T CELL RECEPTOR Vβ GENE USAGE IN THAI CHILDREN WITH DENGUE VIRUS INFECTION

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Abstract. T lymphocyte activation during dengue is thought to contribute to the pathogenesis of dengue hemorrhagic fever (DHF). We examined the T cell receptor Vβ gene usage by a reverse transcriptase-polymerase chain reaction assay during infection and after recovery in 13 children with DHF and 13 children with dengue fever (DF). There was no deletion of specific Vβ gene families. We detected significant expansions in usage of single Vβ families in six subjects with DHF and three subjects with DF over the course of infection, but these did not show an association with clinical diagnosis, viral serotype, or HLA alleles. Differences in Vβ gene usage between subjects with DHF and subjects with DF were of borderline significance. These data suggest that the differences in T cell activation in DHF and DF are quantitative rather than qualitative and that T cells are activated by conventional antigen(s) and not a viral superantigen.

INTRODUCTION

Infection with dengue virus (DV), a mosquito-borne flavivirus, is a major health problem for many tropical and subtropical areas of the world, where as many as 100 million DV infections occur yearly. This virus exists as four distinct serotypes; infection with one DV serotype induces long-lived immunity against reinfection with that serotype. However, epidemiologic and laboratory studies indicate that reinfection with a heterologous DV serotype is more likely to result in the potentially life-threatening form of disease termed dengue hemorrhagic fever (DHF), characterized by increased capillary permeability. Additionally, a high level of CD69 expression has been observed on circulating peripheral blood mononuclear cells (PBMCs) of children with DHF.

The biological basis for the increased T cell activation in DHF has not been determined. Expansion of T cell subsets bearing T cell receptors (TCRs) using particular Vβ gene families has been associated with severe disease in other infectious or autoimmune disorders. Specific examples include toxic shock syndrome, leprosy, infection with human immunodeficiency virus (HIV), Crohn’s disease, and reactive arthritis. In vitro analysis of TCR usage by a panel of DV-specific CD4+ cytotoxic T lymphocyte clones generated from one individual after a primary dengue-4 virus infection demonstrated that eight of 19 clones expressed Vβ17. A similar expansion of Vβ 17-expressing cells was observed following bulk culture stimulation of this donor’s PBMCs with D4 antigen but not following stimulation with an anti-CD3 antibody. These data provided preliminary evidence that DV-infection might result in expansion of T cells bearing a particular Vβ chain.

We therefore sought to determine whether an expansion of T lymphocytes expressing a particular Vβ gene occurs in vivo upon DV infection and whether preferential Vβ region usage correlates with disease severity. We analyzed the Vβ gene usage in PBMCs obtained during and after the acute infection from 13 subjects with DHF and 13 with DF using a semi-quantitative polymerase chain reaction (PCR) method. We confirmed that this method yields highly reproducible results. We detected expansions of T cells expressing particular Vβ genes during acute DV infection in some subjects. However, we did not find consistent expansions of specific Vβ gene families among the group of subjects as a whole or significant differences in Vβ usage between patients with DHF and those with DF.

MATERIALS AND METHODS

Clinical study design and sample collection. Thai children between the ages of six months and 14 years were enrolled in the Dengue Hemorrhagic Fever Project, as previously reported. Written informed consent was obtained from the subjects’ parents or guardians. The study protocol was approved by Institutional Review Boards established by the Ministry of Public Health, Thailand, the Surgeon General’s Office of the Department of the Army, and the University of Massachusetts. We selected at random 13 children with DHF and 13 children with DF from among the study population enrolled between April 1994 and November 1995. Dengue virus infections were confirmed by serologic testing and virus isolation as described. Clinical diagnoses of DHF and DF were assigned according to World Health Organization criteria.

