Persistent expansion and Th1-like skewing of HIV-specific circulating T follicular helper cells during antiretroviral therapy

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ABSTRACT

Background: Untreated HIV infection leads to alterations in HIV-specific CD4+ T cells including increased expression of co-inhibitory receptors (IRs) and skewing toward a T follicular helper cell (Tfh) signature. However, which changes are maintained after suppression of viral replication with antiretroviral therapy (ART) is poorly known.

Methods: We analyzed blood CD4+ T cells specific to HIV and comparative viral antigens in ART-treated people using a cytokine-independent activation-induced marker assay alone or in combination with functional readouts.

Findings: In intra-individual comparisons, HIV-specific CD4+ T cells were characterized by a larger fraction of circulating Tfh (cTfh) cells than CMV- and HBV-specific cells and preferentially expressed multiple IRs and showed elevated production of the Tfh cytokines CXCL13 and IL-21. In addition, HIV-specific cTfh exhibited a predominant Th1-like phenotype and function when compared to cTfh of other specificities, contrasting with a reduction in Th1-functions in HIV-specific non-cTfh. Using longitudinal samples, we demonstrate that this distinct HIV-specific cTfh profile was induced during chronic untreated HIV infection, persisted on ART and correlated with the translation-competent HIV reservoir but not with the total HIV DNA reservoir.

Interpretation: Expansion and altered features of HIV-specific cTfh cells are maintained during ART and may be driven by persistent HIV antigen expression.

Funding: This work was supported by the National Institutes of Health (NIH), the Canadian Institutes of Health Research (CIHR) and the FRQS AIDS and Infectious Diseases Network.

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Studies of peripheral blood responses con-
\[15\] qualitative defects in their capacity to provide help to B cells. CD4+ T cells exhibits a Tfh-like signature that directly correlates with
\[11\] infection, expansion of bulk GC Tfh cells occurs in lymphoid tissue with ART initiation but can stay elevated compared to uninfected controls
\[11-13\]. Bulk cTfh cells from ART-treated donors show a reduced capacity to induce HIV antibody production and B cell differentiation in vitro when compared to uninfected controls \[17,18\]. We observed that cTfh constitute a readily detectable fraction of HIV-specific CD4+ T cells in individuals prior to ART initiation, this irrespective of viral load \[4\]. ART decreased the marked Tfh gene signature present in viremic persons at the transcriptional level \[4\], however, the impact of ART on the differentiation and function of HIV-specific CD4+ T cell has not yet been investigated. Whether immunological features of HIV-specific cTfh would be unique to this virus or shared with other Ag specificities in the same individual, remains to be determined.

Here, we used activation-induced marker (AIM) assays \[4,19,20\], and functional tests to define the frequency, phenotype and function of blood CD4+ T cells, in particular cTfh, specific to HIV, CMV and HBV in ART-treated individuals. We longitudinally investigated partici-
pants pre- and post-ART initiation to delineate features of HIV-spe-
cific CD4+ T cells that were modulated by therapeutic control of viral load from those that persisted despite suppressive ART. Finally, we identified features of these HIV-specific cTfh cells that correlated with the size of the translation-competent HIV reservoir.

2. Material and methods

2.1. Human sample collection and processing

Subject characteristics are summarized in Tables S1-3. HIV-infected, ART-treated participants were on ART for over 12 months with controlled viral load (<50 vRNA copies/ml) for at least 6 months. Donors on ART were not excluded when a single small viral blip (VL > 50 but <200 vRNA copies/ml) occurred with below detection viral load on precedent and subsequent tests. Untreated participants were either treatment naive or untreated for at least 3 months. HIV-uninfected individuals were used as negative controls for HIV antibody and reservoir measurements. PBMCs were isolated from leukapheresis samples by the Ficoll-Hypaque density gradient centrifugation and cryo-preserved in liquid nitrogen until use.

2.2. CD69/CD40L AIM assay

Peripheral blood mononuclear cells (PBMCs) were thawed, washed and put in culture at a concentration of 10 million cells/ml in RPMI 1640 medium (Gibco by Life Technologies, Cat# 11,875-093) supplemented with 0.5% penicillin/streptomycin (Gibco by Life Tech-
nologies, Cat# 15140122) and 10% human serum (Sigma). After a rest of 3 h at 37 °C, a CD40 blocking antibody (Miltenyi Biotec, Cat# 130-094-133, RRID:AB_10839704) was added to the culture to prevent the interaction of CD40L with CD40 and its subsequent downregulation. In addition, antibodies for chemokine receptors CXCRI, CXCRII and CCR5 were added into the culture medium. After 15 min incubation at 37 °C, cells were stimulated with 0.5 μg/ml staphylococcal enterotoxin B (SEB) or 0.5 μg/ml of overlapping peptide pools for CMV pp65 (Cat# PM-PP65), HBV HBsAg (Cat# PM-HBV-IEP), HIV Gag (Cat# PM-HIV-GAG), HIV Env (Cat# PM-HIV-ENV) or HIV Nef (Cat# PM-HIV-NEF) (all JPT) for 9 h at 37 °C. An unstimulated condition served as a negative control. Cells were stained for viability dye

infection, expansion of bulk GC Tfh cells occurs in lymph nodes
\[11-13\]; while technical hurdles have thus far limited studies of the specificity of these cells, part of this expanded GC Tfh population appears to be HIV-specific \[14\]. This quantitative increase in GC Tfh in HIV infection correlates with markers of disease progression and qualitative defects in their capacity to provide help to B cells \[15\]. Studies of peripheral blood responses confirmed an Ag-driven com-
ponent to these alterations: the transcriptome of blood HIV-specific CD4+ T cells exhibits a Tfh-like signature that directly correlates with viral load \[4\]. In contrast to LCMV, however, in HIV infection, this Tfh skewing does not result in virus clearance, perhaps due to the remarkable capacity of HIV to generate escape mutations and elude autologous Ab neutralization.

