Immunotherapy during the acute SHIV infection of macaques confers long-term suppression of viremia

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We report that combination bNAb immunotherapy initiated on day 3 post-infection (PI) maintained durable CD8+ T cell–mediated suppression of SHIVAD8 viremia and preinoculation levels of CD4+ T cells in 9 of 13 treated monkeys during nearly 6 yr of observation, as assessed by successive CD8+ T cell–depletion experiments. In an extension of that study, two treatment interventions (bNAbs alone or cART plus bNAbs) beginning on week 2 PI were conducted and conferred controller status to 7 of 12 monkeys that was also dependent on control mediated by CD8+ cells. However, the median time to suppression of plasma viremia following intervention on week 2 was markedly delayed (85 wk) compared with combination bNAb immunotherapy initiated on day 3 (39 wk). In both cases, the principal correlate of virus control was the induction of CD8+ T cellular immunity.

Introduction
In most untreated individuals, HIV-1 infection leads to high levels of sustained virus replication, unremittent and irreversible depletion of CD4+ T cells, and ultimately symptomatic immunodeficiency with associated opportunistic infections, leading to a fatal clinical outcome. A small number of HIV-1 infected persons (elite controllers) are able to suppress viremia for prolonged periods of time, a phenomenon that is associated with the activity of virus-specific CD8+ T cells (Betts et al., 2006; Deeks and Walker, 2007; Pereyra et al., 2008).

Potent broadly neutralizing antibodies (bNAbs) have been used to prevent and treat lentivirus infections in macaques and humans (Borducchi et al., 2018; Caskey et al., 2015, 2017; Gautam et al., 2016, 2018; Hessel et al., 2016; Ledgerwood et al., 2015; Lu et al., 2016; Lynch et al., 2015; Moldt et al., 2012; Nishimura et al., 2017; Shingai et al., 2013). We previously reported that the administration of combination bNAbs alone during early simian immunodeficiency virus (SIV)/HIV chimeric virus (SHIV) infections of rhesus macaques can induce durable control of virus replication (Nishimura et al., 2017). In that study, 13 macaques were inoculated by the intrarectal (i.r.) or i.v. route with SHIVAD8-EO, and beginning on day 3 after virus challenge, they received a single course of three weekly i.v. infusions (on days 3, 10, and 17) of combination 10–1074 (Mouquet et al., 2012) plus 3BNC117 (Scheid et al., 2011) bNAbs. All of the treated monkeys controlled virus replication for varying periods of time (8–26 wk), and when bNAb levels waned, rebound viremia occurred in 12 of the 13 animals, reaching peak levels of 108 RNA copies/ml in several macaques. During an initial 100–140-wk observation period, six of these animals (three challenged i.r. and three challenged i.v.), designated “controllers,” suppressed plasma viremia to undetectable levels and durably maintained circulating CD4+ T cell numbers (Fig. 1). A CD8+ T cell–depleting mAb, specific for the CD8α chain, was administered to all six controller animals between weeks 48 and 131 after virus challenge (Fig. 1, A–F; left red arrows), resulting in transient bursts of plasma viremia that were rapidly resolved in five of the six monkeys. Quantitative virus outgrowth assays, performed before CD8+ T cell depletion, indicated that <1 cell/106 circulating CD4+ T cells was releasing infectious virions in these macaques. In the final reported experiment, anti-CD8β, which depletes CD8+ T cells but not natural killer, natural killer T, or γδ T cells, was administered to three controller animals between weeks 136 and 150 after SHIVAD8.EO challenge. Although not as potent in vivo as the anti-CD8α mAb, the anti-CD8β mAb infusion induced an immediate increase in plasma viremia (Fig. 1, A, D, and F; blue arrows). In that study, control macaques treated for 15 wk with a combination antiretroviral therapy (cART) regimen also initiated on day 3 after SHIV challenge, exhibited sustained...
plasma viremia upon treatment interruption, and none became virus controllers (Nishimura et al., 2017). These results indicated that passive combination bNAb immunotherapy, initiated during early acute infection, induces potent CD8+ T cell immunity able to durably suppress virus replication.

Here, we have extended these initial findings by (1) monitoring the virologic and CD4+ T cell status of the original “day 3” treated controller monkeys during an additional 2.5–3.5-yr period and (2) initiating a new immunotherapy regimen in which the effects of combination bNAb alone or cART plus bNAb treatment began at the more clinically relevant week 2 after the SHIV infection time point.

