Coinfection with Plasmodium falciparum malaria and Epstein-Barr virus (EBV) is a major risk factor for endemic Burkitt lymphoma (eBL), still one of the most prevalent pediatric cancers in equatorial Africa. Although malaria infection has been associated with immunosuppression, the precise mechanisms that contribute to EBV-associated lymphomagenesis remain unclear. In this study, we used polychromatic flow cytometry to characterize CD8+ T-cell subsets specific for EBV-derived lytic (BMFL1 and BRLF1) and latent (LMP1, LMP2, and EBNA3C) antigens in individuals with divergent malaria exposure. No malaria-associated differences in EBV-specific CD8+ T-cell frequencies were observed. However, based on a multidimensional analysis of CD45RO, CD27, CCR7, CD127, CD57, and PD-1 expression, we found that individuals living in regions with intense and perennial (holoendemic) malaria transmission harbored more differentiated EBV-specific CD8+ T-cell populations that contained fewer central memory cells than individuals living in regions with little or no (hypoendemic) malaria. This profile shift was most marked for EBV-specific CD8+ T-cell populations that targeted latent antigens. Importantly, malaria exposure did not skew the phenotypic properties of either cytomegalovirus (CMV)-specific CD8+ T cells or the global CD8+ memory T-cell pool. These observations define a malaria-associated aberration localized to the EBV-specific CD8+ T-cell compartment that illuminates the etiology of eBL.

First described in 1958 (1), endemic Burkitt lymphoma (eBL) remains one of the most prevalent childhood cancers in equatorial Africa. The average annual incidence is 2 per 100,000 children, with a peak age range of 5 to 9 years (2–4). In 1964, Epstein-Barr virus (EBV) was discovered in a tumor sample obtained from a patient with eBL (5), and EBV DNA has subsequently been detected in tumor cells from 95% of eBL cases (6). Thus, EBV was identified as the first human tumor virus, with ensuing studies revealing the virus-mediated oncogenic processes (7). However, eBL is most common in children residing in areas with the highest malaria transmission intensities (3, 8–10), an enigmatic observation suggesting that malaria-mediated oncogenic processes (7). However, eBL is most common in children residing in areas with the highest malaria transmission intensities (3, 8–10), an enigmatic observation that leaves the malaria-associated mechanisms involved in the etiology of eBL insufficiently established by comparison.

Infection with EBV occurs early in most African populations, and almost 100% of children are EBV seropositive by 3 years of age (11, 12). Primary infection during childhood is typically asymptomatic, whereas infection in young adults can result in acute infectious mononucleosis (AIM), a self-limited lymphoproliferative disorder. To date, most immunologic studies of EBV infection are based on healthy seropositive adults or cases of AIM among adolescents in Europe or the United States (13). Collectively, these studies show that CD8+ cytotoxic T lymphocytes (CTL) are necessary for immune surveillance and control of persistent EBV infection (14, 15). The CTL response to EBV is directed against an array of antigens expressed during the lytic and latent phases of the viral life cycle (13–15), and control is associated with HLA class I-restricted gamma interferon (IFN-γ) responses (16). Previous studies have also demonstrated phenotypic and functional heterogeneity among EBV-specific CD8+ T-cell populations (17). However, little is known about these cells when primary EBV infection occurs during infancy or early childhood.

Beyond the early studies that revealed a geographic overlap between eBL and areas of intense, perennial Plasmodium falciparum malaria transmission (regions of malaria holoendemicity) (8, 9), the malaria-driven mechanisms that contribute to eBL pathogenesis remain obscure. In these regions of equatorial Africa, more than 80% of children are chronically or repeatedly infected with P. falciparum malaria by 5 years of age, and initial malaria exposure occurs within the first few months of life (18, 19). It is established that malaria parasites modulate and evade the host immune system (20). Indeed, these properties underlie the hypothesis that P. falciparum malaria suppresses immunity to EBV during coinfection. In the early 1980s, a series of seminal studies demonstrated that lymphocytes from malaria-infected individuals were unable to control the proliferation of EBV-transformed B cells in relatively crude regression assays (21, 22). Although these observations suggest that P. falciparum malaria infection disrupts EBV-specific immunity, the effector cells or mediators responsible for controlling EBV-infected B-cell growth were not identified, and overall immune competence was not assessed in the small number of individuals studied. More recently, an age-related deficiency in IFN-γ recall responses to EBV lytic and latent antigens was demonstrated in children (i.e., 5 to 9 years of age) with ho-
loendemic malaria exposure compared to those from an area of malaria hypoendemcity (23). In addition, EBV load in African children correlates with malaria exposure (24, 25), further implicating coinfection as a risk factor for eBL tumorigenesis. However, it remains unclear how *P. falciparum* malaria might potentiate a deficit in EBV-specific CD8 T-cell immunity and thus contribute to eBL lymphomagenesis.