While current antiretroviral therapy (ART) regimens are highly effective at suppressing viral replication, resulting in improved immunity against opportunistic infections and remarkable reduction in morbidity and mortality, they do not lead to the restoration of an effective HIV-specific immune response capable of suppressing virus. Consequently, ART interruption generally leads to a rapid viral rebound from a reservoir of latent infected cells \[16\].

Frequencies of total GC Tfh decrease in lymphoid tissue with ART initiation but can stay elevated compared to uninfected controls
\[11-13\]. Bulk cTfh cells from ART-treated donors show a reduced capacity to induce HIV antibody production and B cell differentiation in vitro when compared to uninfected controls \[17,18\]. We observed that cTfh constitute a readily detectable fraction of HIV-specific CD4+ T cells in individuals prior to ART initiation, this irrespective of viral load \[4\]. ART decreased the marked Tfh gene signature present in viremic persons at the transcriptional level \[4\], however, the impact of ART on the differentiation and function of HIV-specific CD4+ T cell has not yet been investigated. Whether immunological features of HIV-specific cTfh would be unique to this virus or shared with other Ag specificities in the same individual, remains to be determined.

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Evidence before this study

Combination antiretroviral therapy (ART) is highly effective in
controlling HIV but requires life-long medication due to the exis-
tence of a latent viral reservoir, and to the fact that, with rare exceptions, ART alone does not restore immune responses capa-
bility of suppressing HIV. T follicular helper cells (Tfh) are of high
interest both as HIV reservoirs and for their critical role in
enabling the development of potent broadly neutralizing anti-
bodies (bNAbs). The identification of a circulating Tfh (cTfh) pop-
ulation in peripheral blood with strong similarities with
lymphoid tissue Tfh has attracted much attention over the past
few years. However, investigations of cTfh at the antigen (Ag)-
specific level have been hampered by the lack of sensitive assays
to detect them. We recently overcame this hurdle by using new
experimental, cytokine-independent approaches to evaluate
HIV-specific CD4+ T cell responses. We found an elevated Tfh
signature in HIV-specific CD4+ T cells of chronically infected,
untreated individuals that correlated with blood HIV viremia.
Although this Tfh signature decreased after ART initiation at the
transcriptional level, detailed phenotypic and functional analyses
of the HIV-specific CD4+ T cell responses during ART are lacking.

Added value of this study

Here, we evaluated HIV-specific CD4+ T cells, and in particular
cTfh, in a cohort of ART-treated individuals. Comparative phe-
notypic and functional analyses with Ag-specific responses
(CMV, HBV) in the same study participant revealed that HIV-
specific cTfh cells are abundant in ART-treated humans and
represent a much larger fraction of the virus-specific CD4+ T cell
response compared to their CMV- and HBV-specific coun-
terparts. HIV-specific cTfh also differ from CMV-specific and
HBV-specific cTfh by multiple phenotypic and functional fea-
tures that are established during chronic viremic infection,
which persist on ART and appear less responsive to viral
suppression than in non-cTfh HIV-specific CD4+ T cells. This
distinctive HIV-specific cTfh profile correlates with the transla-
tion-competent HIV reservoir, but not the total HIV DNA
reservoir, suggesting that persistent HIV antigen expression
maintains these altered features during ART.

Implications of all the available evidence

Increasing evidence suggest that cTfh are ontogenically related
to lymphoid tissue Tfh. As animal models suggest that overabund-
ant, qualitatively impaired Ag-specific Tfh may be detrimental
and lead to low affinity Ab responses, it will be important to
determine if priming of new, effective Tfh responses may be
hampered in therapeutic vaccine trials by competition with such
large pre-existing HIV-specific Tfh populations during ART.
2.5. Transcription factor staining

Table S4 for antibodies). Incubated with antibodies against cytokines for 30 min at 4 °C (see fixation/Permeabilization kit (Thermo Fisher, Cat# 88-8824-00) and Spiec version 5.3 (RRID:SCR_008520) and Spice version 5.3 (RRID:SCR_016603) [21]. For phenotypic analysis of Ag-specific CD4+ T cells, only responses that were >2-fold over unstimulated condition were included to limit the impact of background staining. In contrast, for analysis of Ag-specific CD4+ T cells subsets as percentage of total CD4+ T cells, background-subtracted net values were used, which did not require excluding responses.

2.3. Standard intracellular cytokine staining

PBMCs were thawed, washed and put in culture at a concentration of 4 million cells/ml in RPMI 1640 medium (Gibco by Life Technologies, Cat# 11875-093) supplemented with penicillin/streptomycin (Gibco by Life Technologies, Cat# 15140122) and 10% fetal bovine serum (FBS) (Seradigm, Cat#1500-500). After a rest of 2 h, cells were stimulated with 0.5 µg/ml of overlapping peptide pools for CMVpp65, HBV HBsAg or HIV Gag for 6 h in presence of Brefeldin A (BD Biosciences, Cat# 555029) and monensin (BD Biosciences, Cat# 554724). For some experiments, anti-CD107A-BV785 (Biolegend, Cat#328644, RRID: AB_2565968) was added into culture. Cells were stained for viability marker, surface markers and intracellular cytokines using the IC Fixation/Permeabilization kit (eBioscience, Cat# 88-8824-00) before fixation with 2% PFA and acquisition on the flow cytometer (see Table S4 for antibodies). For the detection of CD107A, granzyme B and perforin within Ag-specific CD4+ T cells, we identified AIM+ cells by intracellular staining for CD69 and CD40L.