**Results**

**Combination bNAb immunotherapy initiated on day 3 postinfection (PI) suppressed SHIV
de replication in 9 of 13 macaques for 6 yr and was mediated by CD8+ T cells**

As shown in Fig. 1, A, B, and D–F, five of the original six controller animals continued to control plasma viremia between weeks 110 and 315 PI. 6 yr after the initiation of bNAb treatment in the day 3 infection cohort, five of the original six controller monkeys also maintained physiological levels of circulating CD4+ T cells (Fig. S1, A–E). As shown in Fig. 1 C, the sixth controller animal (DFIK), which did not exhibit virus rebound following the waning of the initial 3-wk bNAb treatment, was also unable to recontrol virus replication following the administration of the depleting anti-CD8a mAb at week 48 PI. The remaining five day 3 controllers received a final anti-CD8a mAb infusion at PI week 250 or 270, and all experienced another transient burst of viremia that was rapidly suppressed to background levels in four of the five animals (Fig. 1, A, B, and D–F; red arrows on right).

Seven of the original recipients of combination bNAb immunotherapy on PI day 3 had not durably suppressed plasma viremia by week 110 or 140 PI, when the original report was published, and were designated progressors. Interestingly, during the fourth and fifth years after bNAb immunotherapy initiation, three of these seven original progressors (DFFX, DFKX, and DEHW) suppressed virus replication and became late controllers (Fig. 2, A–C). As observed with the original six controller animals, infusion of depleting anti-CD8a mAb into the three late controllers at week 285 or 315 after the virus challenge also resulted in a transient burst of viremia. Three of the other four progressors developed clinical immunodeficiency and were euthanized between years 2 and 5 PI (Fig. 2, D–G). Thus, 9 of the original 13 animals administered combination bNAbs starting on day 3 PI, were able to control viremia in a CD8+ T cell–dependent manner after early antibody therapy.

Phylogenetic analysis of SHIV env gene sequences from the six early controllers collected during peak viremia after the first anti-CD8a antibody treatment (weeks 70–130 PI), from the three late controllers, and from the four progressors (both between weeks 110 and 150 PI) is shown in Fig. S2. Consistent with persistent viremia and viral replication, the env sequences from the four progressor monkeys exhibited the greatest diversity, while the six early controller animals were least diverse. The diversity of env genes from the late controllers was intermediate, an indicator of sustained, but weaker, CD8+ T cell–mediated pressure to suppress the ongoing viral infection.

**Initiation of bNAb only or cART plus bNAb treatment regimens on week 2 PI resulted in controller status in 7 of 12 monkeys**

To assess whether bNAb intervention at day 3 PI, which ultimately resulted in sustained control of virus replication in 9 of 13 macaques, would be effective when initiated at a more clinically relevant time, two groups of SHIV-infected monkeys were treated with bNAbs alone or with cART plus bNAbs starting at week 2 PI. The first cohort of six infected animals received three infusions at biweekly intervals of combination 10–1074 plus 3BNC117 bNAbs (10 mg/kg each) starting 14 d after infection, at the time of peak viremia (102–103 RNA copies/ml; Fig. 3 A). This bNAb regimen modestly suppressed the initial virus replication in five of the six animals by week 10 PI. In one monkey (DG34; Fig. 3 D), bNAb administration reduced plasma viremia below the level of detection, whereas immunotherapy was less effective in the other recipients. Following a virus rebound lasting 15 wk, macaque DG34 controlled SHIV replication to undetectable levels by week 30 PI (Fig. 3 D). Three other recipients of combination bNAb treatment beginning at week 2 also controlled virus replication, but only after 90–150 wk after infection (Fig. 3, B, C,
and E). Infusion of anti-CD8α mAb into these four controller monkeys at week 162 PI elicited transient bursts of plasma viremia, peaking at 10^6–10^7 RNA copies/ml (Fig. 3, B–E). Circulating CD4+ T cell levels in all four controllers were stable during the 3-yr observation period, although transient declines attending the CD8 depletions were observed in these monkeys (Fig. S1, F–I). The two progressor macaques in this treatment group continued to generate progeny virions at set point levels of 10^3–10^4 RNA copies/ml during the post-therapy period (Fig. 3, F and G), and one (DFA3) exhibited a slow continuous reduction of CD4+ T cells that was associated with a rising viral set point (Fig. S1 J).