Blood samples were obtained daily during illness and at a follow-up visit six months after the acute DV infection. The PBMCs were isolated by density gradient centrifugation using Histopaque (Sigma, St. Louis, MO). They were washed and counted, and 2 x 10⁶ cells were then placed into an Eppendorf tube (Brinkman Instruments, Westbury, NY) and washed twice with phosphate-buffered saline. Finally, the cells were pelleted and 1.0 ml of solution D (4 M guanidinium isothiocyanate, 25 mM sodium citrate pH 7.0, 2%
tion. We used 26 cycles of a PCR at 95

The RNA was treated with DNaseI for 2

Analysis of TCR Vβ gene usage by PCR. Analysis of TCR Vβ gene usage was performed as previously de-

Statistical analysis. Pearson correlation was used to compare the expression of all TCR Vβ gene families in serial analyses of the same PBMC samples. For comparison of TCR Vβ expression in different PBMC samples, percentage values were first transformed using the logit function, f(x) = log(x/(1 – x)), using 2% as a minimum value for expression of each TCR Vβ gene family (to avoid skewing of results because of small absolute changes in expression of low abundance TCR Vβ transcripts). Changes in expression of specific TCR Vβ gene families between two time points for an individual subject were considered significant if the differences in transformed values fell outside the mean ± 3 SD of differences for all TCR Vβ gene families between those two time points. We compared the expression of each TCR Vβ gene family between different time points for the study population as a whole using a paired t-test. We compared the expression of each TCR Vβ gene family at each time point and changes in expression of each TCR Vβ gene family between time points in subjects with DHF with the corresponding values in subjects with DF using a t-test. We set P < 0.01 as the criterion for statistical significance, to adjust in part for the number of comparisons.

RESULTS

Characteristics of the study population. Table 1 shows the clinical, serologic, and virologic information on the study subjects. Five of 13 DF patients had primary DV infections, while all of the DHF patients were experiencing secondary infections. All four serotypes of DV were represented in both the DF and DHF groups. Among the subjects studied with DHF, there were six with DHF grade 1, six with DHF grade 2, and one with DHF grade 3. Data on HLA class I alleles were available for 23 subjects and data on HLA class II alleles were available for 24 subjects (Table 1). The mean age of the subjects studied was 8.5 years (95% confidence interval [CI] = 7.3–9.7), and was not significantly different between those children with DHF (mean = 7.9 years) and those with DF (mean = 9.1 years).

Reproducibility of the PCR assay. We first analyzed the reproducibility of the PCR assay for measurement of TCR Vβ gene usage by repeated analysis of cDNA prepared from seven blood samples. The mean correlation coefficient for the comparison of the results from repeated PCR amplification of the same specimens was 0.97 (95% CI = 0.96–0.99).

Changes in TCR Vβ gene usage associated with acute dengue virus infection. To determine whether acute dengue virus infection causes significant shifts in the TCR Vβ gene usage in PBMCs, we examined the TCR Vβ gene usage of PBMCs obtained from each subject on study day 2, fever day +1 (one day after defervescence), and study day 180. For one patient in each group, only the study day 2 specimen was available for analysis. The mean interval between study day 2 and fever day +1 for the study population as a whole was 2.2 days (95% CI = 1.9–2.5), and was not significantly different between those children with DHF (mean = 2.2 days) and those with DF (mean = 2.3 days).
We did not note deletion of cells using any particular TCR Vβ gene family following acute dengue virus infection in the study population. However, we detected TCR transcripts using Vβ20 in only one of the 26 subjects studied, including those with primary dengue virus infection. We were able to amplify TCR Vβ20 transcripts from PBMCs obtained from healthy donors from the United States and from six of the seven control Thai subjects using the same experimental procedures.

The results of our statistical analysis of the changes in TCR Vβ gene usage for the study population as a whole are summarized in Table 2. All analyses for which a paired t-test showed $P < 0.05$ are shown; however, using our selected cutoff of $P < 0.01$, there were few TCR Vβ gene families that showed significant changes in usage. From study day 2 to fever day +1 there was a statistically significant decrease in usage of Vβ5.1. From fever day +1 to study day 180, there were statistically significant increases in usage of Vβ2, Vβ5.1, and Vβ6 and statistically significant decreases in usage of Vβ1 and Vβ13.1. From study day 2 to study day 180, there was a statistically significant decrease in usage of Vβ18.