2.6. CD4+ T cells were isolated from thawed PBMCs using a negative isolation kit (StemCell, Cat# 19052), rested for 2 h at 37 °C, washed and stained for 60 min at room temperature with PE-labeled MHC-II tetramers loaded with DV16 peptide (DRFYKYTLRAEQASQEV) for the DRB1*01:01 allele or VY18 peptide (YVDFRYKTLRAEQASQEV) for the DRB1*11:01 allele (NIH Tetrmer Core Facility at Emory University, Atlanta, GA). These sequences encompass an immunodominant, HLA Class II promiscuous epitope in Gag [22]. Control tetramers loaded with an irrelevant peptide (CLIP: PVSKMRMATPLLMQA) or HIV-uninfected donors with the same HLA-DRB1 genotype served as negative controls. Tetramer+ CD4+ T cells were column enriched using anti-PE beads (Miltenyi, Cat# 130-048-801). Cells were stained for viability marker (Aquaavivid, Thermofisher, Cat# L34957), CXCR5 (45 min, 37 °C), surface markers (30 min, 4 °C) and fixed with 2% PFA before acquisition at the flow cytometer (LSRIII, BD).

2.7. MHCI tetramer staining

CD4+ T cells were isolated from thawed PBMCs using a negative isolation kit (StemCell, Cat# 19052), rested for 2 h at 37 °C, washed and stained for 60 min at room temperature with PE-labeled MHC-II tetramers loaded with DV16 peptide (DRFYKYTLRAEQASQEV) for the DRB1*01:01 allele or VY18 peptide (YVDFRYKTLRAEQASQEV) for the DRB1*11:01 allele (NIH Tetrmer Core Facility at Emory University, Atlanta, GA). These sequences encompass an immunodominant, HLA Class II promiscuous epitope in Gag [22]. Control tetramers loaded with an irrelevant peptide (CLIP: PVSKMRMATPLLMQA) or HIV-uninfected donors with the same HLA-DRB1 genotype served as negative controls. Tetramer+ CD4+ T cells were column enriched using anti-PE beads (Miltenyi, Cat# 130-048-801). Cells were stained for viability marker (Aquaavivid, Thermofisher, Cat# L34957), CXCR5 (45 min, 37 °C), surface markers (30 min, 4 °C) and fixed with 2% PFA before acquisition at the flow cytometer (LSRIII, BD).

2.8. Quantification of total and integrated HIV DNA

Total and integrated HIV DNA were measured in CD4+ T cells isolated from PBMCs by magnetic bead-based negative selection (Stem Cell Technologies, Cat# 19052) by real time nested polymerase chain reaction (PCR) as described previously [23].

2.9. Detection of translation-competent reservoir by RNA flow-FISH

CD4+ T cells harbouring latent translation-competent reservoir were identified using the HIVRNAISH assay as previously described [24,25]. Briefly, CD4+ T cells were isolated by magnetic bead negative selection (StemCell, Cat# 19052) from PBMCs from ART-treated individuals, rested for 3 h and stimulated with PMA (50 ng/ml, Sigma-Aldrich, Cat# P1585) and lonomycin (0.5 µg/ml, Sigma-Aldrich, Cat# I9657) for 12 h. Unstimulated cells and cells from HIV-uninfected individuals served as controls. Cells were stained with surface markers, anti-Gag KC57 (Beckman Coulter) by intracellular staining and labeled for HIV gag RNA with Alexa Fluor 750-coupled probes (ThermoFisher) using the PrimeFlow RNA Assay (ThermoFisher, Cat# 88-18005-210) (see Table S4 for antibodies). Translation-competent CD4+ T cells were identified as cells expressing both HIV Gag protein and gag RNA after PMA/lonomycin stimulation.

2.10. Detection of p24-specific antibodies by ELISA

96 well plates (Thermo Scientific Nunc, FluoroNunc/LumiNunc, MaxiSorp Surface) were coated with 0.1 µg/ml of recombinant p24 (NII AIDS Research and Reference Reagent Program, Cat# 12028) or bovine serum albumin (BSA) (Bioshop, Cat# ALB001.1) in PBS overnight at 4 °C. Plates were blocked for 90 min at RT with blocking buffer (TBS, Tween 0.1%, BSA 2%) and then washed 4 times with washing buffer (TBS, Tween 0.1%). Dilutions of human sera (1:3000) or rabbit anti-HIV p24 antisemur (NII AIDS Reagent Program, Cat# 4250) in washing buffer containing 0.1% of BSA were incubated for 2 h at RT. Plates were washed 4 times with washing buffer before incubation for 90 min at RT with HRP-conjugated secondary Abs goat anti-human IgG HRP (Thermo Fisher Scientific Cat# 31410, RRID: AB_228269) or anti IgG rabbit HRP (Thermo Fisher Scientific Cat# 65-6120, RRID:AB_253967). Plates were then washed 4 times with washing buffer before revealing with standard ECL (Perkin Elmer) with a TriStar luminometer (LB 941, Berthold Technologies).

2.11. Detection of gp120-specific antibodies

Gp120-specific antibodies were detected in plasma samples using a flow cytometry-based assay as described previously [26]. Briefly, CEM.NKr cells were coated with recombinant HIV-1 V32 gp120...
from HIV-infected ART-treated donors or uninfected controls (1:10,000 dilution) for 30 min at 37 °C. Cells were washed with PBS and stained with 1 μg/ml goat anti-human Alexa Fluor 647 (Thermo Fisher Scientific, Cat# A-21445 RRID:AB_2535862) secondary antibody for 15 min in PBS at room temperature. Cells were washed and fixed using 2% PFA before acquisition at the flow cytometer. The geometric mean of the Alexa Fluor 647 signal was used to express plasma gp120-antibody levels.