The second group of six animals received daily cART therapy for 8 wk beginning at week 2 PI and a biweekly, three-infusion course of combination 10–1074 plus 3BNC117 bNAbs (10 mg/kg each) starting at week 9 PI; cART was discontinued at week 10 PI (Fig. 4 A). Peak levels of viremia ranged from 10^6 to 10^8 RNA copies/ml at the time of cART intervention and were rapidly suppressed to background levels between weeks 5 and 8 PI in all animals. The addition of combination bNAbs at week 9 extended the control of SHIVAD8-EO replication until weeks 17–26 PI, after which viral rebound occurred to the 10^4–10^5 viral RNA copies/ml range when antibody levels had declined below effective levels. Viremia was durably suppressed between weeks 70 and 160 PI in three of these macaques, which were designated controllers (Fig. 4, B–D). Administration of the depleting anti-CD8α mAb at week 200 to these three controller monkeys led to transient bursts of virus replication. As was observed with controllers induced by combination bNAb immunotherapy alone (Fig. S1, F–I), modest CD8 declines also accompanied the infusion of the anti-CD88 depleting antibody in the controller animals (Fig. S1, L–N).

The patterns of durable control were different when bNAb immunotherapy was initiated on day 3 or week 2 PI. Although bNAb treatment initiated on either day 3 or day 14 resulted in sustained suppression of SHIVAD8-EO infection in a similar fraction of macaques, there were important differences between the two cohorts. Although the level of viremia on day 3 PI was, at most, 10^3–10^4 RNA copies/ml when combination bNAb immunotherapy was started, the viral loads on week 2 were 10^7–10^8 RNA copies/ml. This difference was reflected in (1) a relatively rapid and nearly complete resolution of rebound viremia for controller monkeys receiving bNAbS beginning on day 3 PI (Fig. 1, A–F) versus (2) a very long and drawn-out process, requiring, in some cases, 2 or 3 yr to achieve control status for both cohorts that received bNAbs starting at week 2 (Fig. 3, B, C, and E; and Fig. 4, B–D). As shown in Fig. 5, this difference is reflected in the median times for bNAb-induced controller status: 38.7 versus 86.9 and 82.5 wk for interventions at day 3 (bNAbs only), week 2 (bNAbs only), or week 2 (cART plus bNAbs), respectively. The three additional day 3 late controller monkeys achieved controller status at a median time of 159 wk after infection.

Assessment of CD8+ T cells in the LNs of SHIVAD8 controller and progressor monkeys

As previously reported, controller or progressor monkeys that started bNAb immunotherapy on day 3 PI generated low levels of gp120 antibodies and minimal or no differences in anti-SIVmac239 Gag protein-mediated peripheral CD8+ T cell responses (Nishimura et al., 2017). To examine CD8+ T cell responses in lymphoid tissues in these treated macaques, we collected available superficial LNs from three controllers, two progressors, and four cART control animals on week 180 PI. As shown in Fig. S3 A, accumulation of follicular (f) CXCR5+ CD8+ T cells (Ferrando-Martinez et al., 2018; Petrovas et al., 2017) was observed in LNs in controller animals. Antiviral-specific CD8+ T cell responses were assessed by cytokine production and degranulation following stimulation with pooled SIVmac239 Gag peptides. The anti-viral responses in both bulk and follicular CXCR5+ CD8+ T cells, residing in LNs from the four controller macaques, were higher than those collected from progressor or cART control monkeys

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Journal of Experimental Medicine
https://doi.org/10.1084/jem.20201214
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However, this was not statistically different because too few animals were analyzed (e.g., $n = 2$ for the progressor macaques). Polyfunctional analysis revealed that the controller animals had higher levels of virus-specific CD69$^+$MIP-1$\beta^+$CD107a$^+$ and CD107a$^+$ single-positive CD8$^+$ T cells than progressors or cART control monkeys (Fig. S4). In contrast to animals administered bNAbs on day 3 PI, follicular CXCR5$^+$ CD8$^+$ T cells did not accumulate in the LNs of the controllers emerging following bNAb treatment.

