Original Article



CRISPR-Mediated Base Conversion Allows Discriminatory Depletion of Endogenous T Cell Receptors for Enhanced Synthetic Immunity

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Emerging base editing technology exploits CRISPR RNA-guided DNA modification effects for highly specific C > T conversion, which has been used to efficiently disrupt gene expression. These tools can enhance synthetic T cell immunity by restricting specificity, addressing histocompatibility leukocyte antigen (HLA) barriers, and promoting persistence. We report lentiviral delivery of a hepatitis B-virus (HBV)-specific recombinant T cell receptor (rTCR) and a linked CRISPR single-guide RNA for simultaneous disruption of endogenous TCRs (eTCRs) when combined with transient cytosine deamination. Discriminatory depletion of eTCR and coupled expression of rTCR resulted in enrichment of HBV-specific populations from 55% (SEM, $\pm 2.4\%$) to 95% (SEM, $\pm 0.5\%$). Intensity of rTCR expression increased 1.8- to 2.9-fold compared to that in cells retaining their competing eTCR, and increased cytokine production and killing of HBV antigen-expressing hepatoma cells in a 3D microfluidic model were exhibited. Molecular signatures confirmed that seamless conversion of C > T (G > A) had created a premature stop codon in TCR beta constant 1/2 loci, with no notable activity at predicted off-target sites. Thus, targeted disruption of eTCR by cytosine deamination and discriminatory enrichment of antigen-specific T cells offers the prospect of enhanced, more specific T cell therapies against HBV-associated hepatocellular carcinoma (HCC) as well as other viral and tumor antigens.

INTRODUCTION

T cells redirected with recombinant T cell receptors (rTCRs) are being investigated in early-phase human studies.^{1–3} Limitations include unpredictable "off-target effects" due to TCR cross-reactivity; for example, cardiac toxicity following therapy with MAGE-A3 rTCR^{4,5} and concerns that endogenous TCR α and β chains may mispair with rTCR chains and give rise to novel dimeric complexes with unpredictable specificities.^{6,7} These limitations have been partially mitigated by predictive modeling of rTCR cross-reactivity and by promoting exclusive rTCR pairing via additional disulfide bonds and other strategies.^{8–11} Also of note is the importance of rTCR assembly on the cell surface as a multi-

meric complex with CD3 chains, as competition from the endogenous TCR (eTCR) for the shared components can limit cell-surface expression.¹² Competition for such cellular components can be addressed either by overexpression of CD3, disruption of eTCR by RNA interference,¹³ or nuclease-mediated genetic disruption of eTCR chains. Previously, zinc-finger nucleases (ZFNs),¹⁴ transcription activator-like effector nucleases (TALENs),¹⁵ and clustered regularly interspaced short palindromic repeat (CRISPR)/CRISPR-associated protein 9 (Cas9) have all been used to disrupt one or both TCR α and β chains.^{16–18} These genome-editing approaches also reduce the likelihood of mispairing, but existing nuclease-based approaches all result in double-stranded DNA breaks and may create large insertions/deletions (indels), trigger translocation events, and increase activation of p53 pathways.¹⁹⁻²³ Recently, a report of autologous anti-tumor therapy with T cells edited using Cas9 to disrupt both TCR and PD1 expression noted readily detectable chromosomal translocations in the infused products,²⁴ and similar aberrations were found after TALEN editing of T cells modified to express anti-CD19 chimeric antigen receptors (CARs).²⁵

Here, we report the application of emerging cytosine deaminase baseediting technology for efficient and seamless base conversion to introduce premature stop codons in homologous regions of TCR beta constant 1 and 2 (*TRBC 1/2*) chains.^{26,27} BE3 (third-generation base editor) is a CRISPR-guided nickase Cas9 (D10A), fused to a rat apolipoprotein B mRNA editing enzyme catalytic polypeptide (rAPOBEC1) deaminase at the N terminus, which operates within a 4- to 8-bp window distal to the protospacer adjacent motif (PAM) sequence. The inclusion of a C terminus fusion comprising a uracil-DNA glycosylase inhibitor (UGI) (derived from *Bacillus subtilis*



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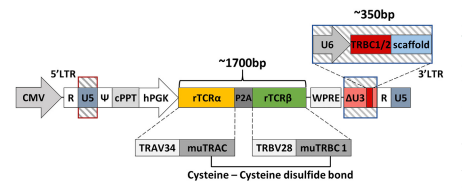


Figure 1. Terminal-CRISPR Lentiviral Vector Configuration Coupling HBV rTCR and CRISPR TRBC1/2 sgRNA Delivery

Lentiviral plasmid configuration, coupling the expression of a recombinant T cell receptor (rTCR) against the hepatitis B virus (HBV) envelope surface antigen 183-91 (S183-91) and a T cell receptor beta constant (TRBC)specific single guide RNA (sgRNA). The S183-91 rTCR is placed under the transcriptional control of an internal human phosphoglycerate kinase 1 (hPGK) promoter, while TRBC1/2 sgRNA is expressed via a human U6 promoter. The rTCR is expressed as a single transcript with the rTCR α chain first, followed by the rTCR β chain separated by a porcine teschovirus-1 2A (P2A) self-cleavage sequence. These recombinant chains are composed of the T cell receptor α variable 34 (TRAV34)

and the T cell receptor β variable 28 (TRBV28) domains, as well as either murine TRAC (muTRAC) or murine TRBC 1 (muTRBC1). The rTCR chains contained an additional cysteine-cysteine disulfide bond between murine constant regions. CMV, cytomegalovirus; cPPT, central polypurine tract; WPRE, woodchuck post-transcriptional regulatory element; LTR, long terminal repeat; ΔU3, deleted unique 3'; R, repeat; U5, unique 5'; ψ, psi; D, diversity region; J, joining region.

bacteriophage PBS1) inhibits uracil-DNA glycosylase and blocks uracil excision, promoting conversion to thymidine as cells replicate. High levels of C > T conversion and low levels of indels have been reported for BE3.²⁸⁻³⁰ Here, we investigate a codon-optimized BE3 (coBE3) in the context of engineering T cells against HBV surface antigen, an important target in the treatment of hepatocellular carcinoma (HCC).^{31,32} HBV viral antigens are processed and presented by major histocompatibility complex (MHC) molecules on the surface of infected cells,^{33,34} and naturally occurring HBV-specific T cells can engage with peptides presented in the context of histocompatibility leukocyte antigen (HLA) to moderate viral and tumor burdens.^{35,36} Nevertheless, such HBV-specific T cell responses can become exhausted during chronic HBV infection,³⁷⁻³⁹ and synthetic HBV-specific T cells can be generated through the expression of rTCRs.⁴⁰⁻⁴⁴ The approach has already been tested clinically in HBV-associated HCC,^{45,46} with further studies planned.