2.14. Data availability

Leukaphereses were obtained from study participants at the McGill University Health Centre, Montreal, Canada, and at the Centre Hospitalier de l'Université de Montréal (CHUM) in Montreal, Canada. The study was approved by the respective IRBs, written informed consent obtained from all participants prior to enrolment.

2.15. Statistical analyses were done using GraphPad Prism version 8 using non-parametric tests. Two-group comparisons were performed using the Mann-Whitney and pairwise comparisons were performed using the Wilcoxon matched pair test. For comparisons between three or more groups, Kruskal–Wallis (for unpaired samples or when values were missing in paired samples) or Friedman one-way ANOVA (for paired samples) with Dunn’s post-test was used. Permutation test (10,000 permutations) was applied for pie-chart comparison using the SPICE software. For correlations, Spearman’s R correlation coefficient was applied. Statistical tests were two-sided and p < 0.05 was considered significant.

2.16. Ethic statement

Leukaphereses were obtained from study participants at the McGill University Health Centre, Montreal, Canada, and at the Centre Hospitalier de l’Université de Montréal (CHUM) in Montreal, Canada. The study was approved by the respective IRBs, written informed consent obtained from all participants prior to enrolment.

2.17. Data availability

Raw experimental data associated with the figures presented in the manuscript are available from the corresponding author upon reasonable request.

3. Results

3.1. AIM assay identifies HIV-specific CD4+ responses with cTfh expansion in ART-treated individuals

To study Ag-specific CD4+ T cells with diverse differentiation and functionality in HIV-infected ART-treated people, we used an approach based on the concurrent detection of activation-induced markers (AIM) on the cell surface after cognate Ag stimulation, as previously described [4,19,20]. PBMCs from a cohort of 27 HIV-infected individuals on ART (Participant characteristics: Table S1 (ART1-27)) were stimulated for 9 h with overlapping peptide pools spanning the sequence of the immunodominant HIV structural protein Gag (Fig. S1a). HIV Gag-specific T cells were identified by concurrent surface expression of AIM CD69 and CD40L (AIM+ cells) (Fig. 1a, S1b). In addition, we examined within the same individual CD4+ T cells specific for other Ags (CMV and HBV) to delineate characteristics that differentiate HIV-specific CD4+ T cells. AIM+ HIV Gag-specific responses were readily detectable in all ART-treated subjects examined, with low background in the absence of exogenous Ag (Fig. 1b, S1c) (responses were considered as positive when more than 2-fold over unstimulated condition). CMV-specific CD4+ T cell responses were detectable in all individuals with positive CMV serology and absent for the 4 CMV-seronegative participants, demonstrating the high specificity of the assay (Fig. 1b, Table S1). Due to the low number of donors with a detectable HBV-specific CD4+ T cell response (6 individuals of the 27 examined; 2 resolved infection, 4 vaccinated) (Fig. 1b, Table S1), we pooled responses elicited by infection or vaccination for further analysis, given the similarities of their profile.

The AIM assay identified significantly higher frequencies of virus-specific CD4+ T cells compared to IFNγ intra-cellular staining (ICS) (Fig. S1d-e), demonstrating that ICS underestimates the frequency of Ag-specific CD4+ T cell responses. This is particularly striking for HIV and HBV, but also apparent for known Th1-skewed responses such as CMV.

Ag-specific cTfh show limited cytokine production and are more likely to be missed by standard ICS, but are detectable by AIM assays [19,27]. Here, we broadly identified cTfh cells as memory (CD45RA–) CD4+ T cells expressing CXCR5 (Fig. S2a, 1c) to avoid the potential confounding factor of PD-1 upregulation in chronic infection [28,29]. Phenotypic analysis was only performed for donors with detectable AIM responses. Compared to CMV- and HBV-specific CD4+ T cells, HIV-specific CD4+ responses were characterized by a significantly higher proportion of CXCR5+ memory cells (Fig. 1d) with comparable CXCR5 MFI for all specificities (Fig. S2b). This skewing was not due to a lower magnitude of HIV Gag expression, but rather was a general feature of HIV-specific CD4+ T cell responses, as analyzed responses against the HIV envelope glycoprotein Env and the accessory protein Nef, which showed a lower magnitude compared to HIV Gag as described previously (Fig. 1e) [22]. Despite lower frequencies, responses were detectable in 13/27 participants for Env and 18/27 participants for Nef. HIV Env- and Nef-specific CD4+ T cell responses were characterized by a similar frequency of CXCR5+ memory cells when compared to Gag-specific CD4+ responses (Fig. 1f).

Accordingly, due to the higher magnitude of HIV Gag-specific CD4+ responses, HIV Gag-specific cTfh were more prevalent in the total CD4+ populations compared to HIV Nef or Env (Fig. 1g). In conclusion, our results demonstrate the expansion of cTfh cells within HIV-specific CD4+ T cell responses compared to other specificities in ART-treated individuals.

3.2. HIV-specific cTfh and non-cTfh cells express multiple co-inhibitory receptors despite viral suppression

IRs are key modulators of T cell signaling for both the regulation of physiologic responses to Ag stimulation and in the context of diseases characterized by persistent Ag exposure. Consistent with a requirement for tight functional control, Tfh cells frequently express IRs such as PD-1, TIGIT and CD200 [30,31]. On the other hand, upregulation of multiple IRs on CD8+ and CD4+ T cells also occurs in cancer or chronic infections including HIV and contribute to exhaustion [reviewed in (32)]. Given this dual role of IRs, we next analyzed IR expression on Ag-specific CD4+ responses in the context of the massive reduction in HIV Ag load due to viral suppression during ART.