Figure 3. Establishment of controller status by a single biweekly three-infusion course of combination bNAb therapy beginning at week 2 PI in rhesus macaques inoculated with SHIVAD8-EO. (A) Six rhesus macaques were inoculated i.r. with 1,000 TCID$_{50}$ of SHIVAD8-EO and treated with 10–1074 plus 3BNC117 (10 mg/kg of each) bNAbs at weeks 2, 4, and 6 PI (blue arrows), and their viral loads were monitored over a 170-wk period. (B–G) Plasma virus loads in four controller macaques (B–E) and two progressors (F and G) are shown. The red arrows indicate the times of anti-CD8$\alpha$-depleting mAb MT807RI infusion.

Figure 4. Establishment of controller status by successive cART therapy for 8 wk beginning at week 2 PI and combination bNAb treatment commencing on week 9 PI in rhesus macaques inoculated with SHIVAD8-EO. (A) Six rhesus macaques were inoculated i.r. with 1,000 TCID$_{50}$ of SHIVAD8-EO and initially received daily cART therapy for 8 wk beginning at week 2 PI (red bar) and were then administered combination 10–1074 plus 3BNC117 bNAbs (10 mg/kg each) at weeks 9, 11, and 13 PI (blue arrows). (B–G) Plasma virus loads in three controller (B–D) and three progressor (E–G) macaques are shown. Red arrows indicate the times of the anti-CD8$\alpha$-depleting mAb MT807RI infusion.
The different CD8+ T cell profiles in LNs from controller status are shown. 

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WK, starting on day 3 PI, completely suppressed virus replication and prolonged rebounds in the second cohort of monkeys, monkeys emerging following bNAb immunotherapy initiated only observed in a single monkey (DG34; Fig. 3 D), which rapidly controlled the initial viremia to this degree but also experienced a nearly 2-yr delay in achieving controller status (Fig. 4). It is worth noting that a latent virus reservoir had been established in several of the day 3 controller monkeys despite the very early initiation of immunotherapy (Nishimura et al., 2017). It is therefore quite likely that such a reservoir was not only present but also had significantly increased in size by week 2 PI, further impeding systemic suppression of virus replication when bNAb treatment was started. That said, the burst of viremia following CD8 depletion at week 200 in all seven controller animals treated with bNAbs ± cART starting at week 9 potently controlled the initial viremia to this degree but also experienced a nearly 2-yr delay in achieving controller status (Fig. 4). It is still too early to tell whether bNAb treatment during the early phase of HIV-1 infections will also mediate sustained virus control in humans. Important differences between both SHIV and HIV-1 pathophysiology (e.g., the median time to immunodeficiency; 2 yr [nonhuman primate] versus 10 yr [human], respectively) as well as variable host responses to immunotherapy make such predictions difficult. Furthermore, direct mechanisms responsible for the development of potent antiviral CD8+ T cell control of virus replication, nearly 4 yr after SHIV challenge, had been achieved.

Although we were unable to identify a distinctive T cell protective signature or mechanism conferring controller status on SHIV-infected rhesus macaques treated with combination anti–HIV-1 bNAbs starting on day 3 or week 2 PI, the consistent correlate of sustained virus suppression was the generation of CD8+ T cells, which persisted for 3–6 yr. In our earlier study, we reported that daily treatment of macaques with cART for only 15 wk, starting on day 3 PI, completely suppressed virus replication (Nishimura et al., 2017). However, all of these monkeys developed high levels of plasma viremia following the cessation of cART and none became controllers, indicating the critical role of bNAbs in conferring sustained repression of viremia.

Based on the data collected from 16 treated animals that became controller monkeys, we speculate that the establishment of long-term controller status in SHIV-infected monkeys may be facilitated and may be dependent on several critical factors. First, upon administration, bNAbs must potently suppress plasma viremia, preferably to undetectable levels, as was the case following their infusion on day 3 PI. When bNAb immunotherapy was initiated at week 2 PI, this degree of control was only observed in a single monkey (DG34; Fig. 3 D), which rapidly and potently suppressed virus replication that was sustained for an additional 2 yr. Second, although bNAb immunotherapy blocks neither virus replication nor progeny virion release from infected lymphoid cells, we previously proposed that bNAb administration in the presence of low levels of antigen production could drive immune complex formation and lead to dendritic cell activation to produce a vaccinial effect. Third, the continuing production of viral antigens, further augmented by the rebounding viremia attending the decline of bNAbs to subtherapeutic levels, could additionally simulate antigen cross-presentation from dendritic cells, stimulating potent and sustained CD4+ and CD8+ T cell responses.