Lentiviral vector delivery of a rTCR specific for HLA-A2/HBV peptide S183-91, incorporating murine constant regions, and coupled to a CRISPR single guide RNA (sgRNA) targeting *TRBC1/2* loci resulted in high levels of targeted cytosine deamination after transient delivery of mRNA encoding coBE3. Thereafter, discriminatory removal of residual eTCR+ cells was achieved by magnetic-beadmediated depletion using the anti-human TCR $\alpha\beta$ monoclonal antibody. Consequently, rTCR expression was enriched, as the murine constant regions lack the specific epitope recognized by this antibody. Phenotypic and functional assessments, including migration and killing in a 3D microfluidic model, verified immunotherapeutic effects following genome editing, and molecular analysis of both DNA and RNA was performed to examine editor effects.

RESULTS

Base Conversion Disrupts eTCR Expression and Allows Enrichment of T Cells Expressing rTCR

A third-generation self-inactivating (SIN) lentiviral vector was generated encoding an HLA-A0201 restricted rTCR (S183-91, FLLTRILTI)

specific for HBV envelope protein⁴⁷ and a linked sgRNA expression cassette targeting TRBC 1/2. The latter was embedded within a deleted unique (Δ U3) region of the 3' long terminal repeat (LTR) under the transcriptional control of an RNA polymerase III human U6 promoter as previously described.48 This configuration is referred to as terminal-TRBC-S183-91 rTCR (TTRBC-S183-91 rTCR) (Figure 1). Upon electroporation of coBE3 mRNA, the sgRNA mediated highly targeted base conversion of two neighboring cytosine nucleotides within exon 1 of TRBC 1/2 loci. Single or double base conversion produces a premature stop codon within a 4- to 8-bp window distal to the nCas9 (D10A) PAM sequence (Figure 2A). Consequently, disruption of endogenous TCR β chain expression eliminated eTCR $\alpha\beta$ assembly, and the inclusion of murine constant regions within the rTCR further addressed any possibility of aberrant cross-pairing between residual recombinant and endogenous chains (Figure 2B). Following the timeline shown in Figure 2C, healthy T cells were readily activated and transduced resulting in 50%-60% rTCR expression (Figures 2D and 2Ei). Exposure to coBE3 led to disruption of eTCR expression and simultaneous emergence of rTCR+ populations, increasing in proportion to approximately 60%-65% of the cultures (Figures 2D and 2Ei). Furthermore, because eTCR was amenable to detection by anti-TCRαβ monoclonal antibody, magnetic bead-mediated depletion of residual eTCRa\beta-expressing cells was possible. Notably, rTCR (constructed with murine C domains) was not susceptible to these reagents; thus, at the end of production, cells could be enriched for endogenous TCR-/recombinant TCR+ (eTCR-/rTCR+), resulting in a highly homogeneous product (>99% eTCR-/95.9% rTCR+) (Figures 2D and 2Ei). There was also a significant increase in the mean florescence intensity (MFI) of rTCR in eTCR-/rTCR+ cells compared to eTCR+/rTCR± , suggesting enhanced cell-surface expression of rTCR in the absence of eTCR, which may otherwise have competed for CD3 chains during assembly (one-way ANOVA, p < 0.02) (Figure 2Eii).

Hepatitis B Antigen-Specific Responses of eTCR-/rTCR+ T Cells

Three different *in vitro* assessments of antigen-specific function were undertaken. First, production of cytokines including interferon- γ

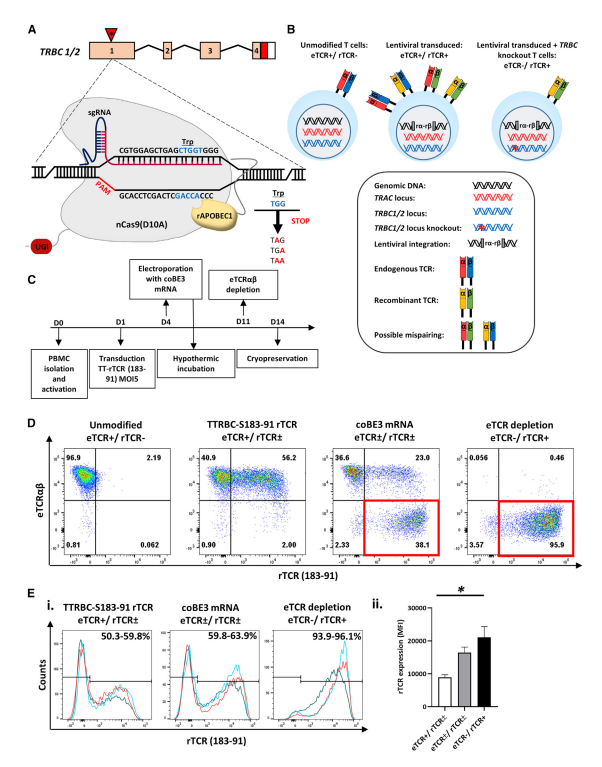


Figure 2. Generation of eTCR-/rTCR+ T cells Using Coupled Cytosine Deaminase Base Editing

(A) Schematic representation of base editor 3 (BE3) targeting exon 1 of the *TRBC 1/2* loci. Editing window (blue) of the BE3 ranged from 4 to 8 bp distal to the PAM (red) with the conversion of tryptophan (Trp) codons to create premature stop codons. (B) Theoretical TCR chain pairing when introducing a rTCR with or without knockout of the endogenous TCR β chain. Incorporation of murine constant regions with an additional disulfide bridge in the recombinant α and β (r α -r β) chains reduced the potential for the mispairing indicated in the middle panel, and disruption of eTCR further reduces the likelihood of mispairing. (C) Schema of cell production. Human peripheral blood lymphocytes were isolated and activated with TransAct (anti-CD3/CD28) (day 0) before transduction (day 1) and electroporation with codon-optimized (co)BE3 mRNA (day *(legend continued on next page)*

(IFN γ), tumor necrosis factor α (TNF- α), interleukin-2 (IL-2), and C-C motif chemokine ligand 4 (CCL4) was determined by flow cytometry in T cells responding to HepG2 cells pulsed with the irrelevant control peptide (HBV core C18-27, FLPSDFFPSV) or a gradient of Hepatitis B target surface envelope peptide (S183-91, FLLTRILTI) concentrations. In all three donors tested, cytokine production was higher in eTCR-/rTCR+ T cells in response to target S183-91 peptide (Figure 3A, i and ii), with absent response to control C18-27 peptide and no-peptide control (Figure S1). Next, we investigated effector function at different effector:target (E:T) ratios in a previously described xCELLigence impedance assay and calculated the relevant normalized cell indices over 72 h after the addition of effector T cells.⁴⁹ An increased index indicated HepG2 target cell proliferation, whereas cell death or apoptosis resulted in a reduced index, signifying higher levels of effector T cell activity (Figure 3B). Control groups included target cells alone (HepG2 alone), and non-transduced effectors (eTCR+/rTCR-), where, as expected, there was a progressive increase and plateau in index. In contrast, both effector groups exhibited a transient rise and then decline in index, with more rapid reductions mediated by eTCR-/rTCR+ cells compared to eTCR+/rTCR± T cells at all E:T ratios (Figure 3Ci). Overall effector function was calculated by area under the curve, as shown in Figure 3Cii, reflecting the increased cytotoxicity by enriched eTCR-/ rTCR+ effector cells compared to their unedited, non-eTCR-depleted counterparts (eTCR+/rTCR±).

