Monofunctional and Polyfunctional CD8$^+$ T Cell Responses to Human Herpesvirus 8 Lytic and Latency Proteins$^{-\dagger}$

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Human herpesvirus 8 (HHV-8) is the etiological agent of Kaposi's sarcoma, primary effusion lymphoma, and multicentric Castleman's disease. It is postulated that CD8$^+$ T cell responses play an important role in controlling HHV-8 infection and preventing development of disease. In this study, we investigated monofunctional and polyfunctional CD8$^+$ T cell responses to HHV-8 lytic proteins gB (glycoprotein B) and K8.1 and latency proteins LANA-1 (latency-associated nuclear antigen-1) and K12. On the basis of our previous findings that dendritic cells (DC) reveal major histocompatibility complex (MHC) class I epitopes in gB, 2 in K8.1, 5 in LANA-1, and 1 in K12. These new HHV-8 epitopes activated monofunctional and polyfunctional CD8$^+$ T cells that produced various combinations of gamma interferon, interleukin 2, tumor necrosis factor alpha, macrophage inhibitory protein 1B, and cytotoxic degradation marker CD107a in healthy HHV-8-seropositive individuals. We were also able to detect HHV-8-specific CD8$^+$ T cells in peripheral blood samples using HLA A*0201 pentamer complexes for one gB epitope, one K8.1 epitope, two LANA-1 epitopes, and one K12 epitope. These immunogenic regions of viral lytic and latency proteins could be important in T cell control of HHV-8 infection.

Human herpesvirus 8 (HHV-8), also referred to as Kaposi’s sarcoma-associated herpesvirus, is a gammaherpesvirus that causes Kaposi’s sarcoma (KS), primary effusion lymphoma, and multicentric Castleman’s disease. The importance of developing effective prevention and treatment for HHV-8 infection is evident in that KS, a neoplasm of endothelial origin, continues to be the most common cancer among human immunodeficiency virus (HIV)-infected patients (8). KS is also the leading cause of cancer in children in sub-Saharan Africa (7). Although the incidence of KS in HIV-infected persons declined with the advent of antiretroviral therapy (ART) (10), KS can occur in persons on ART with suppressed HIV infection and high CD4$^+$ T cell counts (25).

The immune responses responsible for controlling HHV-8 infection and preventing KS are not clear. CD8$^+$ T cell immunity likely plays a significant role in HHV-8 infection, as these cells have been shown to be crucial in controlling infection caused by the other human gammaherpesviruses, i.e., Epstein-Barr virus (EBV) (11, 14). In support of this hypothesis, our laboratory (40–42) and others (4–6, 12, 19, 23, 26–28, 31, 32, 36, 37, 43, 44) have shown that CD8$^+$ T cells produce gamma interferon (IFN-γ) in response to HHV-8 immunodominant epitopes presented by major histocompatibility complex class I (MHC-I) in HHV-8-seropositive individuals. Little is known whether T cells produce other immune mediators in response to HHV-8 infection. Indeed, polyfunctional T cells, i.e., single cells producing two or more immune mediators, have been linked to control of HIV and other persistent infections (1, 24, 29, 33) and could play a role in controlling HHV-8 infection. In one recent study, HHV-8 epitope-specific, polyfunctional T cells were detected in patients with multicentric Castleman’s disease, but these cells did not differ in number from those in healthy controls (13). Another study has found that patients with controlled KS had HHV-8-specific CD8$^+$ T cells that secreted IFN-γ and tumor necrosis factor alpha (TNF-α) but that patients with progressive disease had weaker and less polyfunctional CD8$^+$ T cells (2).

HHV-8 epitope-specific monofunctional and polyfunctional T cell immunity could be important in development of HHV-8 vaccines that induce T cell responses that target these viral epitopes. In the present study, we therefore investigated CD8$^+$ T cell responses to two HHV-8 lytic proteins, gB (glycoprotein B) and K8.1, and two latency proteins, LANA-1 (latency-associated nuclear antigen-1) and K12. We previously showed that optimal induction of T cell reactivity to the HHV-8 protein gB required 1 week of stimulation with peptide-loaded, autologous, mature, monocyte-derived dendritic cells (DC) (40). Using this enhanced DC-T cell stimulation system, we now have revealed several new epitopes for these four lytic and latency HHV-8 proteins in healthy HHV-8-seropositive individuals, which induce both monofunctional and polyfunctional CD8$^+$ T cells. These regions of HHV-8 could be critical in understanding HHV-8 immunopathogenesis and in vaccine development.

