A Restriction Fragment of the C2 Gene Is a Unique Marker for C2 Deficiency and the Uncommon C2 Allele C2*B (A Marker for Type 1 Diabetes)

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Abstract

There are three common C2 protein alleles in Caucasians, C2*C, C2*B, and C2*Q0, with allele frequencies of 0.96, 0.03, and 0.01, as well as Sst I RFLP variants of 2.75, 2.7, 2.65, 2.55, and 2.4 kb, with frequencies of 0.017, 0.533, 0.358, 0.017, and 0.075. Thus, C2*C is informatively split by the RFLP. Of 94 nonrandomly ascertained Caucasian complotypes, 77 contained C2*C, four contained C2*Q0, and 13 had C2*B. None of the C2*C-containing complotypes carried the 2.75 kb Sst I fragment and all of the complotypes with C2*B or C2*Q0 carried it. All of the C2*Q0 alleles were associated with C4A*4, C4B*2 in the complotype S042 as previously reported. C2*B was usually (9/13) in the complotype S042, occasionally (1/13 each) in S045, S041, S04(4,3)0, and S031. Thus, the association of the C2 2.75-kb fragment was with C2*B and C2*Q0, not with C4A*4, C4B*2, or even C4A*4 alone. The complotype SC42 was associated with the 2.65-kb Sst I fragment in four of five instances and in a single example with the 2.7-kb fragment. C2*B and C2*Q0 possibly had a common evolutionary ancestor complotype which carried the 2.75-kb Sst I fragment, and BF*S, C4A*4, and C4B*2. C2*B (particularly as the haplotype HLA-Bw62, SB42, DR4) is associated with type 1 diabetes but C2*Q0 is protective. (J. Clin. Invest. 1991. 88;2142–2145.) Key words: human complement C2·C2 gene·C2 allele·C2 deficiency·restriction fragment length polymorphism

Introduction

There is a single gene for the human complement protein C2,1 situated telomeric to the genes for factor B (BF) and C4 (C4A and C4B) and centromeric to HLA-B (1,2) in the mid-portion of the short arm of the sixth human chromosome. C2 protein shows relatively little polymorphism in Caucasian populations, with ~96% of all chromosomes carrying the common allele C2*C and around 3% carrying the gene for a more basic variant C2*B (3–5). Around 1% of chromosomes have a null gene, C2*Q0, which, in the homozygous state, is associated with a total absence of C2 protein in serum and with an increased risk for systemic lupus erythematosus (6). The C2*Q0 allele has a grossly normal-sized gene in genomic DNA (7). Both C2*B and C2*Q0 are in very strong linkage disequilibrium with the specific alleles BF*S, C4A*4, and C4B*2, forming the complotypes (four allele sets behaving in populations and families as single genetic units) SB42 and S042 (8,9). Less commonly, C2*B was found on SB41, SB45, or SB43.

A multiallelic polymorphism in C2 genomic DNA is detected after digestion with Sst I (or several other enzymes) and Southern blotting with a 300-bp 5' C2 genomic probe (10–12). The variants represent differences in the size of an intron near the 5' end of the C2 gene and include Sst I fragments of 2.4, 2.55, 2.65, 2.7, and 2.75 kb. They are of particular interest because they allow the subdivision of haplotypes with C2*C.

This report concerns the striking linkage disequilibrium of the 2.75-kb Sst I variant in C2 genomic DNA with C2*B and C2*Q0 in the complotypes S042, SB42, SB45, SB41, SB4(4,3)0, and S031. SB42, SB45, SB41 and S042 carry most instances of C2*B and C2*Q0 in Caucasian populations.

Methods

Subjects. Lymphoblastoid lines were established by Epstein-Barr virus transformation of peripheral blood lymphocytes from members of 24 families in which at least one member had type 1 diabetes mellitus. In addition, lymphoblastoid lines were established from peripheral blood lymphocytes of MHC homozygous individuals either by us or by the 10th International Histocompatibility Workshop (13). For the present study, a total of 94 independent normal and diabetic haplotypes were analyzed for the Sst I polymorphism in the 5' portion of the C2 gene. Of these, 54 diabetic and 37 normal haplotypes were randomly ascertained and were used for frequency comparisons. The compilation and analysis of normal and diabetic haplotypes were as previously described (14).

Complotype and HLA determinations. Plasma from whole blood collected into EDTA was used to test for genetic polymorphisms in C4 (C4A and C4B), BF, and C2 proteins. Complotypes were assigned from similar studies in immediate relatives. For C4 typing (15), the plasma was treated with neuraminidase and, in some instances, carboxypeptidase (16) and then subjected to agarose gel electrophoresis and immunofixation with goat anti-human C4 (Atlantic Antibodies, Stillwater, MN). C2 types were determined by isoelectric focusing in thin layer polyacrylamide gel and a C2-sensitive agarose gel overlay incorporat-
ing antibody-sensitized sheep erythrocytes (3). Of the four haplotypes with C2*Q0, two were in a homozygote and all were with HLA-B18 and DR2. For the two heterozygotes, half-normal C2 levels in serum were associated with the haplotype in family studies. BF typing (17) was by agarose gel electrophoresis and immunofixation with anti-human BF (Atlantic Antibodies). The nomenclature for C4 is described previously (15, 18).

