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“INFLAMMESCENT” CX3CR1+CD57+ CD8 T-CELLS ARE GENERATED AND EXPANDED BY IL-15

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ABSTRACT

HIV infection is associated with an increase in the proportion of activated CD8 memory T cells (Tmem) that express CX3CR1, but how these cells are generated and maintained in vivo is unclear. We demonstrate that increased CX3CR1 expression on CD8 Tmem in people living with HIV (PLWH) is dependent on coinfection with human cytomegalovirus (CMV), and CX3CR1+ CD8 Tmem are enriched for a putatively immunosenescent CD57+CD28- phenotype. The cytokine IL-15 promotes the phenotype, survival, and proliferation of CX3CR1+CD57+ CD8 Tmem in vitro, whereas TCR stimulation leads to their death. IL-15-driven survival is dependent on STAT5 and Bcl-2 activity, and IL-15-induced proliferation requires STAT5 and mTORC1. Thus, we identify mechanistic pathways that could explain how “inflammascent” CX3CR1+CD57+ CD8 Tmem dominate the overall memory T cell pool in CMV-seropositive PLWH and that support reevaluation of immune senescence as a nonproliferative dead-end.
INTRODUCTION

High expression of the fractalkine receptor (CX3CR1) on CD8 T cells identifies a population of long-lived effector memory cells with lytic granules and cytolytic capacity (1-3) that do not recirculate through lymphatics (4). Cytolytic CD8 memory T cells (Tmem) also have shortened telomeres and impaired proliferative responses to antigen – a condition termed immune senescence characterized by high expression of CD57 and low expression of the costimulatory receptor CD28 (5-7). CD57+CD28- CD8 T cells are enriched in patients with chronic viral infections, such as human immunodeficiency virus (HIV), and cytomegalovirus (CMV), and are particularly enriched in people with HIV/CMV coinfection (8, 9), who comprise greater than 90% of those infected with HIV (10, 11).

In people living with HIV infection (PLWH), increased CD8 T cell numbers with resultant CD4/CD8 inversion is strongly linked to the development of non-AIDS comorbidities (12, 13). When HIV replication is controlled by antiretroviral therapy (ART), CD4 T cell numbers often recover to near normal levels, but persistently elevated CD8 T cell numbers maintain a low CD4/CD8 ratio (14). CD8 T cell expansion is likely driven by several factors, including pro-inflammatory cytokines such as IL-15, which is elevated during chronic HIV infection (15, 16). We recently showed that CD8 T cell expansion in ART-treated PLWH is linked to CMV coinfection (17). Many of the expanded CD8 T cells are likely to be CMV-reactive (1, 18-20), cytolytic (3, 21), and express CX3CR1 (22, 23), but what drives the sustained numbers of circulating CD8 T cells with ostensibly poor replicative capacity is not well understood.

Here we characterized peripheral blood CD8 T cells in PLWH. The majority of CX3CR1+ CD8 Tmem are CD57+CD28- and are highly enriched in CMV-seropositive (CMV+) donors. These cells have high expression of the transcription factor T-bet and the cytolytic enzymes granzyme B and perforin, and so are poised for effector function. *In vitro* IL-15 exposure promoted the development of the CX3CR1+CD57+ phenotype, induced the proliferation of CX3CR1+CD57+ CD8 T cells, and sustained their viability. IL-15-induced proliferation required STAT5 phosphorylation and mTORC1 signaling, whereas IL-15-enhanced viability required phosphorylation of STAT5 and Bcl-2 activity. Thus, our data call into question the premise of immune
senescence in this durably expanded cell population and provide a mechanistic link between inflammation and the expansion, activation, and persistence of CX3CR1+ CD8 T cells in ART-treated HIV infection.
RESULTS

CX3CR1+ CD8 T cells are mostly CD57+CD28- and are enriched in CMV-seropositive PLWH

Earlier, we characterized a population of CCR7-CX3CR1+ CD8 Tmem that is enriched in the peripheral blood of ART-treated HIV-infected individuals (23). We demonstrated that CMV coinfection was necessary for CD8 T cell expansion in PLWH (17). Here, we found that CCR7-CX3CR1+ CD8 Tmem were significantly expanded in CMV+ PLWH (Figure 1A). Consistent with previous reports (1, 24), we found that a majority of CMV pp65-specific cells expressed CX3CR1 and pp65-specific cells were enriched among CCR7- cells that expressed CX3CR1 (Figure S2A). This was not seen among HIV gag-specific CD8 T cells (Figure S2B). Regardless of CMV serostatus, however, CCR7-CX3CR1+ CD8 Tmem were enriched for CD57+CD28- cells (Figure 1B). CMV+ PLWH donors also had an elevated proportion of CX3CR1hiCD27- CD8 Tmem (Figure S1C) that are thought to be retained in circulation (2, 4, 24). As CX3CR1+CD57+CD28- CD8 Tmem had the highest CX3CR1 density (Figure S1D), CX3CR1hi cells and CX3CR1+CD57+CD28- cells appear to be largely the same population.

