



Chimeric antigen receptor T cells form nonclassical and potent immune synapses driving rapid cytotoxicity

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Chimeric antigen receptor T (CAR-T) cells are effective serial killers with a faster off-rate from dying tumor cells than CAR-T cells binding target cells through their T cell receptor (TCR). Here we explored the functional consequences of CAR-mediated signaling using a dual-specific CAR-T cell, where the same cell was triggered via TCR (trCTL) or CAR (carCTL). The carCTL immune synapse lacked distinct LFA-1 adhesion rings and was less reliant on LFA to form stable conjugates with target cells. carCTL receptors associated with the synapse were found to be disrupted and formed a convoluted multifocal pattern of Lck microclusters. Both proximal and distal receptor signaling pathways were induced more rapidly and subsequently decreased more rapidly in carCTL than in trCTL. The functional consequence of this rapid signaling in carCTL cells included faster lytic granule recruitment to the immune synapse, correlating with faster detachment of the CTL from the target cell. This study provides a mechanism for how CAR-T cells can debulk large tumor burden quickly and may contribute to further refinement of CAR design for enhancing the quality of signaling and programming of the T cell.

chimeric antigen receptor | cell death | cytotoxicity | cytotoxic T lymphocyte | immune synapse

Chimeric antigen receptor T (CAR-T) cells show great promise in the clinic with objective responses and increased relapse free survival in patients with refractory pediatric B-cell acute lymphoblastic leukemia (B-ALL), adult B-ALL, and non-Hodgkin lymphoma (1–4) and very recently in glioblastoma (5). However, clinical responses to date in other hematological and solid cancers have been sporadic (reviewed in ref. 6). There are many potential causes for these observations, including (i) inadequate trafficking of CAR-T cells to the tumor, (ii) lack of CAR-T cell persistence, (iii) tumor microenvironment-induced immune suppression, and (iv) loss of CAR expression in vivo impacting functional response to tumor antigen.

In many patients with refractory disease the CAR-T cell dose used in successful treatments was far smaller than the tumor burden, indicating that CAR-T cells may have proliferated in vivo (7). Our recent data showed that CAR-T cells can mediate rapid and serial killing of tumor cells, indicating that, in addition to CAR-T proliferation, multiple functional pathways may potentially contribute to the dramatic responses observed in some CAR-T cell clinical trials (8). A strong CAR-T cell response in vivo is dependent on stable expression of the CAR, recognition of cognate antigen on tumor cells, and subsequent activation of the effector cell. Activation of a cytotoxic T cell (CTL) is initiated after formation of an immune synapse (IS), a highly organized structure formed at the interface between effector and target cells (9, 10). The TCR-initiated IS comprises a series of concentric rings of clustered molecules (termed supramolecular activating clusters, or SMAC) (10), where each SMAC confers a specific function. The TCR and Lck are clustered in the central SMAC, or cSMAC, while the LFA-1 adhesion molecule and its

adaptor talin are located in the peripheral SMAC, or pSMAC (10, 11). Immunofluorescence imaging revealed that actin accumulates to the distal SMAC (dSMAC), to create the well-known bulls-eye synapse structure (12). Thus, organization of the IS is important as each SMAC confers a function. In the cSMAC, the accumulation of activatory signaling enhances and reinforces the initial response (13, 14). In addition, the cSMAC is the delivery site for cytotoxic granules just before delivery of the lethal hit (12). The adhesion ring present in the pSMAC stabilizes the IS via LFA-1 integrin-mediated binding to ICAM-1 on target cells (15).