Finally, migration and target cell killing by engineered T cells was determined in a 3D microfluidics device. The system captured the migration of effector T cells from a fluidics channel to a collagen gel embedded with target PreS1-GFP-HepG2 cells. Phenotyping of effector T cells confirmed rTCR expression (Figure 4A) and minimal cytokine expression in the absence of stimulation after thawing. Comparable numbers of T cells were observed migrating into the gel between the effector groups (Figure S2) before cytokine expression profiles were compared between cells recovered from inside or outside the gel area (Figure 4B). Both eTCR+/rTCR± and eTCR-/rTCR+ effector groups presented higher levels of IL-2, IFN γ , and TNF- α expression within the gel. Killing of PreS1-GFP-HepG2 cells by eTCR-/rTCR+ cells was confirmed within 24 h, whereas eTCR+/ rTCR± cells at this time point were comparable to control eTCR+/ rTCR- indices and the control PreS1-GFP-HepG2-alone groups (Figure 4C). Direct visualization revealed greater clearance of HepG2 cells after co-culture with eTCR-/rTCR+ T cells (Figure 4D).

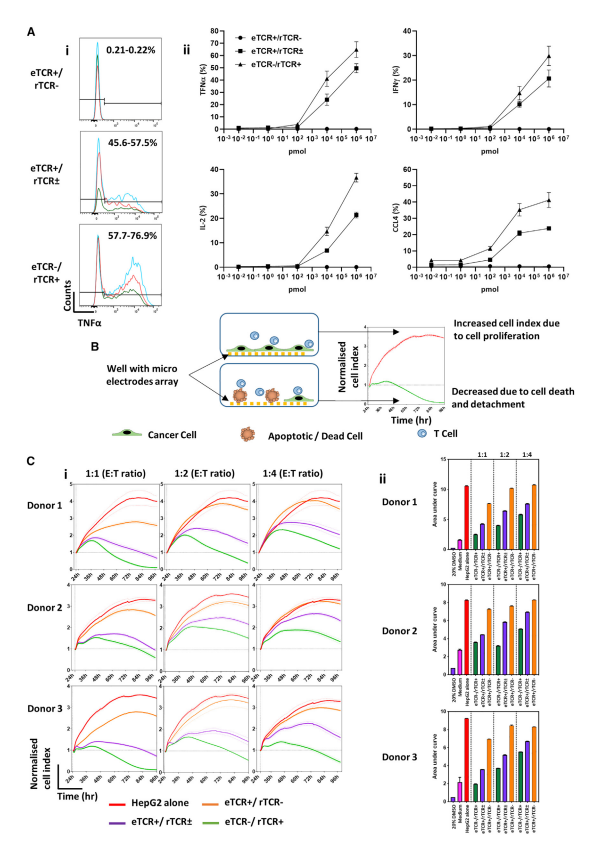
Molecular Characterization of Base Editor Effects

The application of novel genome-editing tools necessitated further investigation of anticipated and unexpected molecular consequences of T cell engineering. There is an established experience of lentiviral mediated effects, including their propensity to integrate into transcriptionally active genes,^{50–52} and we did not re-examine these aspects. However, base conversion effects of coBE3 were characterized in depth, extending comparisons to the effects of SpCas9 disruption in similar experiments disrupting eTCR expression in T cells engineered to express a CAR against CD19. Both modalities had mediated high levels of TCR $\alpha\beta$ disruption (means ± SEM: coBE3, 40.4% ± 5.4%; SpCas9, 52.7% ± 6%) (n = 4; Figure S3), but as anticipated, we found reduced indel frequencies following electroporation of coBE3 (11.9% ± 1.8%) compared to SpCas9 delivery (48.7% ± 6%).

In the context of rTCR delivery, direct sequencing of *TRBC 1/2* in TCR $\alpha\beta$ -depleted eTCR-/rTCR+ T cells was undertaken and analyzed using EditR, with cytosines at positions 5 and 6 distal to the PAM of particular interest (Figure 5A, i and ii). High levels of C > T conversion (G > A sense strand) were captured at these positions (37.3 ± 3.9% and 24.3 ± 2.2% at C5 and C6, respectively), with little activity at other nearby C residues (C1, 5 ± 1.2%; C2, 2.3 ± 1%; and C3, 4.3 ± 1.8%). Next-generation sequencing (NGS) revealed similar levels of C > T conversion at both positions C5 (40 ± 2.9%) and C6 (32.3 ± 3%) (Figure S4). Although mostly seamless, a minority of reads exhibited small (<10 bp; 8.4 ± 1.4%) or large (10–100 bp; 8.2 ± 0.7%) indel signatures (Figure 5Aiii), as others have noted previously.^{28–30}

In silico analysis of sgRNA binding and possible off-target activity was undertaken using the Benchling software platform and presented no exonic off-targets with <3 mismatches. Six genomic loci with the highest scores for off-target activity, all of which contained cytosine bases within the BE3 editing window, were interrogated directly by NGS in three different donors (Figure 5B). We found very low levels (<1%) of conversion activity at these sites, and only one intronic site exhibited C > T changes higher than in its respective non-edited control sample. Recent reports in cell lines have also suggested that promiscuous rAPOBEC1 RNA deamination (including by BE3) can arise following plasmid-mediated expression of base editors.^{54–57} In the T cell context, and with coBE3 transiently expressed by mRNA electroporation, we investigated whether regions directing antigen receptor specificity might be affected. Analysis of RNA from T cells exposed to coBE3 focused on high-throughput interrogation of

^{4).} After overnight hypothermic culture at 30°C, cells were expanded in G-Rex10 flasks for 7 days. Discriminatory depletion of residual endogenous TCR (eTCR)-expressing cells was carried out (day 11) before cryopreservation on day 14. (D) Representative flow cytometry phenotyping of unmodified and terminal TRBC-S183-91 rTCR (TTRBC-S183-91 rTCR)-transduced cells. Delivery of coBE3 mRNA by electroporation caused reduction of eTCR expression (38.1%) and emergence of eTCR-/ rTCR+ cells (red boxes). Magnetic-bead-mediated depletion of residual eTCR+ T cells enriched eTCR- populations, resulting in >99% eTCR-/95.9% rTCR+ (gated on CD45+). (E) Expression of S183-91 rTCR in three healthy donors. (i) Histogram of rTCR (183-91) expression exhibiting transduction ranging from 59.8% to 63.9% in cells exposed to both vector and BE3, which, following TCRαβ bead-mediated depletion, resulted in the enrichment of genome-edited cells, with rTCR levels increased to 93.9%–96.1%. The three colors represent different donors. (ii) Levels of cell-surface rTCR expression measured by mean fluorescence intensity (MFI) (n = 3) showed increased eTCR-/rTCR+ compared to eTCR+/rTCR± cells (gated on CD45+ > rTCR+ population). One-way ANOVA with Tukey's multiple comparison test, p < 0.02. Error bars represent ± 1 standard error of the mean (SEM). nCas9, nickase CRISPR-associated protein 9; UGI, uracil DNA glycosylase inhibitor; rAPOBEC1, rat apolipoprotein B mRNA editing enzyme catalytic polypeptide 1.