HIV Gag-specific cTfh from ART-treated individuals were characterized by a significantly higher frequency of cells expressing the IRs TIGIT and CD200 compared to non-cTfh cells, while the frequency of PD-1+ cells was similar between both subsets (Fig. S3a). When we compared cTfh IR expression between Ags, we observed a high frequency of TIGIT for HIV-specific cTfh cells compared to CMV- and HBV-specific cTfh (Fig. 2a–b). In addition, high levels of CD200 and PD-1 characterized HIV- and HBV-specific cTfh cells (Fig. 2b). The frequency of cTfh cells specific for HIV co-expressing TIGIT, PD-1 and CD200 was significantly higher compared to CMV or HBV (Fig. 2c–d). The differences in IR expression observed for Ag-specific non-cTfh reflected those seen on cTfh, albeit at lower levels (Fig. S3b–c).

IR expression can be transiently induced by activation [33], and may thus be impacted by the 9 h stimulation required for the AIM assay. Therefore, we used MHC Class II tetramers to phenotype HIV-specific cTfh cells in 3 ART-treated donors (Fig. S2b). We detected similar frequencies of TIGIT+ and PD-1+ cells on tetramer+ as on AIM+ Gag-specific cTfh cells, which were increased when compared to the
total cTfh population (Fig. S3d). In contrast, more HIV Gag-specific cTfh cells expressed CD200 when the AIM assay was used. These findings show that a high frequency of HIV-specific cTfh cells in ART-treated donors pre-express multiple IRs, such as PD-1 and TIGIT, before stimulation with the cognate Ag. On the other hand, while a fraction of HIV-specific cTfh cells pre-expresses CD200, this molecule is further rapidly upregulated during stimulation, as shown previously [34]. Therefore, a higher capacity to upregulate CD200 after stimulation might contribute to its increased expression on HIV- vs HBV- and CMV-specific CD4+ T cells.

In summary, our results demonstrate that despite viral suppression on ART, HIV-specific cTfh and non-cTfh CD4+ T cells are characterized by a high frequency of cells expressing multiple IRs.

3.3. HIV-specific cTfh cells produce higher levels of Tfh cytokines than CMV-specific cTfh

To elucidate whether high expression of IRs on HIV-specific cTfh influences Tfh function, we assessed the production of the canonical Tfh-related cytokines CXCL13 and IL-21. Detection of CXCL13 at the protein level is challenging due to their limited expression upon Ag stimulation and unspecific background staining. To overcome this hurdle, we used a modified ICS assay (“delayed ICS”), which included an extended (9 h) stimulation prior to addition of Brefeldin A, thus allowing for upregulation of AIM markers on the cell surface before the phase of intracellular protein trapping (Fig. 3a). We analyzed cytokine expression by a sequential gating strategy: pre-gating on CD69+CD40L+ cTfh cells increased specificity for CXCL13 and IL-21 and enabled robust detection of rare cytokine-expressing cells (Fig. 3b). We detected a significantly higher frequency of CXCL13- and IL-21-producing cells in HIV-specific compared to CMV-specific cTfh cells (Fig. 3c-d). Consistent with the known role of IRs for Tfh function [37], Tfh-cytokine production was highest in cTfh cells expressing multiple IRs (Fig. 3e-f). Therefore, HIV-specific cTfh cells expressing multiple IRs demonstrate robust expression of Tfh-related functions.

3.4. HIV-specific cTfh cells show preferential Th1-like phenotype and function

Co-expression patterns of the chemokine receptors CXCR3 and CCR6 have been associated with T helper cell differentiation and function.
and can also provide an indication of cTfh polarization and of their capacity to provide help to B cells [40]. We thus next examined CXCR3 and CCR6 expression on AIM+ Ag-specific cTfh cells in ART-treated individuals (Fig. 4a). HIV- and CMV-specific cTfh predominantly had a Th1-like (CXCR3+CCR6/C0) phenotype, while HBV-specific cTfh responses showed a mixed cTfh profile with Th2-like (CXCR3/CRC6+) and Th1-like polarizations (Fig. 4a, S4a). HIV-specific cTfh cells expressed in addition higher levels of Eomes and/or T-bet, whereas RORγt or GATA3 was rarely detected (Fig. S4bc), confirming their preferential Th1-like polarization. Importantly, the frequency of Th1-polarized cTfh was significantly higher for HIV than for both CMV and HBV (Fig. 4b).

In contrast, the proportion of CXCR3+CCR6+ Th1-like cells within the HIV-specific non-cTfh subset was comparable to CMV and HBV (Fig. S4d). In addition, HIV-specific non-cTfh cells included a high frequency of CCR6+ Th1Th17-like (CXCR5+CXCX3+) and Th17-like (CXX6+CXX3+) (Fig. S4d). Accordingly, we identified subsets of HIV-specific non-cTfh cells characterized by Eomes/T-bet and RORγt expression (Fig. S4e). Eomes and/or T-bet expression was significantly lower in HIV-specific cTfh than non-cTfh (Fig. S4f). These data suggest that the distinctive Th1-like phenotype skewing of HIV-specific cTfh compared to other specificities might correspond to a mixed differentiation pattern that does not reach the full acquisition of all Th1-related features seen in non-cTfh cells.

