Over the past decade, chimeric antigen receptor (CAR) T cells have emerged as the prototype gene therapy for B cell leukemias. These so-called living drugs are derived from a patient's own cells, reprogrammed to recognize and destroy cancer cells, and then reintroduced into the body. The huge success of this therapy for cancer is rooted in pioneering clinical and preclinical studies, established more than three decades ago, focused on persistent HIV-1 infection. In this issue of the JCI, Bingfeng Liu et al. revisit HIV-specific CAR T cells in an important clinical study that supports broader application of this groundbreaking therapy. Although curative endpoints were not achieved, these findings lay the foundation for augmented approaches applying combinatorial technologies including antigen supplementation.
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**CAR T cells from HIV to cancer and back**

CAR T cell therapies were first described in the early 1990s as a treatment for antiretroviral therapy–suppressed (ART-suppressed) people living with HIV (PLWH). Despite promising initial data in preclinical experiments (1–3), three clinical trials between 1998 and 2005 found minimal impact on virus persistence in PLWH (4, 5). Long-term follow-up in these clinical cohorts established the safety of CAR-modified cell products and suggested that CAR T cells maintain the capacity to expand in response to exogenous activation stimuli (6). In parallel, advents in CAR engineering took this approach to new heights as a potent therapy for patients with hematological malignancies, namely CD19+ B cell leukemias and lymphomas. Among many approaches to potentiate CAR T cell function against malignant targets, various groups have validated more advanced lentiviral vector designs, methods to control ratios of T cell subsets within a CAR product, culture conditions geared toward memory T cell phenotypes, and perhaps most importantly, the addition of costimulatory domains within the CAR molecule, designed to enhance target-dependent killing (7–10).

In this issue of the JCI, Bingfeng Liu et al. took the impressive clinical efficacy of these next-generation CAR products for cancer into account, and built on the safety of first-generation HIV-specific CAR T cells that were established over 20 years ago. Liu and colleagues reported on the results of a clinical trial involving 15 HIV-infected, ART-suppressed participants who received next-generation HIV-specific CAR T cells. The authors’ goal was to assess the safety and impact of CD8+ CAR T cells on virological parameters before and, in the case of six participants, after ART treatment interruption (ATI) (11).

**HIV CAR T cell effects on reactivated viral reservoirs**

Liu et al. enrolled 15 participants with ART-suppressed HIV-1 infection and successfully manufactured a virus-specific CAR T cell product for 14 out of 15 individuals. The CAR featured a single chain variable fragment (scFv) derived from the HIV-1 broadly neutralizing antibody, VRC01, and contained both CD28 and 4-1BB costimulatory domains. Further, the lentiviral vector used to modify CD8+ T cells from each participant coexpressed short hairpin RNA molecules designed to knock down the expression of immune exhaustion proteins PD1, TIM3, and LAG3. In the six participants who underwent ATI, time to viral rebound was substantially longer than in a historical control cohort (12). Correlative measurements further suggested that CAR+ cells expanded coincident with increases in cell-associated viral RNA after ART interruption. Intact provirus detection assays, a current gold standard for virus persistence (13, 14), showed a marked decrease in intact proviruses in this cohort over time. Sequencing experiments suggested that viral diversity decreased following CAR treatments and selection for CAR-resistant variants occurred, although these data were limited in scope. Comparisons to untreated controls (participants who underwent ATI without CAR T cell therapy) were notably limited to historical reports. This limitation is understandable when considering ethical challenges for noninterventional ATI studies (15, 16). The key findings from Liu et al., namely a delay of up to 10 weeks in viral rebound following ATI, offer a tantalizing glimpse of the potential for HIV-specific CAR T cell therapies in ongoing and future studies. One intriguing question is whether posttreatment immune control would have been seen in the six ATI participants if their ART interruption period had been extended beyond a single documented virus-positive time point.

**Next steps for HIV CAR T cell therapies**

Much remains to be learned regarding the requirements for an effective CAR T cell strategy for persistent HIV infection. Many will continue to highlight similarities and opportunities for synergy with CAR T cells for cancer. It is increasingly clear that the local immune environment, the frequency of antigen-expressing target cells, and/or

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1. CD19 + leukemias frequently contain CAR T cell therapies for leukemia (Figure 1). CD19 + T cells (23, 24) and by the low number of latently infected cells (no more than 1 per million CD4+ T cells) in a typical PLWH (13, 25). Several groups including ours have and continue to address these barriers in small and large animal models of HIV persistence (26, 27).

The clinical data from Liu et al. represent a welcome return of HIV-specific CAR T cells to the clinic. Iteratively developing gene therapy and gene editing approaches that augment CAR T cell safety, function, and persistence will be essential to further augment these therapies, with the ultimate goal of enabling lifelong HIV remission without the need for daily ART. Beyond optimizing the best CAR molecules and cell manufacturing approaches in preclinical models, aligning these studies with rationally designed clinical trials will, likewise, be critical. Our previous nonhuman primate study was designed to inform an ongoing clinical trial (Clinicaltrials.gov identifier NCT03617198), in particular focused on the timing between CAR T cell infusion and ATI. Liu et al. interrupted ART 3 to 6 weeks following infusion of CAR-modified cells, and it remains unclear whether a shorter period may be required to ensure remission (i.e., so that recrudescence viral antigen coincides with the peak window of CAR T cell immune surveillance in vivo). Other aspects, including exogenous antigen boosting, enhancement of cell-intrinsic CAR properties, and/or modifications to the local viral reservoir environment, may require further studies in preclinical models. The study by Liu et al. reminds us that CAR T cells were born as a treatment for infectious disease and, following lessons learned in cancer immunotherapy, have come full circle as an HIV-specific clinical therapy. Much work remains to develop this approach into a form that will consistently lead to HIV cure endpoints. A call to action to invest the necessary resources in this extremely promising immunotherapy, not just for HIV, but numerous other diseases, may provide profound clinical benefit.

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