Two mutually compatible theories have been proposed to explain the relationship between EBV and *P. falciparum* malaria in the etiology of eBL (26). The first suggests that malaria coinfection increases the number of latently infected B cells by inducing polyclonal B-cell expansion and consequent lytic EBV reactivation (27). In turn, the greater precursor frequency of EBV-infected B cells increases the likelihood of c-myc translocation, which is a hallmark of all BL tumors (28). The second theory argues that EBV-specific T-cell responses are selectively altered during malaria coinfection, either as a cause or consequence of enhanced EBV replication, leading to impaired viral control and/or immune surveillance (29, 30).

To provide direct evidence for the role of altered EBV-specific T-cell immunity in eBL, we used polychromatic flow cytometry to characterize EBV-specific CD8 T cells in a unique, well-characterized cohort of individuals with divergent malaria exposure. Specifically, 16-parameter flow cytometry panels were developed to quantify differentiation, exhaustion, senescence, and homeostatic potential within six distinct EBV-specific CD8 T-cell populations and one CMV-specific CD8 T-cell population, all restricted by HLA A*0201, and within the CD8 T-cell compartment as a whole. By virtue of the highly multiplexed nature of our measurements, cellular characteristics could be defined in exquisite detail. Furthermore, we employed recently developed data analysis strategies to deconvolute these complex flow cytometry data sets. Accordingly, probability binning (31) and frequency difference gating (32) revealed dramatic differences between cell populations in multidimensional space that eluded conventional analytical approaches. Together, our analyses reveal an immunologic aberration confined to the EBV-specific CD8 T-cell compartment that is associated with *P. falciparum* malaria exposure and further illuminate the relationship between holoendemic malaria and eBL.

**MATERIALS AND METHODS**

**Study participants.** Approval for this study was obtained from the Ethical Review Committee at the Kenya Medical Research Institute (KEMRI) and the Institutional Review Board for Human Studies, University Hospitals of Cleveland, Case Western Reserve University; the latter was the institutional affiliation for A.M.M. at the time of this study. Written informed consent was obtained in all cases from study participants or parents of minors.

Participant recruitment and sample collection were conducted in two epidemiologically distinct areas of western Kenya: (i) Nyanza Province, Kisumu District, in the sublocation of Kayanyangi; (ii) Rift Valley Province, Nandi District, in the sublocation of Kipsamoite. The first study site is situated on the shore of Lake Victoria, 10 km west of Kisumu; malaria transmission in this area is holoendemic (i.e., intense and perennial). The second study site is located in the highlands, 150 km northeast of Kisumu; malaria transmission in this area is hypoendemic (i.e., sporadic with periodic outbreaks of malaria morbidity in a population with low parasite prevalence) (33).

For this study, we collected blood samples from individuals with known HLA class I genotypes and divergent malaria exposure histories who experienced primary EBV infection prior to 3 years of age (25, 34). No evidence of fever or anemia was present in these individuals at the time of venous blood sampling; thus, any cases of malaria parasitemia were asymptomatic.

**Sample collection.** Venous blood was collected in sodium-heparinized tubes and transported to the KEMRI Center for Global Health Research in Kisumu for processing within 3 h. Peripheral blood mononuclear cells (PBMCs) were separated from whole blood by Ficoll-Hypaque density gradient centrifugation, counted, and then resuspended in freezing medium comprising 90% heat-inactivated, filter-sterilized fetal bovine serum and 10% dimethyl sulfoxide (Sigma). Aliquots of 6 million cells/ml per vial were cooled overnight at a rate of −1°C per minute prior to long-term storage in vapor-phase liquid nitrogen. Samples were transported with a Centers for Disease Control and Prevention (CDC) import permit using an MVE IATA-approved vapor-phase shipper to maintain temperatures below −180°C en route.

