Patterns of CD8⁺ Immunodominance May Influence the Ability of *Mamu-B*⁺⁺08-Positive Macaques To Naturally Control Simian Immunodeficiency Virus SIVmac239 Replication

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Certain major histocompatibility complex (MHC) class I alleles are strongly associated with control of human immunodeficiency virus and simian immunodeficiency virus (SIV). CD8⁺ T cells specific for epitopes restricted by these molecules may be particularly effective. Understanding how CD8⁺ T cells contribute to control of viral replication should yield important insights for vaccine design. We have recently identified an Indian rhesus macaque MHC class I allele, *Mamu-B*⁺⁺08, associated with elite control and low plasma viremia after infection with the pathogenic isolate SIVmac239. Here, we infected four *Mamu-B*⁺⁺08-positive macaques with SIVmac239 to investigate why some of these macaques control viral replication. Three of the four macaques controlled SIVmac239 replication with plasma virus concentrations below 20,000 viral RNA copies/ml at 20 weeks postinfection; two of four macaques were elite controllers (ECs). Interestingly, two of the four macaques preserved their CD4⁺ memory T lymphocytes during peak viremia, and all four recovered their CD4⁺ memory T lymphocytes in the chronic phase of infection. *Mamu-B*⁺⁺08-restricted CD8⁺ T-cell responses dominated the acute phase and accounted for 23.3% to 59.6% of the total SIV-specific immune responses. Additionally, the ECs mounted strong and broad CD8⁺ T-cell responses against several epitopes in Vif and Nef. *Mamu-B*⁺⁺08-specific CD8⁺ T cells accounted for the majority of mutations in the virus at 18 weeks postinfection. Interestingly, patterns of viral variation in Nef differed between the ECs and the other two macaques. Natural containment of AIDS virus replication in *Mamu-B*⁺⁺08-positive macaques may, therefore, be related to a combination of immunodominance and viral escape from CD8⁺ T-cell responses.

Understanding the immunological and genetic basis of the natural control of AIDS virus replication should assist in human immunodeficiency virus (HIV) vaccine design. Of particular interest are human “elite controllers” (ECs), rare individuals who spontaneously control HIV viremia to extremely low levels (17). Similarly, a limited number of macaques spontaneously control simian immunodeficiency virus (SIV) replication and become ECs (49, 76).

Several lines of evidence suggest that CD8⁺ T cells play a key role in immune control of immunodeficiency virus replication. The transient in vivo depletion of circulating CD8⁺ lymphocytes in SIV-infected macaques, including EC macaques, results in dramatic increases in plasma viremia (23, 32, 52, 69). CD8⁺ T-cell responses also exert selective pressure on replicating viruses, resulting in the emergence of variants that escape immune detection in both HIV (5, 12, 20, 26, 36, 64, 65) and SIV (3, 7, 19, 35, 51, 59, 60) infection. In addition, it is well established that the expression of specific major histocompatibility complex (MHC) class I alleles is associated with reduced plasma viremia and/or slower disease progression in humans (14, 15, 29, 34, 55) and macaques (49, 57, 61, 63, 76, 77). In particular, human long-term nonprogressor and EC cohorts are enriched for HLA-B27 and HLA-B57. Numerous studies have implicated these molecules in the presentation of epitopes that elicit effective HIV-specific CD8⁺ T-lymphocyte responses (6, 20, 26, 36, 38, 54, 55, 70). This assertion is further supported by studies describing associations between viral escape from the immunodominant HLA-B27-restricted Gag263-272KK10 response and disease progression (8, 20, 26, 36).

Unfortunately, there are many difficulties inherent to studying HIV-infected humans. Viral control appears to be mediated soon after the resolution of acute-phase viremia with the appearance of CD8⁺ T-cell responses in HIV-infected individuals (11, 41), yet HIV is infrequently diagnosed during primary infection (42, 73). Hence, the study of immune responses initially involved in controlling HIV replication is extremely difficult. An additional complication arises due to the diversity of HIV isolates with which individuals might be infected (24, 27). Therefore, complete immunological monitoring would necessitate the sequencing of acute-phase virus and the synthesis of custom peptide sets matched to the infecting virus for every patient.

AIDS research with nonhuman primates provides an animal...
model to complement human studies. In particular, SIV-infected ECs also provide examples of successful immune containment of pathogenic immunodeficiency virus replication. However, unlike in human studies, researchers working with macaques have direct control over key variables such as virus strain, host genotype, and route of infection. Inoculum variability is eliminated because macaques can be infected with a clonal viral stock, e.g., SIVmac239, enabling complete and accurate tracking of early immune responses by use of corresponding peptides in ex vivo immunological assays. Moreover, the timing of immune responses after infection, the associated viral sequence evolution, and plasma virus concentrations may be closely monitored. Hence, the immunology and pathogenesis of acute infection can be more readily studied in macaques than in humans and may aid in our understanding of the correlates of immune protection.

Previously, we assembled a cohort of 192 Indian rhesus macaques, all infected with SIVmac239 (49, 76). Fourteen of these animals were considered ECs and controlled virus replication to fewer than 1,000 viral RNA (vRNA) copies/ml in the chronic phase of SIV infection. Nine of the 14 ECs expressed Mamu-B*17 (76). In addition to Mamu-B*17, we recently discovered that Mamu-B*08 is enriched in EC cohorts and associated with reduced chronic-phase plasma virus concentrations (49). Over 50% of Mamu-B*08-positive macaques become ECs. Interestingly, a preliminary binding motif for Mamu-B*08 appears comparable to that of HLA-B*27, an allele associated with elite control in humans (48). The similarity between Mamu-B*08 and HLA-B*27 makes SIVmac239-infected, Mamu-B*08-positive macaques an ideal system for modeling human ECs.

While not all Mamu-B*08- or Mamu-B*17-positive macaques control viral replication (49, 75, 76), we and others have previously shown that control of SIV replication is not due to genes linked to Mamu-B*17 (75) or to polymorphisms in several host genes, including CCR5, CXCR6, GPR15, RANTES, interleukin-10 (IL-10), APOBEC3G, tumor necrosis factor alpha, and TSG101 (72). Rather, CD8+ T cells appear to play a critical role in the natural containment of SIV replication (23, 61).