Individual alleles are italicized and designated by locus name in capital letters, an asterisk, and a number or "00" if null (e.g., C4A*4, C4B*2, or C4B*Q0). Phenotypes, variants or proteins are given with roman capital letters, a space, and the same number or symbol as the corresponding allele (C4A 4, C4B 2, or C4B Q0). Complotype are designated by their BF, C2, C4A, and C4B alleles, in that arbitrary order (9). Null or Q0 alleles are simply 0. Thus, SB42 indicates the complotype BF*S, C2*B, C4A*4, C4B*2 and S042 is BF*S, C2*Q0, C4A*4, C4B*2.

HLA-A, B, and DR generic typing was performed by standard methods (19). Extended haplotypes were defined previously as HLA-B, complotype, DR allele sets exhibiting significant linkage disequilibrium (20).

Restriction fragment length polymorphism (RELP). DNA was extracted from the B lymphoblastoid lines by the method of Gross-Bellard and colleagues (21). 5 µg of DNA were digested to completion with Sst I restriction endonuclease, using conditions recommended by the manufacturer (Bethesda Research Laboratories, Gaithersburg, MD). Electrophoresis of Sst I-digested DNA was carried out for 96 h at 60 V at 4°C in 2.0% agarose gel in 1X TAE (0.04 M Tris acetate/0.002 M EDTA) with continuous buffer circulation, in a minor modification of the method described previously (11).

DNA cleavage fragments were transferred to Nytran (Schleicher & Schuell, Keene, NH) or Sure-blot membranes (Oncor, Gaithersburg, MD) (22). The 300-bp C2 probe was derived from the genomic clone pG850 and was described previously (10). It was labeled with alpha [32P]dCTP by the random primer method (23). Prehybridization and hybridization were carried out at 45°C, the latter overnight. Membranes were washed twice at room temperature with 0.1x standard saline citrate (SSC), 0.1% SDS, and a third time with the same solution at 52°C for 20-60 min. The membranes were then exposed to x-ray film with an intensifying screen at −70°C for 2-5 d for autoradiography.

Results

In the overall set of chromosomes, including both normal and diabetic haplotypes, there were 77 instances of complotype with C2*C, 13 with C2*B, and four with C2*Q0 (including two from a deliberately selected homozygote). The C2 Sst I restriction fragment length variant of 2.75 kb was found only on haplotypes with the complotypes SB42, SB45, SB41, SB4(3,4), SB31, and S042. The distribution of C2 Sst I variants with respect to C2 protein alleles is shown in Table I. Of five examples of SC42, four were found on haplotypes that carried the 2.65-kb Sst I variant and one SC42 was with the 2.7-kb fragment. All other complotype carried only the previously described C2 variants of 2.4, 2.65, or 2.7 kb. The C2 variant of 2.55 kb was not encountered in this study.

In all instances, the 2.75 kb Sst I variant was inherited in the families in which it occurred in Mendelian fashion and in every case cosegregated with the C2*B or C2*Q0 variant-containing complotype. This is illustrated in Fig. 1.

From the known frequencies of C2*B (0.0282) and C2*Q0 (~0.01) in a large population of normal Caucasian chromosomes (n = 2,180), it can be estimated that the frequency of the 2.75-kb Sst I C2 allele is 0.04, assuming that it occurs only but invariably in association with these alleles. This is in the same range as the frequency of 0.017 estimated directly on a relatively small number of normal caucasian haplotypes.

Of 59 randomly ascertained type I diabetes haplotypes, the 2.75-kb Sst I C2 fragment was found on 5, whereas it was found on two of 37 family control haplotypes. The relative risk for positivity for the 2.75-kb Sst I C2 allele among patients with diabetes compared with family controls was 1.8. The association in the two populations of haplotypes were different. All instances on patient haplotypes were with C2*B, whereas in family normal control haplotypes, two independent instances were with C2*Q0 and one was with C2*B.

To assess whether C2*B is a significant marker for type I diabetes, its frequency among 365 independent Caucasian patient haplotypes was compared with that among 2180 normal Caucasian control haplotypes. C2*B had a gene frequency of 0.0712 in diabetic haplotypes and 0.0271 in normal haplotypes.