Statistically significant changes in usage of one or more TCR Vβ gene families during the acute infection were apparent in 12 subjects. The most common such finding was the expansion in use of a particular TCR Vβ gene between study day 2 and fever day +1. We found expansion in use of a single TCR Vβ gene family in six of 12 subjects with DHF and three of 12 subjects with DF ($P$ not significant). These expansions generally did not persist in the PBMCs obtained six months after the acute infection. For example, Figure 1A shows one subject (C94-050) with DHF and secondary dengue-4 virus infection who demonstrated increased use of Vβ21 on fever day +1. Figure 1B shows one subject (C94-076) with DHF and three of 12 subjects with DF ($P$ not significant).

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fication. The earliest time point studied (study day 2) corresponded to a mean fever day value of −1.2, or slightly more than one day before defervescence. This was the closest approximation available to the viremic phase of infection,18 and 24 of the 26 subjects studied had positive plasma virus culture at that time. The middle time point (fever day +1, one day after defervescence) corresponds closest to the period of plasma leakage in DHF.17 The latest time point (study day 180) is the closest approximation to a baseline for each subject since we did not have access to PBMCs from before dengue virus infection in these subjects. Although the interval between the first two specimens was relatively short, we have evidence that significant immunologic changes occur during this period. In a separate group of study subjects, the expression of CD69 on CD8+ T cells of children with acute dengue virus infection was higher on study day 2 than on fever day +1, particularly in children with DHF.9 On the other hand, the number of atypical lymphocytes in the peripheral blood of children with acute dengue virus infection was markedly higher on fever day +1.9,17 Plasma levels of sCD8 and sIL-2R, markers of T cell activation, were elevated on study day 2 but increased further through fever day +1.7

We used a semi-quantitative PCR assay to measure the usage of 26 TCR Vβ gene families in the PBMC samples. Although some of the data were obtained from repeat analysis of the same samples and interpolation of missing data, we believe that our finding of a high interassay correlation justifies this approach.

Our analysis of the data tested several specific hypotheses. First, we looked for increases in the usage of some TCR Vβ gene families in individual subjects during the acute infection, using the study day 180 specimen for comparison, as an indication of expansion of particular T cell populations. As shown in Figure 1, expansions in certain TCR Vβ subsets during the course of DV infection were observed in some subjects. This finding is in agreement with our in vitro data demonstrating preferential expansion of Vβ17-bearing T cells in the PBMCs of a dengue-immune donor upon stimulation with non-infectious DV antigen.18 The TCR Vβ expansions were most often noted one day following defervescence (fever day +1), which supports the hypothesis that some of the atypical lymphocytes in the peripheral blood are proliferating DV-specific T cells.

The specific TCR Vβ gene families showing expansion during acute DV infection differed among the study subjects. We did not identify any consistent association between the TCR Vβ genes showing expansion and the clinical, virologic, or genetic profile of the subjects. We found statistically significant changes in expression of TCR Vβ1, Vβ2, Vβ5.1, Vβ6, Vβ13.1, and Vβ18 in the study population as a whole. Although we used a relatively conservative criterion for statistical significance of P < 0.01, these findings may still reflect type I error due to the large number of statistical comparisons performed. Therefore, we consider these to be preliminary observations that require confirmation in other study populations.

We did not detect deletions of any specific TCR Vβ gene families related to acute DV infection, regardless of the clinical severity of disease. This finding does not support the

### Table 2

Summary of probability values for comparisons of interval changes in T cell receptor (TCR) gene usage in the study population.

<table>
<thead>
<tr>
<th>Vβ</th>
<th>All subjects*</th>
<th>DFH versus DP*</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Study day 2 to fever day +1</td>
<td>Fever day +1 to study day 180</td>
</tr>
<tr>
<td>1</td>
<td>0.022</td>
<td>(↑) 0.001</td>
</tr>
<tr>
<td>2</td>
<td>0.016</td>
<td>(↑) &lt;0.001</td>
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<tr>
<td>3</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>4</td>
<td>–</td>
<td>0.038</td>
</tr>
<tr>
<td>5.1</td>
<td>(↑) 0.005</td>
<td>(↑) 0.002</td>
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<tr>
<td>5.2</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>6</td>
<td>–</td>
<td>(↑) 0.001</td>
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<td>7</td>
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<td>12</td>
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<tr>
<td>13.1</td>
<td>–</td>
<td>(↓) &lt;0.001</td>
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<tr>
<td>13.2</td>
<td>–</td>
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<tr>
<td>14</td>
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<td>18</td>
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<td>(↓) 0.004</td>
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<td>24</td>
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</table>

*All p values are 2-tailed. Dashes represent comparisons where p values are > 0.05. The direction of change in TCR gene usage is indicated only for comparisons showing p < 0.01.