As noted from the results presented, the different patterns in establishing controller status following initiation of combination bNAb immunotherapy on day 3 and week 2 PI most likely reflect the high levels of circulating and cell-associated virus production at week 2 PI superimposed on a background of more extensive and possibly irreversible injury to the immune system. It should also be noted that identical bNAb dosing regimens (10 mg/kg) were administered to both groups of macaques, rather than infusing larger amounts (e.g., 30 mg/kg) of mAb at week 2 PI. The relatively low bNAb dose given to the monkeys at week 2 PI may have simply been inadequate to effectively control viral loads peaking in the range of $10^7$–$10^8$ viral RNA copies/ml, leading to greater CD4+ T cell killing and the delayed development of a potent CD8+ T cell response. On the other hand, simply controlling viral loads to undetectable levels beginning at week 2 PI was insufficient in achieving controller status since the animals treated with cART between weeks 2 and 10 and then combination bNAbs starting at week 9 potently controlled the initial viremia to this degree but also experienced a nearly 2-yr delay in achieving controller status (Fig. 4). It is therefore quite likely that such a reservoir was not only present but also had significantly increased in size by week 2 PI, further impeding systemic suppression of virus replication when bNAb treatment was started. That said, the burst of viremia following CD8 depletion at week 200 in all seven controller animals treated with bNAbs ± cART beginning at week 2 PI (Fig. 3 and Fig. 4) clearly indicates that potent systemic CD8+ T cell control of virus replication, nearly 4 yr after SHIV challenge, had been achieved.

Discussion

Although we were unable to identify a distinctive T cell protective signature or mechanism conferring controller status on SHIV-infected human recipients of combination anti–HIV-1 bNAbs starting on day 3 or week 2 PI, the consistent correlate of sustained virus suppression was the generation of CD8+ T cells, which persisted for 3–6 yr. In our earlier study, we reported that daily treatment of macaques with cART for only 15 wk, starting on day 3 PI, completely suppressed virus replication (Nishimura et al., 2017). However, all of these monkeys developed high levels of plasma viremia following the cessation of cART and none became controllers, indicating the critical role of bNAbs in conferring sustained repression of viremia.

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It is still too early to tell whether bNAb treatment during the early phase of HIV-1 infections will also mediate sustained virus control in humans. Important differences between both SHIV and HIV-1 pathophysiology (e.g., the median time to immunodeficiency; 2 yr [nonhuman primate] versus 10 yr [human], respectively) as well as variable host responses to immunotherapy make such predictions difficult. Furthermore, direct mechanisms responsible for the development of potent antiviral CD8+ T cell responses remain to be elucidated. Nonetheless, consistent with the requirement for bNAb-induced CD8+ T cells for control of the robust SHIVAD8 infection of macaques, chronically HIV-1–infected human recipients of combination bNAb therapy also exhibited increased CD8+ T cell responses (Niessl et al., 2020). In that study, CD8+ T cell responses peaked at 6–7 wk following the initiation of immunotherapy when bNAb levels in the blood were at their highest levels. Although we were unable to identify a distinctive protective mechanism responsible for subsequent sustained control of virus replication following the administration of combination anti–HIV-1 bNAbs
during the acute SHIV infection of rhesus macaques, the induction of CD8+ T cells was required in controllers from all cohorts.

**Materials and methods**

**Animal experiments**

25 male and female rhesus macaques (Macaca mulatta, 13 from previous study; Nishimura et al., 2017) of Indian genetic origin, ranging from 2 to 4 yr of age at the time of challenge, were maintained in accordance with the guidelines of the Committee on Care and Use of Laboratory Animals and were housed in a biosafety level 2 facility. All animal procedures and experiments were performed according to protocols approved by the Institutional Animal Care and Use Committee of the National Institute of Allergy and Infectious Diseases, National Institutes of Health. Phlebotomies, virus inoculations, mAb and cART infusions, and sample collections were performed as previously described (Endo et al., 2000; Nishimura et al., 2017; Shingai et al., 2012). All animals were negative for the major histocompatibility complex class I Mamu-A*01, Mamu-B*08, and Mamu-B*17 alleles. Animals were not randomized, and the data collected were not blinded.