We next asked whether HIV infection was necessary to generate the CX3CR1+CD57+CD28- phenotype on CD8 Tmem. This was not the case as CX3CR1+CD57+CD28- CD8 T mem were expanded in HIV-uninfected CMV+ donors when compared to findings among HIV-negative, CMV-seronegative (CMV-) donors (Figure 1C). Using t-stochastic neighbor embedding (t-SNE) analysis of CD8 T cells from HIV-uninfected and infected donors stratified by CMV serostatus (Figure 1D), we found that CX3CR1+CD57+CD28- Tmem were enriched in CD8 T cells from CMV+ donors, regardless of HIV infection status, demonstrating that CMV infection alone is sufficient to promote a relatively expanded CX3CR1+CD57+CD28- phenotype on CD8 Tmem.

CX3CR1+CD57+CD28- CD8 Tmem have an effector/cytolytic cell phenotype

Among CCR7-CX3CR1+ CD8 Tmem, CD57+CD28- cells had the highest expression of the effector-phenotype transcription factor T-bet and lowest expression of the memory-phenotype transcription factor eomesodermin (Eomes), resulting in a pro-effector T-bethiEomeslo phenotype (Figure S2A). Consistent with previous reports (21, 25), CD57+CD28- cells were enriched for the cytolytic enzymes granzyme B and perforin (Figure S2B). In general, CX3CR1+ CD8 T cells have greater expression of the checkpoint receptor PD-1 than do CX3CR1-
CD8 T cells (23). However, among CX3CR1+ CD8 Tmem, CD57+CD28- cells had the lowest PD-1 expression (Figure S2C), suggesting that they are less likely to be regulated by PD-1/PD-L1 interactions, and may be distinct from the recently described transitory PD-1+CX3CR1+CD101-TIM-3+ CD8 T cell population that responds to PD-1 blockade in mice (26). Since CD57+ CD8 Tmem lack expression of CD28, we next stimulated CD8 Tmem through the T cell receptor (TCR) without anti-CD28 costimulation and measured cytokine synthesis. CD57+ CD8 Tmem exhibited a significantly different cytokine profile than CD57- CD8 Tmem in response to TCR stimulation in the absence of costimulation (Figure S2D), and this differential cytokine response was characterized by elevated MIP-1β expression (Figure S2E), especially by cells that synthesized only MIP-1β (Figure S2F). Thus, CX3CR1+CD57+CD28- CD8 Tmem have an effector cell transcriptional profile, are poised for cytotoxicity, and are polyfunctional, with enriched MIP-1β expression.

Given that CD57 expression is associated with shorter telomeres, increased activation induced cell death, and poor proliferative potential in response to antigen-mediated stimulation (7, 27), we wondered how CD57+ CD8 Tmem accumulate and persist in CMV+ PLWH. To address this, we examined CCR7-CX3CR1+ CD8 Tmem for intracellular expression of the pro-survival factor Bcl-2 (Figure 2A) and the master transcriptional regulator c-myc (Figure 2B), which is important for stimulating cell cycle progression. Both CD57+CD28- and CD57-CD28+ cells had similar levels of Bcl-2 and c-myc expression, and surprisingly, CD57-CD28- cells had reduced Bcl-2 and c-myc levels when compared to levels in the other two groups – although the functional significance of these differences is currently unknown. We next measured intracellular Ki67 expression to determine what proportion of these cells were actively in cell cycle. Although a lower proportion of Ki67+ CCR7-CX3CR1+ CD8 Tmem were CD57+CD28- than among Ki67- cells, the majority of CCR7-CX3CR1+ CD8 Tmem in cell cycle were CD57+CD28- (Figure 2C). Taken together, our data suggest that these putatively senescent, effector-like CX3CR1+CD57+CD28- CD8 Tmem are in cell cycle or were recently in cell cycle in vivo and are protected from apoptosis.