**Tetrameric peptide-HLA A*0201 complexes.** Soluble, bivalent tetramer-PELA A*0201 (pHLA A*0201) monomers were produced and tetramerized with fluorochrome-conjugated streptavidin as described previously (35). The following peptides (≥95% purity; BioSynthesis) were used for monomer production: (i) GLCTLVAML (GLC; EBV BMFI, residues 280 to 288), (ii) YVLDHLLIVV (YVL; EBV BRF1, residues 109 to 117), (iii) YLLEMLWRL (YLL; EBV LMP1, residues 125 to 133), (iv) FLYALALLL (FLY; EBV LMP2, residues 356 to 364), (v) CGLGGLTMV (CLG; EBV LMP2, residues 426 to 434), (vi) LLDVFRMGV (LLD; EBV EBNA3C, residues 284 to 293), and (vii) NVLPVMATV (NLV; CMV pp65, residues 495 to 503).

**Flow cytometry and data analysis.** Samples were thawed and then stained with pHLA A*0201 tetramers and a panel of fluorochrome-conjugated monoclonal antibodies as described previously (36). Data were analyzed using FlowJo version 9.2 (Treestar Inc.), which contains platforms for probability binning/frequency difference gating and SPICE (31, 32). Adjustments for multiple comparisons were not performed due to considerable overlap between cell subsets (37). Instead, a more stringent threshold for significance was set (*P < 0.01*).

**RESULTS**

Development and validation of polychromatic flow cytometry panels for the evaluation of EBV-specific CD8 T-cell immunity. To evaluate the effects of *P. falciparum* malaria on EBV-specific CD8 T-cell immunity, we conducted a cross-sectional study of children and adults from regions where malaria is either holoendemic or hypoendemic (34). Peripheral blood samples were examined using two 16-parameter flow cytometry panels to determine antigen specificity and molecular coexpression patterns at the single-cell level (Fig. 1A and B). Reagents and staining panels were developed as described previously (36). Assay procedures were validated for rare event analysis using a set of test samples (*n* = 28), with previously defined epitope-specific CD8 T-cell responses corresponding to the HLA A*0201-restricted specificities examined in this study. All pHLA A*0201 tetramers were quality controlled using PBMCs from HLA A*0201− donors and HLA A*0201+ donors with no evidence of CMV or EBV infection; no nonspecific staining was observed in either setting. Individual samples were also evaluated repeatedly to ensure procedural reproducibility. This approach enabled us to interrogate EBV-specific CD8 T-cell populations with unprecedented precision and depth.

**Immunodominance profiles among HLA A*0201-restricted EBV-specific CD8 T-cell populations.** Initially, we examined the frequencies of CD8 T cells directed against six different HLA A*0201-restricted EBV-derived epitopes. Four of these epitopes (YLL, FLY, CLG, and LLD) are derived from latently expressed proteins, and two (GLC and YVL) are derived from lytic-phase
proteins. There are dramatic differences between these epitopes with respect to the milieu and kinetics of antigen presentation to the immune system. Remarkably, however, we found no significant differences between the frequencies of CD8\(^+\) T cells targeting latent and lytic epitopes (Fig. 2A). Moreover, no single epitope-specific CD8\(^+\) T-cell population was consistently immunodominant across the cohort as a whole. These observations held after stratification of study participants for malaria exposure (data not shown) and age (Fig. 2B). The similar distribution of these CD8\(^+\) T-cell populations across all age groups suggests that the nature of the EBV-specific response is set early in life and changes little over time.

**Distinct phenotypes characterize different EBV-derived epitope-specific CD8\(^+\) T-cell populations.** Next, we examined the phenotypic properties of EBV-specific CD8\(^+\) T cells across the entire study population regardless of malaria exposure. Minor phenotypic differences were observed in some EBV-specific CD8\(^+\) T-cell populations by age. However, these were observed only in exceedingly rare cell populations, and there was no clear relationship between the populations that differed; for example, expression of the senescence marker CD57 was not consistently lower in the youngest individuals (data not shown). On this basis, we subsequently aggregated all study participants irrespective of age. In order to minimize the contribution of nonspecific events (“noise”), we stratified the data set according to the number of cells collected on the flow cytometer and selected only those populations that achieved a statistically rigorous minimum number of antigen-specific events (Fig. 3; see also Fig. S1 in the supplemental material).