**Virus challenge**

The origin and preparation of the tissue culture–derived SHIVAD8-EO stock have been previously described (Shingai et al., 2012). 13 animals from the previous study (Nishimura et al., 2017) were challenged with SHIVAD8-EO i.v. (100 or 1,000 median tissue culture infectious dosage [TCID50]) or i.r. (1,000 TCID50) as previously described (Shingai et al., 2014). 12 new animals were challenged with 1,000 TCID50 of SHIVAD8-EO i.r.

**cART regimen**

Six animals were treated with a three-drug (combination) ART regimen comprising two nucleotide RT inhibitors (tenofovir and emtricitabine) and one integrase inhibitor (raltegravir), starting at week 2 for 8 wk. The preformulated cocktail of tenofovir and emtricitabine was administered intramuscularly (complete medium). Cells were frozen in FBS, 100 U/ml penicillin, 100 µg/ml streptomycin, and 2 mM L-glutamine (complete medium). Cells were frozen in FBS containing 10% DMSO and stored in liquid nitrogen until further use.

**Antibody administration**

A combination of 3BNC117 (Scheid et al., 2011) and 10–1074 (Moquet et al., 2012) anti–HIV-1 bNAb (10 mg/kg each) was administered i.v. to individual animals biweekly (at weeks 2, 4, and 6 for bNAb-only animals and at weeks 9, 11, and 13 for cART–plus-bNAb animals).

**Quantitation of plasma viral RNA levels**

Viral RNA levels in plasma were determined by quantitative RT–PCR (ABI Prism 7900HT sequence detection system; Applied Biosystems) as previously described (Endo et al., 2000).

**Lymphocyte immunophenotyping**

EDTA-treated blood samples were stained for flow cytometric analysis for lymphocyte immunophenotyping as previously described (Nishimura et al., 2010).

**Viral RNA Isolation**

Viral RNA extraction from plasma and RT was conducted as described with modifications (Imamichi et al., 2002). Briefly, viral RNA was isolated from macaque plasma using the QIAamp Viral RNA Isolation kit (Qiagen) and converted to cDNA using the Superscript III or Superscript IV First Strand cDNA synthesis kit (Thermo Fisher). cDNA was subject to PCR amplification for the gp120 region using Platinum PCR Supermix HiFi (Thermo Fisher) 5′-TCCTCTCTAGTACCCGC-3′ (nt 6129 forward) and 5′-GTGAGCTCAAGGCCCCAGTC-3′ (nt 9216 reverse) primers, followed by a nested PCR reaction using 5′-AAGGCAGAAAG CAGTTGAC-3′ (nt 6475 forward) and 5′-ATCCACCACAATT GCAGC-3′ (nt 8730 reverse) primers (Integrated DNA Technologies). Initial denaturation was performed at 94°C for 2 min, followed by 32 cycles of 20 s at 94°C, 30 s at 55°C, and 2 min 20 s (nested) or 3 min 20 s (primary) at 68°C, with a final extension for 7 min at 68°C. PCR products were ligated into PCRA-TOPO-TA vectors using the TOPO-TOA Cloning Kit (Thermo Fisher), and each ligation reaction was transformed into TOP10 Chemically competent Escherichia coli (Thermo Fisher) according to manufacturer’s instructions. Plasmid DNA was extracted and purified using the QIAPrep SpinPrep miniprep kit (Qiagen). Sequences were aligned using MacVector sequence analysis software (MacVector Inc.) starting from the first three codons of gp120 (ATGAAAGTG) and stopping after nt 8730 regardless of primers used. Sequence alignments were evaluated using the Model Tool in MEGA X for macOS (Stecher et al., 2020) to find the most appropriate substitution modeling parameters, for which the software recommended the Hasegawa-Kishino-Yano model with Gamma distribution and invariant rate among sites (HKY+G+I; Hasegawa et al., 1985). These recommended parameters were used to reconstruct a maximum likelihood tree in MEGA X inferred via extensive subtree pruning and regrafting (SPR level 5) and 1,000 bootstrap replicates for branch support. The resulting tree was colorized using the Rainbow Tree tool (Los Alamos National Laboratory), and bootstrap values >70% are indicated at the start of each clade. The viral sequences recovered from infected macaques have been deposited in GenBank under accession numbers MT864360–MT864599.

**LN processing**

LN biopsies were placed in RPMI-1640 medium. After removal of fat and connective tissue, the tissue was cut into small pieces and dissociated with the plastic end of a 6-mL syringe plunger. Cell suspensions were passed through a 40-µm cell strainer and washed twice in RPMI-1640 medium supplemented with 10% FBS, 100 U/mL penicillin, 100 µg/mL streptomycin, and 2 mM L-glutamine (complete medium). Cells were frozen in FBS containing 10% DMSO and stored in liquid nitrogen until further use.