**IL-15 promotes the generation of CX3CR1+CD57+ CD8 Tmem**

We next asked what might be driving the generation and expansion of CX3CR1+CD57+CD28- CD8 Tmem, with a specific focus on TCR signals and common γ-chain cytokines. IL-15 exposure in particular has been
shown to promote a CD57+CD28- phenotype on CD8 T cells in vitro (28, 29), enhance mitochondrial biogenesis (30), induce oxidative phosphorylation and c-myc expression (31), and promote antigen-independent proliferation without telomere loss in CD8 Tmem (32, 33). In addition, our group has identified IL-15 as an important driver of bystander CD8 T cell activation and expansion in chronic untreated HIV infection (15). To determine cytokine susceptibility, we measured CCR7-CX3CR1+ CD8 Tmem subpopulations for expression of common γ-chain cytokine receptors (Figure S3A). None of the subpopulations expressed much CD25, the α-chain of the IL-2 receptor, and so are not likely to respond to low-level IL-2 exposure. All three subpopulations expressed robust levels of CD122, the IL-2/IL-15 receptor β-chain, suggesting they could be responsive to IL-15 or high concentrations of IL-2. There was a large variability in expression of CD127, the α-chain of the IL-7 receptor; CD57-CD28+ cells often expressed CD127 whereas CD57+CD28- cells rarely (median 11.7%) did, consistent with the more effector-like phenotype of CD57+CD28- cells and with previous reports regarding CX3CR1hi CD8 Tmem (2, 34). We next stimulated PBMCs from CMV+ PLWH with IL-2, IL-15, or plate-bound anti-CD3 and soluble anti-CD28 mAbs (TCR) for 2 days to determine if these factors induced the CX3CR1+CD57+CD28- phenotype in CD8 Tmem. Treatment with IL-15 or TCR stimulation resulted in an increase in the expression of CX3CR1, but only IL-15 also sustained CD57 expression, which was downregulated on CD8 Tmem after TCR stimulation (Figure 3A), possibly due in part to activation-induced cell death of the CD57+ cells. All three subpopulations of CCR7-CX3CR1+ Tmem responded to IL-15 and to TCR stimulation by increasing granzyme B and perforin co-expression (Figure S3B). Thus, in a mixed pool of CD8 T cells, IL-15 induced expression of CX3CR1, CD57, and granzyme B and perforin, whereas TCR stimulation induced expression of CX3CR1, granzyme B and perforin, but not CD57. IL-2 had minimal effect on CD8 Tmem phenotype. To determine if the IL-15-induced increases in CX3CR1 and CD57 reflected de novo acquisition of these molecules, we sorted CCR7+CX3CR1-CD57-CD28+CD45RO+ central memory CD8 T cells (Tcm) from HIV/CMV coinfected donors and stimulated them with IL-15 for 7 days (Figure 3B). We found a substantial increase in CX3CR1 and CD57 expression on the sorted cells, coupled with a decrease in CD28 expression. Our data therefore support the concept that the persistent accumulation of CX3CR1+CD57+ CD8 Tmem in CMV+ PLWH could be due at least in part to antigen-nonspecific mechanisms.

**IL-15 promotes CX3CR1+CD57+ CD8 Tmem viability via STAT5 and Bcl-2 activity**
IL-15 has been shown to induce mitochondrial biogenesis and promote oxidative phosphorylation (OXPHOS) in T cells (30, 35). Therefore, we next measured forward scatter (i.e. cell size), mitochondrial mass, OXPHOS, and c-myc expression in stimulated CX3CR1+CD57+ CD8 Tmem (Figure 3C-F). While CX3CR1+CD57+ CD8 Tmem responded similarly to IL-15 and TCR stimulation in terms of cell size, mitochondrial biogenesis, and OXPHOS activity, only IL-15 elicited sustained c-myc expression in these cells.

CD57 expression has been proposed to mark T cells near the end of their replicative capacity (7, 27). Indeed, we found that sorted CX3CR1+CD57+CD28- CD8 Tmem had decreased viability after 7 days of TCR stimulation (Figure 4A,B). Conversely, viability was preserved in CD57+CD28- cells by both IL-2 and IL-15. Because IL-15 promotes expression of the anti-apoptosis protein Bcl-2, which supports cell survival upon activation (36, 37), we measured Bcl-2 levels after stimulation (Figure 4C). While both IL-15 treatment and TCR stimulation elicited an early upregulation of Bcl-2 levels, only IL-15 signals durably maintained high Bcl-2 expression. Thus, our data suggest that sustained Bcl-2 expression might be required to maintain cell viability of CX3CR1+CD57+ CD8 Tmem in the presence of the robust cell growth and mitochondrial activity induced by IL-15 and TCR signals.