Given the large number of measured parameters and the challenges associated with the interpretation of such complex multivariate data, our initial analyses focused on the differentiation stage of EBV-specific CD8\(^+\) T cells, defined by the coordinate expression of CD45RO, CCR7, CD27, CD127, and CD57 (see Fig. S2 in the supplemental material). Although CD57 typically defines senescent cells (38), we have observed in previous analyses that expression is not exclusively linked to advanced differentiation (39), which complicates the categorization and nomenclature of...
maturational subsets. In this light, we examined other phenotypic categories separately within the CD57$^+$ and CD57$^-$ subsets. This analysis showed that many of the CD57$^-$ cells directed against both latent and lytic EBV-derived epitopes were central memory-like cells, expressing various combinations of CD45RO, CCR7, CD27, and CD127. Notably, EBV-specific CD8$^+$ T cells with a naïve-like phenotype (CD45RO$^-$ CCR7$^+$ CD27$^+$ CD127$^+$ CD57$^-$) were frequently observed, in line with previous reports in other systems (40, 41). This is consistent with the recent description of stem cell-like memory T cells, which express many other surface markers typically associated with antigen-inexperienced naive cells and likely represent a very early stage of memory T-cell differentiation (42).

In further analyses, we examined the phenotypic properties of EBV-specific CD8$^+$ T-cell populations in terms of homeostatic potential (CD127), exhaustion (PD-1), and senescence (CD57). Clear phenotypic differences were observed between CD8$^+$ T-cell populations that targeted distinct EBV-derived antigens (Fig. 4). Thus, substantial diversity exists within the CD8$^+$ T-cell response to EBV, which likely reflects the complexity of antigen expression and immunosurveillance. Despite this complexity, however, CD8$^+$ T cells directed against the latent antigens FLY, CLG, and YLL generally expressed markers consistent with the retention of homeostatic potential in the relative absence of exhaustion/senescence. In contrast, CD8$^+$ T cells directed against the lytic antigens GLC and YVL more frequently displayed an exhausted/senescent phenotype lacking homeostatic potential. Interestingly, CD8$^+$ T cells specific for the latent epitope LLD (EBNA3C) exhibited similar proportions of exhausted/senescent cells compared to the antilytic populations, thereby suggesting that the biology of EBNA3C expression may differ from that of other latent antigens.

**Phenotypic differences between EBV-specific CD8$^+$ T cells associated with malaria endemcity.** Next, we stratified study participants by malaria exposure and examined the phenotype of EBV-specific, CMV-specific, and total CD8$^+$ T-cell populations. Malaria exposure was determined as described previously (43), based on well-defined epidemiologic parameters, and confirmed with clinical or laboratory data. Specifically, blood-stage infection in children occurs with $<10\%$ prevalence in regions where malaria is hypoendemic, whereas childhood infection rates in regions

![FIG 2](image.png)
of malaria holoendemicity are consistently >50% throughout the year (44). Individuals from regions with strikingly divergent cumulative malaria exposure (hypendemic versus holoendemic) were compared for this study, and two distinct approaches to data analysis were employed.

FIG 3 Distribution of phenotypes within each antigen-specific CD8+ T-cell population, grouped by the number of events collected on the flow cytometer. Data are shown for CD8+ T cells directed against EBV-derived latent antigens. Pies depict data for samples where 1 to 5, 6 to 10, 11 to 20, or 20+ antigen-specific events were collected. Slices within each pie represent the proportion of CD8+ T cells expressing a particular combination of the markers analyzed. Pies for each event category were compared by permutation analysis to determine if the distribution of phenotypes differed according to the number of events collected. For example, among CD8+ T cells specific for the FLY epitope, the distribution of phenotypes observed when 1 to 5 events were collected (pie 1) was significantly different from the distribution observed when 20+ events were collected (P = 0.037). Similarly, the distribution differed significantly when 6 to 10 (pie 2) and 11 to 20 (pie 3) events were collected (P = 0.074 and P = 0.009, respectively). Thus, the phenotype of FLY-specific CD8+ T cells was not considered in samples with less than 20 such events. For all other latent-phase antigens, the distribution of phenotypes was not significantly different when more than 5 events were collected; these pies are highlighted with a black border. Therefore, for these specificities, only study participants with fewer than 5 antigen-specific events were excluded from the analysis. The numbers in the center of the pie represent the number of study participants with the specific phenotype.