We next investigated whether the phenotypic polarization of HIV-specific cTfh correlated with function. We analyzed expression of classical Th1-cytokines in HIV- or CMV-specific cTfh cells (Fig. 4c). We detected a significantly higher frequency of IFNγ, TNFα and IL-2-expressing cells in HIV- vs CMV-specific cTfh (Fig. 4c-d). This profile of elevated Th1-cytokine expression was not seen for non-cTfh cells; indeed, IFNγ secretion was significantly reduced in HIV-specific non-cTfh compared to CMV-specific, while IL-2 production was increased (Fig. S4g). Consistent with previous findings at the bulk cTfh level [30], CXCR3+ HIV-specific cTfh populations (CXCR3+CXCX6-Th1-like, and to a lesser extent CXCR3+CXCX6-Th1/Th17-like) produced IFNγ following stimulation (Fig. 4e). However, a large fraction of Ag-specific CXCR3+ cTfh cells also produced the Th1-cytokine expression was not seen for non-cTfh cells; indeed, IFNγ secretion was significantly reduced in HIV-specific non-cTfh compared to CMV-specific, while IL-2 production was increased (Fig. 4e). Consistent with previous findings at the bulk cTfh level [30], CXCR3+ HIV-specific cTfh populations (CXCR3+CXCX6-Th1-like, and to a lesser extent CXCR3+CXCX6-Th1/Th17-like) produced IFNγ following stimulation (Fig. 4e). However, a large fraction of Ag-specific CXCR3+ cTfh cells also produced the Th1-cytokine expression was not seen for non-cTfh cells; indeed, IFNγ secretion was significantly reduced in HIV-specific non-cTfh compared to CMV-specific, while IL-2 production was increased (Fig. 4e). 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In contrast and as shown previously [41], cTfh cells, independent of their specificity, rarely expressed cytolytic markers such as CD107A, granzyme B or perforin (Fig. S5ab). These functions were, however, detectable in Ag-specific non-cTfh cells, especially for CMV (Fig. S5ab). Collectively, these observations demonstrate a preferential Th1-associated phenotype and function of HIV-specific cTfh cells in ART-treated individuals, contrasting with lack of cytotoxic molecule expression.

3.5. Suppression of HIV viremia by ART does not reverse the phenotype of HIV-specific cTfh cells

Having identified several distinct features of HIV-specific CD4+ T cell responses in ART-treated individuals, we next investigated whether these characteristics were present prior to ART initiation, and if they were modulated by therapeutic control of Ag load. For this, we examined samples from 7 HIV-infected donors obtained...
during the chronic phase of untreated infection before ART initiation and after 1 to 3.2 years of therapy (Participant characteristics: Table S2). We first determined the magnitude of HIV-specific CD4+ T cell responses and detected little change after starting therapy using the AIM assay, contrasting with a clear contraction of the HIV-specific CD4+ T cell response when the IFN-γ+ ICS was used (Fig. 5a).

In a transcriptional analysis of HIV-specific CD4+ T cells, we recently demonstrated that ART decreased the Tfh gene signature present in viremic individuals off therapy [4]. As this analysis was performed at the RNA level on total AIM+ HIV-specific CD4+ cells, we next investigated whether HIV-specific cTfh and non-cTfh were affected differentially at the phenotypic level by ART.

Although the frequency of cTfh cells within the total memory CD4+ T cell pool did not change following ART initiation, we observed an increase in the frequency of cTfh cells within the HIV-specific CD4+ T cell population (Fig. 5b). We detected a significant decrease in ICOS+ HIV-specific cTfh and non-cTfh cells (Fig. 5c), consistent with a general reduction of activation of HIV-specific CD4+ T cells. In addition, we detected a significant decrease in the frequencies of PD-1+, CD200+ and PD-1+TIGIT+CD200+ HIV-specific non-cTfh cells (Fig. 5d).

**Fig. 4.** HIV-specific cTfh are predominantly Th1-polarized. (a) Example plot showing CXCR3/CCR6 expression on HIV-specific cTfh. (b) Comparison of Ag-specific cTfh subsets based on CXCR3/CCR6 coexpression. (c) Example plots showing expression of IFN-γ in HIV- and CMV-specific cTfh. (d) Frequency of Th1-cytokine+ HIV- vs CMV-specific cTfh. (e) Frequency of HIV Gag-specific cTfh subsets expressing IFN-γ, CXCL13 or IL-21. (f) Co-expression analysis of Ag-specific cTfh cells based on CXCL13, IFN-γ and IL-21 expression. $n = 27$ (HIV), $n = 23$ (CMV) or $n = 6$ (HBV) for (b); $n = 12$ for (d), $n = 9$ for (e,f). P-values shown in figure were calculated by Kruskal–Wallis test with Dunn’s post test (b), Wilcoxon test (d, f) or Friedman test with Dunn’s post test (e).
In contrast, viral suppression on ART had a limited impact on expression and co-expression of IRs on HIV-specific cTfh cells (Fig. 5d). Similarly to IR expression, we did not detect changes in the CXCR3/CCR6 phenotype of HIV-specific cTfh cells following ART initiation (Fig. 5e), contrasting with a significant decrease in Th1-polarization of HIV-specific non-cTfh cells.

In summary, our results demonstrate that the phenotype of cTfh, characterized by a preferential Th1-polarized and high expression of multiple IRs, is established during chronic HIV infection and maintained despite suppressed viremia, this at least for the first three years after ART initiation.

3.6. Persistent levels of plasma HIV-specific antibodies in ART-treated individuals correlate with HIV-specific cTfh

Untreated HIV infection is associated with GC Tfh expansion and hypergammaglobulinemia [11]. Although HIV-specific Ab plasma levels decrease after ART initiation, they are detectable in the plasma of long-term treated individuals [42]. It has been suggested that ongoing Ag stimulation in lymphoid tissue may contribute to elevated HIV-specific plasma concentrations [43].

We therefore next assessed plasma levels of p24- and gp120-specific antibodies in our cohort of ART-treated individuals and how they relate to persistent Tfh responses. Despite their decrease after viral suppression on ART (Fig. 6a), HIV-specific Ab levels were detectable in nearly all ART-treated participants (Fig. 6b) and remained stable over time during ART (Fig. 6c). In addition, levels of p24- and gp120-specific Abs in plasma of ART-treated individuals were directly correlated (Fig. 6d). There was no association between total HIV-specific CD4+ T cell or HIV-specific non-cTfh responses and HIV-specific Abs (Fig. 6e-f). However, we detected a significant correlation between HIV-specific cTfh and persistent plasma HIV-specific Ab levels in ART-treated donors (Fig. 6g). Thus, our results suggest ongoing stimulation of HIV-specific B and T cells, which is associated with augmented Ab responses in ART-treated people in vivo.

