Attacking Latent HIV with *convertible* CAR-T Cells, a Highly Adaptable Killing Platform

**Graphical Abstract**

**Highlights**
- Inert MICA-fused Ab and *convertible* CAR-T cells (cCAR-T) only kill when combined
- cCAR-T decreases tumor size in a mouse lymphoma model as efficiently as scFv CAR-T
- cCAR-T kills HIV-infected primary cells with high efficiency and specificity
- cCAR-T reduces the inducible reservoir in blood of HIV+ individuals by 50%–60% in 48 h

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**In Brief**
An adaptable CAR-T cell platform based on cytotoxic lymphocytes engineered to bind a variety of broadly neutralizing anti-HIV antibodies can effectively kill HIV-infected primary cells and reduce viral reservoirs in the blood of infected individuals on antiretroviral therapy.
Attacking Latent HIV with convertibleCAR-T Cells, a Highly Adaptable Killing Platform

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SUMMARY

Current approaches to reducing the latent HIV reservoir entail first reactivating virus-containing cells to become visible to the immune system. A critical second step is killing these cells to reduce reservoir size. Endogenous cytotoxic T-lymphocytes (CTLs) may not be adequate because of cellular exhaustion and the evolution of CTL-resistant viruses. We have designed a universal CAR-T cell platform based on CTLs engineered to bind a variety of broadly neutralizing anti-HIV antibodies. We show that this platform, convertibleCAR-T cells, effectively kills HIV-infected, but not uninfected, CD4 T cells from blood, tonsil, or spleen and only when armed with anti-HIV antibodies. convertibleCAR-T cells also kill within 48 h more than half of the inducible reservoir found in blood of HIV-infected individuals on antiretroviral therapy. The modularity of convertibleCAR-T cell system, which allows multiplexing with several anti-HIV antibodies yielding greater breadth and control, makes it a promising tool for attacking the latent HIV reservoir.

INTRODUCTION

The main obstacle to curing HIV-infected individuals is the existence of a reservoir of latently infected cells that persists despite long-term antiretroviral therapy (ART). These rare cells harbor an integrated HIV provirus but generally do not express viral proteins, making them invisible to the immune system and difficult to eliminate (Chun et al., 1997; Finzi et al., 1997; Richman et al., 2009; Wong et al., 1997). The establishment of latency may have various causes, including the infection of partially active cells transitioning to a resting state (Chun et al., 1995; Siliciano and Greene, 2011), the activation of a virus-encoded program (Razooky et al., 2015), or even the infection of resting CD4 T cells (Cameron et al., 2010). Regardless of the origin of the latent reservoir, the greater its size, the harder it is for the immune system to control viral levels (Lori et al., 1999), and the faster the virus rebounds in the event of ART interruption (Li et al., 2016). Although the advent of combination ART has revolutionized management of HIV infection and prevents development of AIDS, HIV-positive individuals must adhere to lifelong treatment, not without adverse side effects (Renju et al., 2017). Despite falling short of complete viral eradication, a consistent ability to reduce the size of the viral reservoir and control its activity to achieve a sustained viral remission in the absence of ART would radically alter approaches to the treatment of infection around the world (Goulder and Deeks, 2018; Lori et al., 1999).

Some HIV-positive individuals who received treatment early during acute infection exert long-lasting control negating viral rebound after withdrawal of ART. In these rare individuals, termed post-treatment controllers (PTCs) (Hocqueloux et al., 2010; Sáez-Cirión et al., 2013), the immune system manages to keep the latent reservoir small and sometimes even reduces the reservoir size and controls viremia without drug intervention (Sáez-Cirión et al., 2013). The reasons for PTCs’ better control of HIV load are still under investigation, but levels of CD4 T cell activation are lower in these individuals than in non-controllers. CD4 T and natural killer (NK) cell responses are also higher and levels of inflammation are low (Sáez-Cirión et al., 2013; Samri et al., 2016). We seek an intervention that routinely establishes post-treatment control in all HIV-infected people from both developed and developing countries.

One of the main approaches to reducing the size of the reservoir is “shock and kill” (Archin et al., 2012). This approach entails exposing cells to one or more latency reversing agents (LRAs) to induce viral gene expression, ideally with little or no toxicity to the host (Jean et al., 2019). Once successfully shocked, reservoir cells begin producing the viral protein Env, which is inserted on the cell surface. These cells may now be killed either due to a viral cytopathic effect or by immune cells recognizing as foreign the viral Env protein/peptides expressed on the surface of the
reservoir cells (Jones and Walker, 2016; Leonard et al., 1988). However, only strong LRAs lead to cell killing via viral-induced cytopathic effects, and their clinical utility is quite limited because of their toxic side effects, including potential triggering of a cytokine storm (Chun et al., 1999; Prins et al., 1999). More subtle, non-toxic compounds would be desirable, but they may only stimulate a small fraction of the latent reservoir necessitating serial administration (Cillo et al., 2014; Jean et al., 2019). Regardless of the way reservoir cells are reactivated, the success of the shock-and-kill strategy crucially depends on an efficient means of killing the reactivated cells.

HIV infection is commonly associated with the exhaustion of cytotoxic T lymphocytes (CTLs) manifested by loss of both effector function and proliferative capacity (Cella et al., 2010; Hersperger et al., 2010; Kalams et al., 1999; Shin and Wherry, 2007). In addition, CTL-resistant viral strains emerge, especially if treatment with ART is delayed beyond acute infection (Deng et al., 2015; Jain et al., 2013; Shan et al., 2012; Yang et al., 2002). Any attempt to boost the cell-mediated immune response of HIV-infected individuals must overcome these two hurdles.

Our work focuses on designing a more reliable way to kill the reactivated cells. Such killing is pivotal for reducing the size of the reservoir and critical for implementing a “reduce-and-control” strategy that, together with an immune intervention, might lead to post-treatment control. Two options for killing include: (1) improve the activity of the resident CTLs (Migueles et al., 2009), or (2) introduce new CTLs (Sung et al., 2018). Several broadly neutralizing antibodies (bNAbs) are now available that react with infected cells (Julg and Barouch, 2019; Mayer et al., 2017; McCoy, 2018; Sok et al., 2016). These bNAbs have the ability to recognize HIV-infected cells and contribute to their neutralization but still need a functional immune system to execute the bNAb-directed killing (Bar-On et al., 2018; Caskey et al., 2017). New CTLs have also been introduced by first removing patient T cells, inserting into these cells a chimeric antigen receptor (CAR) against a specific surface target along with costimulatory and signaling domains (e.g., 4-1BB and CD3ζ) (Guedan et al., 2018) and then reinfusing the CAR-T cells into the patient (Pule et al., 2003). In the most successful application of this CAR-T cell approach, cells were engineered to express an anti-CD19 single chain fragment variable (scFv), which specifically recognizes the CD19 antigen (Onea and Jazirehi, 2016) expressed on malignant cells from patients with refractory B cell lymphoma (Jacobson, 2019; Onea and Jazirehi, 2016; Zheng et al., 2018). Although successful, one of the problems of current CAR-T cells is that once administered, the cells are always in an “on-state” thus control is limited, and there is little to no way to halt the activity of these cells should serious autoimmune side effects emerge (Bonifant et al., 2016; Zheng et al., 2018).

Similar strategies have been attempted to engineer CAR-T cells against HIV, by fusing scFv part of an anti-HIVEnv antibody (Sung et al., 2018) (see Figure 1E) or the extracellular domains of the CD4 molecule that recognize HIV Env, to the zeta chain of the T cell receptor creating a CAR-T cell (Deeks et al., 2002). Although well-tolerated in humans, these CAR-T cells failed to produce clinical effects potentially due to lack of an adequate host response to LRAs (Wagner, 2018). The use of a single targeting motif on these CAR-Ts significantly limits the ability of these cells to address HIV’s diversity and high rate of viral epitope drift in patients. A more versatile platform that is inherently more responsive to viral fluidity is therefore needed to maximize the impact of a CAR-T-based strategy.