MATERIALS AND METHODS

**Study subjects.** Healthy, HIV-1-negative subjects were selected based on their HHV-8 antibody status and MHC-I genotype, and written informed consent was obtained. Detection of HHV-8 serum antibody specific for viral lytic antigens was done using an indirect immunofluorescence assay (40). High-resolution HLA molecular typing was conducted by the University of Pittsburgh Medical Center Tissue Typing Laboratory. The donors were classified into HLA A*0201-positive HHV-8-seropositive and HLA A*0201-positive HHV-8-seronegative groups.
Synthetic peptides. For the initial studies, libraries of 15-mer peptides overlapping by 11 amino acids (aa) derived from K12, gB, and K8.1 protein sequences were synthesized (PEPscreen; Sigma, St. Louis, MO). For the larger protein LANA-1, a library of 15- to 20-mer peptides overlapping by 11 aa was used. Protein sequences were obtained from the National Center for Biotechnology Information (NCBI) database, with accession number AAD46501 for LANA-1, accession number AAD46499 for K12, accession number ABD28851 for gB, and accession number ABD28902 for K8.1. As LANA-1 contains a large repeat region, we used one set of representative peptides to span this region (peptides 59 to 75) (Fig. 1A). For epitope mapping studies, putative optimal 9-mer peptides were synthesized based on anchor residues for HLA A*0201 (30) as well as peptides No rC terminus amino acid shorter or longer than optimal (15). The following previously published 9-mer, HLA A*0201-restricted HHV-8 epitopes were also used: LANA-1238-246 (WATESPIYV) (12), LANA-11116-1124 (QMARLAWEA).

FIG. 1. IFN-γ ELISPOT responses to four HHV-8 latency and lytic proteins. CD14+ PBMC from healthy HLA A*0201-positive HHV-8-seropositive donors were stimulated with autologous, mature DC that were loaded with overlapping 15- to 20-mer peptides derived from each of the HHV-8 proteins LANA-1, K12, gB, and K8.1. IFN-γ production was measured by a DC-enhanced ELISPOT assay, and the number of spots produced by cells without peptide was subtracted from the number of spots produced by cells with peptide to give net values for spots. The donors (three or four donors used for each protein) are listed in each row, while the representative peptide numbers are listed in each column. The colors for the boxes represent the net number of IFN-γ spots per 10^6 cells as follows: white, <1 spot; light gray, 1 to 199 spots; dark gray, 200 to 399 spots; black, 400 to 599 spots; and red, ≥600 spots. Hot spots are circled for LANA-1 (A), K12 (B), gB (C), and K8.1 (D).
Novel T cell epitopes in HHV-8 lytic and latency proteins. To define MHC-I epitopes that generate CD8+ T cell responses to HHV-8 proteins, a DC-enhanced IFN-γ ELISPOT assay was used. This was based on our previous finding that this method was necessary to reveal an epitope in HHV-8 gB due to the nonrobust nature of the immune response to HHV-8 in healthy HHV-8-seropositive persons (40). T cells were stimulated with autologous mature DC that were loaded with overlapping peptides derived from HHV-8 LANA-1, K12, gB, and K8.1, and single-cell IFN-γ production was measured by the ELISPOT assay.

