Genetic Analysis of Low Vβ3 Expression in Humans

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Summary

While studying the T cell receptor (TCR) repertoire of normal individuals, we found that more than 20% of adults have low levels of circulating Vβ3.1+ T cells in both CD4 and CD8 populations. A similar frequency was found in fetal cord blood samples, suggesting that in most cases, the Vβ3.1low phenotype is inherited. In support of this conclusion, children expressing low levels were only found in families where one of the parents expressed this phenotype. In two large families, genetic studies showed that low expression was a recessive trait and dependent on inheritance of particular TCR VB gene complexes. Family members with the low phenotype, however, expressed Vβ3.1 genes with normal sequences and expressed normal levels of receptor per cell. Results from these families suggest that up to 50% of normal individuals may carry a Vβ3.1 allele that is defective in its ability to rearrange effectively. In another large family, low expression in one individual was shown not to be determined by genes within the TCR VB gene or major histocompatibility complexes, suggesting a different mechanism for low Vβ3.1+ T cells. Overall, our results describe novel mechanisms that result in low levels of Vβ3.1+ T cells in a relatively large subset of the normal human population.

Materials and Methods

Sample Preparation. Mononuclear cells were isolated from peripheral blood or cord blood samples by Ficoll-Hypaque density gradient separation, and used for analysis of TCR VB expression and preparation of RNA (11). For genetic studies in families, neutrophils were also isolated from the pellet during gradient separation of mononuclear cells and used for preparation of genomic DNA.

Immunofluorescence Analysis. Mononuclear cells were stained using biotinylated mAb directed to Vβ2, Vβ3.1, Vβ5.1, Vβ5.2/5.3, Vβ6.7a, Vβ8.1/8.2, Vβ12, Vβ13.1, Vβ13.2, and Vβ17 (12–15). Streptavidin-PE (Fisher Biotech, Pittsburgh, PA) was used as a second-step reagent. Samples were also stained with FITC-conjugated mAbs directed to CD4 and CD8 (both from Becton Dickinson & Co., Mountain View, CA). Fluorescence intensity was analyzed with a cytofluorograph (Epics Profile; Coulter Corp., Hialeah, FL), and at least 10^4 cells were analyzed for each antibody combination.

The specificity of the 8F10 mAb was assigned based on its ability to stain T cell hybridomas transfected with a DNA construct containing Vβ3.1 as described (15), but not their untransfected parents or hybridoma cells expressing other human Vβ elements (data not shown).
shown). A similar perfect correlation was seen when a panel of T cell clones was typed for surface expression with the 8F10 mAb and for VB3.1 mRNA expression by PCR (11). Additionally, stimulation of peripheral blood cells with the 8F10 mAb resulted in a selective increase of VB3.1 mRNA expression (13, and Donahue, J. P., P. Marrack, J. Kappler, and B. L. Kotzin, unpublished observations).

Sequencing of TCRBV3S1 Genes. TCRBV3S1 (hereafter called VB3.1) sequences were obtained by amplifying cDNA derived from unstimulated cells using PCR with oligonucleotides specific for the VB3.1 leader (GTGAGAATTCCACCTGAAAGTAACC-TCCTCTGTC) and TCRBC (CGGGTGGGAACACCTTGTTC-GGATCCTC). Amplified DNA was cloned into pT7Z18R (Pharmacia Fine Chemicals, Piscataway, NJ) using EcoRI and BamHI sites, or into the pCR™ cloning vector as specified by the manufacturer (Invitrogen, San Diego, CA). The ligation product was transformed into competent Escherichia coli cells (Invitrogen), and colonies containing appropriate inserts were chosen randomly for sequencing. DNA sequencing was performed by the dideoxy method using Sequenase (United States Biochemical Corp., Cleveland, OH). At least five cDNA clones derived from each sample were sequenced.

Identification of Polymorphisms within the TCRB Gene Complex. Previously described VB gene polymorphisms were used primarily to genetically type family members and to determine the inheritance pattern in families. For example, in Families A and B, we utilized polymorphisms in simple sequence repeats (e.g., simple sequence length polymorphisms [SSLPs]) contained within the VB6 subfamily of gene segments as described (16). Sequences of the oligonucleotides used for PCR amplification were as described (16). PCR employed a thermal cycler (Cetus/Perkin Elmer; Emeryville, CA). In each reaction, 100 ng of genomic DNA was mixed with primers (0.3 μM final concentration), 1.0 U of Taq polymerase (Perkin Elmer), and 20 μM each of dNTPs in a total volume of 50 μl. Amplification was performed for 35–40 cycles, with 1 min of denaturation at 95°C, 1 min of annealing at 55°C, and 2 min of extension at 72°C. PCR products were separated on a 15% polyacrylamide gel, and visualized with ethidium bromide and UV irradiation.