CTLs and NK cells express on their surface the NK group 2D receptors (NKG2D) that recognize a family of ligands overexpressed on cells stressed by viral infection or transformation (Cosman et al., 2001). The ζ1-ζ2 domains of these major histocompatibility complex class I chain-related (MIC) ligands—that include MICA, MICB, and UL16 binding proteins (ULBPs) 1–6—bind to NKG2D that activates the cytolytic function of these cells (Bauer et al., 1999; Cosman et al., 2001; Steinle et al., 2001) (see Figure 1A). We leveraged the natural binding of MIC/ULBP ligands to NKG2D to develop an exclusive orthogonal ligand-receptor interaction to generate the components of a highly modular universal CAR-T cell platform (for more thorough biochemical characterization and application of the platform in other fields see U.S. patent 10,259,858 and Landgraf et al. [2019]). The engineered extracellular domain of the NKG2D variant was tethered to the intracellular 4-1BB and CD3ζ co-signaling domains to generate the CAR while the mutant ligand domain was fused to an antibody to generate a bispecific molecule termed a MicAbody. This two-part system, which we term convertibleCAR-T (cCAR-T) cells, can readily be multiplexed with several distinct MicAbodies distinguished by their ability to engage unique epitopes.

We reasoned that by combining bNAbs and cCAR-T, we might harness their individual benefits to expand the breadth of killing of HIV-infected targets while minimizing the emergence of viral resistance. We have explored the killing properties of these cCAR-T cells against both productively HIV-infected cells from human blood, tonsil, and spleen and reactivated blood CD4 T cells obtained from ART-suppressed HIV-positive individuals. We describe our findings, concluding this platform shows great potential as a robust approach to reduce the reservoir size.

RESULTS

HIV bNAbs Fused to an Orthogonal MIC Ligand Direct T Cells Expressing an Inert NKG2D-Based CAR Construct to Recognize HIV-1-Infected Primary Cells

To combine bNAbs and CAR-T cells, we took advantage of a natural ligand-receptor system that normally participates in the body’s surveillance for malignant and virally infected cells. This system uses a ligand from the MIC/ULBP family, expressed on stressed cells, and its receptor NKG2D, expressed on CTL and NK cells (Figure 1A). The binding pocket of the NKG2D extracellular domain was mutated to render it inert and incapable of binding its natural ligands (Figure 1B). Compensatory mutations in the ζ1-ζ2 domain of a MIC ligand were introduced to eliminate binding to wild-type NKG2D and promote sub-nanomolar engagement of the mutant NKG2D (Figure 1F). This established a unique and exclusive orthogonal pairing system in which the mutant NKG2D was subsequently formatted as a CAR for expression on CD8 T cells (cCAR-T) and the mutant ζ1-ζ2 fused to HIV bNAbs (Figure 1C; STAR Methods) to generate bispecific
agents called MicAbodies. The mutant z1-α2 exclusively engages the mutant NKG2D-CAR expressed on the CAR-T cells (Figure 1G; Table S1), while the Fv domains engage the cognate antigen of the HIV-Env protein on the surface of tonsil-derived infected cells (Figure 1H). This high avidity and specific interaction generates an immunologic synapse that activates the cytotoxic effector functions of the CAR-T cells to kill the targeted cell.

Four MicAbodies from two types of HIV-specific bNAbs were constructed. The first two bNAbs, 3BNC60 and 3BNC117, recognize the CD4 binding site on the virus envelope (Malbec et al., 2013; Scheid et al., 2011, 2016); the other two, PGT121 and 10-1074, are HIV-V3 glycan loop domain-binding bNAbs (Malbec et al., 2013; Mouquet et al., 2012; Walker et al., 2011) (Table S2). Both types of bNAbs were shown to be potent in clinical trials when used alone or in combination (Bar-On et al., 2018; Caskey et al., 2017, 2019; McCoy, 2018).

To test the specificity of CAR-T killing, we constructed two additional MicAbodies as negative controls: one based on the B cell-specific, anti-CD20 monoclonal antibody, rituximab (Maloney et al., 1997), and the other on the breast cancer-specific monoclonal antibody, anti-HER2 or trastuzumab (Herbst and Hong, 2002) (Table S2). Using fluorophore-conjugated MicAbodies, we demonstrated specific binding of MicAbodies to the HIV-infected (GFP+) tonsil cells in a comparable manner to natural bNAb (Figure 1H).

Finally, to ensure that the effect of our system on infected cells would reflect specific binding to the CAR-T cells, we introduced two mutations (D265A/N297A) in the MicAbody's heavy chain (Table S2). These mutations abolish the MicAbodies' ability to trigger cell killing via antibody-dependent cellular cytotoxicity (ADCC) often involving NK cells as effectors (Shields et al., 2001). Adding these mutations helps to ensure that killing is only triggered by the MicAbody-CAR-T interaction, resulting in a safe, modular, and regulatable killing platform.

We first tested the platform in vivo in a subcutaneous lymphoma mouse model, where NSG mice were injected with Raji lymphoma cells. After tumor implantation, the mice were injected with rituximab MicAbodies (20 µg) every 2 days for 6 doses, and...
A

Relative CD4 Viability

E:T Ratio: 0 1 5 10 20

500pM HIV MicAbody mix

B

Relative CD4 Viability

E:T Ratio: 0 1 5 10 20

10pM HIV MicAbody mix

C

Relative CD4 Viability

No CAR-T 0.1 +cCAR-T 0.1 No CAR-T 0.1 +cCAR-T 0.1

3BNC60 (nM) +cCAR-T 3BNC117 (nM) +cCAR-T PGT121 (nM) +cCAR-T 10-1074 (nM) +cCAR-T

D

Relative CD4 Viability

No CAR-T 0.1 +cCAR-T 0.1 No CAR-T 0.1 +cCAR-T 0.1

3BNC60 (nM) +cCAR-T 3BNC117 (nM) +cCAR-T PGT121 (nM) +cCAR-T 10-1074 (nM) +cCAR-T

(legend on next page)
cCAR-T cells were injected 1 day after the first MicAbody dose. Tumor volumes were regularly monitored by caliper measurements. Conventional CAR-T (CD19 scFv) and cCAR-T armed with rituximab MicAbodies displayed similar levels of control of tumor cell growth. Delivery of cCAR-T without MicAbody produced no significant anti-tumor effects (Figure 1).

**cCAR-T Cells Combined with HIV MicAbodies Specifically Kill HIV-Infected Primary CD4 T Cells**

Although viral load is commonly tested in blood, and a large number of latency studies have been performed using blood cells (Shacklett et al., 2019), it is important to explore HIV infection and latency in lymphoid tissues because most of the reservoir resides there, and replication of the virus mainly occurs in these tissues (Chun et al., 2008; Haase et al., 1996; Pantaleo et al., 1993; Yukl et al., 2010). We have principally used tonsil-derived cells for infection with HIV-GFP in this study (Doitsch et al., 2010) (Figure S1; STAR Methods). Principal findings have also been confirmed in both spleen and peripheral blood cells.