The IFN-γ ELISPOT assay results for four HLA A*0201-positive HHV-8-seropositive donors (A29, A34, A52, and A57) to peptides from each viral protein are displayed in Fig. 1. For the larger proteins LANA-1 and gB, peptides containing possible epitopes, termed “hot spots” (Fig. 1, circled areas), were defined as peptides that generated a positive response above the background level of mock-stimulated cells for all donors tested, with at least one donor responding in the highest category (red boxes). For the smaller proteins K12 and K8.1, we considered hot spots to be those peptides that the majority of donors responded to above the background level, with at least one donor responding in the highest category. For LANA-1, out of 106 peptides tested, we found 8 hot spots (Fig. 1A). For K12, out of 13 peptides tested, we found 1 hot spot (Fig. 1B). For gB, out of 235 peptides tested, we found 9 hot spots, including 1 containing our previously described epitope (40) (Fig. 1C). For K8.1, out of 60 peptides tested, we found 3 hot spots, including 2 containing previously described epitopes (5, 43) (Fig. 1D). Two HLA A*0201-positive HHV-8-seronegative donors did not respond above the background level to any of these hot spots from the four viral proteins (data not shown). Thus, these hot spots represented regions of the four HHV-8 proteins that displayed positive reactivity associated with HHV-8 seropositivity.
We next mapped minimal epitope sequences for 10 of these hot-spot peptides. To determine the minimal, optimal epitope, we used peptide families consisting of the putative optimal 9-mer based on anchor residues for HLA A*0201, along with peptides with N or C terminus amino acid truncation (8-mers) or extension (10-mers). Initially, we adapted the conventional method to define optimal T cell epitopes, i.e., T cell response to different concentrations of N- and C-terminal extensions and truncations of peptide in a 16-h ELISPOT assay, to our 7-day extended assay. However, stimulating PBMC for 7 days with DC loaded with 5 different 10-fold concentrations of these peptide families did not differentiate a dominant peptide response (data not shown). Therefore, we tested the peptide families for the optimal epitopes by stimulating the cells for 7 days with a single concentration of peptide (2 μg/ml), followed by a conventional 16-h ELISPOT assay using the same peptide. We assessed five peptide families from LANA-1, one from K12, two from gB, and two from K8.1. One of the hot spots that we identified (K8.1 peptide 38 [Fig. 1D]) contained a published 15-mer epitope (43). Therefore, we included this peptide family to determine the minimal epitope sequence. Two of the hot spots (gB peptide 139 [Fig. 1C] and K8.1 peptide 56 [Fig. 1D]) contained previously published 9-mer epitopes (5, 40) that we included as controls. As shown in Fig. 2, the DC-enhanced assay revealed positive IFN-γ responses for the known HHV-8 epitopes as well as for the peptide families of each protein. We defined the optimal epitope as the peptide from each family that generated the highest level of IFN-γ production above the level for the background (i.e., mock-stimulated cell cultures). By these approaches, we were able to define five novel, minimal epitope sequences for LANA-1, one for K12, two for gB, and two for K8.1 (Table 1).

**MHC-I restriction and binding of novel HHV-8 epitopes.** To verify that the novel epitopes were HLA A*0201 restricted, we generated epitope-specific T cell lines and used B cells that express only HLA A*0201 (A2 cells) or do not express any

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**TABLE 1. Novel HHV-8 HLA A*0201 epitopes identified in LANA-1, K12, gB, and K8.1**

| Protein | Peptide(s) | Amino acid positions | Sequence |
|---------|------------|----------------------|----------|
| LANA-1  | 17         | 140–148              | PESSQRPPP |
| LANA-1  | 33, 34     | 281–289              | AMLVLLAEI |
| LANA-1  | 48         | 417–425              | DGGDGKNTL |
| LANA-1  | 63         | 688–697              | OQDEQQQOD |
| LANA-1  | 83         | 920–928              | PVVSTHEQI |
| K12     | 7, 8       | 23–32                | WRLGAIPLL |
| gB      | 44         | 159–168              | SSKMVNVNG |
| gB      | 207        | 736–745              | MLMIIIIVAI |
| K8.1    | 21         | 73–81                | RLAAGSPSS |
| K8.1    | 38         | 135–143              | ALISAFSGS |

* Representative peptide numbers from Fig. 1.
HLA (null cells) as APC in the standard ELISPOT assay. Positive IFN-γ responses with the use of A2 cells as APC were evident for our five novel LANA-1 epitopes, one novel K12 epitope, two novel gB epitopes, and two novel K8.1 epitopes (Fig. 3, black bars). IFN-γ responses were also detected with the use of A2 cells as APC for our previously determined HLA A*0201-restricted gB epitope (40) (Fig. 3, black bars). For all peptides tested, response levels were lower when the null B cells were used as APC (Fig. 3, gray bars). Taken together, these results support that our novel epitopes from the four HHV-8 proteins are HLA A*0201 restricted.

Having demonstrated that these peptides are presented in the context of HLA A*0201, we then tested them in an MHC-I binding assay to confirm their specificity. To this end, ProImmune performed their class I REVEAL binding assay on our novel minimal epitopes as well as several known epitopes from the four HHV-8 proteins used in our study. The binding rates are reported as percentages relative to the binding of a known, strong HLA A*0201 T cell epitope. The peptides that showed high binding rates (i.e., above the intermediate control of a known, weak HLA A*0201 T cell epitope) were three known epitopes (gB492-500, K1217-25, and LANA-11116-1124) as well as two of our novel epitopes (LANA-1238-246 and K8.1 1209-1217) which supports that all of these peptides are HLA A*0201-restricted gB epitope (40) (Fig. 3, black bars). For all peptides tested, response levels were lower when the null B cells were used as APC (Fig. 3, gray bars). Taken together, these results support that our novel epitopes from the four HHV-8 proteins are HLA A*0201 restricted.