**Functional assays of CD8+ T cells in LN**

Thawed LN cells were suspended in complete medium for 1 h at 37°C/5% CO2. The cells were then stimulated for 6 h at 37°C/5% CO2 with 4 µg/mL SIV Gag peptide pool in the presence of monensin (GolgiStop; BD Biosciences), brefeldin A (Sigma), and anti-CD107a antibody (clone H4A3; Biolegend) for the
assessments of cellular degranulation. The unstimulated condition (with DMSO instead of SIV Gag peptide pool) was included for all samples. For each functional readout, the frequency of SIV Gag-specific stimulated cells was background (unstimulated condition) subtracted.

**Flow cytometry for intracellular cytokine staining**

Cells were surface stained with Aqua amine viability dye (In-vitrogen) for 10 min at room temperature, followed by surface staining with directly conjugated and biotinylated antibodies for 15 min at room temperature and streptavidin-BV421 (Biorad) incubation for 15 min at room temperature. Washes between and following the surface staining steps were performed in FACS buffer (PBS containing 2% FBS). Cells were fixed and permeabilized in Cytofix/Cytoperm (BD Biosciences) for 30 min at 4°C, washed in Perm/Wash buffer (BD Biosciences), and stained intracellularly with directly conjugated antibodies for 30 min at 4°C.

The cells were washed in Perm/Wash buffer and re-suspended in FACS buffer before data acquisition. Samples were acquired on a modified BD FACSymphony (BD Biosciences) equipped with 355-, 405-, 488-, 532-, and 628-nm lasers. Data were analyzed using FlowJo software version 9.9.6 (BD Biosciences). The antibodies used for intracellular cytokine staining are shown in Fig. S5. The gating strategy for functional readouts was evaluated in either total CD8+ or fCD8+ T cells (identified as live CD3+CD8+CCR7-CD95hiCXCR5hi fCD8+ cells) as shown in Fig. S5. Within each of these cell populations, the functional readouts were evaluated as either coexpression of CD69 and cytokine (IFN-γ, TNF, or MIP-1β) or total CD107a expression. The polyfunctionality analyses were performed using the Simple Presentation of Incredibly Complex Evaluations software v.6.0 (Vaccine Research Center, National Institute of Allergy and Infectious Diseases, National Institutes of Health).

**Online supplemental material**

Fig. S1 shows CD4+ T cell levels in macaques treated with bNabs alone or cART plus bNabs. Fig. S2 describes env gene diversity in SHIV-infected macaques following initiation of bNab immunotherapy on day 3 Pl. Fig. S3 demonstrates that follicular CXCR5+ CD8+ T cell accumulation and SIV-specific CD8+ T cell responses in LNs from the controller macaques occurred following combination bNab therapy beginning on day 3 Pl, but not in the controller macaques following bNab ± cART treatment beginning at week 2 Pl. Fig. S4 displays polyfunctional profiles of bulk CD8+ and follicular CXCR5+ CD8+ T cell responses to SIV Gag in the LNs of day 3 controller, progressor, and cART-only treated monkeys. Fig. S5 lists the antibodies used for intracellular cytokine staining and the gating strategy used to define CCR7hiCD95biCXCR5bi fCD8+ cells in flow cytometry datasets.

**Acknowledgments**

We thank A. Peach, P. King, and B. Yankulova for determining plasma viral RNA loads and sequencing and R. Petros, C. Ramera, A Bruce, K Werku, and E. Sanford for diligently assisting in the maintenance of animals and assisting with procedures.

This work was supported by the Intramural Research Program of the National Institute of Allergy and Infectious Diseases, National Institutes of Health. The research was also funded in part by the Bill and Melinda Gates Foundation Collaboration for AIDS Vaccine Discovery grants OPP1092074 and OPP124068. In addition, National Institute of Allergy and Infectious Diseases of the National Institutes of Health grant HIVRAD P01 AI100148, National Institutes of Health Center for HIV/AIDS Vaccine Immunology and Immunogen Discovery grant 1UM1 AI100663-01, National Institutes of Health/National Institute of Allergy and Infectious Diseases grant P01 AI138212, and the Robertson Foundation of Rockefeller University (all to M.C. Nussenzweig) funded this research. M.C. Nussenzweig is a Howard Hughes Medical Institute investigator.