Here, we investigated early immune responses and viral evolution in four Mamu-B*08-positive Indian rhesus macaques during primary SIVmac239 infection. From previous studies (49), we estimated that two of the four macaques would control viral replication. We hypothesized that this control would be a function of the Mamu-B*08-specific CD8+ T-cell responses and viral escape. We found that containment of SIV replication began before 10 weeks postinfection in three of the four Mamu-B*08-positive macaques. During peak viremia, the CD4+ memory T-cell subset decreased in only two of the four Macaca mulatta (367). In these cases, freshly isolated or cryopreserved PBMC were depleted of CD8+ cells by using a CD8 microbead kit for nonhuman primates (Miltenyi, Auburn, CA) along with LS columns (Miltenyi) according to the manufacturer’s protocols. Labeled cells were removed by magnetic separation, and the remaining cells were used as described above in IFN-γ ELISPOT assays. Fluorescence-activated cell sorting analysis using CD3 FITC (clone SP34-2; BD Biosciences) and CD4 allophycocyanin (clone SK3; BD Biosciences) cell surface markers confirmed that the CD8+ cell depletions removed >99% of the CD8+ lymphocytes. Wells were imaged and counted with AIDS EliSpot reader version 3.40 or 4.0 (AID, Strassberg, Germany) and analyzed as previously described (46, 47). A response was considered positive if the mean number of spot-forming cells (SFC)
from the duplicate sample wells exceeded the background level (mean of wells without peptide stimulation) plus 2 standard deviations. Background levels were subtracted from each well, and assay results are shown as numbers of SFC per 1 x 10^6 PBMC. Responses of <50 SFC per 1 x 10^6 PBMC were not considered positive.

MHC class I tetramer and surface staining. Ex vivo MHC class I tetramer stains were performed on freshly isolated or cryopreserved PBMC as previously described (46). MHC class I tetramers were constructed with minor modifications as previously described (31, 46). Cryopreserved PBMC were thawed at 37°C and washed twice in R10 (RPMI 1640 medium [HyClone, Logan, UT] supplemented with 10% fetal calf serum [HyClone], 2 mM l-glutamine [HyClone], and 1x antibiotic-antimycotic solution [HyClone]) before staining.

Briefly, ~5 x 10^6 cells were stained with 5 μl of 0.1 mg/ml tetramer stocks, 3 μl of CD3 FITC (clone SP34-2; BD Biosciences), and 5 μl of CD95 PerCP (clone SK1; BD Biosciences) in ~100 μl of R10. After the cells were fixed in 1% paraformaldehyde (Electron Microscopy Sciences, Hatfield, PA), approximately 0.5 x 10^6 to 1 x 10^6 lymphocyte-gated events were acquired on a FACS Calibur (BD Biosciences) and analyzed using FlowJo 8.4.5 (TreeStar, Inc.). Percentages represent the numbers of lymphocyte-gated events that were CD3^+ CD95^+ and MHC class I tetramer positive. The threshold of detection in these assays was 0.02% CD3^+ CD95^+ MHC class I tetramer-gated lymphocytes.

ICS assay. IL-2 and IFN-γ intracellular cytokine staining (ICS) assays were performed on freshly isolated PBMC as previously described (23, 71). Briefly, each test contained ~5 x 10^5 PBMC. Peptide pools were used at a concentration of 10 μM. SIV peptide pools each contained 10 15-mer peptides overlapping by 11 amino acids. Fifteen-mer peptides were provided by the NIH AIDS Research and Reference Reagent Program (Germantown, MD). Approximately 0.5 x 10^6 lymphocyte-gated events were acquired on a FACS Calibur (BD Biosciences) and analyzed using FlowJo 8.4.5 (TreeStar, Inc.). All values were normalized by subtracting the background staining level (negative control of PBMC in media without stimulation).

Sequencing of plasma vRNA. Viral sequencing was performed based on methods previously described (23, 60). Briefly, vRNA was extracted from plasma by using a Qiagen MinElute kit (Valencia, CA). We used a Qiagen One Step RT-PCR kit to amplify overlapping regions between approximately 300 and 800 nucleotides in length that spanned the entire SIVmac239 genome. The RT-PCR conditions for all amplicons were as follows: 50°C for 30 min; 95°C for 15 min; 45 cycles of 94°C for 30 s, 53°C for 1 min, and 72°C for 15 s; and 68°C for 20 min. Cycling ramp rates were 2°C per second. The amplified cDNA was purified using a Qiagen PCR purification kit. Plasmids containing cloned sequences were purified using a QIAprep Spin Miniprep kit (Qiagen). Both strands of each amplicon were sequenced on a 3730 DNA analyzer (Applied Biosystems, Foster City, CA). Sequences were assembled using Aligner version 1.6.3 (CodonCode, Dedham, MA). DNA sequences were conceptually translated and aligned to the wild-type SIVmac239 sequence with the MacVector 9.0 trial version (Accelrys, Burlington, MA).

Statistical analysis. Statistical analyses consisted of two sample t tests for CD4+ T-cell (CD3^+ CD4^+ gaged lymphocytes) and total CD4+ T-cell (CD3^+ CD4^+ gaged lymphocytes) counts in Mamu-B*08-positive and Mamu-B*08-negative SIVmac239-infected macaques at seven different time points. The time points were categorized into seven intervals, as 0, 2, 6, 8, 10, 12, and 18 (or 28) weeks postinfection. Mean values for comparison between Mamu-B*08-positive (n = 4) and Mamu-B*08-negative (n = 4) macaques at each of the time points were obtained using the TTEST procedure in version 9.1 of SAS (Cary, NC).

The Z-score test was employed to compare the viral loads observed in each of the four Mamu-B*08-positive macaques against the means of the progressor (n = 175) and EC (n = 10) cohorts at 12 different time points. None of the SIVmac239-infected macaques in the progressor and EC cohorts expressed Mamu-B*08. The viral load data from the 187 animals within these two cohorts were obtained from a prior investigation (49). The time points were defined here as the numbers of weeks, between 0 and 20, after SIVmac239 infection, subdivided into 12 intervals. The goal was to ascertain whether the viral loads in Mamu-B*08-positive macaques were significantly different from the mean of the EC cohort or significantly lower than the mean of the progressor cohort.