Direct identification of circulating CD8+ T cells specific for HHV-8 epitopes. On the basis of the highest scores of our peptide-MHC class I binding results, we synthesized five HLA A*0201 pentamers for two novel epitopes (LANA-1238-246 and K8.1 1209-1217) and three known epitopes (gB492-500, K1217-25, and LANA-11116-1124). As displayed in Fig. 4, the mean ± standard error (SE) percentages of epitope-specific CD8+ T cells in eight healthy HLA A*0201-positive HHV-8-seropositive individuals were 0.048% ± 0.010% for LANA-1238-246, 0.050% ± 0.012% for LANA-11116-1124, 0.053% ± 0.018% for K12-17-25, 0.050% ± 0.009% for gB492-500, and 0.096% ± 0.020% for K8.1 1209-1217. The mean levels of pentamer-positive CD8+ T cells were 0.020% ± 0.003% in three healthy HLA A*0201-positive HHV-8-seropositive individuals (Fig. 4). Overall, the level of pentamer-positive CD8+ T cells in the seropositive donors was significantly greater than that in the seronegative donors (P < 0.05).

Monofunctional and polyfunctional CD8+ T cell responses to novel HHV-8 epitopes. There are many cytokines in addition to IFN-γ, as well as chemokines and cytotoxic molecules, that are important in antiviral T cell responses. Moreover, polyfunctional T cells, which are defined as a single cell producing two or more such immune mediators, are an important immune correlate of protection against HIV-1 disease progression (1). As it is currently not clear whether monofunctional and polyfunctional CD8+ T cells play a role in controlling HHV-8 infection, we used a panel of markers for immune mediators to examine polyfunctional CD8+ T cell responses. This panel included the cytokines IFN-γ, TNF-α, and IL-2, the chemokine MIP-1b, and the degranulation mobilization marker CD107a. We used a modified procedure with DC-enhanced T cell cultures for the ICS assay developed in our laboratory (16) to improve detection of production of these immune mediators (for a representative analysis, see Fig. S2 in the supplemental material).

As shown in Fig. 5, positive responses for a variety of immune mediators were detected for our novel minimal epitopes and several known epitopes. The data are shown as bars representing the frequency of the listed mediator combination, with each color representing a different viral peptide. Each pie chart represents total responses to the listed peptide, with each color representing the number of cytokines produced. The ICS results revealed that in healthy HHV-8-seropositive individuals controlling infection, HHV-8 latency and lytic protein epitopes induced both monofunctional and polyfunctional CD8+ T cell responses to novel HHV-8 epitopes.
responses (Fig. 5). Although all of the peptides displayed a trend of more monofunctional responses than polyfunctional responses, the differences were not significant. With all of the HHV-8 epitopes tested taken into consideration, all five immunemediators were produced by monofunctional T cells, with a predominance of IFN-γ, IL-2, and CD107a (Fig. 5). Polyfunctional T cells producing two immune mediators were notable for the combinations of IFN-γ and IL-2 and for MIP-1β and IL-2. Predominant patterns of polyfunctional responses for three immune mediators included IFN-γ, TNF-α, and CD107a or MIP-1β, IL-2, and CD107a. Polyfunctional T cells producing four immune mediators included IFN-γ, TNF-α, MIP-1β, and CD107a or IFN-γ, TNF-α, IL-2, and CD107a.

Different HHV-8 epitopes induced different patterns of both monofunctional and polyfunctional responses. For example, the known epitope from gB (gB492-500) induced notable monofunctional responses consisting of IL-2 or CD107a and polyfunctional responses consisting of MIP-1β and IL-2 or IFN-γ, TNF-α, and CD107a (Fig. 5A). The known epitope K8.1 (K8.1135-143) displayed notable monofunctional responses consisting of IFN-γ or TNF-α and polyfunctional responses consisting of the combination of (i) CD107a and TNF-α, (ii) MIP-1β, TNF-α, and IFN-γ, or (iii) MIP-1β, TNF-α, IFN-γ, and CD107a (Fig. 5A).

The traditional method for mapping epitopes is through an IFN-γ ELISPOT assay. However, while analysis using one marker allows for the determination of response magnitude, the inclusion of several markers provides insight regarding the quality of the response. Therefore, along with our novel and known HHV-8 epitopes, we also investigated polyfunctional responses to one peptide family from each of the four HHV-8 proteins included in our study (Fig. 6). While trends were evident, no single peptide had a significantly higher magnitude of responses or more polyfunctional responses than the other peptides within the peptide families examined.