To determine the optimal effector-to-target ratio, we combined cCAR-T cells and tonsil-derived cells at various effector: target (E:T) ratios in the presence of a mix of the four HIV-specific MicAbodies and assessed killing efficiency by measuring the depletion of HIV-infected CD4 T cells (HIV-positive cells were monitored by the expression of the GFP reporter; see also Figure S2 for gating strategy). For each experiment, one million infected tonsil cells (containing \( \sim 1 \times 10^5 \) GFP+ HIV-infected cells) were incubated for 48 h with a range of cCAR-T cells from no-CAR-T to 2 \( \times 10^5 \) CAR-T cells (in the no-CAR-T controls, untransduced CD8 T cells from the same donor were added). All experiments were conducted in the presence of ART (saquinavir, 5 \( \mu \)M) to prevent a spreading infection that could potentially confound the killing results.

We found that the killing efficiencies of cells infected with HIV-1 CCR5 (R5) tropic virus (BaL/GFP) correlated with the number of effector cCAR-T cells present. These results were consistent regardless of the concentration of MicAbodies tested (Figures 2A and 2B; GFP+ bars). Although specific killing of HIV-infected cells (GFP+) improved as we added more effector cells by an E:T ratio of 20:1, the viability of the uninfected bystander CD4 T cells began to decrease both with high and low MicAbody concentrations (Figures 2A and 2B; GFP− bars). Based on these results, we selected an E:T ratio of 10:1 for all of subsequent experiments.

Next, a number of control experiments were performed to assess the specificity of cCAR-T killing HIV-infected cells. To confirm that the killing of target cells depended on HIV recognition, we tested unarmed cCAR-T or cCAR-T armed with the B cell-specific MicAbody rituximab or the anti-HER2 MicAbody, trastuzumab. No detectable killing of HIV-infected CD4 T cells was observed with these control MicAbodies (Figure 2C, left). Only when an HIV-specific MicAbody was added to the cCAR-T was a reduction in the number of HIV-infected, GFP+ tonsil cells observed (Figure 2C, right). Infected CD4 T cell number was reduced by \( \sim 60\% \) and specific killing was observed with all four HIV MicAbodies at doses spanning two orders of magnitude (0.1–10 nM). We conclude that cCAR-T cells armed with HIV-specific MicAbodies can successfully kill HIV-infected tonsil cells.

To determine whether in-well toxicity or non-specific cCAR-T toxicity was contributing to reducing cell viability, we monitored the non-HIV-infected CD4 T cells (GFP− cells) in each well. No decrease in viability of these uninfected CD4 T cells was observed in any of the wells (Figures 2A–2C, gray bars, GFP−). We conclude that cCAR-T cells armed with an HIV-specific MicAbody selectively kills HIV-1-infected cells without affecting uninfected bystander cells. To test a second lymphoid tissue, we performed killing experiments on cells derived from the spleens of healthy donors, finding similar results (Figures S3).

In ~50% of AIDS patients, the R5 tropic virus converts to an X4 (CXCR4) tropic virus (Schulz et al., 1992, 2011). We therefore tested the ability of cCAR-T cells to kill tonsil-derived CD4 T cells infected with X4-NL4-3, a GFP-tagged X4-tropic virus (Leyv et al., 2004). We found that cCAR-T killed these cells efficiently and selectively when combined with the Env/C4D4-binding MicAbodies (3BNC60/117), which are known to neutralize both R5-tropic and X4-tropic viruses (Bruel et al., 2016). However, when cCAR-T cells were combined with the Env/V3-loop binding MicAbodies, known to preferentially neutralize R5-tropic over X4-tropic viruses (Bruel et al., 2016), the killing was reduced (Figures 2D and S4). We conclude that the type of bNAb selected for MicAbody construction is critical in the context of X4-tropic viral infections. These observations support the importance of being able to multiplex the arming of the cCAR-T cells.

**Figure 2. Specific Killing of Primary CD4 T Cells Infected with HIV by cCAR-T Combined with HIV Env-Specific MicAbodies**

To determine the optimal effector to target cell ratio, one million tonsil-derived cells (\( \sim 1 \times 10^5 \) HIV/GFP-infected target cells) were incubated with a range of cCAR-T effector cells from zero (0:1) to \( 2 \times 10^5 \) (20:1) cCAR-T cells: target cells for 48 h with the mix of four HIV Env-specific MicAbodies (Mix). In the absence of cCAR-T cells (0:1); the donor-matched untransduced CD8 T cells were present. Live GFP+/CD3+/CD8+ cells were counted to assess off-target killing (GFP−). (A and B) HIV-MicAbody mix was tested with high (500 pM) (A) or low (10 pM) (B) concentration of each individual MicAbody. Data derived from three independent experiments; mean ± SEM. (C and D) R5 tropic HIV-1 (BaL) (C) or X4 tropic HIV-1 (NL4-3) (D) were used to infect tonsil cells followed by testing of individual HIV-specific MicAbody for specific killing by cCAR-T cells. One million tonsil-derived cells (\( \sim 1 \times 10^5 \) infected cells) were incubated with \( 1 \times 10^5 \) cCAR-T cells for 48 h, in the presence of different concentrations (0.1–10 nM) of HIV Env-specific MicAbodies. B cell-specific MicAbody (Ritux) and anti-HER2-specific MicAbody (HER2) were used as negative control MicAbodies. Results are presented relative to the no MicAbody control. For each individual MicAbody, an internal control of no cCAR-T supplemented with the highest MicAbody concentration tested is presented. To assess off-target killing or generalized in-well toxicity, viability of GFP+ CD4 T cells was assessed. This experiment was performed four times using cells from independent donors. Data are represented as mean ± SEM. *p \( \leq 0.05 \), **p \( \leq 0.01 \), ***p \( \leq 0.001 \) (compared to no cCAR-T presence).

See also Figures S1, S2, S3, and S4.
Laboratory-adapted HIV strains, such as R5-BaL and X4-NL4-3 used above, are less predictive of in vivo outcomes than strains mediating transmission between HIV-positive individuals, also known as transmitted/founder (T/F) viruses, which consistently display CCR5 co-receptor tropism (Keele et al., 2008; Li and Chen, 2019; Parrish et al., 2013; Schwartz et al., 2018). To test our system in a more clinically relevant setting, we infected primary cells with a T/F virus, 109FPB4 (F4) (Cavrois et al., 2017; Neidleman et al., 2017). Our previous experiments with the R5-BaL and X4-NL4-3 strains used MicAbodies in the nanomolar (nM) concentration range, all of which induced similar levels of killing (Figure 2). To identify the minimum amount of MicAbody sufficient to trigger CAR-T-mediated killing, we assessed the killing of T/F infected cells in the presence of picomolar (pM) MicAbody concentrations. After 48 h of incubation, dose-dependent killing of up to 65% of infected cells was observed with three of the four HIV-specific MicAbodies over a 10–500 pM concentration range (Figure 3A). In the case of the 10-1074-based MicAbody, no detectable killing of T/F HIV-infected cells occurred at any of the MicAbody concentrations. These results show that the cCAR-T cell platform can recognize and efficiently kill primary cells infected with both laboratory-adapted and T/F HIV strains, but depending on which virus the CD4 T cell harbors, killing may vary depending on the specificity of the selected MicAbody. Moreover, our results show that picomolar MicAbody concentrations are sufficient to trigger robust killing of infected cells by the cCAR-T cell platform.