Before performing inferential testing, we checked for key underlying assumptions of the t tests and Z-score tests (i.e., normality of residuals and homoscedasticity). The failure of the data to support these assumptions led us to transform the data for both CD4+ and viral loads via the natural logarithm to improve conformity to the assumptions. To circumvent the assumption of normality of residuals, we employed one-way nonparametric analysis using an exact Wilcoxon rank sum test with the continuity correction for the CD4+ T-cell counts only to further verify our results. Because statistical significance and direction of effects were the same across these analytic approaches, we based our interpretations on results from t tests only.

Nucleotide sequence accession numbers. The SIV genome sequences from 18 weeks postinfection were the following GenBank accession numbers: EU280803 (r91003-SIVmm239), EU280805 (r1027-SIVmm239), EU280804 (r0032-SIVmm239), and EU280806 (r0219-SIVmm239).

RESULTS

Mamu-B*08-positive, SIV-infected macaques control SIVmac239 viral replication and recover their CD4+ memory T lymphocytes after the acute phase of infection. We infected four Mamu-B*08-positive Indian rhesus macaques in an attempt to understand the dynamics of early infection and the way that the majority of macaques expressing this protective allele control viral replication. After challenging the macaques intravenously with the cloned viral isolate SIVmac239, we followed plasma viral concentrations, peripheral blood CD4+ counts (including total and memory subsets), antigen-specific responses (CD8+ and CD4+), and viral evolution during the first 20 weeks after SIV infection.

Three of the four macaques controlled their viral set points of SIVmac239 replication below 20,000 vRNA copies/ml and two of these had viral set points of ~1,000 vRNA copies/ml (Fig. 1). Only one macaque, animal r91003 (Mamu-B*01-positive and -B*08 positive), displayed plasma virus concentrations no different from those seen in 175 SIVmac239-infected macaques that progressed to AIDS (~500,000 vRNA copies/ml) and was thus termed a progressor. In contrast, animal r1027 (Mamu-A*01-positive and -B*08-positive) had a viral set point of <20,000 vRNA copies/ml and was termed a slow progressor/controller. Controller macaques maintain viral set points >10-fold lower than our previously studied SIV-infected macaque cohort (49). Macaques r00032 (Mamu-A*02-positive and -B*08 positive) and r0219 (Mamu-B*08-positive) controlled replication of this pathogenic SIVmac239 isolate to ~1,000 vRNA copies/ml and were classified as ECs. When comparing the plasma virus concentrations of the three Mamu-B*08-positive macaques that controlled SIV viremia with those of 175 macaques that progressed to AIDS, we found that the EC macaques r00032 and r0219 had viral set points significantly below those of 175 macaques that progressed to AIDS (P = 0.0066 and P = 0.0076, respectively). The difference between the viral set point of controller r1027 and those of the progressor macaques was near statistical significance (P = 0.0578). We also found that the plasma virus concentrations in r1027, r0032, and r0219 first began to differentiate themselves from those in the progressors between weeks 6 and 8 postinfection and trended toward significance as early as 4 weeks postinfection (data not shown). These data further suggest that the initial immunopathogenic events of immunodeficiency virus infection are crucial in controlling viral replication.

We also measured concentrations of circulating CD4+ T cells in our four SIV-infected Mamu-B*08-positive macaques. High levels of acute-phase immunodeficiency virus replication leads to the destruction of the CD4+ memory T-cell compartment (44, 53), thereby crippling the immune system during the critical stage of HIV/SIV infection. As expected, four Mamu-B*08-negative macaques (r95107, r98059, r0041, and r01035) that progressed to AIDS lost their CD4+ memory T cells (defined as CD3+ CD4+ CD95+ lymphocytes) during the
acute phase of infection and never recovered their CD4+ memory T lymphocytes (Fig. 2). Two Mamu-B*08-negative animals, r98059 and r00041, were infected with SIVmac239 at the same time as the four Mamu-B*08-positive macaques, while r95107 and r01035 were two Mamu-B*08-negative historical controls. By comparison, only two of the four Mamu-B*08-positive macaques experienced an acute-phase loss in their CD4+ memory T-cell levels during peak viremia. CD4+ memory T-cell levels rebounded in all four macaques (Fig. 2). For each time point between weeks 6 and 18 postinfection, the Mamu-B*08-positive macaques preserved a significantly higher number of CD4+ memory T cells than the four Mamu-B*08-negative macaques ($P < 0.014$). Interestingly, the Mamu-B*08-positive progressor macaque r91003 began to show a decline in its CD4+ memory T-cell count only at 18 weeks postinfection, perhaps due to high plasma virus concentrations. The total CD4+ T-cell counts (defined as CD3+ CD4+ lymphocytes) were not statistically different between the groups of Mamu-B*08-positive and B*08-negative macaques (data not shown).

**Dominance of Mamu-B*08-restricted CD8+ T-cell responses during the acute phase of SIVmac239 infection.** We initially examined the entire repertoire of SIV-specific immune responses by using ex vivo IFN-γ ELISPOT with 15-mer peptides that overlapped by 11 amino acids for the entire viral proteome (Fig. 3). These assays also included eight minimal optimal epitopes restricted by Mamu-B*08 (48) as well as known epitopes and other peptides that bind to Mamu-A*01 (2, 4) and Mamu-A*02 (46, 67, 71). In addition, we performed MHC class I tetramer staining on the immunodominant epitopes restricted by Mamu-A*01 (Tat28-35SL8 and Gag181-189CM9) and Mamu-A*02 (Gag151-159GY9 and Nef159-167YY9) during primary infection in macaques that expressed these alleles (data not shown). Since CD8+ T-cell responses usually peak at 3 weeks postinfection (66), we chose this time point to extensively investigate the possible role of SIV-specific CD8+ T cells in disease protection or progression during the acute phase.