3.7. The translation-competent HIV reservoir correlates with Th1-like phenotype and function of HIV-specific cTfh

As the phenotypic profiles of HIV-specific cTfh cells during untreated infection and during ART were similar, we next explored possible links between these maintained features detected in our
analyses and markers of HIV persistence during ART. We assessed the size of the HIV reservoir in CD4+ T cells using PCR for total viral DNA (vDNA), PCR for integrated vDNA and by measuring the inducible translation-competent reservoir by an RNA flow cytometric fluorescent in situ hybridization (RNA flow-FISH) assay. With this method, the translation-competent reservoir is defined as CD4+ T cells capable of co-expressing HIV gag RNA and Gag protein after in vitro stimulation (Fig. 7a) [24,25]. Translation-competent HIV RNA+/Gag+ CD4+ cells were undetectable in most ART-treated individuals in the absence of stimulation and in HIV-uninfected controls (Fig. 7b, Table S3), but could be identified in ART-treated donors after stimulation with PMA/Ionomycin with varying frequencies (Fig. 7b). DNA PCR-based methods detect defective viral genomes or proviruses unable to express viral proteins, and as expected the two PCR assays gave much larger estimates than the RNA flow-FISH method (Fig S6a). As low Ag persistence might
contribute to maintain the pool of virus-specific T cells on ART, we examined the relationship between reservoir size and magnitude of HIV-specific CD4$^+$ T cell responses. We observed no significant association between the frequency of HIV-specific CD4$^+$ T cells or cTfh responses and vDNA reservoir measurements but a non-significant correlation between the magnitude of the HIV-specific CD4$^+$ T cell response and the translation-competent reservoir (Fig. S6b).
We next explored potential relationships between the size of the total or integrated vDNA reservoir and phenotypic or functional features of HIV-specific cTfh or non-cTfh responses and found no significant association (Fig. 7c). In contrast, the size of the translation-competent HIV reservoir during ART was significantly positively associated with TIGIT, ICOS expression and the Th1-like phenotype and -function of HIV-specific cTfh (Fig. 7c, 56c-d). Of note, no association was detected between the size of the translation-competent reservoir and the phenotype or function of HIV-specific non-cTfh in ART-treated individuals (Fig. 7c). Therefore, our results indicate that the expression of IRs and Th1 functions of HIV Gag-specific cTfh cells correlate with a marker of the functional viral reservoir in individuals on ART. This suggests that the low but continuous levels of HIV Ag production may contribute to the maintenance of this population after prolonged therapy.

4. Discussion

Tfh cells are of high interest in HIV infection: at the total subset level, they serve as important sites of viral replication pre-ART [12] and reservoirs post-ART [13]. Studies of the total cTfh compartment show positive associations between memory Tfh cells and generation of HIV-specific broadly neutralizing antibodies (bNAb) [30,44,45], but investigations of HIV-specific cTfh have been hampered by the lack of sensitive tools to assess them. It remained unclear whether frequencies of HIV-specific Tfh cells, and/or altered functions, set them apart from Tfh specific for other viruses. Here, we used cytokine-independent AIM assays, alone or in combination with functional readouts, to characterize HIV-specific cTfh. We found that compared to their CMV- and HBV-specific counterparts in the same donors, HIV-specific cTfh, defined as CCR5+ memory cells, were more abundant and that this difference was exacerbated after ART initiation. On suppressive therapy, HIV-specific cTfh maintained a distinct phenotype with high expression of IRs, high production of cytokines, and a Th1-like skewing. Positive correlations between the magnitude, phenotype and function of HIV-specific cTfh responses with HIV-specific antibody levels and with the size of the translation-competent reservoir suggest that persistent exposure to viral products contribute to ongoing stimulation of HIV-specific Tfh despite undetectable HIV viremia on ART.

Our experimental approach gives a different picture of the impact of ART on HIV-specific CD4+ T cell responses compared to traditional assays such as IFN-γ ICS. HIV-specific CD4+ T cell responses were detectable in all ART-treated individuals examined by the AIM assay, in contrast to IFN-γ ICS. Matched samples obtained before and after ART initiation demonstrated a clear contraction of IFN-γ HIV-specific CD4+ T cell responses, whereas AIM+ responses were maintained. Therefore, the impact of ART on the magnitude of HIV-specific CD4+ T cells may have been overestimated in previous reports using IFN-γ as a readout [46]. Rather than massive attrition, smaller changes in frequency associated with more dramatic changes in polarization, differentiation and function of HIV-specific CD4+ T cells occur after viral suppression.