FIG 4 Relative frequencies of EBV-specific CD8+ T cells expressing various combinations of CD45RO, CD57, CD127, and PD-1 within each tetramer+ population. Data are arranged in order of increasing differentiation, such that the bar charts on the left side of the figure depict CD127+ events and the bar charts on the right side of the figure depict CD127− events. Within each section, increasingly exhausted or senescent populations are found to the right. For example, CD8+ T cells expressing CD127 without markers of exhaustion or senescence are represented on the far left; these are most evident within the populations specific for YLL, LLD, GLC, and YVL. In contrast, exhausted or senescent CD8+ T cells without homeostatic potential are represented on the far right; these are most evident within the populations specific for LLD, GLC, and YVL. Bars represent median values for the depicted cell types (x axis key) within CD8+ T-cell populations specific for the listed EBV-derived epitopes (right y axis labels). Hatched green (latent) and red (lytic) lines connect the medians for each distinct phenotype to aid visual comparison within each specificity.

In the first approach, the proportion of CD8+ T cells that expressed each single marker was compared across both study groups and any statistically significant differences were noted. Subsequently, the proportion of CD8+ T cells that expressed every combination of two, three, four, or five markers was tested across the study groups. These analyses were conducted for each EBV-specific CD8+ T-cell population and for the CMV-specific and total CD8+ T-cell populations. Figure 5 shows examples of the differences observed when dual-marker expression was examined within the EBV-derived GLC epitope-specific CD8+ T-cell population. Study participants from regions where malaria is hypendemic (blue) had more CD45RO−CD127+ and CD45RO+CD57− T cells than individuals from holoendemic (red) areas (P = 0.003 and P = 0.009, respectively) and fewer CD45RO−CD127− T cells (P = 0.008).
Similar analyses were performed for all combinations of phenotypic markers. Tables 1 and 2 list the phenotypically defined CD8<sup>+</sup> T-cell subsets across all specificities that were significantly elevated in study participants from regions where malaria is hypoendemic and holoendemic, respectively. In general, the numbers of cell types associated with less differentiated memory T-cell populations (e.g., CD127<sup>+</sup> cells) were elevated in individuals from regions with hypoendemic malaria, whereas the numbers of more differentiated (e.g., CD27<sup>+</sup>, CCR7<sup>+</sup>) cells were elevated in those exposed to holoendemic malaria. Although no adjustments were made for multiple comparisons, several comparisons were highly significant (P < 0.01), and taken together, the data present a uniform picture. Specifically, individuals with chronic/recurrent malaria exposure have elevated levels of highly differentiated EBV-specific CD8<sup>+</sup> T cells compared to those of individuals with little or no malaria exposure. Importantly, no statistically significant phenotypic differences between study groups were observed within either the CMV-specific or the total CD8<sup>+</sup> T-cell compartment. Horizontal bars represent median values, and boxes delineate the 25th and 75th percentiles. Box colors depict malaria endemicity (hypoendemic, blue; holoendemic, red).

**TABLE 1** Phenotypically defined CD8<sup>+</sup> T-cell subsets that differed by malaria exposure. Horizontal bars represent median values, and boxes delineate the 25th and 75th percentiles. Box colors depict malaria endemicity (hypoendemic, blue; holoendemic, red).