To define mechanisms of IL-15-supported survival of CX3CR1+CD57+ CD8 Tmem, we next stimulated sorted CX3CR1+CD57+CD28- cells with IL-15 in the presence or absence of specific inhibitors (Figures 4D, S4A). Inhibiting Bcl-2 with venetoclax reduced the viability of the sorted cells underscoring an important role for Bcl-2 here. IL-15 signals through STAT5 to activate the nutrient sensor target of rapamycin complex 1 (mTORC1)(38), and mTORC1 is both promoted by and reciprocally promotes c-myc activity (39, 40). However, neither blocking mTORC1 activity with rapamycin nor blocking the combined activities of mTORC1 and mTORC2 with KU-0063794 affected IL-15-induced survival benefit. Blocking STAT5 phosphorylation prevented IL-15-induced survival, as did blocking Bcl-2. Thus, IL-15 promotes survival of sorted CX3CR1+CD57+CD28- CD8 Tmem via STAT5 and Bcl-2, but independent of mTORC signaling.

**IL-15-induced CX3CR1+CD57+ CD8 Tmem proliferation is dependent on STAT5 and mTORC1**
Given both IL-15 and TCR stimulation could affect cell size, c-myc expression, and mitochondrial biogenesis of CX3CR1+CD57+CD28- CD8 Tmem, we next asked if these exposures could drive proliferation of this reportedly senescent T cell population (Figure 5A). Because of the differences we observed in sorted cell viability after stimulation, we gated only on viable cells for proliferation analyses. IL-15 and TCR signals, but not IL-2 treatment, resulted in robust proliferation of viable cells. As TCR stimulation did not promote survival, and IL-2 did not drive proliferation of these cells, while IL-15 did both, we focused on IL-15 induced proliferation. In contrast to its impact on viability, inhibiting Bcl-2 with venetoclax did not affect proliferation induced by IL-15. However, blocking mTORC1 with either rapamycin or KU-0063794 or inhibiting STAT5 phosphorylation reduced IL-15-induced proliferation (Figures 5B, S4B). Thus, in contrast to stimulation through the TCR or exposure to IL-2, IL-15 stimulation uniquely promoted both the viability and proliferation of CX3CR1+CD57+CD28- CD8 Tmem via STAT5 signals.

In summary, in sorted CX3CR1+CD57+CD28- CD8 Tmem, the response to IL-15 stimulation in the absence of mTOR signals mimics the response to IL-2 treatment (enhanced viability, poor proliferation), whereas the response to IL-15 stimulation in the absence of Bcl-2 activity was similar to the response to TCR stimulation (poor viability, robust proliferation), and inhibiting STAT5 abrogated the entire IL-15 treatment response (Figure 5C). To confirm that IL-15-induced STAT5 activation was upstream of mTOR activity, we measured the phosphorylation of STAT5 and ribosomal S6 protein in CD57+ CD8 Tmem after 45 minutes of IL-15 exposure in the presence or absence of a STAT5 phosphorylation inhibitor or rapamycin (Figures 5D, S4C). As expected, blocking mTORC1 activity with rapamycin inhibited the phosphorylation of S6 but not STAT5. Blocking STAT5 phosphorylation, however, also impaired S6 phosphorylation – confirming that STAT5 activation is upstream of mTORC1 activity in IL-15-stimulated CD57+ CD8 Tmem. In contrast, inhibiting STAT5 phosphorylation had no effect on S6 phosphorylation in CD4 T cells stimulated with IL-15 (Figure S4D). We next determined whether inhibiting mTOR signaling eliminated the IL-15-induced increase in mitochondrial activity. Blocking mTORC1 activity with rapamycin significantly reduced the IL-15-induced increases in mitochondrial mass and OXPHOS in CX3CR1+CD57+ CD8 Tmem after 4 days of exposure (Figures 5E, S4E). Thus, our data suggest that IL-15 promotes proliferation of ostensibly senescent CD8 Tmem by driving mTORC1-dependent mitochondrial activity.
DISCUSSION

The drivers and determinants of memory CD8 T cell expansion in HIV infection are incompletely described and the distinct effects of HIV and CMV infections on CD8 Tmem phenotypes are only beginning to be understood. Recent studies have suggested that CD8 Tmem that express the vascular endothelium homing receptor CX3CR1 are an important population of cells that promote tumor and virus clearance, and which may provide a source of cells that respond to immunotherapies, such as blockade of the PD-1 pathway (1, 2, 26, 41, 42). Here, we investigated the memory CD8 T cell pool in people with or without HIV infection and found that CMV+ individuals had enrichment of CX3CR1+ CD8 T cells that display surface markers of senescence (CD57+CD28-) and intracellular accumulation of cytolytic molecules (e.g. granzyme B and perforin), regardless of HIV infection status. These cells can be maintained and expanded in an antigen-independent manner in vitro by IL-15, via the combined activities of STAT5, Bcl-2, and mTORC1. Whereas IL-15 promoted both CX3CR1 and CD57 expression and reduced CD28 expression (43, 44), even on sorted Tcm (Figure 3B), TCR stimulation resulted in upregulation of CX3CR1 but not CD57 on CD8 Tmem in vitro (Figure 3A), suggesting that TCR stimulation drives cells toward a CX3CR1+CD57-CD28- phenotype. Whether CX3CR1+CD57-CD28+ cells serve as stimulation-dependent precursors for the other two groups is unknown, and further study will be required to discern the developmental interactions among CX3CR1+ CD8 Tmem subpopulations in PLWH and in HIV-uninfected controls.