CD8+ T-cell responses against the majority of the SIV proteins were detected in all four SIV-infected macaques. While these macaques all expressed Mamu-B*08, their immune responses should also be influenced by other known and unknown MHC class I alleles that they expressed. As expected, the progressor macaque r91003 (also Mamu-A*01 positive) made robust immune responses against the immunodominant Mamu-A*01-restricted CD8+ T-cell epitopes Gag181-189CM9 and Tat28-35SL8 (Fig. 3 and data not shown). These CD8+ T-cell responses peaked at 3 weeks postinfection, with $\sim 0.39\%$ (Gag181-189CM9) and $\sim 2.24\%$ (Tat28-35SL8) of CD3+ CD8+ lymphocytes staining with the relevant tetramer. Aside from these two Mamu-A*01-restricted responses, along with the Mamu-A*01-restricted response against Gag151-159GY9, the majority of the CD8+ T-cell response generated by r91003 targeted a single Mamu-B*08-restricted epitope in Vif and an epitope of unknown MHC class I restriction in Rev. Progressor macaque r91003 made no substantial CD8+ T-cell responses against Nef, although three known Mamu-B*08-restricted CD8+ T-cell epitopes are located in this protein (48).

The controller macaque r01027 (also expressing Mamu-A'01) mounted SIV-specific responses of a similar magnitude against the Mamu-A'01-restricted CD8+ T-cell epitopes Tat28-35SL8 and Gag181-189CM9 (Fig. 3 and data not shown).
MHC class I tetramer staining showed that the CD8+ T-cell response against Gag<sub>181-189</sub>CM9 peaked at 2 weeks postinfection (0.77%), while the Tat<sub>28-35</sub>SL8-specific CD8+ T cells reached 1.95% at 3 weeks postinfection. Along with a robust CD8+ T-cell response against Vif<sub>172-179</sub>RL8, several Mamu-B*08-restricted responses of lower magnitude (<200 SFC/10<sup>6</sup> PBMC) were detected in r01027 (Fig. 3). A sizeable CD8+ T-cell response of unknown MHC restriction to Rev was also identified in this macaque.

At 3 weeks postinfection, the two EC macaques r00032 and r02019 mounted total SIV-specific immune responses near or exceeding 10,000 SFC/10<sup>6</sup> PBMC (Fig. 4). These included several robust CD8+ T-cell responses to both the Vif and Nef epitopes restricted by Mamu-B*08 (Fig. 3 and 4). Additionally, animal r00032 (also Mamu-A*02 positive) made a response against the immunodominant Mamu-A*02-restricted CD8+ T-cell epitope Nef<sub>159-167</sub>YY9 (~2.4% at 2 and 3 weeks postinfection) (data not shown). Notably, we did not detect a response to Gag<sub>71-79</sub>GY9, which is frequently codominant with Nef<sub>159-167</sub>YY9 in Mamu-A*02-positive animals (46, 71). However, two responses, of >500 SFC/10<sup>6</sup> PBMC and of unknown MHC class I restriction, were directed against Gag<sub>241-291</sub>(G) and Gag<sub>361-411</sub>(J) peptide pools in EC r00032 (Fig. 3). While EC macaque r02019 did not express any other characterized MHC class I alleles that restrict known minimal optimal epitopes, a substantial response(s) of unknown restriction against Vpr was identified.

Using the week 3 ELISPOT pool response data along with responses to known minimal optimal epitopes and MHC class I tetramer stains, we defined the proportion of the total SIV-specific immune responses restricted by Mamu-B*08 and other MHC class I molecules (Fig. 4). Mamu-A*01-positive macaques mount two immunodominant, acute-phase CD8+ T-cell responses, Gag<sub>181-189</sub>CM9 and Tat<sub>28-35</sub>SL8 (3, 61), that typically account for ~50% of the total SIV-specific CD8+ responses (56). However, the normally dominant Mamu-A*01-restricted immune responses contributed only a slightly larger percentage of the total SIV-specific immune response than the Mamu-B*08-restricted immune responses in progressor macaque r91003 and controller r01027. Mamu-A*01-restricted CD8+ T cells accounted for 44% of the total SIV-specific immune response in r91003 and 26.5% of the total SIV-specific

**FIG. 2.** CD4+ memory T cells are maintained in Mamu-B*08-positive macaques infected with SIVmac239. (A) Absolute counts of CD4+ memory T cells were obtained from the four SIVmac239-infected, Mamu-B*08-positive macaques (blue lines) in addition to the four SIVmac239-infected, Mamu-B*08-negative macaques (red lines). Mamu-B*08-negative macaques r00059 and r00041 were infected along with the four Mamu-B*08-positive macaques. CD4+ memory T-cell counts from two additional Mamu-B*08-negative, -B*17-negative macaques (animals r95107 and r01035) were acquired from cryopreserved PBMC. Archived PBMC were not available at week 2 from r01035 or at week 18 from r01035 and r00041. In place of the week 18 time point, absolute counts of CD4+ memory T cells were acquired from the closest available time point (week 28), indicated in parentheses. (B) Geometric means of absolute counts of CD4+ memory T cells. The blue line represents the four Mamu-B*08-positive macaques, and the red line represents the four Mamu-B*08-negative macaques. By use of log-transformed arithmetic means, statistically significant differences were found between the absolute CD4+ memory T-cell counts of the Mamu-B*08-positive and -B*08-negative groups at the following time points: week 6 (P = 0.001), week 8 (P < 0.001), week 10 (P < 0.001), week 12 (P = 0.014), and week 18 (or 28) (P = 0.001).
FIG. 3. Ex vivo whole PBMC IFN-γ ELISPOT using peptides spanning the entire SIVmac239 proteome and the relevant MHC class I-restricted minimal optimal CD8+ T-cell epitopes at 3 weeks postinfection. Eighty-one peptide pools (10 15-mer peptides overlapping by 11 amino acids) were tested in IFN-γ ELISPOT assays spanning the complete SIVmac239 proteome. Total responses for each protein were calculated by adding the mean values of the individual peptide pools for each SIV protein. Individual minimal optimal peptides were included to detect responses.
immune response in r01027, compared to 32.7% and 23.3%, respectively, for Mamu-B*08-restricted CD8+ T cells (Fig. 4A). Animal r00032 expressed both Mamu-A*02 and Mamu-B*08. While Mamu-A*02-restricted responses accounted for 14.4% of the SIV-specific response, Mamu-B*08-restricted responses accounted for 48.7% of the total number of SIV-specific CD8+ T cells. Macaque r02019 did not express any other characterized MHC class I alleles aside from Mamu-B*08, and Mamu-B*08-restricted CD8+ T cells accounted for 59.6% of its total SIV-specific response.