In contrast to the IS formed via TCR activation, the IS formed by ligation of the CAR in T cells is completely understudied. In addition, the ensuing signaling events leading to CAR-T cell activation are only partially described. A recent study reported phosphorylation of ZAP70 when HEK-CAR19 cells were conjugated with the CD19⁺ Raji cell line (16). Other studies have demonstrated increased PI3K activity, phospho-AKT levels (17), and T cell survival proteins (18) by third-generation CARs compared with second-generation CARs following antigen stimulation. However, the kinetics and duration of CAR-mediated signaling and the implications for CAR-T cytotoxicity efficiency is not known. To address these issues we used a dual-receptor CAR-T cell model, where the OTI TCR specific for SIINFEKL peptide in association with

Significance

Davenport et al. discovered that the chimeric antigen receptor (CAR) immune synapse structure is different from the T cell receptor (TCR) synapse. The CAR immune synapse formed a disorganized pattern of Lck and more rapidly recruited lytic granules compared with the TCR. The differing immune synapse correlated with faster killing of tumor target cells and detachment from dying tumor cells by CAR-T cells. These findings provide a mechanism whereby CAR-T cells can effectively reduce large tumor burden in patients. This study will form a basis upon which to compare future receptor design to modulate signaling and programming of cytotoxic CAR-T cells to improve treatment of solid cancers.

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H-2K^b and a second-generation anti-HER2 CAR (CD28-CD3 ζ) were expressed in the same CD8⁺ T cell (8). This enabled us to compare, within the same cell, TCR (trCTLs) versus CAR-initiated (carCTLs) IS structure, signaling, and the downstream effect on tumor target killing efficiency.

Results and Discussion

The carCTL IS Is Not Conventional. We previously showed that carCTLs detached faster than trCTL from dying target cells (8). To explore the mechanisms for this observation we first examined the ability of CAR-T cells to form conjugates with OVA₂₅₇- or HER2-expressing EL4 tumor targets. In a standard conjugate assay a significantly higher percentage of trCTL, compared with carCTL, formed conjugates with tumor cells from 10 to 40 min (Fig. 1A). These results are consistent with more rapid carCTL detachment from tumor cells up to 40 min (Fig. 1A), supporting our previously published observations of a faster “off-rate” when viewed by live cell microscopy (8). These off-rates are associated with the avidity of CTL-target interactions. In our model, the number of target cell antigen (HER2 or H-2K^b MHC) molecules were equivalent (8); therefore, the interactions described in this paper compare interactions with a similar number of interacting ligands.

It is well established that TCR signals activate LFA-1, which in turn promotes T cell signaling in the synapse and T cell activation. In vivo, LFA-1 is important for T cell homing and trafficking, with variable levels of LFA-1 having been found on tumor-infiltrating lymphocytes in melanoma tumors (19). Flow cytometry-based conjugate assays are able to detect stable conjugate formation, formed by CTL TCR/CAR interactions as well as LFA-1 binding to its ligand (ICAM-1) on tumor cells.

Using a conjugate assay we determined the level of stable immune synapse formed by trCTLs or carCTLs and the role of LFA-1 in forming these stable ISs (Fig. 1A and C). The trCTLs formed significantly more conjugates than carCTLs from 10 to 40 min, suggesting that trCTL conjugates were more stable than those formed by carCTL (Fig. 1A). trCTL conjugate formation was significantly more dependent on the presence of LFA-1 than carCTL (Fig. 1A and C, *i*). This indicates that carCTL are not as dependent on LFA-1 for synapse formation. Furthermore, while the trCTL IS had a distinct LFA-1 adhesion ring the carCTL IS was characterized by a disorganized LFA-1 pattern (Fig. 1B).

It has previously been shown that three TCR–MHC interacting molecules are sufficient to trigger cytotoxicity by OTI CTLs; however, LFA-1 clustering (as measured by ICAM-1–GFP) required 8 to 10 peptide-loaded MHC (pMHC) complexes for stable ring formation (20). This indicates that formation of a classical stable IS is not always required for cytotoxicity. Furthermore, in our model system carCTLs may bypass the formation of the stable IS structure to induce tumor-cell killing. Single-molecule imaging techniques have shown that a single pMHC ligand is sufficient to trigger TNF α and IL-2 secretion by CD4⁺ T cells (21). However, it is not known whether CD4⁺ or CD8⁺ carCTLs require formation of a stable synapse structure for cytokine production.