<table>
<thead>
<tr>
<th>Epitope</th>
<th>CD45RO&lt;sup&gt;+&lt;/sup&gt;</th>
<th>CD45RO&lt;sup&gt;+&lt;/sup&gt;,CCR7&lt;sup&gt;+&lt;/sup&gt;</th>
<th>CD57&lt;sup&gt;-&lt;/sup&gt;,CD127&lt;sup&gt;-&lt;/sup&gt;</th>
<th>CD57&lt;sup&gt;-&lt;/sup&gt;,CD127&lt;sup&gt;-&lt;/sup&gt;,CD45RO&lt;sup&gt;+&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>CLG</td>
<td>0.003</td>
<td>0.009</td>
<td></td>
<td></td>
</tr>
<tr>
<td>GLC</td>
<td>0.808</td>
<td>0.151</td>
<td>0.230</td>
<td></td>
</tr>
<tr>
<td>CMV</td>
<td>0.151</td>
<td>0.837</td>
<td>0.250</td>
<td></td>
</tr>
<tr>
<td>YVL</td>
<td>0.031</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LLD</td>
<td>0.003</td>
<td>0.009</td>
<td></td>
<td></td>
</tr>
<tr>
<td>FLY</td>
<td>0.003</td>
<td>0.009</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**FIG 5** Representative examples of phenotypic patterns within EBV-derived GLC-specific CD8<sup>+</sup> T-cell populations that differed by malaria exposure. Horizontal bars represent median values, and boxes delineate the 25th and 75th percentiles. Box colors depict malaria endemicity (hypoendemic, blue; holoendemic, red).

Malaria-associated phenotypic differences in EBV-specific CD8<sup>+</sup> T-cell populations pinpointed in multidimensional space. The manual gating approach described above has two significant limitations. First, any population differences that are only manifest in multiple dimensions will be more difficult to resolve, and appropriate gating strategies that discriminate such subsets optimally may not be evident from examination of the dual-parameter histograms. Second, a large number of phenotypic populations are necessarily identified, leading to loss of significance when correcting for multiple comparisons. To overcome these limitations, we adopted a second approach to confirm and extend our observations. Specifically, we used frequency difference gating (32), a method that does not rely on serial bivariate analysis and therefore has the potential to reveal population differences that occur across multidimensional space.

Using this approach, we initially examined EBV-derived GLC epitope-specific CD8<sup>+</sup> T cells (Fig. 6). The results confirmed and extended our previous analysis, in which CD8<sup>+</sup> T cells with a central memory-like phenotype (CD45RO<sup>+</sup>, CD127<sup>+</sup> and CD45RO<sup>+</sup>,CD57<sup>+</sup>,CD127<sup>+</sup>) were associated with hypoendemic malaria exposure (Table 1 and Fig. 5). Frequency difference gating revealed a phenotypic space defined by the CD45RO<sup>+</sup>,CD27<sup>+</sup>,CCR7<sup>-</sup>,CD127<sup>-</sup>,CD57<sup>-</sup>,PD-1<sup>+</sup> expression pattern that contained 9.9% of GLC-specific CD8<sup>+</sup> T cells from individuals with hypoendemic malaria exposure and only 2.9% of the corresponding CD8<sup>+</sup> T-cell population from individuals with holoendemic malaria exposure (P < 0.05). Thus, as in the previous analysis, highly differentiated cells were enriched within the GLC-specific CD8<sup>+</sup> T-cell compartment in individuals with holoendemic malaria exposure.

Examination of the other specificities revealed more striking differences (Fig. 6). In individuals with hypoendemic malaria exposure, frequency difference gating identified an EBV-derived FLY epitope-specific CD8<sup>+</sup> T-cell population that was highly enriched for central memory-like cells compared to the holoendemic group (27.7% versus 5.7%, P < 0.0001). This difference was not apparent with the first approach to data analysis. Furthermore, we found that individuals with hypoendemic malaria exposure also had higher levels of central memory-like cells when responses to any latent epitope were pooled (P < 0.001). Differences according to malaria endemicity across pooled CD8<sup>+</sup> T-cell populations specific for EBV-derived latent epitopes are summarized in Fig. S3 in the supplemental material. No statistically significant
phenotypic differences between study groups were observed for pooled EBV-derived lytic epitope-specific, CMV-specific, or total CD8\(^+\) T cells.

DISCUSSION
This is the first comprehensive study of EBV-specific CD8\(^+\) T-cell immunity in individuals, including children, who reside in a region of malaria endemicity that imparts a significantly elevated risk for the development of eBL. Accordingly, the results presented here reveal previously unappreciated characteristics of the human immune response to this common, chronic viral infection and the potential role of altered immunity in susceptibility to cancer.