Despite the presence of as many as 12 MHC class I alleles expressed in a given macaque (13, 16, 62), Mamu-B*08-restricted CD8+ T cells accounted for a large proportion (23.3 to 59.6%) of the total SIV-specific response. Furthermore, the contribution of Mamu-B*08-restricted SIV-specific immune responses is likely an underestimate because, unlike for Mamu-A*01 (2, 4) and Mamu-A*02 (46), a comprehensive peptide binding motif and epitope mapping study has not yet been completed for Mamu-B*08.

Increased breadth of Mamu-B*08-restricted CD8+ T-cell responses against epitopes in Vif and Nef was associated with elite control. Despite the fact that all four macaques made strong Mamu-B*08-restricted CD8+ T-cell responses, only two animals were ECs, controlling viral replication to approximately 1,000 vRNA copies/ml (49, 76). We next analyzed the breadth of the Mamu-B*08-restricted immune responses to examine the possible role that multiple epitope-specific, Mamu-B*08-restricted CD8+ T-cell responses might play in the control of SIV replication. At 3 weeks postinfection, the progressor macaque r91003 made a narrowly focused immunodominant response against Vif172-179RL8 that constituted 87% of the total Mamu-B*08-restricted CD8+ T cells detected in this macaque (Fig. 4B). Controller macaque r01027 also made an immunodominant response to this same epitope (53.7%, Vif172-179RL8). However, three other Mamu-B*08-restricted CD8+ T-cell responses (Vif23-131RL9, Env573-581KL9, and Nef137-146RL10) each accounted for >10% of the Mamu-B*08-restricted immune breadth in r01027. By contrast, the EC macaques r00032 and r02019 made robust CD8+ T-cell responses to several of the eight Mamu-B*08-bound peptides, none of which accounted for >50% of the Mamu-B*08-restricted CD8+ T-cell response. At 3 weeks postinfection, five of the eight Mamu-B*08-restricted responses were >1,000 SFC/10^6 PBMC for the EC r00032, while the other EC r02019 had three responses exceeding this magnitude (Fig. 4B).

To study changes in the Mamu-B*08-restricted epitope breadth over time, we performed MHC class I tetramer stains on cryopreserved PBMC during the acute to early chronic phase of SIV infection for all eight Mamu-B*08-restricted CD8+ T-cell epitopes (Fig. 5). We found that the MHC class I tetramer stains largely agreed with the magnitude of the IFN-γ ELISPOT responses at 3 weeks postinfection (Fig. 4) and subsequent time points (data not shown). At 3 weeks postinfection, ECs r00032 and r02019 typically displayed the largest percentages of SIV-specific CD8+ T cells for each of the eight Mamu-B*08-restricted CD8+ T-cell responses. These two ECs targeted three epitopes, two in Vif (RL9 and RL8) and one in Nef (RL10), with massive CD8+ T-cell responses (0.76 to 7.31%) (Fig. 5). Controller macaque r01027 displayed several low-level responses aside from its immunodominant Vif172-179RL8 that were maintained over time. In contrast, progressor r91003 mounted a Mamu-B*08-restricted immune response focused primarily against the Vif172-179RL8 epitope at 3 weeks postinfection, although a sizeable response against Nef137-146RL10 was detected later after infection (Fig. 5). Interestingly, only r91003 failed to make a substantial response against the Vif23-131RL9 epitope during any of the time points, potentially implicating this response in Mamu-B*08-restricted control. Not surprisingly, only low-level frequencies for the majority of the Mamu-B*08-restricted CD8+ T cells were found in the three macaques controlling viremia at 18 weeks postinfection. This diminution of their robust acute-phase responses was likely due to small amounts of antigen stimulation due to low SIV plasma virus concentrations after primary infection.

Stronger and broad CD4+ T-cell responses in the Mamu-B*08-positive EC macaques. Several previous studies have shown an association between strong and broad CD4+ T-cell responses and control of HIV/SIV replication (10, 23, 25, 28, 50, 68). CD4+ T cells may also play a role in the prevention of CD8+ T-cell exhaustion (45). Therefore, we investigated the involvement of CD4- mediated immune responses in the four Mamu-B*08-positive macaques. At 6 weeks postinfection, we performed ex vivo IL-2 and IFN-γ ICS assays to determine whether responses to peptide pools detected in the PBMC IFN-γ ELISPOTs at 3 weeks postinfection were mediated by CD4+ or CD8+ T cells. Progressor r91003 and controller r01027 did not make any appreciable CD4+ responses at this time point (data not shown). In contrast, EC macaque r00032 made three responses directed against the Gag121-171(D), Rev15,54(A), and Nef15,51(A) pools, while EC macaque r02019 responded to the Gag241-291(G) pool. All of these responses were at or below 0.1% of the level for CD4+ lymphocytes (data not shown).

We next determined whether CD4+ T-cell responses broadened after resolution of primary viremia, as a durable, long-
FIG. 4. Involvement of Mamu-B*08-restricted CD8+ T-cell responses in the overall CD8+ T-cell-mediated immune response against SIVmac239 at 3 weeks postinfection. (A) Mamu-B*08-restricted CD8+ T cells make a major contribution to the total SIV-specific immune response during the acute phase of infection. The contributions of known responses are shown for Mamu-A*01, Mamu-A*02, and Mamu-B*08 by attributing peptide pools to particular MHC class I molecules based on the locations of known SIV epitopes in these pools. (B) Mamu-B*08-restricted CD8+ T cells recognize a broader epitope repertoire in EC macaques early in SIV infection. CD8+ T-cell responses accounting for >10% of the total Mamu-B*08-restricted immune response are indicated above the appropriate bar. Minimal optimal peptides were used for the
lasting control may also be aided by CD4+ T-cell responses (45). IFN-γ ELISPOT assays were performed on PBMC depleted of CD8+ cells at 20 weeks after SIV infection (Fig. 6). Controller r01027 made only one detectable CD4+ T-cell response to a peptide pool in Gag. Surprisingly, despite high levels of SIV viremia, the number of CD4+ T-cell responses made by progressor r91003 was comparable to those made by the EC macaques r00032 and r02019. However, all of the following Mamu-B*08-restricted epitopes: Vif 123-131RL9, Vif 172-179RL8, Rev 44-51RL8, Env 573-581KL9, and Nef 246-254RL9. Mamu-B*08 epitopes annotated with an asterisk were represented with peptides slightly larger than the minimal optimal as these responses were in the process of being fine mapped (Rev 12-20KL9* represents two overlapping 15-mer peptides positions 5 to 23, Nef 8-16RL9* represents a 10-mer peptide at positions 7 to 16, and Nef137-146RL10* represents an 11-mer peptide at positions 136 to 146). Data for both panel A and panel B were derived from ex vivo IFN-γ ELISPOT assays at 3 weeks postinfection (Fig. 3). Background levels were subtracted from each well. Mean responses of <50 SFC per 1 × 10^6 cells (white bars) were not considered positive.