Earlier studies proposed that CD57 expression marked terminally-differentiated Tmem that underwent activation-induced apoptosis in response to TCR stimulation (7, 27), and our results confirm the findings that TCR engagement results in the death of these cells. However, we show here that terminally-differentiated CX3CR1+CD57+CD28- CD8 Tmem are capable of prolonged survival (via Bcl-2) and proliferation (via mTORC1) in response to IL-15. These effects are each dependent on the STAT5 signal transduction pathway, consistent with mechanistic studies in mice that demonstrated that antigen-independent inflammatory IL-15 stimulation of CD8 Tmem enhances cycling via STAT5 and mTORC1 (38) and survival via Bcl-2 (45). We found that IL-15 stimulates cell growth, mitochondrial biogenesis, and OXPHOS in CX3CR1+CD57+CD28- CD8 Tmem, consistent with previous reports in CD8 and CD4 T cells (30, 35). IL-15 likely elicits these effects, at least in part, through its upregulation of c-myc, since c-myc activity is strongly linked to cell proliferation (46),
mitochondrial biogenesis (47), and cell metabolism (48), and c-myc has been shown to be important for IL-15-mediated homeostatic proliferation of CD8 Tmem (31). Conversely, mTORC1 activity has also been shown to promote c-myc expression, suggesting the presence of a feed-forward loop (40, 49). Here, mTORC1 activity was necessary for the optimum induction of mitochondrial biogenesis and OXPHOS following IL-15 exposure in CX3CR1+CD57+CD28- CD8 Tmem. IL-15 stimulation has also been shown to upregulate carnitine palmitoyltransferase 1a (Cpt1a) (30, 50, 51), a crucial rate-limiting enzyme for the β-oxidation of long chain fatty acids (51, 52) that is downstream of c-myc (53). CD28-mediated costimulation drives Cpt1a in naïve T cells primed by anti-CD3 (54), but other evidence suggests that anti-CD3/anti-CD28 may reduce Cpt1a expression (52, 55). In either case, it is possible that in the absence of CD28 costimulatory signals, TCR activation of CX3CR1+CD57+CD28- CD8 Tmem is pro-apoptotic as a result of insufficient c-myc expression, dysfunctional metabolism, and a lack of Cpt1a-mediated fatty acid β-oxidation (FAO). Stimulation with IL-15, on the other hand, may promote Cpt1a expression via c-myc, leading to sustained metabolic activity and enhancing cell survival. We note, however, that a recent study showed that continuous IL-15 exposure drove less Cpt1a expression and FAO in NK cells than did intermittent IL-15 exposure (56). Whether these effects occur in IL-15-exposed CX3CR1+CD57+CD28- CD8 Tmem, which share many functional and phenotypic attributes with NK cells, remains to be elucidated, and further research is needed to determine if and how Cpt1a and FAO are necessary for the optimal viability of stimulated CX3CR1+CD57+CD28- CD8 Tmem.

Recently IL-15 has been shown to be necessary for the generation and function of innate-like, virtual memory CD8 T cells (Tvm), which exhibit a memory phenotype despite never encountering specific cognate antigen, and which may provide a cross-reactive, innate-like protection from pathogens (57, 58). In humans, Tvm express a terminally-differentiated Tmem phenotype (e.g. CD57+), but can be distinguished from antigen-experienced terminally-differentiated memory T cells by expression of killer-cell immunoglobulin-like receptors (KIRs) and/or NKG2A (57, 59, 60). Although we cannot exclude a contribution of Tvm to the IL-15-induced proliferation that we observed here, our preliminary data of NKG2A expression in a separate cohort of PLWH suggest that most CX3CR1+CD57+CD28- CD8 Tmem that proliferate in response to IL-15 are presumably antigen-experienced cells (data not shown).
As the immune system ages, and as infection episodes and comorbidities accumulate, memory T cells continue to differentiate until there is an abundance of cells expressing markers associated with poor proliferative capacity, reduced lifespan, vascular homing potential, and robust cytotoxicity (CD57, CX3CR1, and granzyme B). CMV infection in particular appears to be a potent contributor to the generation of these cells – yet it is not clear how these terminally-differentiated CD8 T cells are sustained. We show here that under conditions of inflammatory IL-15 stimulation, possibly related to CMV persistence (61), ostensibly “senescent” CX3CR1+CD57+CD28- CD8 Tmem are generated, armed with effector molecules, proliferate, and retain viability in an antigen-independent manner. Our findings indicate that these highly differentiated CD8 Tmem cells are not invariably senescent and that they can be driven to proliferate when stimulated by an inflammatory cytokine – IL-15. We propose the term “inflamescent” to describe CX3CR1+CD57+CD28- CD8 Tmem, which express traditional markers of senescence yet retain survival and proliferation capacities, and which are both driven by and contribute to inflammation.
METHODS

Human Donors and Tissues.