In Family A, we also utilized PCR and polymorphisms within the TCRBC loci as described (17). Polymorphisms in two regions identified as CB21 and CB25 (17) could be identified by changes in BglII and KpnI restriction sites, respectively. Primers utilized for PCR were as described (17) and PCR conditions were identical to those described above. After precipitation of 20 μl of the PCR product, digestion was carried out with 2 U/μg DNA of enzyme at 37°C overnight. Undigested and digested aliquots were separated on a 2% agarose gel and visualized with ethidium bromide and UV irradiation.

In Family B, previously reported polymorphisms in the VB6.1 structural genes were also utilized (8). One of the mutations at amino acid position 62, eliminates a BsiE1 restriction enzyme site. Primers utilized were 5'-GTTCTCAACTTGTTGTCTCCCG (+ strand) and 5'-GGATACACGGCGTGACGTC (- strand). 25 μl of amplified product was digested with 2 U/μg DNA of BsiE1 (New England Biolabs Inc., Beverly, MA) overnight at 55°C. Undigested and digested products were separated on a 15% polyacrylamide gel and visualized with ethidium bromide staining.

In Family C, typing of maternal TCRB haplotypes utilized Southern analysis and a previously described RFLP described by a BgIII restriction site in the TCRBC2 gene segment (18). In addition, a VB3.1 polymorphism was utilized to determine inheritance of paternal VB gene complexes. To determine the presence of the paternal allele, cDNA from unstimulated PBMC was amplified using VB3.1 and TCRBC specific primers utilized above for sequencing. This fragment was then reamplified using internal oligonucleotides specific for VB3.1 (GTGAGAATTCCACCTGAAAGTAACC-CAGAGCTCG) and TCRBC (TTCTGATGGCTCAAACACG). Fragments were digested with AatII overnight before analysis on a gel.

Results

Variable Percentages of Vß3.1 + T Cells in Adult and Fetal Cord Blood. We examined the peripheral blood TCR repertoire in a group of unrelated normal adults to look for possible deletions of specific Vß subsets. These studies were accomplished first using a quantitative PCR technique (11) and subsequently using flow cytometry with mAb directed to T cells bearing Vß2, Vß3.1, Vß5.1, Vß5.2, Vß6.7, Vß8.1/8.2, Vß12, Vß13.1, Vß13.2, and Vß17. By PCR analysis and with a mAb directed to Vß3.1 (clone 8F10), we noted a broad range in the percentage of Vß3.1 + T cells in the normal population, and discovered several individuals with distinctly low levels (staining data are shown in Fig. 1 a). Unlike results with some of the other anti-Vß antibodies, low levels of Vß3.1 + cells, when present, occurred in both the CD4 + and CD8 + populations. 7 of 31 unrelated adults studied (or 23%) had low percentages of Vß3.1-bearing T cells, defined as <2.0% of CD4 + and CD8 + T cells. Most of these individuals were studied multiple times over several months and the percentage of Vß3.1 + T cells remained remarkably stable in all cases (data not shown).

We also examined the TCR repertoire of fetal cord blood samples (Fig. 1 b). 5 of 25 samples contained low percent-
ages of Vβ3.1+ T cells in both CD4+ and CD8+ subsets. The frequency of low values (∼20%) was similar to that observed for adults (Fig. 1a), suggesting that infection after birth does not account for most of the cases with the Vβ3.1low phenotype.

We carefully examined the density of Vβ3.1 expression per cell, as determined by mean channel of fluorescence intensity, in a large number of different individuals with both low and high percentages of Vβ3.1+ cells in blood. The density was identical regardless of the percentage of positive cells (data not shown).

Inheritance Pattern for the Vβ3.1low Phenotype and Contribution of TCR and MHC genes. We searched for families in which the mother and father expressed divergent levels of Vβ3.1+ T cells. Of the first 13 sets of parents studied, 7 had one spouse with a low level of Vβ3.1-expressing T cells. In these seven families, the frequency of children with low Vβ3.1 levels (<2% in both CD4+ and CD8+ subsets) was 12 of 30 compared with 0 of 23 in the other six families (p < 0.001 by Fisher’s exact test). These results strongly suggest that low levels of Vβ3.1+ T cells are determined genetically.