For epitopes presented by HLA A*0201, we found that EBV-specific CD8\(^+\) T cells directed against lytic and latent antigens were equally represented across the study population as a whole, even though the proteins are presumably expressed with different kinetics and under different conditions in vivo. We also found that the signatures of EBV-specific T-cell immunity are largely set following primary infection and appear to change little over time, as no phenotypic differences were linked to age. Furthermore, although the immune system is thought to "see" latent antigens constantly, CD8\(^+\) T cells directed against these epitopes generally retained homeostatic potential and were rarely exhausted/senescent. In contrast, CD8\(^+\) T cells directed against lytic antigens, the expression of which is thought to be episodic and infrequent, were more likely to display an exhausted/senescent phenotype lacking homeostatic potential. Given the prevalent hypothesis that such differences are related to antigen exposure, these observations suggest the occurrence of frequent EBV reactivations in the overall study population.

### TABLE 2
Phenotypically defined CD8\(^+\) T-cell subsets that were significantly elevated in regions with holoendemic malaria

<table>
<thead>
<tr>
<th>Epitope</th>
<th>CD45RO(^+)/CD127(^-)</th>
<th>CD45RO(^+)/CCR7(^-)</th>
<th>CD45RO(^+)/CD127(^-)/PD1(^+)</th>
<th>CD45RO(^+)/CD127(^-)/CD27(^-)/CD57(^-)</th>
<th>CD27(^-)/CD127(^-)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CLG</td>
<td>0.015</td>
<td>0.087</td>
<td>0.016</td>
<td>0.008</td>
<td>0.003</td>
</tr>
<tr>
<td>GLC</td>
<td>0.566</td>
<td>0.621</td>
<td>0.791</td>
<td>0.446</td>
<td>0.296</td>
</tr>
<tr>
<td>LLD</td>
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<td>0.013</td>
<td>0.988</td>
<td>0.650</td>
<td>0.013</td>
</tr>
<tr>
<td>YLL</td>
<td>0.383</td>
<td>0.608</td>
<td>0.192</td>
<td>0.192</td>
<td>0.192</td>
</tr>
<tr>
<td>CMV</td>
<td>0.566</td>
<td>0.621</td>
<td>0.791</td>
<td>0.446</td>
<td>0.296</td>
</tr>
<tr>
<td>CD8</td>
<td>0.520</td>
<td>0.710</td>
<td>0.264</td>
<td>0.446</td>
<td>0.296</td>
</tr>
</tbody>
</table>

**FIG 6** Frequency difference gating reveals CD8\(^+\) T-cell subsets that are altered by malaria exposure. A subset of CD45RO\(^+\)/CD127\(^-\)/CD27\(^-\)/CCR7\(^-\) PD-1\(^+\) (central memory-like) EBV-derived GLC-specific (top) and FLY-specific (middle) CD8\(^+\) T cells is diminished in individuals from regions with holoendemic malaria. In the pooled data from all CD8\(^+\) T cells that target latent antigens, a subset that includes central memory-like cells is dramatically reduced in individuals with holoendemic malaria exposure.
Interestingly, CD8\(^+\) T cells directed against the latent epitope LLD (EBNA3C) displayed phenotypic characteristics similar to those observed for the corresponding lytic antigen-specific populations; this finding is in keeping with previous studies that detail the typical immunodominance of EBNA3C (13) and suggests quantitatively enhanced or qualitatively distinct presentation of this protein among the latent antigens studied.

An intriguing observation from our data is the identification of EBV-specific CD8\(^+\) T cells with a naïve-like (CD45RO\(^-\) CD27\(^-\) CCK7\(^-\) CD127\(^-\) CD57\(^-\)) phenotype. Within the context of this study, the functional capacity of these EBV-experienced naïve-like CD8\(^+\) T cells is as yet unknown. It is likely that such cells populate a very early differentiated memory pool (41) and perhaps even constitute an antigen-specific "stem cell-like memory" population (42). However, further work is required to clarify the biological significance of this observation.

