the DR, DQ subregions of the extended haplotypes bearing DR3. In the DR subregion two extended haplotypes bearing DR3 were identifiable as follows (fig 1):

1. B8,DR3 and non-B8,non-B18,DR3: DRA1 (Bgl II=4-20 kb; Eco RV=8-20 kb), DRB1 (Hinc II=2-50 kb), DRB3.1 (Bam HI=4-00 kb; Hind III=2-60 kb; Pvu II=5-60 kb; Pst I=1-07 kb).

2. B18,DR3: DRA2 (Bgl II=4-50 kb, Eco RV=12-50 kb), DRB1 (Hinc II=2-50 kb), DRB3.2 (Pvu II=5-40 kb, Eco RI=2-90 kb).

Similarly, these two extended haplotypes bearing DR3 were demonstrable for DQA1 and DQB1 chain genes (fig 2). The distribution of these two types of extended haplotypes bearing DR3 (B8,DR3 and non-B8,non-B18,DR3: B18,DR3) in normal subjects (55:6), patients with RA with thrombocytopenia or proteinuria (23:1), patients with RA without these side effects (42:3) was essentially similar ($\chi^2=0.18$ to 0.73; NS).

Southern blots were then analysed for distribution of RFLP for DQA2 and DQB2 genes. Two DQA2 genes were identifiable; one DQA2.1 (Taq I=2-25 kb; Pst I=8-00 kb) correlated with the B8,DR3 haplotype, and the other DQA2.2 (Taq I=2-15 kb; Pst I=6-84 kb) was associated with B18,DR3 and non-B8,non-B18,DR3 alleles (fig 2). Similarly, two DQB2 genes, DQB2.1 associated with B8,DR3 (Pst I=2-90 kb) and DQB2.2 with B18,DR3 and non-B8,non-B18,DR3 (Pst I=4-10 kb), were demonstrable (fig 2). Thus the combined analysis for DR and DQ (α and β chain) genes distinguished three extended haplotypes bearing DR3: B8,DR3; B18,DR3; and non-B8,non-B18,DR3.

Based on RFLP for DQA2 and DQB2 chain genes, two extended haplotypes bearing DR3, B8,DR3 and non-B8,DR3 (including B18,DR3 and non-B8,non-B18,DR3), were demonstrable. The distribution of the extended haplotype bearing B8,DR3, which carries the DQA2.1 and DQB2.1 genes in patients with RA with thrombocytopenia or proteinuria (92%), was significantly different from its distribution in patients without these side effects (71%; p<0.05), and in normal healthy subjects (66%; p<0.015).

Discussion

The data in this study show that the incidence of DR3 was significantly higher in patients with RA who developed gold induced thrombocytopenia or proteinuria than in normal healthy controls or patients with RA who did not develop these side effects after gold treatment. In addition, patients with RA without these side effects had a significantly higher prevalence of DR4 than the normal controls or patients with RA with thrombocytopenia or proteinuria. The higher prevalence of DR4 in patients with RA with thrombocytopenia or proteinuria was not statistically different from that in normal controls. The distribution of three extended haplotypes bearing DR3 in patients with RA with thrombocytopenia or proteinuria was significantly different from that in normal healthy controls, but not from that in patients with RA without these toxic reactions. Southern blot analysis of HLA-D region genes with α and β chain cDNA probes showed that the B8,DR3 extended haplotype, which carries DQA2.1 and DQB2.1 genes, was present in a significantly higher proportion in patients with RA with thrombocytopenia or proteinuria than in normal subjects or patients with RA without these gold induced side effects. The data suggest that the genomic region on chromosome 6 involved in susceptibility to gold induced thrombocytopenia.
or proteinuria should be extended to the DQA2, DQB2 gene loci.

A number of investigators have found a correlation between DR3 and gold induced thrombocytopenia or proteinuria. On the other hand, three reports could not confirm that the patients with DR3 were more likely to have toxic reactions to gold than the patients without these antigens. These reported differences may be related to patient selection (racial factors, seropositive, or seronegative) and to the gold preparation used. The results in our study are in agreement with earlier reports, which showed patients with RA carrying DR3 are at a higher risk of developing gold induced thrombocytopenia or proteinuria. In addition, the present data extend the earlier results by showing that the distribution of extended haplotypes bearing DR3 in patients with RA with thrombocytopenia or proteinuria was significantly different from that in normal healthy controls.

The HLA-D region of human major histocompatibility complex consists of multiple gene loci encoding α and β subunits that form αβ heterodimers. The HLA-D region genes are arranged in a block consisting of several subregions including DR, DQ, and DP, with each subregion containing at least one α chain and one β chain gene. In recent years RFLP analysis of the HLA-D region genes (DR (A1,B1,B3,B4); DQ (A1,A2,B1,B2)) has developed into a powerful tool for genomic typing. A number of investigators have examined the polymorphism of extended haplotypes bearing DR3 by Southern blot analysis, and the polymorphism detected by RFLP analysis of DR, DQ genes in these DR3 extended haplotypes correlated with phenotypic polymorphism recognised by serology. These data strongly suggest that it is possible to identify three different extended haplotypes bearing DR3 from phenotypes by RFLP. In this study we examined members of 35 DR3 positive families, including 15 families of patients with RA. In the remaining cases we used Southern blot analysis of genomic DNA, as suggested by a number of investigators, for the identification of extended haplotypes bearing DR3. The distribution of different extended haplotypes bearing DR3 in normal subjects and in patients with RA in this study was the same as that found by other investigators in normal subjects and in patients with RA.

In addition, in an earlier study all (15 of 15) DR3 positive patients with RA who developed gold induced nephropathy carried the B8,DR3 extended haplotype. Our data, therefore, confirm the earlier results and extend them as the distribution of DQA2 and DQB2 chain genes on haplotypes bearing DR3 in patients with RA with thrombocytopenia or proteinuria was significantly different from that in normal healthy controls or in patients with RA without these gold induced toxic reactions.

Because of linkage disequilibrium between polymorphic alleles of HLA genes, certain combinations of alleles are inherited as extended haplotypes. Several independent examples of extended haplotypes bearing DR3 have recently been examined. The independent examples belong to an extended haplotype bearing DR3 and are similar in RFLP patterns, suggesting that if an allele can be divided into antigens most of the extended haplotypes belonging to a subtype are similar. These data suggest that alleles in B, DR, DQ regions of extended haplotypes bearing DR3 are fixed and thus support the extended haplotype theory.

HLA-D region antigens play a crucial part in regulation of immune response and in pathogenesis of diseases. The results in our study suggest that subjects carrying a particular extended haplotype bearing DR3 may display disease associations that differ from those of subjects positive for other DR3 extended haplotypes. Similar results have been reported for gluten sensitive enteropathy, where only the B8,DR3 haplotype, and not the other DR3 haplotypes, is increased in patients. In contrast, both B8,DR3 and B18,DR3 extended haplotypes are increased in insulin dependent diabetes mellitus.

This work was supported by research grants from the Arthritis Society and the Medical Research Council of Canada. The DRα (pII-p-I), DRβ (pII-p-II), and DQ (pII-p-I) cDNA probes were kindly given by Dr P A Peterson, Uppsala, Sweden and the DQβ (pDCox107) probe by Dr K Tsui, Kanagawa, Japan.


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doi: 10.1136/ard.49.8.582

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