Peripheral blood was acquired in Vacutainer tubes containing EDTA (BD) from persons living with HIV infection (PLWH) receiving combination ART with plasma HIV RNA <40 copies/mL (CMV-seronegative, n = 9; CMV-seropositive, n = 42) and HIV-uninfected controls (n = 27). Participant characteristics stratified by HIV infection status are shown in Table S1. CMV serostatus was determined clinically by IMMULITE 2000 CMV IgG Ab immunoassay in plasma (Siemens). HIV-infected participant characteristics stratified by CMV infection status are shown in Table S2. Although there were significant differences between the cohorts of PLWH, we did observe trends toward more CD8 T cells and a lower CD4/CD8 ratio in the CMV+ donors – consistent with our previous observations (17). Peripheral blood mononuclear cells (PBMCs) were purified by centrifugation over a Ficoll-Paque cushion (GE Healthcare).

Flow Cytometry.

Lymphocytes were identified by forward and side scatter, and phenotype was assessed using the following fluorochrome-conjugated antibodies: anti-CD3 BUV737 (UCHT1; BD), anti-CD4 BUV395 (SK3; BD), anti-CD8 BV605 (SK1; BD), anti-CCR7 PE-Cy7 (3D12; BD), anti-CX3CR1 PerCP-Cy5.5 (2A9-1; BioLegend), anti-CD25 APC-H7 (M-A251, BD), anti-CD27 AlexaFluor 700 (M-T271, BD), anti-CD28 BV421 or BB515 (CD28.2; BioLegend), anti-CD45RO BV650 (UCHL1, BD), anti-CD57 AlexaFluor 647 or Pacific Blue (HNK-1; BioLegend), anti-CD69 FITC (Bcl-2/100, BD), anti-c-myc PE (9E10, R&D Systems), anti-Bcl-2 FITC (Bcl-2/100, BD), T-bet PE (4B10, eBioscience), eomes FITC (WD1928, eBioscience), anti-granzyme B FITC (GB11; BD) and anti-perforin PE (B-D48; BioLegend). For detection of
intracellular phosphorylated proteins, cells were fixed with 16% formaldehyde for 10 minutes at 37°C, then washed and treated with 90% methanol for 20 minutes at -20°C, then washed and incubated for 20 minutes with anti-STAT5 pY694 PE (47/Stat5[pY694], BD) and anti-S6 p240 APC (REA420, Miltenyi Biotec). To identify CMV or HIV antigen-specificity, PBMCs were incubated with BV421-conjugated MHC class I HLA-A*02-restricted CMV pp65 (NLVPMVATV) or HIV gag (SLYNTVATL) peptide tetramers (NIH Tetramer Facility) for 1 h at room temperature prior to surface antibody staining. Mitochondrial mass was determined by labeling cells with MitoTracker Green dye (Molecular Probes), and mitochondrial oxidative phosphorylation activity was determined using MitoTracker Orange dye (Molecular Probes) per manufacturer’s instructions. Memory T cells (Tmem) were determined phenotypically by excluding phenotypically naïve (CD45RO-CCR7+) cells. High dimensionality t-stochastic neighbor embedding (t-SNE) analysis was performed using 12 parameters: CCR7, CD27, CD28, CD45RO, CD57, CD95, CD101, CD244, CX3CR1, PD-1, SLAMF6, and TIM-3. For some experiments, CD8 T cells enriched by magnetic beads (AutoMACS, Miltenyi Biotec) were then antibody labeled and sorted into subpopulations (CCR7-CX3CR1+CD57+CD28-, CCR7-CX3CR1+CD57-CD28-, CCR7-CX3CR1+CD57+CD28+, or CCR7+CX3CR1-CD57-CD28+; average sort purity 89.4%) using a FACSAlia (BD) or S3 sorter (BioRad). All flow cytometry data were acquired on a LSRFortessa flow cytometer (BD) and analysis including t-SNE, performed using FlowJo software (FlowJo).