The features of EBV-specific CD8\(^+\) T-cell immunity described above were independent of malaria exposure yet were observed in individuals who typically acquire the virus during early childhood (25). Nonetheless, malaria coinfections were associated with additional phenotypic differences between CD8\(^+\) T-cell populations specific for latent epitopes and the lytic epitope GLC (BMLF1), which displayed characteristics of a more differentiated stage in individuals from regions of malaria holoendemicity. Thus, despite the fact that all study participants were healthy, distinct malaria-associated irregularities in EBV-specific CD8\(^+\) T-cell immunity were apparent. Overall, these results support a combined role for early primary EBV infection and cumulative malaria exposure in EBV-specific T-cell immune dysregulation.

Due to the cross-sectional nature of this study and the relatively low incidence of eBL, we were not able to address directly the causal relationship between EBV, \textit{P. falciparum}, malaria, and eBL. However, our data offer new evidence that addresses two longstanding questions in the field.

First, as with any immunologic study in humans that attempts to inform pathogenic mechanisms, it is critical to understand whether the observed deficits represent cause or effect in the disease process. Our previous study revealed a deficiency in IFN-γ production by EBV-specific CD8\(^+\) T cells in children residing in a region with holoendemic malaria (23). However, the events that precede this immune deficiency are unclear. On that basis, we postulated that more detailed studies of EBV-specific CD8\(^+\) T-cell immunity in areas with divergent malaria endermicity might provide a better window into the complex etiology of eBL. Given the remarkable diversity of T cells, simple immunophenotypic analyses could miss fine subsets critical to the understanding of this relationship. In addition, although EBNA1 is the only EBV-derived protein expressed in eBL (45), the risk for disease may be set when T cells specific for a wide variety of lytic and latent antigens are generated. For these reasons, we used a sophisticated flow cytometric approach optimized for rare event analysis to demonstrate that a loss of central memory-like EBV-specific CD8\(^+\) T cells is associated with malaria exposure. Such immunologic perturbations within the EBV-specific CD8\(^+\) T-cell compartment could predispose to the functional deficits that accompany eBL (46).

Second, our study addresses the long-standing question of whether eBL is associated with a generalized, malaria-induced suppression of T-cell immunity (21, 22, 47). Our data demonstrate that EBV-specific, but not CMV-specific or total, CD8\(^+\) T-cell populations show significant differences associated with malaria exposure. These findings are consistent with previous studies showing that T-cell responses to malaria antigens and nonspecific mitogens are equally robust across study populations and age groups (46). Thus, malaria exposure uniquely impacts the EBV-specific CD8\(^+\) T-cell compartment, which argues against a role for generalized immune suppression in the pathogenesis of eBL.

In the context of a cross-sectional study, however, it is not possible to discern how the observed immunologic differences arise. Recently, a malarial antigen capable of inducing the reactivation of latent EBV was identified (48), possibly providing a mechanistic explanation for our findings. In this scenario, repeated malaria infections induce EBV reactivation, thereby seeding new, potentially highly activated and dividing B cells (27), which consequently increase the risk of B-cell transformation. As EBV-specific CD8\(^+\) T cells attempt to control these frequent reactivations, their differentiation is accelerated and the pool of central memory cells for such specificities is selectively depleted. This possibility is supported by seroprofiling studies, which demonstrate that elevated antibody titers to viral capsid antigen (VCA) and the Z Epstein-Barr replication activator (ZEBRA) protein are associated with holoendemic malaria exposure (49). An elevation in these antibody titers signifies viral reactivation (50), and higher VCA antibody titers were observed prior to the development of eBL in a study of Ugandan children (51, 52). In addition, malaria infection in concert with EBV reactivation may modulate elements of the innate immune system that influence T-cell maturation and differentiation (53, 54).

In summary, we have demonstrated that a select set of EBV-derived antigen-specific CD8\(^+\) T cells is altered in individuals coinfected with \textit{P. falciparum} malaria and at increased risk for eBL. These findings inform our studies of eBL pathogenesis and suggest that EBV-targeted immune interventions in the setting of holoendemic malaria might hold promise for the prevention of this devastating pediatric malignancy.

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REFERENCES

41. Pirillo E, Kimmel R, Chelimo K, Midi dorp JM, Odada PS, Ploutz-


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