Many studies suggest that Tfh cells and GC play a critical role in the ability of the immune system to generate bNAb, which are uncommon, leading to the hypothesis that rarity and/or qualitative defects of HIV-specific Tfh could be limiting factors in the bNAb generation in infected humans [47]. Here, we demonstrate that, surprisingly, the median fraction of cTfh within the HIV-specific CD4+ T cell population in ART-treated people is large, 3.7-fold greater than in CMV-specific and 2.8-fold greater than in HBV-specific CD4+ T cells. The frequencies of HIV-specific cTfh observed by AIM assay were similar to the frequencies measured by tetramer staining in our experiments and to those reported by another group [18]. These results also confirm the capacity of the CD69/CD40L AIM assay to sensitively detect cTfh to chronic pathogens, besides its capacity to detect acute cTfh responses, as has been recently shown for the flu vaccine [20]. As these comparisons were made on an intra-individual basis, they control for the frequent persistence of an inflammatory environment despite suppressive ART and potential for bystander changes in T cell responses, therefore demonstrating actual virus-specific differences in cTfh responses. These expanded HIV-specific cTfh also present distinct phenotypic and functional features: they express higher levels of the IRs PD-1, TIGIT and CD200 compared to their CMV- and HBV-specific counterparts yet produce more cytokines. This augmented production includes not only the Th1 cytokines CXCL13 and IL-21, but also the Th1 cytokines IFN-γ, TNFα, and IL-2, consistent with the Th1-like phenotype of high CXCR3 expression. The production of Th1-associated cytokines CXCL13 and IL-21 was highest in Ag-specific cTfh cells expressing multiple IRs. Previous studied demonstrated the importance of PD-1 for optimal Tfh cell positioning and function including IL-21 production in vivo [37]. In this manuscript, we compare our findings related to HIV-specific CD4+ T cells to responses specific to CMV, which also causes a chronic infection albeit with some differences. The length of CMV infection in our cohort is likely greater when compared to HIV. However, CMV viral loads are usually undetectable in ART-treated non-immunocompromised individuals [48]. Life-long CMV Ag exposure can induce a particularly large T cell response, which is characterized by high Th1-related cytokine expression including cytolytic markers and expansion of senescent cells [49]. However, we detected these functions only in the CMV-specific non-cTfh population. No cTfh expansion nor preferential IR upregulation in CMV-specific CD4+ T cell responses was detected in our cohort, similar to HBV-specific responses induced by vaccination or resolved infections. In contrast, other studies observed a Th1 expansion and Th1-phenotype in other chronic infections that share some features with HIV, such as LCMV clone 13 in mice [5], SIV in non-human primates [50], HCV [8], and malaria [51], in humans. The underlying mechanisms are poorly understood, but several studies suggested that increased duration of Ag exposure, high levels of Ag and inflammatory cytokines, common for these infections, favor this polarization (reviewed in [52]).

Two non-mutually exclusive mechanisms may be responsible for maintaining the characteristics of virus-specific CD4+ T cell responses during viral suppression on ART. First, persistent low-level HIV Ag expression could continuously stimulate Tfh responses on ART and maintain HIV-specific cTfh phenotype and function. Importantly, such a mechanism does not imply residual replication of the full virus, as abortive expression of some viral genes or trapped Ag, for example on follicular dendritic cells, may have similar consequences [53]. HIV Ags can be detected in lymphoid tissue of ART-treated donors [54,55]. In support of this hypothesis, we detected a significant correlation between the size of the translation-competent reservoir and expression of TIGIT, ICOS and Th1-like phenotype and function of HIV-specific cTfh but not non-cTfh cells. In contrast, HIV DNA measurements, which include a large fraction of defective genomes unable to produce viral Ags [56], were not associated. One possible explanation to our observations is that occasional reactivation of persistently infected cells leads to stochastic production of viral proteins within the GCs of lymphoid tissues and stimulation of HIV-specific Tfh cells. These cells eventually exit the GC when the burst of HIV products resolves, and subsequently become cTfh cells. A second possibility of persistent characteristics of HIV-specific CD4+ T cells is permanent epigenetic programming. Previous studies demonstrated that during chronic viral exposure, virus-specific CD8+ T cells in mice and humans are epigenetically altered so that PD-1 expression was persistently elevated even after Ag withdrawal [57–60]. Thus, maintenance of certain phenotypic profiles in HIV-specific cTfh cells during ART could be related to an epigenetic reprogramming due to high levels of chronic Ag exposure during untreated infection.

We recently showed that HIV-specific CD4+ T cells in untreated HIV infection are characterized by an atypical Tfh signature in
Acknowledgments

We thank Joséé Girouard, the clinical staff at the McGill University Health Centre in Montreal and all study participants for their invaluable role in this project; Dr. Dominique Gauchat, Philippe St-Onge and the CRCHUM Flow Cytometry Platform as well as Dr. Olfa Debbeche and the CRCHUM BSL3 platform for technical assistance. The following reagent was obtained through the NIH Tetramer Core Facility: HLA Class II tetramers DRB1*01:01 DRYFKTLRAEQASQEV-PE and DRB1*11:01:YVDFFKTLRAEQASQEV-PE. Fig. S1A was prepared using images from Servier Medical Art by Servier, which is licensed under a Creative Commons Attribution 3.0 Unported License.

Funding Sources

This study was supported by the National Institutes of Health (no. HL02565 to D.E.K., no. AI100863 and no. AI144462 CHAVI-ID to D.E.K. (Consortium PI: Dennis Burton); the Canadian Institutes of Health Research grants (no. 137694 and no. 320721 to D.E.K.; foundation grant no. 352417 to A.F.; no. 36440 to N.C.); the Canada Foundation for Innovation Program Leader grant (no. 31756 to D.E.K.); and the FRQS AIDS and Infectious Diseases Network. D.E.K. is a FRQS Merit Research Scholar. N.C. is supported by a FRQS Research Scholar Career Award. J.N. is supported by scholarships from the FRQS, the Bavarian Research Alliance (BayFor) and the Department of Microbiology, Infectious Diseases and Immunology of the University of Montreal. A.E.B. is supported by a CIHR post-doctoral fellowship. J.R. is the recipient of a Mathilde Krim Fellowship in Basic Biomedical Research from AmfAR. J.T. is supported by a CIHR scholarship. J.P.R. is the holder of the Louis Lowenstein Chair in Hematology & Oncology, McGill University. A.F. is supported by a Canada Research Chair Award. The funding sources had no role in: study design; in the collection, analysis, and interpretation of data; in the writing of the report; and in the decision to submit the paper for publication.

Supplementary materials

Supplementary material associated with this article can be found in the online version at doi:10.1016/j.ebiom.2020.102727.

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