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TCR hypervariable regions (TCRVa and TCRVB CDR3 regions). Analysis of samples collected at serial time points, from 1-8 days after BE3 mRNA delivery, found no obvious evidence of aberrant deamination compared to controls (99%-100% cysteines unmodified), and intact sequence integrity of HBs183-91 rTCR was verified (Figure 5C). In addition, transcriptomic analysis on these samples detected anticipated effects of T cell activation and transduction over time (Figure S5). Thus, the first principal component (PC1) accounted for 76% of variance when comparing day 5 and day 12. As the second principal component (PC2) accounted for only 13% of variance, no major transcriptional changes between edited and non-edited cells were noted. In silico analysis had identified a further 24 unique sites of possible off-target BE activity in exonic regions. However, these were all found to have low transcriptional activity (averaged <100 reads) in both edited and non-edited T cells and, therefore, unlikely to be of importance.

Thus, while on-target deamination and creation of TCR-stop codons were highly efficient, there was no notable activity at sites of potential interest at either the DNA or RNA level for coBE3.

DISCUSSION

T cell immunotherapy against conventional tumor-associated targets such as NY-ESO-1 are being widely investigated, and recent reports indicate that autologous T cells with additional CRISPR-Cas9 modifications designed to improve persistence and efficacy can be safely infused.^{24,58} Emerging base editor technologies offer the prospect of highly specific C > T (G > A) base conversion that can be harnessed to create seamless premature stop codons or modify splice sites to disrupt gene expression for advanced T cell engineering.

We previously reported the first therapeutic use of autologous T cells modified to express HBsAg-specific T cell receptors in a subject with chemoresistant, extrahepatic, metastatic disease. In that case, HBV antigens were detectable in HCC metastases but not in donor-derived liver (following cadaveric liver transplantation), thereby reducing the risk of T cell-mediated hepatitis. Gene-modified T cells survived, expanded, and mediated a reduction in HBsAg levels, and while efficacy was not established, there was no significant on- or off-target toxicity.² A small number of additional subjects have been treated

subsequently, although the approach remains highly patient tailored, and extending to larger numbers of patients is logistically challenging and costly. Similar hurdles are being addressed in the arena of hematological malignancies through the generation of "universal" T cells expressing CARs from non-HLA-matched healthy donors. As such, depletion of endogenous TCR and other antigens by genome editing has allowed HLA barriers to be circumvented, and ongoing trials suggest that such universal CAR T cells can expand and persist sufficiently to induce molecular remission.²⁵ The editing tools applied in clinic have included TALENS and CRISPR-Cas9, and they rely on targeted DNA cleavage and repair by non-homologous end joining (NHEJ), which results in the creation of indels leading to gene disruption. Application of CRISPR-guided base conversion to create stop codons or alter the critical splice site to disrupt gene expression offers the possibility of seamless gene disruption with greatly reduced likelihood of translocations or toxicity. We report the application of APO-BEC deaminase technology for the generation of engineered T cells, which are then rendered devoid of endogenous TCRs and uniformly express rTCR specific for an epitope of HBsAg. The resulting product was homogeneous and exhibited enhanced rTCR intensity, greater levels of cytokine production, and antigen-specific functional integrity in models of HCC elimination. An ability to discriminate, selectively process, and deplete eTCR T cells while rTCR populations are untouched provides critical advantages, especially for strategies when allogeneic donor cells bearing potentially alloreactive eTCRs can be eliminated. Non-human protein sequences within constructs have the potential to be immunogenic, although murine TCR constant regions are considered unlikely determinants in the generation of human anti-mouse antibodies.⁵⁹ Likewise, the BE configurations use bacterial and rodent-derived elements, but expression is transient during ex vivo culture and unlikely to be problematic in vivo.

The rapid development of tools enabling highly targeted base conversion through deamination effects promises tantalizing opportunities, although the in-depth characterization of desirable and unwanted effects in subsequent therapeutic applications has to be mapped. Existing CRISPR-Cas base editors using rAPOBEC1 (including coBE3) are known to mediate off-target DNA edits and transcriptome-wide RNA deamination in both protein-coding and non-coding regions.^{54–56} While these could be problematic, newer variants with more precise

Figure 3. Anti-HBV Responsiveness of eTCR+/rTCR±, compared to Base-Edited eTCR-/rTCR+ Effector T Cells

(A) Cytokine responses of effector T cells to the HepG2 cell line pulsed with target HBV surface peptide (S183-91, FLLTRILTI). n = 3. (i) Histograms of tumor necrosis factor α (TNF- α) responses to HepG2 target cells pulsed with 1 μ M target peptide. Both the eTCR+/rTCR± and eTCR-/rTCR+ effector groups are gated on CD45+ > CD3+ > rTCR+ > CD8+, whereas unmodified eTCR+/rTCR- effectors are gated on CD45+ > CD3+ > rTCR- > CD8+. The three different colors represent results from three donors. (ii) Cytokine responsiveness at different concentrations of target peptide (S183-91). HepG2 target cells were pulsed with 1 μ M control peptide (C18-27) to ensure specificity of response, showing cytokine responsiveness comparable to that of the no-peptide control. Effector groups eTCR+/rTCR± and eTCR-/rTCR+ are gated on CD45+ > CD3+ > rTCR+ > CD8+, whereas unmodified cells are gated on CD45+ > CD3+ > rTCR- > CD8+. Error bars represent ±1 SEM. (B) Schematic depiction of the xCELLigence impedance assay showing cancer cells (green) seeded in wells with micro-electrode array (yellow) in the presence of effector T cells (blue). Where T cells recognize cancer cells, this leads to cell death (brown) and reduced impedance, resulting in lower cell index values and area under the curve (AUC). (C) xCELLigence data across different effector:target (E:T) ratios (1:1, 1:2, and 1:4). (i) Visualization of normalized cell index (NCI) over time; all donors showed an increased NCI with decreased E:T ratio. Both HepG2 alone (red) and eTCR-/rTCR+ (green) groups show an initially increased NCI followed by a marked decline. Normalized to time point prior to effector T cell addition. (ii) Summary data of AUC. Increased AUC values were observed at the lower E:T ratios, with eTCR-/rTCR+ consistently presenting with the lowest AUC values. Error bars represent ±1 SEM. IFN γ , interferon γ ; IL-2, interleukin-2; CCL4, C-C motif chemokine ligand 4.