Author contributions: Y. Nishimura, M.C. Nussenzweig, and M.A. Martin conceived of the study and wrote the manuscript. Y. Nishimura, M.C. Nussenzweig, R. A. Koup, R. Gautam, and M.A. Martin designed the research and analyzed the data. O.K. Donau, J. Dias, S. Ferrando-Martinez, E. Jesteadt, R. Sadjapour, and A. Buckler-White performed the experiments. R. Gelezuiunas provided antiretroviral drugs.

Disclosures: The authors declare no competing interests exist.

Submitted: 11 June 2020
Revised: 6 August 2020
Accepted: 18 August 2020

**References**


Figure S1. CD4+ T cell levels in macaques treated with bNAbs alone or cART plus bNAbs. (A–Q) Absolute numbers of circulating CD4+ T cells in five controller macaques following bNAb therapy beginning on day 3 PI (A–E); in four controller (F–I) and two progressor (J and K) macaques following bNAb therapy alone beginning at week 2 PI; and in three controller (L–N) and three progressor (O–Q) macaques after cART plus bNAb therapy beginning at week 2 PI are shown.
Figure S2. **Env gene diversity in SHIV-infected macaques following initiation of bNAb immunotherapy on day 3 PI.** An unrooted phylogram of env gene sequences displays phylogenetic diversity in controllers (red) at weeks 70–130 PI, in late controllers (green rectangles) at weeks 110–150 PI, and in progressors (blue) at weeks 110–150 PI. Diversity of 0.01 is indicated. Branch lengths were measured by the number of substitutions per site. Bootstrap numbers >70% are indicated at the start of each clade.
Figure S3. Follicular CXCR5+ CD8+ T cell accumulation and SIV-specific CD8+ T cell responses in LNs from the controller macaques occur following combination bNAb therapy beginning on day 3 PI, but not in the controller macaques following bNAb ± cART treatment beginning at week 2 PI. (A–D) Treatments were initiated on day 3 (A and B) or at week 2 (C and D). (A) Follicular CXCR5+ CD8+ T cells accumulated in LNs of day 3 controller (C) animals (n = 4) compared with the same T cell subset collected from progressor (P) macaques (n = 2) or control monkeys treated with cART only, starting at day 3 (n = 4). Comparable fractions of bulk CD8+ T cells were present in LNs from the three groups of monkeys. The Kruskal-Wallis test followed by Dunn’s multiple comparisons test showed a statistically significant difference in levels of fCD8+ T cells between controller and progressor monkeys (P = 0.0228). (B) Antiviral responses in both bulk and follicular CXCR5+ CD8+ T cells in LNs from controller macaques were higher than in LNs from progressor or cART-only control macaques. The CD8+ T cell responses to SIVmac239 Gag were measured by intracellular cytokine staining (IFN-γ, MIP-1β, TNF, and CD107a). (C) The fractions of follicular CXCR5+ CD8+ T cells in LNs from controller (n = 7) and progressor (n = 4) animals were comparable. (D) Antiviral responses in both bulk and follicular CXCR5+ CD8+ T cells in LNs from controller macaques were comparable to the responses measured in progressor animals. The CD8+ T cell responses to SIVmac239 Gag were measured by intracellular cytokine staining (IFN-γ, MIP-1β, TNF, and CD107a). Error bars indicate SD of data from multiple animals.
Figure S4. Polyfunctional profile of bulk CD8+ and follicular CXCR5+ CD8+ T cell responses to SIV Gag in the LNs of day 3 controller, progressor, and cART-only monkeys. The pie charts and graphs show the mean frequency of cells performing all different combinations of effector functions. Graphs also show the individual data points. C, controller macaques; P, progressor macaques.
Antibodies for intracellular cytokine assays

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Gating strategy (representative example)

Figure S5. Antibodies used for intracellular cytokine staining and gating strategy used to define CCR7loCD95hiCXCR5hi fCD8+ cells in flow cytometry datasets. Representative flow cytometry plots from one representative animal are shown. Functional readouts were evaluated as either coexpression of CD69 and cytokine (IFN-γ, TNF, or MIP-1β) or total CD107a expression. Ab, antibody; ECD, electron coupled dye; FSC, forward scatter; SSC, side scatter.