**cCAR-T Cell-Mediated Killing Correlates with HIV Gene Expression Levels in Target Cells**

Although highly specific, cCAR-T cell-mediated killing consistently plateaued at ~65% (Figure 3A). We sought to determine whether this plateau reflects a differential sensitivity of the infected cells to cCAR-T cells, possibly correlating with the levels of viral gene expression occurring in the infected cells. In these studies, cells were infected with HIV strains that express GFP as a multiply spliced RNA under control of the viral LTR (Kutsch et al., 2002; Neidleman et al., 2017). In these viruses, GFP is inserted in the Nef coding region. Nef is expressed at physiological levels (Neidleman et al., 2017) and mediates CD4 downregulation of infected cells (Figure S2C). Therefore, GFP expression is a quantitative marker of overall HIV gene expression. To assess a potential correlation between levels of HIV expression and sensitivity to cCAR-T killing, we gated...
HIV-infected cells based on their GFP fluorescence level (low or high) and tested their response to cCAR-T killing. While CD4 T cells with high GFP levels were readily eliminated by the cCAR-T system (killing efficiency up to \( \approx 90\% \)) (Figure 3B), many CD4 T cells with lower GFP levels survived exposure to the cCAR-T cells (killing efficiencies of \( \approx 30\% \)) (Figure 3C). We conclude that killing is correlated with the amount of HIV gene products the infected cells produce and most likely with the amount of Env protein they express on their cell surface. This is consistent with the observation that targeted antigen density is a major determinant in CAR-T cell efficacy (Walker et al., 2017; Watanabe et al., 2015).

**Single-Cell Time-Lapse Microscopy Shows a Delay in bNAb-Armed cCAR-T Cell-Killing Kinetics**

The cCAR-T system requires a “three-body collision”: the cCAR-T cell, the MicAbody and the infected cell to create the cytolytic immunologic synapse. In order to monitor the killing kinetics, we used time-lapse microscopy. We incubated \( 3 \times 10^6 \) primary cells infected with HIV-F4 with a mix of the four HIV-specific MicAbodies in the presence of \( 3 \times 10^5 \) cCAR-T cells on a \( \mu \)-Dish slide. We then monitored the number of HIV-positive cells every 30 min for 48 h. As observed in our end-point experiments (Figures 3 and S5), cCAR-T cells killed 90% of the HIV-infected cells when the pool of MicAbodies was added (Figure 4). Interestingly, the killing did not commence until 10–15 h after the initiation of the cultures. These findings suggest a requisite period of time for the effector cells to arm with MicAbodies and successfully collide with their target cells.

**Multiplexing Two MicAbody Types Results in Specific Killing of Two Different Types of Target Cells**

HIV-1 has proven to evade every single-drug or single-antibody therapy, regardless of their level of initial efficacy. Thus, the use of more than one bNAb-based MicAbody will likely be required to prevent the emergence of resistant viruses over time. Because
the cCAR-T system is a two-part modular system, one of its advantages is the ability to multiplex with different MicAbodies, thus attacking the virus-infected cells with more than one type of bNAb. To confirm effective multiplexing with cCAR-T, we armed these cells with two different types of MicAbodies, one specific for B cells (rituximab) and the other corresponding to a mix of HIV-specific MicAbodies and assessed their killing properties in cultures containing a mixture of HIV-1-infected primary cells and ~10% B cells. The killing of B cells was not diminished by co-arming with HIV-specific MicAbodies (Figure 5A). Similarly, the presence of the B cell-specific MicAbody did not significantly impair killing of HIV-infected cells (Figure 5B). We conclude that multiplexing MicAbodies is possible within the cCAR-T system.

**Ex Vivo cCAR-T Cell Killing of Reactivated Reservoir Cells Present in the Blood of HIV-Infected Individuals on Long-Term ART**

We next turned to the effects of cCAR-T cells in the context of the latent HIV reservoir. In the prior in vitro primary-cell-based experiments, we used cells from the lymphoid tissues of healthy individuals productively infected ex vivo with the various HIV strains. Because CD4 T cells from blood can only be productively infected after activation (Muñoz-Arias et al., 2015), we compared cCAR-T cell killing of tonsil-derived and activated blood-derived CD4 T cells, and found that both were killed with a similar efficiency (Figure S5). To investigate potential effects of cCAR-T on the latent HIV reservoir, CD4 T cells were isolated from the blood of six HIV-positive individuals on suppressive ART (characteristics of the study participants are shown in Table S3). These cells were treated with a strong LRA (PMA + ionomycin) and then cultured with cCAR-T cells armed with HIV MicAbodies at two concentrations. Because the specific virus genotype was not known, we used a mixture of the four HIV-specific MicAbodies. Two days after co-culture, RNA was extracted from the cells (cell-associated RNA [caRNA]) and HIV-RNA levels were assayed by droplet digital PCR (ddPCR) using an HIV-specific probe (Shan et al., 2013) (Figure 6A). We observed a significant reduction in the amount of HIV RNA in cultures including cCAR-T cells and MicAbodies, but not in cultures including MicAbodies and untransduced parental CD8 T cells or unarmed CAR-T cells (Figure 6B). Our results, showing superior killing with lower MicAbody concentration (0.1 nM), compared to 1 nM, are probably the result of the pro-zone effect reducing killing efficiency with higher antibody concentrations (Vaidya et al., 2017). These results indicate that it is possible to attack the inducible latent reservoir with bNAb-armed cCAR-T cells. The use of this ex vivo system provides a proof-of-concept validation for the ability of cCAR-T cells armed with HIV-specific MicAbodies to reduce the reactivated latent reservoir.

**DISCUSSION**

A Modular CAR-T Cell Platform Armed with HIV bNAbs that Recognizes and Kills HIV-Infected Cells

Many attempts to cure HIV are focusing on “shock and kill” with the goal of activating virus expression in the latent reservoir followed by immune clearance of the reservoir cells, all performed under the cover of ART to prevent viral spread. Results with the first generation of LRAs, including histone deacetylase inhibitors (HDACi) and protein kinase C (PKC) activators are disappointing due mainly to a lack of potency and/or unacceptable toxicity (Prins et al., 1999; Rasmussen and Lewin, 2016). The fact that CTLs present in HIV-infected individuals exhibit an exhausted phenotype and diminished killing capacity is also of concern as is the potential presence of CTL-resistant viruses within the latent reservoir. A delayed start of ART certainly sets the stage for the emergence of such resistance.

To ensure highly effective killing of reactivated reservoir cells, we developed and tested a platform, cCAR-T cells, based on the binding of a mutant NKG2D receptor to an orthogonal MIC ligand fused to broadly neutralizing HIV antibodies (MicAbodies). The exclusive interaction engineered into this ligand-receptor pair ensures that on its own each component is functionally inert as was shown both in vitro and in vivo. As a consequence, neither the cCAR-T cells nor the MicAbody are able to kill target cells until they have bound to each other and the MicAbody has specifically engaged its epitope on an HIV-infected target cell to create an immunologic synapse. This system robustly incorporates many features that not only provide targeting flexibility and
specificity but also enhanced controllability and safety. During HIV infection, soluble ligands (e.g., MICA and ULBP) for the natural NKG2D receptor are expressed but cleaved from the surface of infected cells, thereby compromising the ability of CTL and NK cells to kill infected cells (Matusali et al., 2013). The mutation of the MIC ligand and NKG2D receptor such that each is unable to bind their natural substrates circumvents this problem. Additionally, the MicAbody has been rendered ADCC-deficient with mutations in the Fc region that disrupt the binding to Fcγ receptors present on NK cells (Shields et al., 2001). In summary, these modular features ensure that the cCAR-T cells and the MicAbodies are unable to kill target cells until they are bound to each other and the MicAbodies have specifically contacted an HIV-infected target cell to create the lethal immunologic synapse.

We are able to infect different lymphoid tissues and blood CD4 T cells with a variety of HIV strains and then reduce the number of productively infected cells using cCAR-T cells armed with an HIV-specific MicAbody. Importantly, we can also use these cCAR-T cells to attack latent reservoir cells reactivated ex vivo, removing 50%–60% of the inducible reservoir within 48 h. Furthermore, killing by cCAR-T is specific for infected cells and occurs only when an HIV-specific MicAbody is present. Although killing efficiency correlates with viral protein expression, we predict that levels of Env present in vivo will be sufficient for recognition by the cCAR-T platform because these reactivated reservoir cells where shown to be bound to and be enriched using the same HIV bNAbs we are using (Cohn et al., 2018).