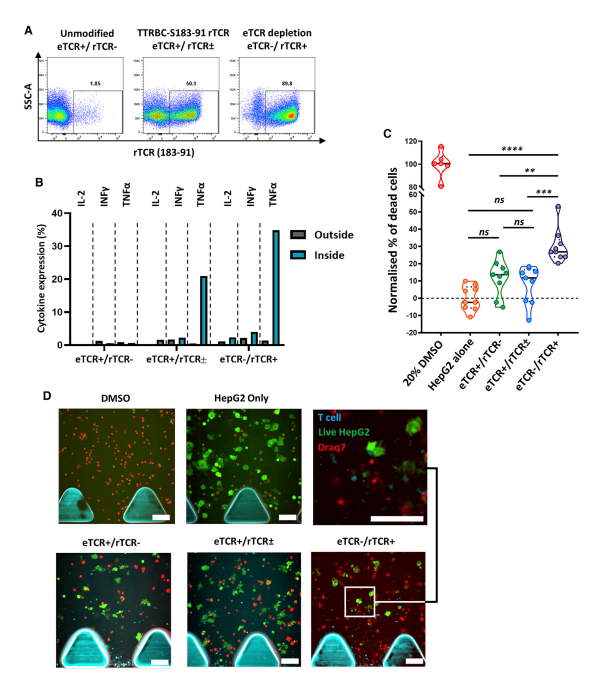


Figure 4. Effector T Cell Cytokine Responsiveness and Target Cell Killing in a 3D Microfluidics Device

(A) Flow-cytometry-based phenotyping of effector T cells used in 3D killing assay post-cryopreservation (gated on CD45+). (B) Histogram depicting cytokine responsiveness of effector T cells isolated from either outside the collagen gel (gray) or inside the collagen gel (blue) after 24 h (cells pooled from 3 replicate devices, gated on rTCR+CD8+; unmodified cells gated on rTCR-CD8+). (C) Normalized killing of target PreS1-GFP-HepG2 cells in response to effector T cell groups presented as violin plot with median (solid black line) and $25^{th}/75^{th}$ quartiles (dotted black lines). PreS1-GFP-HepG2 alone (orange) and 20% DMSO (red) were used as negative and positive controls, respectively. Increased cytotoxicity was observed with eTCR-/rTCR+ (purple) effectors, compared to PreS1-GFP-HepG2 alone (p < 0.0001), eTCR+/rTCR- (green, p < 0.002), and eTCR+/rTCR± (blue, p = 0.0001). Each point represents a section of a 3D microfluidics device from n = 3 technical replicates (3 sections analyzed per device). One-way ANOVA with Tukey's multiple comparison test. (D) Visualization of a region within the collagen gel. Addition of 20% DMSO resulted in cell death (red), while PreS1-GFP-HepG2 target cells alone resulted in high viability (green). Addition of effector T cells (blue) resulted in different degrees of target cell killing between different effector groups. Scale bars, 100 µm.

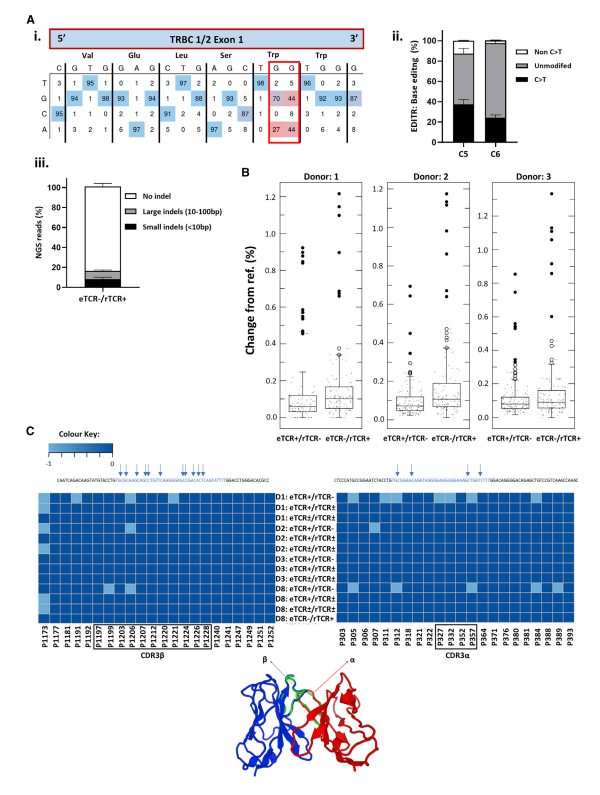


Figure 5. Molecular Analysis of On-/Off-Target DNA Editing and Fidelity of CDR3 α and β Regions within rTCR mRNA Transcripts

(A) Sanger sequencing of on-target editing at *TRBC 1/2* loci in eTCR-/rTCR+ cells. (i) Representative EditR analysis with wild-type sequence (top) and four possible bases (side) shown at each position. Target G > A conversions (red box) generate a premature stop codon (Trp > *). (ii) Summary of EditR data for three donors at cytosine positions 5 *(legend continued on next page)*

DNA restricted editing are already in development and should continue to evolve as ever more efficient, specific, and non-toxic editing tools. Our analysis of possible off-target sgRNA activity in three donors found minimal base conversion effects at predicted DNA sites. Importantly, examination of RNA detected no major differences in gene expression levels between base edited and non-edited T cells, with only very minor perturbations and C > U conversions of the CDR3 variable regions, no greater than in control cells. Such changes could otherwise redirect the specificity of the introduced TCR and would risk causing autoimmunity or off-target T cell effects.

Conclusions

Removal of eTCR enhances the expression of introduced rTCR, reduces the risk of aberrant cross-pairing, and allows discriminatory enrichment of engineered T cells. The strategy also opens the door to generating "universal" allogeneic T cells from healthy HLA-mismatched donors by reducing the risk of graft versus host disease. In the case of the rTCR specific for HBs183-91, blood from healthy HLA-A201 donors could readily be further edited to disrupt mismatched HLA molecules, creating immunologically stealthy cells. Additional multiplexed editing of T cell exhaustion markers may promote enhanced persistence and anti-tumor effects. Ultimately, pre-manufactured banks of eTCR-/rTCR+ T cells specific for groups of dominant HLA/peptide combinations could provide treatment options for large numbers of subjects.

MATERIALS AND METHODS

CRISPR Guide RNA

Guide sequences compatible with coBE3 targeting homologous sequences in *TRBC1* and -2 were designed using the CRISPR design tool Benchling (https://www.benchling.com) and provided an ontarget editing score for predicted activity at each cytosine around the editing window;²⁸ TRBC1/2: $C_{0.8}C_{11}C_{5.7}AC_{21.9}$ $C_{21.4}AGCUCAG$ CUCCACG (anti-sense, numbers indicate predicted editing scores for the specific cytosine base). Predicted exonic off-target binding required at least 3 mismatches within the protospacer.

Lentiviral Construct for rTCR and sgRNA Delivery

Lentiviral design for coupled transgene and guide RNA expression has been previously described.⁴⁸ Briefly, rTCR HLA-A0201/ HBs183-91 was cloned under the control of an internal human phosphoglycerate kinase 1 (hPGK) promoter, and a CRISPR guide expression cassette was embedded in the lentiviral 3' LTR. This comprised a 5' RNA polymerase III promoter (U6) and a sgRNA specific for TCRB1/2 with a 5' G for improved transcription. Vector stocks were produced in 293T cells by transient transfection with third-generation packaging plasmids and concentrated by ultracentrifugation prior to storage at -80° C.