We analyzed three of the above families with the largest number of individuals for the contribution of VB or MHC genes to the Vβ3.1 phenotype. The analysis of Family A is shown in Fig. 2a. The mother expressed low levels of Vβ3.1+ T cells in both CD4+ and CD8+ subsets, whereas the father did not express this phenotype. Six children were analyzed, of which three expressed the Vβ3.1low phenotype. Other T cell subsets quantitated by monoclonal anti-Vβ antibodies were not strikingly different among family members (only data for Vβ8.1/8.2 are shown). As shown in Fig. 2a, the inheritance of MHC haplotypes in this family was independent of Vβ3.1+ T cell levels. It is interesting to note that expression of the low phenotype correlated with inheritance of one of the TCRB complexes from the father, who did not express the low phenotype (Fig. 2a). The statistical likelihood of this happening by chance is p = 0.05 (by Fisher’s exact test). Either of the mother’s TCRB complexes appeared to allow for the low phenotype in the children.

This analysis of Family B is shown in Fig. 2b. Again, the mother expressed the low phenotype that was expressed by four of the seven children. Vβ3.1 levels in the other three children were actually higher than those expressed by the father. 5 wk later, repeat studies of three children, including two with the low phenotype, showed identical Vβ3.1 percentages. As shown in Fig. 2b, MHC haplotypes inherited from the mother or father did not correlate with Vβ3.1+ T cell levels. However, similar to Family A, this family also showed a concordance of the Vβ3.1low phenotype with one of the paternal TCRB gene complexes (p = 0.03 by Fisher’s exact test). Again, either TCRB complex from the mother, who expressed the low phenotype, allowed for low levels in her offspring.

One additional family (Family C) with the largest number of available individuals was studied to determine the importance of TCRB or MHC genes (Fig. 2c). The mother (II-2) exhibited a Vβ3.1low phenotype as did one sister (II-3). Of the eight children examined, only one child (III-4) was Vβ3.1low. These phenotypes remained stable during the 12-mo period in which the family was studied. For example, percentages of peripheral blood CD4+ T cells expressing Vβ3.1 in samples taken several months apart from the father (II-1) were 4.9, 5.7, and 5.0%. In contrast, values for the mother (II-2) were 1.5, 1.7, and 1.3%, and values for child III-4 were 2.0 and 1.9%.

Inheritance of MHC haplotypes in Family C is shown in Fig. 2c. In several individuals, the same MHC type was associated with disparate Vβ3.1 expression, indicating that a particular MHC type was not sufficient to explain the Vβ3.1low phenotype.

During the sequencing of Vβ3.1 genes expressed in Family C, a new Vβ3.1 allele (designated TCRBV3S1*4[N]) was identified in the father, which differs from previously published sequences (designated TCRBV3S1*1 [19, 20]) by a single nonproductive change (A → G) in the third position of the threonine codon at position 84. Although it does not alter the amino acid sequence, this change does generate a unique AatII restriction site, which was useful for identifying the paternal VB complex inherited among family members. Using this AatII polymorphism and a previously characterized genomic TCRBC2 BglII polymorphism (18), we identified two individuals who inherited the same VB complexes but who had different Vβ3.1 phenotypes (III-4 and III-8).

Vβ3.1 Gene Sequences Expressed by Individuals with the Low Phenotype. Expressed Vβ3.1 gene sequences were analyzed in a number of different individuals to determine if coding region polymorphisms contributed to low levels of Vβ3.1+ T cells. Table 1 shows that structural Vβ3.1 gene alterations do not account for differences in Vβ3.1 expression. Table 1 also shows that a relatively large fraction of the cDNA clones from Vβ3.1low individuals represent nonfunctional rearrangements (p < 0.005, compared with sequences from individuals not expressing the low phenotype). It is important to note that the coding region of the nonfunctional clones also matched the TCRBV3S1*1 sequence.

Discussion

We initiated these studies of human TCR repertoire formation by analyzing lymphocytes from normal individuals for Vβ expression using both quantitative PCR and different anti-Vβ mAb. Although considerable variation was found for many Vβs, only a few, such as Vβ3, were found to be expressed at distinctly low levels in both CD4+ and CD8+ subsets in some individuals. Our results show that >20% of normal adults express the Vβ3.1low phenotype. A similar frequency in fetal cord blood samples and a strong inheritance pattern in family studies strongly support the hypothesis that the Vβ3.1low phenotype is genetically determined in most cases.

In two families studied, the Vβ3.1low phenotype correlated perfectly with inheritance of one TCRB complex from the parent who was not low. Either allele from the Vβ3.1low parent was sufficient for expression of the low phenotype.
The most straightforward explanation for these findings is that the low phenotype is determined in a recessive fashion in these families. If this is the major mechanism for the low phenotype in the human population, then nearly 20% of individuals must be homozygous for the \( V\beta^3.1 \text{low} \) allele, and as predicted by the Hardy-Weinberg equilibrium for allele frequency, nearly 50% of individuals would then be expected to carry such an allele in a heterozygous fashion.