FIG. 5. Comparison of Mamu-B*08-restricted CD8+ T-cell responses detected by MHC class I tetramers during the first 18 weeks postinfection. Cryopreserved PBMC from the four Mamu-B*08-positive macaques (black, r91003; green, r01027; blue, r00032; and red, r02019) were thawed and stained at the indicated time points for the eight Mamu-B*08-restricted epitopes. Results of MHC class I tetramer stains are shown here as percentages of CD3+CD8+ MHC class I tetramer-positive gated lymphocytes. The threshold of detection in these assays was 0.02% CD3+CD8+ MHC class I tetramer-positive gated lymphocytes.
CD4⁺ T-cell responses detected in r91003 were low frequency, with between 50 and 150 SFC/10⁶ CD8⁺ cell-depleted PBMC. In comparison, the ECs exhibited stronger CD4⁺ immune responses. EC r00032 mounted three CD4⁺ T-cell responses, while EC r02019 had six CD4⁺ T-cell responses at 150 SFC/10⁶ CD8⁺ cell-depleted PBMC. Almost half of the CD4⁺ T-cell responses generated by r91003, r00032, and r02019 were against Gag, a protein that is frequently recognized by CD4⁺ T cells in successful vaccinees and ECs (10, 23, 25, 28, 68, 74). A number of the low-level responses (100 SFC/10⁶ CD8⁺ cell-depleted PBMC) from these three macaques were also found in frozen IFN-γ ELISPOT assays between weeks 14 and 26 postinfection (data not shown).

FIG. 6. Detection of CD4⁺ T-cell responses at 20 weeks postinfection by ex vivo CD8⁺ cell-depleted IFN-γ ELISPOT. Thirty-five peptide pools, with 10 15-mer peptides overlapping by 11 amino acids, spanning the entire SIVmac239 proteome (except for Pol and Env due to limited availability of PBMC) were tested in IFN-γ ELISPOT assays on PBMC depleted of CD8⁺ cells. Each column represents the number of SFC per 10⁶ CD8⁺ cell-depleted PBMC directed against a single peptide pool at 20 weeks postinfection. Only positive responses are indicated. Background levels were subtracted from each well. Mean responses of <50 SFC per 1 × 10⁶ cells (dashed line) were not considered positive.
TABLE 1. Amino acid replacements in the circulating plasma virus of the four Mamu-B*08-positive macaques at 18 weeks after SIVmac239 infection

| Type of substitution  | No. of amino acid replacements |
|----------------------|--------------------------------|
|                      | Complete          | Mixed base | Total     |
| MHC class I associated |                                |            |
| Mamu-B*08<sup>a</sup> | 6                 | 4         | 10        |
| Mamu-A*01<sup>b</sup> | 2<sup>c</sup>     | 4<sup>e</sup> | 6         |
| Mamu-A*02<sup>c</sup> | 0                 | 0         | 0         |
| Not MHC associated   |                                |            |
| Suboptimal           | 6                 | 1         | 7         |
| Other<sup>d</sup>    | 2                 | 0         | 2         |
| Unknown              | 1                 | 7         | 8         |
| Total                | 17                | 16        | 33        |

<sup>a</sup>Expressed by r91003 (progressor), r01027 (controller), r00032 (EC), and r02019 (EC).
<sup>b</sup>Expressed by r91003 (progressor) and r01027 (controller).
<sup>c</sup>Expressed by r00032 (EC).
<sup>d</sup>Substitutions at position 67 in Env were previously detected in 33 of 35 SIV-infected macaques regardless of MHC class I genotype (60).
<sup>e</sup>This number includes concomitant changes in the overlapping open reading frame for Vpr (see Fig. S1C in the supplemental material).

Early viral variation in several Mamu-B*08-restricted CD8<sup>+</sup> T-cell epitopes. It has been shown that CD8<sup>+</sup> T cells exert enormous selective pressures and can lead immunodeficiency virus escape as early as 4 weeks postinfection, (3, 5, 7, 8, 12, 19, 20, 26, 35, 36, 51, 59, 60, 64, 65). We therefore investigated whether viral escape was a factor in determining whether a Mamu-B*08-positive macaque develops high viral loads or controls SIV replication. Sequencing the entire SIVmac239 genome from plasma virus at 18 weeks postinfection revealed only a few substitutions in the clonal SIVmac239 virus (see Fig. S1 in the supplemental material). In total, mutations resulting in 33 amino acid substitutions (17 complete amino acid substitutions and 16 incomplete replacements) were found in plasma virus from the four SIV-infected macaques (Table 1). A deletion in the RNA that encodes the gp120 Env protein was also found in progressor macaque r91003 (positions 418 to 424) (see Fig. S1 in the supplemental material). Of these 33 amino acid substitutions, 7 were previously documented to be the result of suboptimal nucleotides that routinely mutate in the majority of Indian rhesus macaques in vivo, likely increasing the fitness of SIVmac239 (1). An additional two substitutions located in Env also do not appear to be associated with HMC class I expression (Table 1). Of the remaining 24 substitutions, 16 were within defined MHC class I epitopes, while 8 cannot be accounted for at this time.