**T cell stimulation assays.**

PBMCs or sorted CD8 T cell subpopulations were treated for 45 minutes (phosflow assay), or one, two, four, and seven days with medium control, recombinant IL-2 (100U/ml; Novartis), recombinant IL-15 (20ng/ml; R&D Systems), or anti-CD3 (10µg/ml; HIT3a, BD) and anti-CD28 (5µg/ml, CD28.2, BD). After treatment, T cells were immunostained and analyzed by flow cytometry. In some experiments, cells were labeled with CellTrace Violet (Molecular Probes) per manufacturer’s instructions prior to culture to measure proliferation. Determination of proliferation precursor frequency (percent divided) was done using FlowJo software. In other assays, cells were treated with rapamycin (250ng/ml, Millipore Sigma), KU-0063794 (500nM, SelleckChem), venetoclax (500nM, SelleckChem), or N’-((4-Oxo-4H-chromen-3-yl)methylene)nicotinohydrazide (500µM, Millipore Sigma), an inhibitor of STAT5 activity. For intracellular cytokine staining, PBMCs were stimulated with plate-bound anti-CD3 (10µg/ml; HIT3a, BD) or medium control in the presence of brefeldin-A (GolgiPlug, BD).
After 6h, cells were harvested, washed, labeled with Live/Dead and surface Ab and permeabilized as above. Cells were then stained with anti-IFNγ FITC (B27, BD), anti-IL-2 PE-CF594 (5344.111, BD), anti-MIP-1β PE (D21-1351, BD), and anti-TNF APC (Mab11, BD) for 40 minutes on ice.

Statistics.
Comparisons between unrelated groups used nonparametric two-tailed Mann Whitney U tests. Comparisons among three or more groups were performed with nonparametric Kruskal-Wall tests with Dunn’s multiple comparison post-tests. If data passed D’agostino and Pearson omnibus normality test then multiple groups were compared by one-way ANOVA. Contingency analyses used Fisher’s Exact test. Paired group analyses used Wilcoxon matched-pairs signed rank test. All statistics were performed using Prism software (version 6 or 8, GraphPad). Significance thresholds were set at P-values less than 0.05.

Study Approval.
All human experiments were approved by the Institutional Review Board of University Hospitals, Cleveland Medical Center (Protocol #01-98-55). Blood samples were acquired with informed consent, and in accordance with the Declaration of Helsinki.
AUTHOR CONTRIBUTIONS

S.R.M. designed, performed, and analyzed experiments, and wrote the manuscript. B.C., J.C.M., and S.P. performed and analyzed experiments. C.L.S., S.F.S., C.C., D.A.Z., N.T.F., and S-A.Y. provided reagents and materials, and contributed to the study design and analysis. B.R. and S.G. provided patient samples and contributed to the study design. M.M.L. provided patient samples, contributed to study design and analysis, and helped to write the manuscript. M.L.F. conceived the study design, performed and analyzed experiments, wrote the manuscript, and guided the project. All authors reviewed and approved the manuscript.

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Figure 1. CX3CR1+ CD8 Tmem are expanded in HIV/CMV coinfected individuals.

Representative staining and quantification of (A) CX3CR1 and CCR7 expression on CD8 Tmem and (B) CD57 and CD28 expression on CCR7-CX3CR1- and CCR7-CX3CR1+ CD8 Tmem subpopulations in CMV- (n=8) and CMV+ (n=12) PLWH. Data represent median ± IQR. Significance determined by Mann-Whitney test. (C) Percentage of CD8 Tmem that are CX3CR1+CD57+CD28- in CMV-seronegative (n=9) and CMV-seropositive (n=10) HIV-uninfected donors. Data represent median ± IQR. Significance determined by Mann-Whitney test. (D) t-stochastic neighbor embedding (t-SNE) analysis of 12-parameter data of CD8 T cells from combined HIV-CMV- (n=4), HIV-CMV+ (n=4), HIV+CMV- (n=4), and HIV+CMV+ (n=4) donors. CX3CR1+CD57+CD28- Tmem are shown in red. Proportion of donor populations that are CX3CR1+CD57+CD28- CD8 Tmem, stratified by HIV and CMV infection status.
Figure 2. CX3CR1+CD57+CD28- CD8 Tmem express Bcl-2, c-myc, and are in cell cycle *in vivo*.

Intracellular expression of Bcl-2 (A) and c-myc (B) in subsets of CCR7-CX3CR1+ CD8 Tmem stratified by CD57 and CD28 expression from CMV+ PLWH (n=6). Data represent median ± IQR. Significance determined by Kruskal-Wallis test with Dunn’s correction for multiple comparisons. (C) Percentage of Ki67- and Ki67+ CCR7-CX3CR1+ CD8 Tmem that are CD57+CD28- from CMV+ PLWH (n=10). Data represent median ± IQR. Significance determined by Wilcoxon matched-pairs test.
Figure 3. IL-15 promotes CX3CR1 and CD57 expression and mitochondrial activity in CD8 Tmem.