The distribution of \( V\beta^3.1 \) expression in the population overall suggested that there may be three phenotypes corresponding to the three possible genotypes: \( V\beta^3.1 \text{low/low} \), \( V\beta^3.1 \text{low/high} \), and \( V\beta^3.1 \text{high/high} \). However, although being...
Table 1. V$\beta$3.1 Sequences Expressed in Peripheral Blood by Different Individuals

| Study group | No. cDNA clones sequenced | No. wild type | No. in-frame |
|-------------|---------------------------|---------------|--------------|
| General population |                         |               |              |
| V$\beta$3.1
$^{\text{low}}$ ($n = 3$)$^*$ | 16 | 16$^+$ | 15 |
| V$\beta$3.1
$^{\text{low}}$ ($n = 4$)$^*$ | 22 | 22 | 13 |
| Family A |                         |               |              |
| II-2 (V$\beta$3.1$^{\text{high}}$) | 5 | 5 | 4 |
| III-6 (V$\beta$3.1$^{\text{low}}$) | 5 | 5 | 3 |
| Family B |                         |               |              |
| II-1 (V$\beta$3.1$^{\text{high}}$) | 5 | 5 | 5 |
| II-2 (V$\beta$3.1$^{\text{low}}$) | 10 | 10 | 9 |
| II-7 (V$\beta$3.1$^{\text{low}}$) | 7 | 7$^+$ | 4 |
| Family C |                         |               |              |
| II-1 (V$\beta$3.1$^{\text{low}}$) | 5 | 5$^+$ | 5 |
| II-2 (V$\beta$3.1$^{\text{low}}$) | 5 | 5 | 3 |

$^*$ A minimum of five sequences were analyzed per individual.
$^+$ Nucleotide sequences matched the published TCRBV31$^{\text{low}}$ allele sequence (19, 20).
$^+$ Two sequences carried different single nucleotide changes that were not present in the mother or father, and were felt to be errors introduced during PCR amplification.
$^+$ These sequences contained a single nucleotide difference (see Fig. 2 c), which does not result in a change of the amino acid sequence.

homozygous for the V$\beta$3.1$^{\text{low}}$ allele appeared to determine low expression in the families studied. Heterozygous expression did not necessarily determine a middle range of expression. This is best illustrated in Family B, in which three heterozygous children were identified. Their levels of V$\beta$3.1$^+$ cells, however, were clearly at the high end of the total distribution.

The polymorphism(s) in the V$\beta$3.1 allele that causes low expression is currently unknown. However, it appears to be unique compared with other previously described VB gene polymorphisms (7–10). Thus, the V$\beta$3.1$^{\text{low}}$ allele is not abnormal in its coding region. Functional V$\beta$3.1 mRNA and normal V$\beta$3.1 gene sequences were expressed in all individuals studied with the low phenotype. In addition, a relatively large number of nonfunctional V$\beta$3.1 rearrangements were found in individuals with the low phenotype, and in every case the V$\beta$3.1 coding sequence was normal. Finally, it should also be pointed out that despite the lower percentage of V$\beta$3.1$^+$ cells in certain individuals, the amount of TCR protein expressed per cell is normal. We conclude that the V$\beta$3.1 allele in these families is defective in its ability to rearrange normally. Although the increased frequency of nonfunctional rearrangements might be expected based on the low percentage of V$\beta$3.1$^+$ T cells alone, this finding may also be secondary to the V$\beta$3.1 allele defect itself.

Using techniques that quantitate V$\beta$ mRNA, variability in V$\beta$3 expression has been observed by others (9, 21, 22). Vandekerckhove et al. (22) reported that after implantation of fetal liver and thymus from V$\beta$3-discordant individuals into immunodeficient (SCID) mice, the V$\beta$3.1$^{\text{low}}$ phenotype was determined before negative selection in the thymus and was dependent on the fetal liver rather than thymus donor. These results are consistent with a mechanism whereby the V$\beta$3.1$^{\text{low}}$ phenotype is determined by the TCRB gene complex.

In one individual of a separate family, we found that the V$\beta$3.1$^{\text{low}}$ phenotype was not determined solely by either VB or MHC genes. Although none of the children with disparate phenotypes was identical for both VB and MHC genotypes, it is unlikely that a combination of TCR and MHC genes is required for the low phenotype, since the amino acid coding region sequence of the V$\beta$3.1 genes in the family members did not differ. These results suggest a different mechanism for low V$\beta$3.1$^+$ T cells in this individual, including the possible existence of a V$\beta$3.1-specific superantigen. However, it should be emphasized that the inheritance pattern in this family was unexpected in that only one of eight children expressed the low phenotype. Because our conclusions rest on this one individual, we cannot exclude an infectious exposure after birth that led to the low phenotype. Furthermore, since only one other sibling (III-8) was matched for inheritance of the paternal TCRB haplotype, we also cannot exclude a genetic recombination event that would explain higher levels in this child.
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