Mamu-B*08-restricted CD8<sup>+</sup> T cells were responsible for selecting the majority of the viral variation in the replicating plasma virus at 18 weeks postinfection. We found 10 amino acid replacements in or near Mamu-B*08-restricted epitopes (Table 1; also see Fig. S1 in the supplemental material). Six of these were the result of complete nucleotide substitutions, while four were identified as mixed-base substitutions. Surprisingly, viral variation was observed in four of the eight known Mamu-B*08-restricted CD8<sup>+</sup> T-cell epitopes (Vif<sub>123-131</sub>RL9, Vif<sub>172-179</sub>RL8, Nef<sub>136-147</sub>RL10, and Nef<sub>246-254</sub>RL9), with at least one mutation detected in each of the four macaques at 18 weeks postinfection.

Selective pressures exerted by Mamu-A*01-restricted CD8<sup>+</sup> T cells accounted for six amino acid replacements found at 18 weeks postinfection (Table 1; also see Fig. S1 in the supplemental material). Escape mutations were detected in both Mamu-A*01-positive macaques (r91003 and r01027) in the Tat<sub>28-35</sub>SL8 epitope, consistent with the rapid evolution of this sequence (3, 59). Three concomitant changes in the overlapping open reading frame for Vpr were also identified. The only other substitution that was likely due to Mamu-A*01-restricted CD8<sup>+</sup> T cells was in the Env<sub>726-735</sub>ST10 epitope in controller macaque r01027. No variation was identified in Mamu-A*02-restricted epitopes in EC macaque r00032.

We further defined the viral ontogeny in these four Mamu-B*08-restricted epitopes (Vif<sub>123-131</sub>RL9, Vif<sub>172-179</sub>RL8, Nef<sub>136-147</sub>RL10, and Nef<sub>246-254</sub>RL9) by population sequencing of replicating plasma virus at seven time points between weeks 4 and 18 postinfection (Fig. 7). Epitope mutations were detected as early as 10 weeks after SIV infection in Nef<sub>136-147</sub>RL10 in the EC r02019. The majority of the viral variants detected at 18 weeks postinfection appeared between weeks 12 and 13 postinfection. Nonetheless, we saw viral variation in four of the eight Mamu-B*08-restricted epitopes, indicating selective pressure mediated by Mamu-B*08-restricted CD8<sup>+</sup> T cells on multiple epitopes. A majority of these mutations were observed previously in the chronic phase, and selection was correlated with expression of Mamu-B*08 (48).

Viral variation in the Mamu-B*08-restricted CD8<sup>+</sup> T-cell epitopes appeared to be more pronounced in the progressor (r91003) and controller (r01027) macaques (Fig. 7). The Vif<sub>123-131</sub>RL9 epitope showed the same pattern of viral variation (alanine to valine) in all four macaques. By contrast, mutations in the second Vif epitope, Vif<sub>172-179</sub>RL8, were detected only in the controller r01027. The Nef<sub>136-146</sub>RL10 epitopes displayed two patterns of viral variation. The first pattern was seen in the two ECs and involved an isoleucine (I)-to-threonine (T) change at position eight of the epitope. The second pattern was found in the other two animals and involved a change of alanine (A) to proline (P) in the amino acid N-terminal to Nef<sub>136-146</sub>RL10 that may alter antigen processing (48). Interestingly, a similar A-to-P substitution immediately N-terminal of an HLA-B57-restricted HIV epitope in Gag is known to alter antigen processing and to abrogate recognition (18). The second Nef epitope, Nef<sub>246-254</sub>RL9, accumulated substitutions only in progressor r91003 and controller r01027. In both cases, there was a leucine (L)-to-proline (P) substitution at the C terminus of the epitope.

DISCUSSION

Previous investigations have linked particular MHC class I alleles to control of HIV replication (14, 15, 29, 34, 55), yet understanding how CD8<sup>+</sup> T cells restricted by these protective alleles contribute to viral control remains a mystery. Recently, we identified an association between the Indian rhesus macaque MHC class I allele Mamu-B*08 and control of SIVmac239 replication (49). While identifying CD8<sup>+</sup> T-cell responses restricted by this allele in EC macaques, we also discovered that the preliminary peptide binding motif of Mamu-B*08 appears...
to be similar to the peptide binding motif of HLA-B27 (48), an MHC class I allele associated with control of HIV replication in humans. The high percentage of Mamu-B*08-positive macaques that become ECs (50%) and the functional similarity of Mamu-B*08 to HLA-B27 make these MHC class I-defined macaques ideal for modeling human ECs. We therefore studied the immunopathogenic events of acute-phase SIV infection in four Mamu-B*08-positive macaques in an attempt to further understand CD8-mediated viral control in ECs.

In this observational study, we show that three of four Mamu-B*08-positive macaques controlled replication of the pathogenic SIVmac239 isolate (Fig. 1). Two of these macaques (r00032 and r02019) were ECs with plasma viral concentrations of approximately 1,000 vRNA copies/ml at 20 weeks postinfection. The third macaque (r01027) showed some measure of control of this highly pathogenic virus, with a plasma viral concentration of 20,000 vRNA copies/ml at 20 weeks postinfection. Only macaque r91003 had plasma viremia similar to the viral set points in the majority of animals that progressed to AIDS (5 × 10^5 vRNA copies/ml).

We then investigated how the majority of Mamu-B*08-positive macaques controlled viral replication. Recent experiments have demonstrated that the rapid depletion of the CD4 memory T cells during the acute phase of HIV/SIV infection might be an important factor contributing to disease progression (44, 53). Interestingly, while the CD4 memory T cells in the PBMC were depleted during primary SIV infection in two of the four Mamu-B*08-positive macaques, all four of
the Mamu-B*08-positive macaques recovered their CD4+ memory T cells by the chronic phase of infection (Fig. 2). By contrast, four Mamu-B*08-negative macaques infected with SIVmac239 experienced acute-phase loss of their CD4+ memory T-cell subset and none of the Mamu-B*08-negative macaques recovered this important CD4+ T-cell subset. At 18 weeks postinfection, the progressor macaque (r91003) started to show signs of CD4+ memory T-cell depletion, likely related to the high plasma virus concentrations in this animal.