### Table I. Sst I C2 DNA Variants in Relation to C2 Alleles

<table>
<thead>
<tr>
<th>C2 Sst I variant</th>
<th>C2*C</th>
<th>C2*B</th>
<th>C2*Q0</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.75</td>
<td>0</td>
<td>13</td>
<td>4</td>
</tr>
<tr>
<td>2.7</td>
<td>35</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2.65</td>
<td>36</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2.4</td>
<td>6</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Common Origin of C2 Deficiency Gene and C2*B

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Figure 1. (A) Family C with C2*B in the complotype SB42. The mother of this family is homozygous for SB42 (BF*S, C2*B, C4A*4, C4B*2) and the 2.75-kb Sst I C2 fragment. The father is heterozygous for these genetic markers. Two children, II-2 and II-4, are homozygous who have coinherted SB42 and the C2 Sst I 2.75-kb fragment from both parents. The other two children, II-1 and II-3, are heterozygotes, having inherited the a haplotype (FC(3,2,0), 2.7 kb) from their father as well as the c or d SB42, 2.75-kb haplotype (distinguished by other MHC genes, not shown) from their mother. (B) Family H with C2 deficiency. The C2 deficiency complotype S042 (BF*S, C2*Q0, C4A*4, C4B*2) segregates with the 2.75-kb C2 Sst I fragment from the father and mother. Both children are homozygous for C2 deficiency and the 2.75-kb Sst I C2 fragment.


Table II. Possible Extended Haplotypes with C2* B on Independent Type I Diabetes and Normal Haplotypes

<table>
<thead>
<tr>
<th>Haplotype</th>
<th>Normal n = 2180</th>
<th>Diabetes n = 365</th>
<th>Haplotype ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>HLA-Bw62, SB42, DR4</td>
<td>9</td>
<td>0.0041</td>
<td>9</td>
</tr>
<tr>
<td>HLA-Bw55, SB45, DRw6</td>
<td>6</td>
<td>0.0027</td>
<td>1</td>
</tr>
<tr>
<td>HLA-Bw60, SB42, DR4</td>
<td>3</td>
<td>0.0014</td>
<td>2</td>
</tr>
<tr>
<td>HLA-B7, SB42, DR4</td>
<td>3</td>
<td>0.0014</td>
<td>2</td>
</tr>
<tr>
<td>HLA-B37, SB42, DR4</td>
<td>2</td>
<td>0.0009</td>
<td>0</td>
</tr>
<tr>
<td>HLA-B35, SB42, DR4</td>
<td>2</td>
<td>0.0009</td>
<td>0</td>
</tr>
</tbody>
</table>

There were 59 normal and 26 diabetes haplotypes with C2* B. Fraction refers to the number of such haplotypes shown divided by the total normal or diabetes haplotypes. D/N refers to the ratio of frequencies among diabetes and normal haplotypes as specified.

(P < 0.0001), relative risk for positivity for C2* B = 2.8). On the other hand, C2* Q0, almost always on the extended haplotype [HLA-B18, SO42, DR2] or its fragments, was found on ~1% of normal haplotypes but on no diabetic haplotype.

The MHC haplotypes on which C2* B occurred more than once on either independent patient or independent normal haplotypes are shown in Table II. These haplotypes represent possible extended or conserved haplotypes. In both populations, the haplotype HLA-Bw62, SB42, DR4 was the most common. This haplotype was significantly increased among diabetic haplotypes (P < 0.0004). The haplotype HLA-Bw55, SB45, DRw6, previously suspected of being an extended haplotype, was noted on six normal haplotypes but only one patient haplotype. Other C2* B-bearing haplotypes were not increased among diabetics.

Discussion

Because the 2.75-kb Sst I C2 restriction fragment is always associated with the SB42, SB45, SB41, SB31, SB4(3), or SO42 complotypes and reflects an unusual size intron common to these complotypes, it is likely that C2* B and C2* Q0 had a common ancestor. Perhaps surprisingly, because SB42 and SB42 may have arisen from SC42, the association does not include SC42 in caucasians. Thus, the extensive polymorphism in the C2 gene detected with Sst I (10–12) or a number of other restriction enzymes (24), already shown to be capable of dividing haplotypes carrying C2* C and BF* F from those with C2* C and BF* S (25), is also useful in distinguishing C2* B, and C2* Q0 from C2* C.

This is particularly important for C2* Q0, for several reasons. Heterozygotes for C2 deficiency can be difficult to detect by measuring C2 serum levels alone (26). Homozygotes have increased susceptibility to systemic lupus erythematosus and the availability of an additional indicator of the C2 null allele is clearly useful.

Our data confirm that C2* B is increased in frequency among type I diabetics (27) and we now note a marked decrease in frequency or absence of C2* Q0, undoubtedly related to the fact that the extended haplotype which usually carries it also carries HLA-DR2. Because the 2.75-kb RFLP variant is associated with both C2* B (a marker for susceptibility to diabetes) and C2* Q0 (a marker for resistance), it is not as good a marker for disease susceptibility or resistance as the protein variants in C2.

We have found a number of C2* B-bearing haplotypes, some of which may be extended or conserved, particularly HLA-Bw62, SB42, DR4, and HLA-Bw55, SB45, DRw6. HLA-Bw62, SB42, DR4 is a probable conserved or extended haplotype with a markedly increased frequency in diabetics. It clearly contributes to the associations of both C2* B and DR4 with diabetes. These findings are consistent with previous observations of linkage disequilibrium between C2* B and HLA-B15(w62) (5, 28, 29). The haplotype HLA-Bw55, SB45, DRw6 has been noted earlier to carry susceptibility for pemphigus vulgaris (30) and is probably a carrier of a defective CYP21 gene causing congenital adrenal hyperplasia (31).

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References


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