Primary Human Lymphocyte Culture and Modification

Blood was obtained from consented healthy volunteers compliat with ethic (REC number 0141/001). Peripheral blood mononuclear cells (PBMCs) were isolated by Ficoll density gradient and subsequently activated with TransAct Reagent (130-111-160, Miltenyi Biotec) at 10 µL/mL. TexMACS medium (130-097-196, Miltenyi Biotec) with 3% human AB serum (GEM-100-512-HI, Seralabs) and 100 U/mL IL-2 was used for all lymphocyte cell culture. Transduction with lentiviral vector was performed 24 h post-activation at a multiplicity of infection (MOI) of 5. Electroporation of coBE3 mRNA was performed at day 4 post-activation, after which cells were cultured in a G-Rex 10 (P/N 80040S, Wilsonwolf). Lymphocytes were cultured for 11 days post-activation and magnetically depleted using anti-TCR α/β -biotin (130-098-219, Miltenyi Biotec) followed by incubation with Anti-Biotin Microbeads Ultrapure (130-105-637, Miltenvi Biotec) and separation through LD Columns (130-042-901, Miltenyi Biotec). Cells were rested overnight before flow-cytometry-based phenotyping and cryopreservation.

Phenotyping Flow Cytometry

Flow cytometry was performed on a 4-laser BD LSRII (BD Biosciences), with subsequent analysis executed using FlowJo v.10 (TreeStar). Cells were stained according to the manufacturer's instructions with Mouse TCR β constant-APC (clone H57-597, BioLegend, catalog no. 109211), Human TCR α/β PerCP-Vio 700 (clone REA652, Miltenyi Biotec, catalog no. 130-113-540), PD1-PE (clone PD1.3.1.3, Miltenyi Biotec, catalog no. 130-117-384), CD4-VioBlue (clone REA623, Miltenyi Biotec, catalog no. 130-114-534), and CD45-VioGreen (clone REA747, Miltenyi Biotec, catalog no. 130-110-638).

Antigen-Specific Responses

Target HepG2 cells were pulsed with HBV surface envelope peptide S183-91 (FLLTRILTI, JPT Peptide Technologies) and irrelevant control HBV core peptide C18-27 (FLPSDFFPSV, JPT Peptide Technologies) at gradient concentrations for 1 h at 37°C. Cryopreserved effector T cells (eTCR+/rTCR-, eTCR+/rTCR±, and eTCR-/rTCR+) were thawed and cultured at an E:T ratio of 1:1, and 0.1 µg/mL Brefeldin A (Sigma) was added before overnight

and 6 distal to the PAM, presented as C > T changes (black), non-C > T changes (gray), and no editing (white). Error bars represent ±1 SEM. (iii) NGS analysis of on-target editing of *TRBC 1/2* loci and quantification and characterization of indels after BE3 editing found only low levels of small (<10 bp, black) or large (10–100 bp, gray) indels, with the majority of reads presenting with no indels (white). Error bars represent ± 1 SEM. (B) Boxplots showing off-target editing detected by NGS analysis at the top 6 *in silico* predicted off-target sites for the TRBC 1/2 sgRNA, with comparison of unedited eTCR+/rTCR- and edited eTCR-/rTCR+ groups (n = 3). Larger dots represent outliers; in all cases, $\leq 1.3\%$ conversion. Two-tailed independent t test between unmodified (eTCR+/rTCR-) and edited (eTCR-/rTCR+) samples shown for donor 1 (p > 0.5), donor 2 (p = 0.001), and donor 3 (p > 0.1). (C) Serial examination of RNA from rTCR HBs183-91 for 8 days post-coBE3 mRNA delivery (days 5–12 post-activation) found no evidence of promiscuous deamination, with fidelity of CDR3 α and CDR3 β regions maintained. Amplicon positions are marked above for C residues, and schematic highlights hypervariable CDR3 α and CDR3 β regions that confer HLA-peptide specificity. CDR3 regions were mapped as a heatmap in R using the *gplots* library for C > T conversion rates at the marked sites (TCR Clone software: TCRmodel).⁵³

co-culture. A BD LSR Fortessa X20 flow cytometer (BD Biosciences) was used for cell acquisition, with FlowJo v10 (TreeStar) used to analyze phenotype and function of effector T cell groups. Phenotyping included intracellular staining with TNF- α FITC (clone MAb11, BD Biosciences, catalog no. 502906), MIP-1b PE (clone D21-1351, BD Biosciences, catalog no. 550078), IL-2 PerCP-eFluor 710 (clone MQ1-17H12, eBioscience, catalog no. 46-7029-42), GranzymeB AF700 (clone GB11, BD Biosciences, catalog no. 560213), IFN γ V450 (clone B27, BD Biosciences, catalog no. 560371), and surface staining with CD3 BUV395 (clone UCHT1, BD Biosciences, catalog no. 563546) and mouse TCR β constant-APC (clone H57-597, Bio-Legend, catalog no. 109211).

Electroporation of Base Editor mRNA

The BE3 amino-acid sequence was sourced from previously published work containing a single C terminus nuclear localization signal.²⁸ Additionally, the DNA sequence has been codon optimized by GeneArt (Thermo Fisher Scientific). coBE3 mRNA was produced by TriLink and was clean-capped (Cap 1), polyadenylated, and purified by high-performance liquid chromatography (HPLC). Electroporation used a 100- μ L tip kit and the Neon Transfection System (Thermo Fisher Scientific). Cells were electroporated at 20 × 10⁶ cells per milliliter in buffer T, using protocol 24 (1,600 V, 10 ms, 3 pulses) with 50 µg/mL coBE3 mRNA.

Following electroporation, T cells were incubated overnight at 30° C before restoration to 37° C.

Molecular Characterization of On-Target DNA Editing

Genomic DNA extraction was performed using the DNeasy Blood and Tissue Kit (69504, QIAGEN) and PCR sequencing was undertaken using primers for *TRBC1/2* loci: *TRBC* forward, 5'-AGGTCGCTGTGTGTTTGAGC-3'; and *TRBC* reverse, 5'-CTATCC TGGGTCCACTCGTC-3'. Sanger sequencing data (Eurofins Genomics) was analyzed using EditR (https://moriaritylab.shinyapps.io/ editr_v10/).⁶⁰ In addition, amplified products were library prepped for NGS using a Nextera XT Kit (Illumina, Cambridge, UK). After the library preparation, individually barcoded samples were pooled and run in MiSeq using a 500-V2 nano-cartridge. Demultiplexed fastq files were uploaded to Galaxy⁶¹ for trimming and alignment. NHEJ signatures were analyzed using Pindel,⁶² and haplotypes were analyzed using Freebayes.⁶³ Figures were created in R.