Up to 90% of primary human cells infected at high levels with HIV can be killed using the cCAR-T cell system. Killing is dose-dependent and requires the addition of only picomolar quantities of MicAbodies. This killing rate is similar or superior to that previously reported for in vitro CAR-T killing of HIV-infected cells CD3ε-,CD28-,CD19-, and 1 μM ionomycin. After incubation for 2 days with cCAR-T cells or matched untransduced CD8 T cells and a mix of four different HIV-specific MicAbodies (either 0.1 or 1 nM of each MicAbody), cell-associated RNA (caRNA) was extracted, and HIV RNA was quantitated by droplet digital PCR (ddPCR).

(B) HIV RNA quantification results normalized to HIV RNA levels present in CD8 T cells and 1 nM HIV antibody mix. This experiment represents studies of 6 HIV-infected individuals. Data are represented as mean ± SEM. NS = p > 0.05, *p ≤ 0.05, **p ≤ 0.01. See also Figure S5 and Table S3.

The cCAR-T Cell Platform Increases the Flexibility and Safety

Treatment of HIV-infected individuals with a single anti-retroviral drug almost invariably leads to the development of drug resistance, an observation that prompted the introduction of combination antiviral therapy. Administering a CAR-T specific for a single antigenic target (Sotillo et al., 2015) or a single bNAb (Caskey et al., 2017) could also select for emergence of resistant virus and loss of therapeutic benefit. To overcome this limitation, recent clinical trials with bNAb-based therapies are combining two strong bNAbs (10-1074 and 3BNC117) (Bar-On et al., 2018). However, classic CAR-T cells, as currently implemented, carry only a single anti-HIV scFv domain. Deploying a second targeting domain requires designing a second CAR-T cell, which is cumbersome and costly. However, in the cCAR-T platform, multiplexing with two or more HIV-specific MicAbodies is easily accomplished. Indeed, our studies indicate that cCAR-T cells can be multiplexed with different antibodies promoting an effective attack on two different types of cells simultaneously.

cCAR-T cells containing the 10-1074 bNAb were unable to kill F4-infected cells but efficiently killed R5-BaL-infected cells. Our
HIV-specific CAR-T platform is based on broadly neutralizing antibodies that can recognize and neutralize hundreds of strains, although a single antibody falls short of recognizing all strains (Eroshkin et al., 2014). This observation again strongly argues for arming cCAR-T cells with more than one bNAb-based MicAbody to increase the breadth of killing. The general ability to multiplex antibodies in the cCAR-T platform provides for great flexibility and an increased level of effectiveness. Additionally, the picomolar quantities needed for cytolyis supports the possibility of even higher levels of multiplexing if needed, leading to the possibility of implementing a single, defined cocktail of MicAbodies to achieve extremely broad coverage using a single CAR construct.

Another advantage of the cCAR-T system is that as a two-part system, it can be introduced into patients in an inert state. Only after the administration of the specific MicAbody or MicAbody mix will the cCAR-T start killing. This feature should allow clinicians to control both the dose and timing of administered HIV-specific MicAbody. By contrast, classical CAR-T cells are “on” all the time. Using a platform that is inert unless both parts are present promises to enhance safety and avoid some of the problems limiting the applicability of the current CAR-T systems in oncology, including increased cytokine production and CAR-T killing of unintended healthy host cells expressing low levels of the tumor target (Fedorov et al., 2013; Kochenderfer et al., 2012; Ma et al., 2019).

To further increase the safety of cCAR-T cells, one could envision specifically activating or suppressing the cCAR-T cells in vivo, taking advantage of their exclusive ligand–receptor interaction. In theory, any molecule fused to the mutated MIC-ligand binding domain will deliver the heterologous molecule specifically to its cognate NKG2D partner on cCAR-T cells. An example would be linking interleukins to the MIC ligand to preferentially promote cCAR-T growth and survival (McGill et al., 2010; Richer et al., 2015; Younes et al., 2016). Of note, CAR-T cells can persist in vivo in HIV-infected individuals for more than 10 years after administration (Scholler et al., 2012). If cCAR-T cells prove equally durable, they could become a silent reservoir that clinicians could tap into when the need arises by simply delivering exclusively to the cCAR-T cells activation and proliferation signals along with the condition-specific MicAbody. Similarly, it should be possible to deliver specific suppressing molecules to rapidly or transiently silence the cCAR-T cells in case of adverse side effects or deliver a kill signal if rapid termination of cCAR-T activity is desired.

Developing Affordable and Universal CAR-T Cells

In spite of the therapeutic potential of cCAR-T cells, its application in the clinic still faces important hurdles. The first one is the cost of such a treatment for a single patient. Whereas producing the MicAbodies is affordable and easily scalable, removing cells from a patient followed by isolation, expansion, and transduction of these autologous cells with the mutated NKG2D-CAR receptor followed by their expansion and reinfusion is both expensive and requires an additional degree of technical expertise (Sarkar et al., 2018). In addition, the long process from bedside to laboratory and then back to bedside makes deploying the current cCAR-T treatment difficult to envision in developing countries, many of which carry crippling burdens of HIV disease (Kharsany and Karim, 2016; Wang et al., 2016).

Several efforts are underway to reduce the cost of CAR-T therapies and make it more scalable. One high priority is to create allogeneic “universal donor” cells (Ruella and Kenderian, 2017; Torikai and Cooper, 2016) to provide off-the-shelf cells that can be used for all patients (Graham et al., 2018). To achieve such universal donor cells, the T cell receptor (TCR) complex can be disabled to prevent graft-versus-host disease (Yang et al., 2015) and host-versus-graft elimination can be blocked by mutations within β2-microglobulin that prevent MHC-I surface expression (Ren et al., 2017). Modern techniques for cell editing, including CRISPR, transcription activator-like effector nucleas (Talen), and Zinc finger nucleases (ZFNs), offer potential ways to knock out those key cell-surface receptors responsible for graft-versus-host and host-versus-graft reactions (Osborn et al., 2016; Ren et al., 2017). Another promising approach involves the use of induced pluripotent stem cells (iPSCs) (Takahashi et al., 2007) as the source of the CAR-T cells. In this method, iPSCs are engineered to express the CAR prior to being differentiated into T cells (Themeli et al., 2013). These cells will act as effector cells in vivo, as has already been observed for hematopoietic stem/progenitor cells (HSPCs), which go through T cell development and selection in vivo (Zhen et al., 2017). It is now possible to convert iPSCs to various immune cells including T cells that contain chimeric antigen receptors (Lee, 2019; Themeli et al., 2013). Such cells can be engineered to be HLA-compatible for large populations thus dramatically reducing cost and increasing scalability of the CAR-T approach (Xu et al., 2019). Such universal donor cells could be readily adapted to the cCAR-T platform in view of its simplicity and modular design.