Changes in logit-transformed values for TCR gene usage were compared between subjects with dengue hemorrhagic fever (DHF) and those with dengue fever (DF) by t-test.

The direction of change in TCR gene usage is indicated only for comparisons showing p < 0.01.
Figure 1. T cell receptor (TCR) \( V/\beta \) gene usage during and after acute dengue virus infection in individual subjects. Fever day +1 is one day after defervescence. A, subject C94-050 (dengue hemorrhagic fever [DHF]) had a significant expansion of \( V/\beta21 \) gene usage on fever day +1. B, subject C94-076 (dengue fever [DF]) had a significant expansion of \( V/\beta24 \) gene usage on fever day +1.
Figure 2. T cell receptor (TCR) Vβ gene usage during and after acute dengue virus infection in subjects with dengue hemorrhagic fever (DHF) and dengue fever (DF). Polymerase chain reaction of TCR Vβ genes was performed on cDNA from 13 subjects with DHF and 13 subjects with DF. Values plotted are the mean ± SD percent usage of 26 TCR Vβ gene families at the three time points examined. Fever day +1 is one day after defervescence.

suggestion that a DV-encoded superantigen might be responsible for the increased T cell activation observed during DV infection, particularly in patients with DHF.7,8 Superantigens can induce marked activation followed by deletion of T cells expressing a particular TCR Vβ gene.24–26 Our failure to detect such an effect in this study population is subject to several limitations. Only one of the subjects studied had grade 3 DHF, and none of the cases of DHF were associated with a primary DV infection; massive activation of T cells not specific for classical DV antigens caused by interaction with a superantigen might be more likely to play a role in those clinical situations. Also, we detected TCR Vβ20 transcripts in only one of the 26 subjects. This may reflect genetic differences between the Thai and United States populations, but we cannot exclude the possibility that T cells using the Vβ20 gene were deleted earlier in infection or by a prior DV infection. Analysis of TCR Vβ gene usage in the PBMCs of DV-naive Thai children would distinguish between these possibilities.

Lastly, we compared the TCR Vβ gene usage in subjects with severe illness (DHF) with that in subjects with milder illness (DF). Although we found a significant difference in the usage of Vβ22 during the acute infection, the magnitude of this difference was small. Similarly, the statistically significant difference in the change in TCR Vβ1 gene usage from study day 2 to study day 180 between the two groups of subjects was small in magnitude. These differences may well reflect type I error. In general, our findings suggest that there are not qualitative differences in the T cell repertoire responding to acute DV infection between subjects with and
without plasma leakage, and that the increased T cell activation in DHF reflects quantitative differences in the T cell response.

Other groups have similarly failed to find differences between the TCR Vβ repertoire in the PBMCs of HIV-infected subjects and that of control subjects. However, skewing of the TCR Vβ repertoire has been observed in cells obtained from either the lungs14 or lymph nodes27 of HIV-infected patients. Others have also noted that alterations of TCR Vβ gene usage occur much less frequently in PBMCs compared to the organs or tissues in which disease pathology is directly observed.28,29 Dengue virus is believed to infect primarily monocytes and macrophages in vivo,30–32 and a specific organ or tissue that preferentially supports DV replication has not been identified. However it is possible that during acute DV infection, notable biases in Vβ expression may occur in tissues not examined in this study, such as lymph nodes or liver.

The pathogenesis of DHF is still poorly understood, but data suggest that increased levels of T cell activation play an important role. The cause of this heightened T cell activation is an active topic of investigation. To our knowledge, this study provides the first detailed analysis of TCR Vβ gene usage in the PBMCs of patients with acute DV infections. Our results provide additional insight into the nature and timing of T cell responses that should be helpful in directing future studies into the pathophysiologic events in acute DV infection.

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REFERENCES