Molecular Characterization of Off-Target DNA Editing

The online software Benchling was used to predict off-targets for the *TRBC* guide. Libraries were prepared on the top six off-targets using the same methodology as described earlier (NGS for on-target DNA editing) and combinations of target-specific primers (Supplemental Materials and Methods).

Characterization and Analysis of the Transcriptome

Sequential RNA samples from engineered T cells were prepared for RNA sequencing (RNA-seq) using the KAPA mRNA HyperPrep Kit (Roche) at UCL Genomics. Initial analysis was performed on a customized Galaxy workflow followed by transcriptomics analysis on iDEP 9.1 (workflow using R packages). The online software tool CRISPR RGEN "Cas-OFFinder" predicted 1,071 off-target sites for the TRBC guide, with parameters set for up to 3 mismatches and a 1-nt bulge). A pipeline was developed for further investigation of these sites in RNA-seq data (Supplemental Materials and Methods).

Screening for rTCR RNA Editing Effects

Total RNA was extracted using a QIAamp RNA Blood Mini Kit (QIAGEN, 52304) for TCR library preparation and sequencing as previously described.^{64,65} rTCR RNA was reverse transcribed using a murine TRBC-specific primer (5'-TGGACTTCTTTGCCGTTG AC-3'). Following ligation of an oligonucleotide containing the Illumina SP2 primer and unique molecular identifiers, products were amplified using primers specific to the murine constant α and β chains (5'-CGTTGATCTGGCTGTCGAAG-3' and 5'-TTGACCC ACCAAGACAGCTC-3', respectively). Finally, libraries were built in two further steps of amplification during which the SP1 sequencing primer, indices, and Illumina adaptors were added. Part of the primers used in these were also specific for the constant regions (5'-ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNG CCAATGCACGTTGATCTGGCTGTCGAAG-3' and 5'-ACACTCT TTCCCTACACGACGCTCTTCCGATCTNNNNNNGCCAATCC GTTGACCCACCAAGACAGCTC-3'). The final purified libraries were verified using TapeStation (Agilent Technologies) and Qubit (Thermo Fisher Scientific) and were multiplexed and sequenced on a MiSeq system (Illumina) using 500-V2 cartridges (Illumina). Fastq files were demultiplexed using Demultiplexor (https://github.com/ innate2adaptive/Decombinator).66 Using Galaxy tools,61 the demultiplexed fastq files were trimmed (Trim Galore and Trimmomatic) and aligned (Bowtie2) to the relative TCR HBV gene map. Aligned files were interrogated for the frequency of the reference sequence per base around the complementarity-determining region 3 (CDR3) (100-bp total window).

Data Availability

All fastq files will be available on the NCBI Sequence Read Archive upon publication of this article (BioProject: PRJNA637371).

xCELLigence Impedance Assay

Target HepG2 cells were seeded (1×10^5 per well) in the dedicated device (E-Plate VIEW 16, ACEA Biosciences) and cultured for 24 h. Impedance measurement was acquired with an interval of 15 min by an array of electrodes located at the bottom of the plate. Different T cell preparations and E:T ratios were added in the well after 24 h, and the impedance signal was recorded for the subsequent 72 h. Three different donors were tested in triplicate conditions.

3D Microfluidics Device

Briefly, dissociated PreS1-GFP-HepG2 target cells were mixed with Collagen Type I gel (Rat Tail, Corning) and injected into the dedicated region of the 3D cell-culture chip (DAX-1, AIM Biotech), before gel polymerization, following a previously developed protocol.^{38,43,49} R10 media with 3 μM DRAQ7 (BioLegend) cell-impermeable nuclear dye was then added to the media channels to hydrate the gel, and chips were incubated at 37° C. T cells were stained with 3 μ M CellTracker Violet BMQC (Thermo Fisher Scientific) and were injected into one of two media channels flanking the gel region before overnight incubation. 3D confocal images were acquired daily with a high-content imaging system (Phenix, PerkinElmer). T cells from the liquid channel were collected by manual pipetting; then, collagenase solution was injected into the device to retrieve the immune cells migrating in the hydrogel region for flow cytometry analysis on a 4-laser BD LSRII (BD Biosciences).

Statistics

Statistical analysis was performed using GraphPad Prism software, v.8.0.0.

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10. 1016/j.omtm.2020.09.002.

AUTHOR CONTRIBUTIONS

W.Q. developed the concept, supervised the project, designed experiments, and reviewed the manuscript. R.P. designed and performed experiments, analysed data, and wrote the manuscript. A.P., Z.M., J.Y.J., S.A.G., and K.A.S. designed and performed experiments, analysing data, and reviewed the manuscript. C.G. was involved in experimental design, and reviewing the manuscript. A.P., A.B., and M.K.M. supervised scientist performing experiments and reviewed the manuscript.

CONFLICTS OF INTERESTS

W.Q. holds interests unrelated to this project in Autolus. W.Q. received unrelated research funding from Cellectis, Servier, Miltenyi Biotec, and Bellicum. The other authors declare no competing interests.

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Supplemental Information

CRISPR-Mediated Base Conversion Allows

Discriminatory Depletion of Endogenous T Cell

Receptors for Enhanced Synthetic Immunity

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Supplementary Bioinformatics Methods:

Screening RNA-seq data for *in silico* predicted *TRBC* off-target:

Using the RNA-seq data, a customised bioinformatics methodology was used to investigate the frequency of 1,071 off-target predictions produced by Cas-OFFinder:

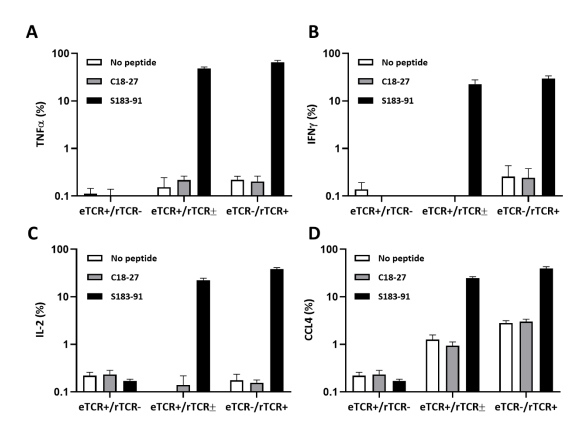
1) Off-targets sites were padded to cover 250bp around each site; 2) The file was converted to a .bed file including the chromosome number, with start and end positions, and strand orientation. 3) Using Homer annotatepeaks the .bed file was annotated to include information on genomic positions (intronic, exonic, intergenic, 3' UTR, 5' UTR etc), gene names, Ensembl numbers and other related information. 4) File was abbreviated for further processing, and a header was created in file. 5) After including exonic regions only, 51 sites remained. 6) By removing duplicates already identified by Cas-OFFinder, 24 exonic sites were advanced. 7) Naïve Variant Caller was performed on RNAseq data previously aligned to human genome (version hg38) and .bed file including the 24 sites. 8) The resulting .vcf files were annotated for any variants using Variant Annotator with a minor allele frequency of 1. 9) Resulting tabular files were finalized on excel. 10) Step 7 resulted with 0 variants annotated. 11) To verify the results, the .vcf files from step 6 were cross-referenced to Integrated Genome Viewer for the aligned .bam files. 12) All files were filtered to include only C to T (or G to A if on anti-sense strand) changes. 13) Any variants found in untransduced (eTCR+/rTCR-) samples were excluded. 14) Remaining variants exhibited coverage of less than 100bp and in most cases less than 10bp, with low MAPQ. This suggested these 24 sites had low transcriptional activity and not likely to be critical to function.