(A) Representative histograms and quantification of CX3CR1 and CD57 expression 48h after indicated stimulation expressed as fold change (FC) over medium control on CD8 Tmem from CMV+ PLWH (n=6). Data represent median ± IQR. Significance determined by Kruskal-Wallis test with Dunn’s correction for multiple comparisons. (B) Representative histograms and quantification of CX3CR1, CD57, and CD28 expression on sorted CD8 central memory T cells (Tcm) 7 days from CMV+ PLWH (n=6) after IL-15 stimulation expressed as
FC over medium control. Data represent median ± IQR. Significance determined by Kruskal-Wallis test with Dunn’s correction for multiple comparisons. (C-F) Representative histograms from day 4 of stimulation and quantification of forward scatter (FSC-A)(C), MitoTracker Green (D), MitoTracker Orange (E), and intracellular c-myc (F) after stimulation expressed as FC over medium control in CCR7-CX3CR1+CD57+CD28- CD8 Tmem from HIV-uninfected controls (n=7). Data represent median ± IQR. Significance among groups determined by Kruskal-Wallis test with Dunn’s correction for multiple comparisons: *$P \leq 0.05$, **$P \leq 0.01$, IL-15 vs IL-2; #$P \leq 0.05$, ##$P \leq 0.01$, TCR vs IL-2; %$P \leq 0.05$, IL-15 vs TCR. Significance within groups (7d vs. 1d) determined by Wilcoxon matched-pairs test: *$P \leq 0.05$. 
Figure 4. IL-15 promotes viability of CCR7-CX3CR1+CD57+CD28- CD8 Tmem via STAT-5 and Bcl-2 activity.

(A) Schematic of flow sorting strategy. (B) Representative labeling with Live/Dead Aqua and quantification of viability after 7d stimulation expressed as FC over medium control in sorted CCR7-CX3CR1+CD57+CD28-CD8 Tmem from CMV+ PLWH (n=5-17/stim). Data represent median ± IQR. Significance determined by Kruskal-Wallis test with Dunn’s correction for multiple comparisons. (C) Representative histograms from day 4 of stimulation and quantification of intracellular Bcl-2 staining after stimulation expressed as FC over medium control in CCR7-CX3CR1+CD57+CD28- CD8 Tmem from HIV-uninfected controls (n=7). Data represent median ± IQR. Significance among groups determined by Kruskal-Wallis test with Dunn’s correction for multiple comparisons: *P ≤ 0.05, **P ≤ 0.01, ***P ≤ 0.005, IL-15 vs. IL-2; ##P ≤ 0.01, TCR vs IL-2; %P ≤ 0.05,
$P \leq 0.01$, IL-15 vs TCR. Significance within groups (7d vs. 1d) determined by Wilcoxon matched-pairs test:

* $P \leq 0.05$. (D) Viability after 7d stimulation with medium control or IL-15 with or without indicated inhibitors in sorted CCR7-CX3CR1+CD57+CD28- CD8 Tmem from CMV+ PLWH (n=8-14/stim). Data represent median ± IQR. Significance determined by Kruskal-Wallis test with Dunn’s correction for multiple comparisons.
Figure 5. IL-15 promotes proliferation of CCR7-CX3CR1+CD57+CD28- CD8 Tmem via STAT-5 and mTORC1 activity.

(A) Representative labeling with CellTrace Violet and quantification of proliferation after 7d stimulation expressed FC over medium control in sorted CCR7-CX3CR1+CD57+CD28- CD8 Tmem from CMV+ PLWH (n=4-15/stim), gated on viable cells. Data represent median ± IQR. Significance determined by Kruskal-Wallis test with Dunn’s correction for multiple comparisons. (B) Proliferation after 7d stimulation with medium control or IL-15 with or without indicated inhibitors in sorted CCR7-CX3CR1+CD57+CD28- CD8 Tmem gated on viable cells from CMV+ PLWH (n=8-14/stim). Data represent median ± IQR. Significance determined by Kruskal-Wallis test with Dunn’s correction for multiple comparisons. (C) Summary of viability and proliferation outcomes. (D) STAT5 pY694 and S6 pS240 expression 45min after stimulation with medium control or IL-15
with or without indicated inhibitors in gated CD57+ CD8 T cells from CMV+ PLWH (n=9). Data represent median ± IQR. Significance determined by Kruskal-Wallis test with Dunn’s correction for multiple comparisons. (E) MitoTracker Green and MitoTracker Orange labeling expressed as FC over medium control in CD57+ CD8 T cells from CMV+ PLWH (n=9) following 4 days stimulation with IL-15 with or without rapamycin (Rapa). Significance determined by Wilcoxon matched-pairs test.