Mamu-B*08-restricted CD8+ T-cell responses contributed substantially to the acute phase of SIVmac239 infection (Fig. 4). However, the pattern of immunodominance varied from animal to animal, and breadth seemed to correlate with successful control of immunodeficiency virus replication. At 3 weeks postinfection, progressor r91003 focused its Mamu-B*08-restricted CD8+ T-cell responses primarily on only one of the eight mapped Mamu-B*08 CD8+ T-cell epitopes, Vif172-179 RL8. Vif172-179 RL8-specific cells accounted for 87% of the eight mapped Mamu-B*08 CD8+ T-cell responses (Table 1). By comparison, viral variation was not detected in any of the Mamu-A*02-restricted epitopes, and amino acid replacements were present in only two Mamu-A*01-restricted epitopes (Tat28-35 SL8 and Env276-283 ST10) (see Fig. S1 in the supplemental material). Four of the eight Mamu-B*08-restricted epitopes (Vif23-31 RL9, Vif172-179 RL8, Nef136-141 RL10, and Nef246-254 RL9) exhibited viral variation consistent with mutations identified in a previous study (48). These results may suggest that Mamu-B*08-restricted CD8+ T cells exert more selective pressure than other MHC class I-restricted responses during the critical early phase of infection. Alternatively, the mutations observed in these epitope sequences could occur in regions of the viral genome that are not under strong evolutionary constraints. Our group and others have previously shown that fitness costs may play a role in determining the rate at which escape mutations accumulate and revert in vivo (21, 22, 43).

We also followed the ontogeny of mutations by population sequencing these four Mamu-B*08-restricted epitopes at various time points. We discovered the first evidence of viral variation in circulating plasma virus at 10 weeks postinfection and mutations within several Mamu-B*08-restricted epitopes by 13 weeks postinfection (Fig. 7). Interestingly, the patterns of amino acid substitutions in the Nef137-146 RL10 and Nef246-254 RL9 epitopes differentiated the four Mamu-B*08-positive macaques into two separate groups, the ECs and the non-ECs. It will be intriguing to see if viral control is lost through the accumulation of additional mutations as the three macaques that control SIV replication progress further into the chronic phase of infection.

Similar to those in HLA-B27 and HLA-B57-positive individuals (6, 9, 38), the acute-phase CD8+ T-cell responses in Mamu-B*08-positive macaques were dominated by Mamu-B*08-specific CD8+ T-cell responses (Fig. 3 and Fig. 4). Surprisingly, Mamu-B*08 appeared to reduce the dominant influence of Mamu-A*01 on CD8+ T-cell responses. Previously, such immunodominance was described to occur in macaques expressing both Mamu-A*01 and Mamu-A*02 during the acute phase of SIVmac251 infection (58). Typically, in Mamu-A*01-positive macaques, the Mamu-A*01-restricted Tat28-35 SL8 and Gag161-179 CM9-specific CD8+ T cells account for >50% of the acute-phase SIV-specific immune responses (56). However, for both r91003 and r1027, the Mamu-A*01-restricted contribution was below 50%. Responses restricted by Mamu-A*02 in r00032 also appeared to be diminished in this Mamu-B*08-positive macaque compared to those in Mamu-A*02-positive macaques that do not express Mamu-B*08. It should be emphasized, however, that these data are derived from only three animals and that larger cohorts of Mamu-B*08-positive macaques expressing other MHC class I molecules should be studied to clarify this issue.

HIV disease progression in HLA-B27-positive humans appears to be associated with viral escape from the immunodominant Gag263-272 KK10 response during the chronic phase of infection (8, 20, 26, 36). However, additional HIV-specific CD8+ T-cell responses may be involved in the initial control of viremia. Because immune responses in HIV-infected individuals are normally characterized using IFN-γ ELISPOT with
consensus peptides, often after primary infection, definition of the entire breadth of the HLA-B27-restricted CD8+ T-cell responses may be incomplete. While preliminary, our current findings appear to indicate that the breadth of CD8+ T-cell responses restricted by protective MHC class I alleles, rather than a single immunodominant response, may be important in determining the control of replication of the AIDS virus.

Surprisingly, while Mamu-B*08 appears to bind similar peptides to HLA-B27, an immunodominant CD8+ T-cell response directed against Gag has not been identified for Mamu-B*08. Rather, Mamu-B*08 restricts robust CD8+ T-cell responses largely from Vif and Nef. Moreover, we are currently attempting to identify all of the SIV epitopes restricted by Mamu-B*08. At this time, we have identified only a single Gag-specific CD8+ T-cell response, and it is both low frequency and recognized in only a few Mamu-B*08-positive macaques (J. T. Loffredo et al., unpublished data). Hence, while CD8+ T cells targeting Gag have been shown to be extremely effective in controlling immunodeficiency virus replication (7, 8, 20, 26, 35, 36, 39, 51, 70), it may be possible for individuals to control viremia by directing responses against other proteins as well. Mamu-B*08-positive, SIV-infected macaques may offer an intriguing model for studying such a mechanism. Interestingly, both HLA-B27 and Mamu-B*08 present many epitopes containing two N-terminal basic amino acids. These peptides are relatively resistant to peptidase activity, and thus, those peptides may be more stable, and this may result in more efficient MHC class I antigen presentation (30).

Approximately 50% of Mamu-B*08-positive macaques become ECs, controlling replication of the pathogenic SIVmac239 isolate to <1,000 vRNA copies/ml (49). Given that the chronic-phase viral set points in 175 macaques that progressed to AIDS were ~500,000 vRNA copies/ml, and approximately two-thirds of macaques die by 1 year postinfection (40), this level of control is remarkable. However, not all Mamu-B*08-positive macaques become ECs after SIVmac239 infection, and the immune system’s role in successful viral containment remains difficult to define. Understanding how some of these macaques become progressors or controllers might give us key insights into how to make an effective HIV vaccine. Here, we provide evidence from a small yet provocative study that the immunodominance and breadth of CD8+ T-cell responses restricted by protective MHC class I alleles may facilitate the development of elite control. Further experiments are needed to explore this intriguing idea, especially given the fundamental implications that this might have for future vaccines.

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