Supplementary Table:

Table S1. Target-specific primer combinations for NGS on top 6 Benchling off-targets

Notes	Off-target Number	Sequence	Genomic Positions Analysed
Fwd primer	1	GAGGCAGGTCTTTCACATTCA	chr19:47,157,708-47,157,727
Rev primer	1	CAAAACGGAAGGAGGCATGT	chr19:47,157,708-47,157,727
Fwd primer	2	AATCTTCACAGCCCCTCCTC	chr9:124,276,868-124,276,887
Rev primer	2	AGTGCGGCCTCAGGTAGC	chr9:124,276,868-124,276,887
Fwd primer	3	GAGAGTGAACGAGGGTGTGG	chr20:33,016,052-33,016,071
Rev primer	3	CCACCTTCACCCCTCGAG	chr20:33,016,052-33,016,071
Fwd primer	4	GTGGTAAGTGTGAGCATTGCAG	chr6:151,587,120-151,587,139
Rev primer	4	TCAGTCCATACTTTCCCCGC	chr6:151,587,120-151,587,139
Fwd primer	5	GATACCTGCTGTCCCCTCTC	chr5:153,537,378-153,537,397
Rev primer	5	CAGAGGTAGATTGCTATGGACCT	chr5:153,537,378-153,537,397
Fwd primer	6	CTTATCCCAGCAGCACTTGG	chr17:45,118,095-45,118,114
Rev primer	6	TCTCTTCCTCCCACAACACC	chr17:45,118,095-45,118,114

Following an internally-built workflow, a 20bp window around the predicted off-target nucleotide position, was analysed to a minor allele frequency of 0.01%. Primer sequences are given in the 5'-3' orientation. Fwd: forward, Rev: reverse.



Supplementary Figures:

Figure S1: Cytokine responsiveness of effector T cells to HepG2 targets alone or pulses with either target (S183-91)/ control (C18-27) peptide.

Flow cytometry data quantifying TNFα (**A**), IFNγ (**B**), IL-2 (**C**), and CCL4 (**D**) cytokine responses of effector T cells against HepG2 targets alone (white) or pulsed with 1µM of either target HBV surface peptide (S183-91, FLLTRILTI, black), or irrelevant control HBV core peptide (C18-27 FLPSDFFPSV, grey) (n=3 biological replicates). Effector groups eTCR+/rTCR± and eTCR-/rTCR+ are gated on CD45+>CD3+>rTCR+CD8+; whereas unmodified eTCR+/rTCR- cells are gated on CD45+>CD3+>rTCR->CD8+. Error bars ± 1 SEM.

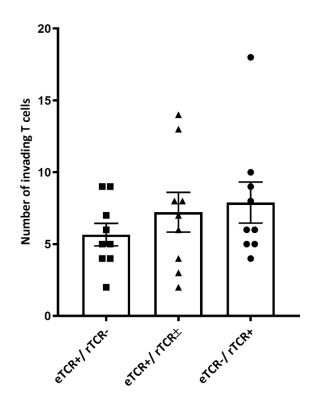


Figure S2: Effector T cell migration within a 3D microfluidics device. Quantification of effector T cells entering the gel region using imaris software. Each point represents a section of a 3D microfluidics device from n=3 technical replicate (3 sections analysed per device). Error bars ± 1 SEM.

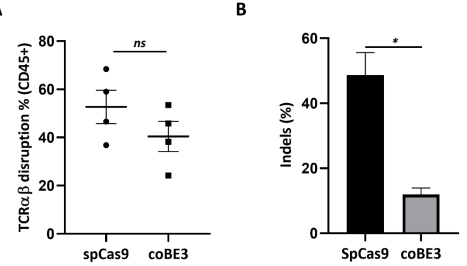
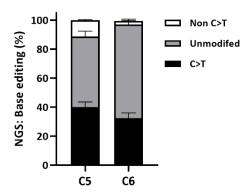


Figure S3: On-target genome editing comparison of *TRBC1/2* editing with coBE3 and SpCas9. **A)** T cells isolated from n=4 healthy donors were transduced at MOI 5 with a lentiviral terminal-CRISPR vector expressing TRBC1/2 sgRNA, and an anti-CD19 chimeric antigen receptor (CAR) transgene. T cells were subsequently electroporated with either coBE3 or SpCas9 mRNA at 50µg/ml. Flow cytometry based analysis of TCR $\alpha\beta$ disruption was gated on CD45+ and no significant difference between SpCas9 vs coBE3 disruption was uncovered. Error bars ± 1 SEM. Mann-Whitney U test, *p* > 0.3). **B)** NGS analysis on the whole cell population found notably higher indel formation at the *TRBC1/2* loci after SpCas9 compared to coBE3 editing (*p* < 0.03, Mann-Whitney U test; Error bars ± 1 SEM. Analysis run on unmodified control revealed an indel frequency of 1.9%.



Protospacer Sequence: GCACCTCGACTCGAC₆C₅ACCC

Figure S4: Next generation sequencing data analysis of on-target coBE3 C>T conversions. Summary of NGS data from n=3 healthy donors at cytosine positions 5 and 6 distal to the PAM, presented as C>T changes (black), non C>T changes (white) and unmodified (grey). Error bars ± 1 SEM.

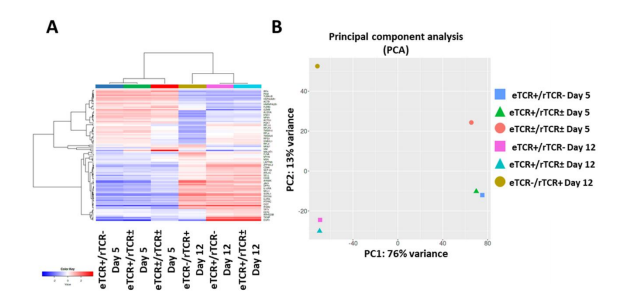


Figure S5: RNA transcriptomics of base edited T cells. PBMCs activated on day 0, T cells transduced with lentiviral TTRBC-S183-91 rTCR on day 1 and electroporated with coBE3 mRNA on day 4. RNA was extracted on days 5 and 12, including from samples enriched by depletion of T cells expressing residual eTCR A) Heatmap of top 100 most variable genes (iDEP9.1) at days 5 and 12 in untransduced cells (eTCR+/rTCR-), transduced and not-edited (eTCR+/rTCR±) and transduced, edited and purified (eTCR-/rTCR+) **B)** Principal Component Analysis demonstrating correlation between PC1 (76% variance) and tested samples over time, capturing the effects of activation and transduction and suggesting little perturbance from base editing activity.