The Urgent Need for Safe and Effective LRAs

The other important hurdle to overcome before cCAR-T cells can be effectively tested in HIV-infected patients is the lack of safe and effective LRAs. To use cCAR-T in the kill phase of a “shock and kill” strategy, an effective shock must be delivered to render the latently infected cells visible to the immune system. The most commonly used LRAs include HDACi and PKC or PTEFb activators (Prins et al., 1999; Rasmussen and Lewin, 2016). Furthermore, several types of LRAs, such as HDACi and PKC activators, exert suppressive effects on the cytolytic function of CTLs and by analogy would probably suppress activity of cCAR-T as well (Walker-Sperling et al., 2016). Nonetheless, approaches to improve shocking strategies, by lowering toxicity of some LRAs or administrating a weak LRA in multiple dosing, have shown some promise (Ke et al., 2018). Recently, combining new regimens of LRAs including toll-like receptor 7 (TLR-7) agonists (Jiang et al., 2018) and the PGT121 broadly neutralizing HIV antibody in SHIV-infected rhesus macaques resulted in prolonged times to viral rebound or in some cases no rebound (Borducchi et al., 2018). Of note, the TLR-7 agonist appears able to activate...
HIV-specific CTLs (Tsai et al., 2017), making it a preferable LRA for cCAR-T applications. Another emerging type of LRAs is the second mitochondria-derived activator of caspases (SMAC) mimetics that act through activation of the non-canonical nuclear factor-κB (NF-κB) pathway and show promising LRA capabilities without the pleiotropic effect seen with other LRAs (Pache et al., 2015; Sampey et al., 2018). One apparent negative feature of all of these suggested LRAs is the small fractional response they elicit (Battivelli et al., 2018; Cillo et al., 2014; Jean et al., 2019). In order to reach a robust level of reactivation, repetitive administration of the LRAs will be required. Using a controllable platform such as the cCAR-T could ensure high activity of the killer cells at the right time and inert circulating killer cells between LRA and MicAbody administrations.

cCAR-T Cells as a Component of a Reduce-and-Control Strategy
Absent a means for the safe and complete eradication of HIV in infected individuals, a reduce-and-control strategy seems to be an attractive option. This strategy involves both a reduction in the size of the reservoir and the introduction of an immune intervention that allows the infected host to control the virus despite the removal of ART. In its purest form, this strategy would imitate the natural, but rare state found in post-treatment controllers (PTCs). One of the hallmarks of post-treatment control is the presence of a small reservoir. A smaller reservoir size has been shown to be a good predictor of the ability of HIV-positive individuals to become PTCs (Sáez-Cirión et al., 2013). Our ex vivo results show that we can reduce the reactivated reservoir size by 50% over the course of a 2-day experiment. Introducing cCAR-T cells and refueling their killing activity with multiple, spaced doses of MicAbodies might provide a powerful way to prevent reservoir expansion in vivo, while maintaining the patients on ART would prevent the spread of infection by the reactivated cells. Future studies will concentrate on developing MicAbodies with different HIV-specific bNAbS, testing combinations of MicAbodies and evaluating the cCAR-T platform in vivo. Studies are being carried out now to test cCAR-T cells and MicAbodies in mice, and future experiments are envisioned in experimentally infected macaques prior to moving to human testing. In summary, the cCAR-T cell system draws together exciting advances in both broadly neutralizing HIV antibody biology and CAR-T cell technology, to create a promising killing platform for attacking the latent HIV reservoir.

STAR*METHODS
Detailed methods are provided in the online version of this paper and include the following:

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  - Mice
  - Primary-cell cultures
  - Cell line
  - Virus strains
- **METHOD DETAILS**
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  - cCAR-T production
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  - Culture and infection of primary cells
  - HIV-1 infection
  - Reactivation of cells from HIV-positive individuals
  - cCAR-T cell killing assay
  - Measurement of cell-associated RNA by ddPCR
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  - Flow cytometry
  - in vivo assay
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- **DATA AND CODE AVAILABILITY**

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.cell.2019.10.002.

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AUTHOR CONTRIBUTIONS

DECLARATION OF INTERESTS
None of the Gladstone/UCSF scientists have a commercial relationship with Xyphos Biosciences. K.C.K. is an employee of Xyphos Biosciences, S.R.W. is an employee and shareholder of Xyphos Biosciences, K.L. and N.K. are members of the Xyphos Biosciences scientific advisory board, and D.W.M. is a founder of Xyphos Biosciences, a shareholder, and member of its scientific advisory board. A patent application has been filed related to this work.

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# STAR METHODS

## KEY RESOURCES TABLE

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**Experimental Models: Cell Lines**

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**Experimental Models: Organisms/Strains**

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LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for reagents should be directed to and will be fulfilled by Dr. Warner C. Greene (warner.greene@gladstone.ucsf.edu). Plasmid sequences for the cCAR-T construct (which includes the mutant NKG2D receptor detailed in this manuscript) and the bNAb-MicAbodies (including the orthogonal ULBP2 variant) will be made available upon request. Purified MicAbodies can be generated upon execution of a material transfer agreement (MTA) with inquiries directed to Dr. Kaman Kim (kaman@xyphosinc.com).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Human samples
Blood from HIV-infected individuals was obtained from volunteers participating in the SCOPE cohort (Hunt et al., 2003). Participants gave their informed consents as part of the SCOPE cohort. Specific characteristics of these participants and their ART regimens are summarized in Table S3.

Mice
Six-week old female NSG mice (Jackson Laboratories, Bar Harbor, ME) were housed and studied in strict accordance with the Institutional Animal Care and Use Committee (IACUC). Studies were performed by ProMab Biotechnologies, Inc. (Richmond, CA).

Primary-cell cultures
Human healthy tonsils and spleens were obtained from the Cooperative Human Tissue Network (CHTN, https://www.chtn.org). Human lymphoid aggregate culture (HLAC) prepared from tonsil or spleen was cultured in HLAC medium: RPMI supplemented with 15% heat-inactivated fetal bovine serum (FBS), 100 μg/ml gentamicin, 200 μg/ml ampicillin, 1 mM sodium pyruvate, 1% nonessential amino acids, 2 mM L-glutamine, and 1% fungizone, at 37°C in 5% CO₂ incubator.

Concentrated white blood cell preparations from healthy volunteers were obtained from Vitalant (https://www.vitalant.org) or from STEMCELL Technologies. PBMCs were cultured in RPMI supplemented with 10% FBS, 1000 U/ml Penicillin and 1 mg/ml Streptomycin and 2 mM L-glutamine, at 37°C in 5% CO₂. cCAR-T and parental CD8 cells were cultured in T cell medium (X-Vivo 15 media, 5% human AB serum; 10 mM neutralized N-acetyl-L-Cysteine, 0.1% 2-mercaptoethanol, supplemented with 40U/ml IL-2). No personal identifiers were provided for either the uninfected lymphoid tissues or blood samples.

Cell line
Female HEK293T cells were transfected with various molecular clones of HIV to produce high titer virus preparations. Female Exp293 cells were used for protein expression. Lenti-X 293T cells were used for lentivirus production. Cells were cultured in DMEM supplemented with 10% FBS, 1000 U/ml Penicillin and 1 mg/ml Streptomycin and 2 mM L-glutamine, at 37°C in 5% CO₂.

Virus strains
HIV molecular clones (pNL4-3-GFP, pBaL-GFP, and pF4-GFP) were purified from E.coli and used to transfect HEK293T cells.

METHOD DETAILS

Protein Expression and Purification
The orthogonal MIC variant was cloned as a C-terminal fusion to the human kappa light-chain via an APTSSSGGGGS linker. Additionally, D265A/N297A (Kabat numbering) mutations were introduced into the CH2 domain of the heavy chain of all antibody and MicAbody clones to reduce binding to all FcγR receptors in order to eliminate antibody-dependent cell cytotoxicity (ADCC) function. Cognate heavy- and light-chains for each bNAb clone were generated by swapping out the VH and VL domains (Table S2). Heavy- and light-chain plasmid DNAs (in the mammalian expression vector pD2610-V12, ATUM) for a given antibody clone were co-transfected into Exp293™ cells and purified by Protein A resin.

cCAR-T production
Leukopaks from healthy anonymous donors (STEMCELL Technologies) were used for primary T cell isolation. After a brief rinse in PBS + 2% FBS, cells were resuspended at 5x10⁷ cells/ml in PBS + 2% FBS and CD8⁺ cells were enriched by negative selection (STEMCELL Technologies Human CD8 T Cell Isolation Kit). 50 μl of isolation cocktail was added per ml of cells and, after a 5 min room temperature incubation, 50 μl of RapidSpheres™ (Stemcell) were added per ml of cells and total volume adjusted with PBS (to 35 ml total for each 21 ml of cells). Cells were isolated by applying an EasySep™ magnet for 10 min followed by transfer of buffer containing negatively enriched cells to new tubes for a second round of purification. Cells were then resuspended, counted, and cryopreserved at 10-15x10⁶ cells/cryovial.