Taken together, these results show that the CD8+ T cell epitopes from the four HHV-8 proteins presented by DC induced a variety of both monofunctional and polyfunctional T cell responses in HHV-8-seropositive healthy donors. The specificity of this T cell response is less evident in polyfunctional T cell determinations.

DISCUSSION

It is postulated that CD8+ T cell responses play a significant role in controlling HHV-8 infection and preventing development of KS (2, 12). A central issue in understanding the CD8+ T cell immunity in HHV-8 infection is the identification of MHC-I epitopes for the more than 80 open reading frames (ORFs) of the virus. This has been difficult to do mainly because of relatively nonrobust CD8+ T cell responses to the virus (22). In this study, we employed a DC-based enhancement of CD8+ T cell responses to four HHV-8 lytic and latency proteins modified from our previous approach (40) to provide a greater insight into immunopathogenesis of the virus. Using this DC-enhanced IFN-γ ELISPOT assay, we have revealed five novel HLA A*0201-restricted CD8+ T cell epitopes in LANA-1, one in K12, two in gB, and two in K8.1. We were also able to detect epitope-specific CD8+ T cells directly in peripheral blood samples using HLA A*0201 pen-
tamer complexes specific for one gB epitope, one K8.1 epitope, two LANA-1 epitopes, and one K12 epitope. These results were expanded and confirmed by using a DC-enhanced ICS assay with polychromatic flow cytometry to detect multiple immune mediators. By this approach, we revealed that HHV-8-seropositive healthy donors controlling infection have circulating monofunctional and polyfunctional CD8\(^+\) T cells specific for an array of HHV-8 lytic and latency protein epitopes. The novel epitopes in our study did not all correspond to published motifs for the preferred HLA A*0201 anchor residues at positions 2 and 9. Such lack of correspondence to MHC-I allele-specific peptide motifs has been recognized for T cell epitopes of other viruses. Thus, the immunodominant HLA A*0201-restricted influenza A virus M1\,_{55-66} epitope GILGFVFTL does not have the preferred HLA A*0201 anchor residue L or M at position 2 (35). Moreover, over half of the HLA A*0201-restricted epitopes for vaccinia virus do not fit the optimal peptide binding motif (35). Aside from binding to MHC-I molecules, there are other important factors that determine the T cell response to a peptide, such as the presence CD8\(^+\) T cell precursors, interactions with T cell receptors and peptide transporters, and generation of peptides by different protease cleavage pathways (3, 26). Furthermore, our epitopes were determined by two different functional assays (ELISPOT and ICS assays) and represent CD8\(^+\) T cell positive reactivity in HLA A*0201-positive HHV-8-seropositive individuals.

FIG. 5. Polyfunctional CD8\(^+\) T cell responses to known and novel HHV-8 epitopes. CD14\(^-\) PBMC from healthy HLA A*0201-positive HHV-8-seropositive donors were stimulated for 1 week with autologous mature DC loaded with known HHV-8 epitopes (5, 6, 12, 40) (A) or novel epitopes from LANA-1 (B) or K12, gB, or K8.1 (C). Responses were measured by polychromatic flow cytometry, and net response values were averaged for three donors (mean \(\pm\) SE). Diagrams were generated using SPICE.
Given the large number of ORFs in HHV-8, we focused on ORFs that code for lytic and latency proteins that are considered important in HHV-8 pathogenesis and oncogenesis. We selected two latency cycle proteins, LANA-1 and K12, and two lytic cycle proteins, gB and K8.1. The lytic protein gB is a virion glycoprotein that binds cell surface heparan sulfate and induces signaling pathways, and the lytic protein K8.1 is a highly immunogenic (antibody-inducing) virion glycoprotein that also
binds cell surface heparan sulfate (40, 43). During latency, LANA-1 is a cell cycle regulatory protein important in anti-apoptotic functions and episome maintenance. The K12 (Kaposi) latency protein has roles in B cell signaling, apoptosis, and cell transformation induction (12, 26). While we do acknowledge that a limitation of our study is the small sample size, we found T cell reactivity to peptides of these lytic and latency proteins, indicating that immunity to each could be important in control of HHV-8 infection. We are currently applying a battery of our newly identified and previously documented epitopes to assess T cell immunity to HHV-8 infection and progression to KS in the Multicenter AIDS Cohort Study. Remarkably, given the need to use DC to reveal this T cell function, direct staining of unstimulated PBMC with HLA A*0201 pentamer complexes for two lytic and three latency protein epitopes identified these antigen-specific CD8+ T cells. HHV-8 specific CD8+ T cells have previously been detected in blood samples of HHV-8-seropositive transplant recipients and patients with AIDS-related and classical KS (19). However, to our knowledge, this is the first direct evidence of HHV-8-specific CD8+ T cells in blood samples of healthy HHV-8-seropositive individuals. We are currently determining the memory phenotype of these circulating T cells and if there is a functional downregulation of these cells.