The extracellular domain of the mutant NKG2D was cloned with the CD8α signal sequence, hinge and transmembrane domains from CD8α, and the intracellular signaling domains from 4-1BB and CD3ζ into the pHR-PGK transfer plasmid for second generation
Pantropic VSV-G pseudotyped lentivirus production along with packaging plasmids pCMVdR8.91 and pMD2.G as previously described (Roybal et al., 2016). For each batch of lentivirus produced, the three plasmids (pHR-PGK, 7.2 μg; pCMVdR8.91, 12.9 μg; and pMD2.G, 2.5 μg) were combined with 720 μl Opti-MEM™ (Fisher) then mixed with Fugene-HD (Promega) before adding to 6x10⁶ Lenti-X 293T cells (Takara Bio) that had been seeded one day prior in a 10 cm dish. Two days post-transfection, supernatants were collected by centrifugation and passed through 0.22 μm filters. SX concentrated PEG-6000 and NaCl are added to achieve final concentrations of 8.5% PEG-6000 (Hampton Research) and 0.3 M NaCl, incubated on ice for 2 h, then spun at 3500 rpm at 4°C for 20 min. Concentrated viral particles were resuspended in 0.01 volume of PBS, and stored frozen at −80°C.

Prior to lentiviral transduction, one vial of cryopreserved CD8 T cells was thawed and diluted into 10 ml of T cell medium “TCM” (X-Vivo 15 media, Lonza; 5% human AB serum, Corning; 10 mM neutralized N-acetyl-L-Cysteine, Sigma-Aldrich; 1X 2-mercaptoethanol, Thermo Fisher; 30 IU/ml human IL-2, R&D Systems). Cells were centrifuged at 400 x g for 5 min, resuspended in 10 ml TCM, adjusted to 1x10⁶ cells/ml, 1 ml dispensed into each well of a 24-well plate, and allowed to rest overnight. Cells were then activated for 24 h with Dynabeads Human T-Activator CD3/CD28 (Thermo Fisher) per manufacturer’s protocol. Concentrated lentiviral particles (50 μl) were added per well, cells incubated overnight, then transferred to T25 flasks with an added 6 ml TCM. After three days of expansion, Dynabeads were removed and cells back-diluted to 5x10⁵ cells/ml with daily monitoring to ensure they did not exceed 4x10⁶ cells/ml. Transduction efficiency was assessed by flow cytometry using a Rituximab-MicAbody that had been directly conjugated to Alexa Fluor 647 (Alexa Fluor Protein Labeling Kit, Thermo Fisher) per manufacturer’s protocol. Transduction efficiencies were above 70% for all cCAR-T tested.

**Spinning disc time-lapse Microscopy**

3x10⁶ tonsil-derived cells were seeded in a 4-well imaging chamber and placed under the microscope for subsequent imaging. Imaging was performed using an Axiovert inverted fluorescence microscope (Carl Zeiss), equipped with a Yokogawa spinning disk, a CoolSNAP HQ2 14-bit camera (PhotoMetrics), and laser lines for 488 nm (40% laser power, 400-ms excitation) and 561 nm (40% laser power, 200-ms excitation). To facilitate time-lapse imaging, the microscope has a programmable stage with definite focus. GFP fluorescence measured by flow cytometry.

**Culture and infection of primary cells**

Spleen and tonsil tissues were minced into small pieces, then passed through a 70-μm cell strainer into FACS buffer (PBS supplemented with 2% FCS and 2% EDTA). The cell suspensions were then passed through 40-μm cell strainer to prepare a single cell culture. Preprocessed HLAC cells or blood, were mounted on top of Ficoll-Paque and centrifuged at 400 g for 30 min. The middle mononuclear cell layer was washed twice with FACS buffer and cells were resuspended in HLAC medium. Prior to HIV infection, PBMC cells were activated by 100 IU/ml IL-2 in the presence of 10 μg/ml Phytohemagglutinin (PHA) for 3 days.

**HIV-1 infection**

HEK293T cells were transfected using Fugene-HD in 24 well flat-bottom-plates with plasmid DNA (100 ng) corresponding to molecular clones of HIV-1 expressing a GFP reporter and Nef using an internal ribosomal entry site (Neidleman et al., 2017). The medium was replaced after 16 h, and cells were tested for GFP expression. HLAC/PBMC were over-laid on the adherent GFP-expressing HEK293T cells for 24 h. Suspension HLAC/PBMC cells were separated from the adherent cells after the 24 h incubation, and spreading infection was allowed to proceed until 4%-10% of the HLAC CD4 T cells were infected as determined by GFP epifluorescence measured by flow cytometry.

**Reactivation of cells from HIV-positive individuals**

Leukopacks from HIV-positive individuals on ART were processed using Ficoll-Paque gradients similar to that for healthy donor blood. CD4 T cells were enriched by negative depletion with an EasySep Human CD4+ T Cell Enrichment Kit (STEMCELL). Subsequently cells were activated with 50 ng/ml Phorbol 12-myristate 13-acetate (PMA) and 1 μM ionomycin for 3 days to induce reactivation of latent proviruses within reservoir cells.

**cCAR-T cell killing assay**

Primary target cells were cultured until the level of infection rose above 4% in a Live/CD3+/CD8+ gate as determined by FACS analysis. One million target cells (HLAC or PBMC) with ~10⁴ infected cells were plated in a 96 V-bottom plate. 10⁴ cCAR-T cells or donor-matched untransduced CD8 cells were incubated with different concentrations or types of MicAbodies for 5 min and then added to the target cells for 48 h incubation. All experiments were performed in the presence of 5 μM saquinavir to prevent spreading infection. For reactivated CD4 T cells from HIV-positive individuals, 5x10⁵ CD4 T cells were plated in the presence of 5x10⁵ cCAR-T cells or matched untransduced CD8 T cells in the presence or absence of a mix of the four HIV-specific MicAbodies. Cultures were incubated for 48 h.
Measurement of cell-associated RNA by ddPCR

Two days after co-culture of reactivated CD4 T cells from HIV-positive individuals with cCAR-T cells and MicAbody, cells were collected and centrifuged for 10 min and RLT lysis buffer (Qiagen) was immediately added to cell pellets. Cell associated RNA (caRNA) was extracted with the RNeasy kit (Qiagen) following manufacturer’s protocol. Extracted RNA was reverse transcribed and pre-amplified using previously described HIV-specific primers (Laird et al., 2015) (see also Key Resources Table) using the Superscript III One-Step RT-PCR system (Life Technologies) with 10 μL purified RNA in 25 μL final volume. RT-PCR was carried out using the following steps: reverse transcription at 50°C for 30 min, denaturation at 95°C for 2 min, 10 cycles of amplification (94°C 15 s, 55°C 30 s, 68°C 30 s) and a final amplification step at 68°C for 5 min on a ThermoFisher PCR instrument. Subsequently, ddPCR was applied to quantify pre-amplified cDNA. For ddPCR droplet generation, reactions were loaded into the Bio-Rad QX-100 emulsification device following the manufacturer’s instructions. Samples were transferred to a 96-well reaction plate and sealed with a pre-heated Eppendorf 96-well heat sealer (Bio-Rad). Finally, samples were amplified on a BioRad C1000 Thermocycler and analyzed on BioRad QX100 ddPCR Reader using QuantaSoft Software (Bio-Rad). Each 25 μL ddPCR mix comprised the ddPCR Probe Supermix (no dUTP), 900nM primers, 250nM probe (Laird et al., 2015), and 5 μL cDNA. The following conditions were used: 10 min at 95°C, and 40 amplification cycles (30 s denaturation at 94°C followed by 59.4°C extension for 60 s) and a final 10 min at 98°C.