Previous studies have shown that the central repeat region of LANA-1 inhibits proteasomal degradation and slows protein synthesis in order to inhibit cellular surveillance for CD8+ T cell epitopes (17). This model predicts that host T cell reactivity to LANA-1 would be minimal. However, we detected CD8+ T cell reactivity to several regions of LANA-1 in these healthy HHV-8-seropositive subjects. Others have shown anti-LANA-1 CD8+ T cell responses in patients with KS (12). Similarly, CD8+ T cell recognition has been found for EBV-encoded nuclear antigen 1 (21, 39), which had previously been proposed to escape CD8+ T cell recognition through cis inhibition of synthesis or blockade of proteasomal degradation by its glycine-alanine repeat domain. Thus, it appears that CD8+ T cells can mount functional responses to certain regions of LANA-1 regardless of cis effects on its production.

Polyfunctional CD8+ T cells, i.e., cells producing more than one immune mediator, are associated with superior control of persistent viral infections such as those with HIV-1 (1, 24). A recent investigation found similar patterns of polyfunctional T cell responses in patients with multicentric Castleman’s disease and healthy controls (13). Other investigators have reported that patients with nonprogressive KS have stronger and more frequent polyfunctional CD8+ T cell responses than those with progressive KS (2). In the present study, we established that polyfunctional T cells specific for known and novel epitopes are present in the PBMC of healthy HHV-8-seropositive individuals. We found that in these healthy, nonimmunosuppressed subjects controlling HHV-8 infection, there was a trend of more monofunctional CD8+ T cell responses than polyfunctional responses for all of the epitopes. Overall, our data suggest that the tight control of HHV-8 infection in healthy individuals may not require a predominance of circulating, virus-specific, polyfunctional CD8+ T cell reactivity. Furthermore, we found that different epitopes induced different patterns of monofunctional and polyfunctional responses. Similar results were found for T cell responses to influenza virus, where CD8+ T cells specific for some but not all viral proteins produced a wide range of cytokines (18). This could be explained by differences in T cell avidity for the peptide-MHC complex.

Traditionally, the standard in monitoring CD8+ T cell responses has been measuring single-cell IFN-γ production in response to antigenic stimuli by an overnight ELISPOT assay. While analysis using one marker allows for the determination of response magnitude, the inclusion of several markers provides insight regarding response quality. Therefore, we examined ICS responses to families of putative optimal epitopes consisting of peptides with N- and C-terminal extensions and truncations from each viral protein to further delineate the minimal epitope sequences. We found that there were different patterns of T cell reactivity revealed by ICS compared to those revealed by the ELISPOT assay. This stresses the need to move beyond IFN-γ ELISPOT assays in defining T cell immunity to viral infections (1, 38). The data also suggest that CD8+ T cells may mount diverse, functional reactivity to 8- to 10-mer variants of an MHC-I 9-mer epitope when presented by DC. This is supported by a recent finding in our laboratory that DC are able to generate efficient monofunctional and polyfunctional T cell responses against 8- to 10-mer N- and C-terminal variants of HIV Gag and Nef epitopes (15a). This enhancement of T cell responses by DC to peptide variations could also explain the broad spectrum of ELISPOT responses evident in our HHV-8 epitope mapping study.

In conclusion, we have revealed several novel HLA A*0201 minimal epitopes in both HHV-8 lytic and latency proteins. We have also shown the presence of antigen-specific CD8+ T cells in peripheral blood samples specific for two lytic and three latency protein epitopes. We have demonstrated that epitopes within these HHV-8 lytic and latency proteins induced both single and multiple immune mediators in CD8+ T cells from healthy HHV-8-seropositive individuals. The likely involvement of CD8+ T cells in the immune response to HHV-8 infection has implications for the prevention and treatment of HHV-8-associated cancers. These targeted regions of the virus that induce immune responses by CD8+ T cells could be critical in HHV-8 immunopathogenesis and the progression to KS.

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