ELISA binding assay

The extracellular domain of wild-type NKG2D or mutant NKG2D was fused to the C terminus of human IgG1-Fc, DNA constructs for Fc-NKG2D molecules were expressed in Expi293™ cells, secreted protein purified by Protein-A affinity chromatography and eluted material fractionated by size-exclusion chromatography (SEC) on an ÅKTA Pure system using Superdex 200 columns. For ELISA binding assays, 1 μg/ml of Fc-NKG2D reagents were coated onto microtiter plates and a dilution series of MicAbody introduced followed by detection with HRP-conjugated mouse-anti-human kappa chain antibody then developed with 1-Step Ultra TMB ELISA.

Flow cytometry

Equal volumes from all treatment wells were spun down and the cells were stained with fluorochrome-conjugated antibodies (see Key Resources Table). Fixable Viability Kit Zombie Violet (BioLegend) was used to exclude dead cells while simultaneously staining for surface markers (CD3/CD4/CD8/CD19). AccuCount counting beads (Spherotech) were added after the last wash to control for sampling errors and cells were fixed in 1% paraformaldehyde. Data were acquired on LSR-II (BD Bioscience), and FlowJo software was used for analysis (TreeStar). Killing of HIV-infected cells was assessed by measuring the reduction in the number of GFP+ cells relative to control (see Figure S2 for gating strategy).

In vivo assay

Female NSG mice were implanted with 1X10⁶ Raji-Luc cells subcutaneously and reached tumor volumes of < 100 mm³ on day 12, at which point 20 μg of Rituximab-MicAbody were injected intraperitoneally in a 100 μL volume and repeated every two days for a total of six doses. 1X10⁷ cCAR-T cells, comprised of a 1:1 absolute CD4:CD8 ratio, was injected intravenously on day 13 in a 100 μL volume. Untreated animals received PBS. Tumor volumes were regularly monitored by caliper measurements and weights were also tracked to monitor overall health of the animals. n = 3 mice per cohort.

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical details of individual experiments, including number of independent donors, mean values, standard error of the mean (SEM), and p values derived from two-tailed t tests are described in the figure legends and specified in the figures. Statistical analyses were performed using Microsoft Excel software. p values ≤ 0.05 were considered statistically significant. For comparison between two treatments, a Student’s two-tailed t test was used. For the in vivo mice assays a two-way ANOVA test was used to assess statistically significance of the results (p values ≤ 0.05). Asterisk coding in figures is as follows: * p ≤ 0.05; ** p ≤ 0.01; *** p ≤ 0.001. Data are presented as means with error bars indicating SEM unless otherwise stated.

DATA AND CODE AVAILABILITY

This study did not generate datasets.
Figure S1. Lymphoid Tissue-Based Killing Assay, Related to Figure 2
Tonsil or spleen cells were purchased from CHTN and processed (see STAR Methods). These cells were infected using HEK293T transfected with various HIV expression plasmid DNAs. The infected cells were then combined with cCAR-T cells and different MicAbodies for 48 h. Killing was determined by reduction in the number of infected cells expressing GFP (live CD3+/CD8^+ GFP^+) compared to the No cCAR-T cell control assessed by flow cytometry. In each experiment potential in-well toxicity or off-target killing by MicAbodies and cCAR-T cells was measured by tracking the number of live CD3^+ CD8^- GFP^- cells.
Figure S2. Gating Strategy for Flow Cytometric Assessment of cCAR-T Cell Killing in Presence of MicAbodies, Related to Figures 2 and 3

Following antibody staining, cells were analyzed on an LSR2 flow cytometer (Becton Dickinson). (A) Cells in the lymphocyte gate were analyzed in two singlets gates (B). Next an exclusion live gate was made dividing the live cells into CD3+ or CD3- gates by using Fixable Viability Kit Zombie Violet. (C) Cells were further gated on CD3+/GFP+ (infected cells) or CD4+/GFP- (uninfected cells). Further gating based of GFP expression allowed separation into high and low GFP expressing cells. Relative viability of uninfected control cells (CD3+/CD4+/GFP-) in each well was assessed in parallel. (D) Gating on effector cells was made by a live CD3+/CD8+ gate. (E) For B cell killing, the gating was made on CD3-/CD19+ live cells.
Figure S3. Specific Killing of HIV-Infected Primary Spleen CD4 T Cells by cCAR-T Cells Combined with HIV Env-Specific MicAbodies, Related to Figure 2

Specific killing of R5 tropic HIV-1 (BaL)-infected spleen cells by cCAR-T cells armed with 4 different HIV Env-specific MicAbodies. One million splenocytes containing approximately 1x10^4 infected CD4 T cells were incubated with 1x10^5 cCAR-T cells for 48 h, in the presence of different concentrations (10-500 pM) of HIV Env-specific MicAbodies. B cell-specific MicAbody (Ritux) and anti-HER2 MicAbody (HER2) were incorporated as negative controls. Results are presented relative to the No cCAR-T cell control. For each individual MicAbody, an internal control of no cCAR-T cell supplemented with the highest MicAbody concentration tested is presented. Results are cumulated from four independent experiments. Data are represented as mean ± SEM. * = p ≤ 0.05, ** = p ≤ 0.01, *** = p ≤ 0.001, compared to no MicAbody.
Figure S4. Specific Killing of CXCR4 Tropic HIV-Infected Primary CD4 T Cells by cCAR-T Cells Combined with Specific HIV MicAbodies, Related to Figure 2

Specific killing of X4-tropic HIV-1 (NL4-3) infected tonsil (A), spleen (B), or blood cells (C) by 4 single HIV-specific MicAbodies with cCAR-T cells was assessed. One million primary cells (~1x10^5 infected cells) were incubated with 1x10^5 cCAR-T cells for 48 h in the presence of different concentrations (10-500 pM) of the HIV Env-specific MicAbodies. B cell-specific MicAbody (Ritux) and anti-HER2 MicAbody (HER2) were used as negative MicAbody controls. Results are presented for each individual MicAbody, no cCAR-T cells and donor-matched untransduced CD8 cells supplemented with the highest MicAbody concentration tested are included as controls. Results are cumulated from three independent experiments for each tissue. Data are represented as mean + SEM relative to the CD8 control. * = p ≤ 0.05, ** = p ≤ 0.01, *** = p ≤ 0.001.
Figure S5. Comparison of cCAR-T Killing of HIV-Infected Cells Present in Activated PBMC versus Tonsil-Derived Cells, Related to Figure 6

F4-HIV-infected cells from tonsil (HLAC) or activated blood cells (PBMC) were cultured with cCAR-T cells at a 10:1 effector-to-target ratio for 48 h in the presence of a mix of four HIV MicAbodies, or HER-2 MicAbody. GFP+ cell number was measured by flow cytometry and data are presented relative to cCAR-T with no MicAbody present. Data are represented as mean ± SEM relative to no MicAbody. NS = p > 0.05, * = p ≤ 0.05, *** = p ≤ 0.001. n = 3.