Despite equivalent killing by CAR.OTI CTL of either OVA- or HER2-expressing target cells there was no effect on the percentage of target cell killing through the TCR or CAR when LFA-1 was blocked, indicating efficient cytotoxic granule delivery (Fig. 1C, *ii*). Consistent with this data, live-cell imaging demonstrated the capacity of carCTLs to induce cytotoxicity of MC57-HER2 target cells in the presence of LFA-1 blocking antibody (Fig. S1 and Movie S1). Therefore, despite data suggesting that LFA-1 is involved in the polarization of lytic granules in natural killer cells (22), LFA appears to be redundant for both trCTL- and carCTL-mediated cytotoxicity by CAR.OTI CTLs (Fig. 1C, *ii*).

A striking observation in this study was that the gross molecular structure of the carCTL IS was found to be different from the conventional trCTL synapse (Fig. 1D). Lck is a proximal signaling molecule recruited to the TCR for activation. In a conventional trCTL IS, Lck clusters to the cSMAC, surrounded by an LFA-1 ring, that is usually important for CTL adhesion. In contrast, the IS structure of the carCTL was characterized by a lack of Lck clustering and displayed Lck patches in a disorganized pattern (Fig. 1D). To determine whether this lack of Lck clustering was specific to the CAR or was dependent on antigen specificity we examined Lck at the IS of T cell synapses triggered by a CAR of alternate specificity. A similar second-generation CAR construct, differing only in the single chain specificity for carcinoembryonic antigen (CEA), was used to transduce OTI T cells and Lck examined by immunofluorescence. CEA-specific carCTLs displayed the same lack of organized Lck clustering at the IS (Fig. S2), indicating that carCTLs display an inherent difference from trCTLs in the IS structure.

Following quantitation the proportion of carCTLs with accumulation of Lck microclusters and actin rings was found to be significantly reduced, compared with trCTLs (Fig. 1E). Measurements across the IS of the actin clearance inner diameter revealed that trCTLs and carCTLs both formed equivalent synapse sizes at 5 min; however, the carCTL synapse size was significantly smaller than that formed by trCTLs at 20 min (Fig. 1E). These data indicated that the carCTL IS duration was shorter than the trCTL IS. This correlated with the faster carCTL off-rate (8), a smaller IS characterized by actin-rich accumulation and protrusions, as well as microclusters of proximal Lck signaling (Fig. 1D and E and Fig. S2). Our data are supported by previous studies including a report by James and Vale (16) who demonstrated clustering of CD19-specific CARs in a reconstituted HEK cell line recognizing CD19⁺ Raji B target cells with a highly convoluted membrane surface at the IS. This is also consistent with another paper reporting CD19-CAR clustering and actin accumulation at the IS (23).

The TCR IS initiates phosphorylation of proximal (Lck and ZAP-70) and distal (ERK) signaling proteins (13) and is critical for CTL assembly of the cytotoxic machinery (12) and killing of target cells. However, these characteristics of the CAR IS signaling network and its effect on cytotoxicity are unknown.

carCTLs Induce Rapid Proximal Signaling of Shorter Duration than trCTLs. Given the differences observed in Lck clustering at the IS we next examined Lck phosphorylation in carCTLs and trCTLs in response to cognate antigen. Western blot analysis revealed a rapid twofold increase in phospho-Lck (pLck) protein expression by 2 min in carCTL following antigen stimulation that decreased to the level induced in trCTLs by 10 min (Fig. 2A). The distal phospho-ERK (pERK) protein expression was also found to be increased in carCTLs at 2 min and 10 min following antigen stimulation and remained stable for 30 min (Fig. 2B). Calcium increase in the cytoplasm of CTLs is an early indicator of proximal TCR signaling and antigen recognition. We consistently found that carCTLs exhibited fewer Ca²⁺ fluxes before the propidium iodide (PI) blush (Fig. 2C, *Left*), consistent with our previous findings (24).

To determine whether increased pLCK and pERK led to any change in the intensity of the Ca²⁺ flux we next compared this in carCTLs and trCTLs following antigen-specific activation (Fig. 2C). Interestingly, there was no difference observed in the Ca²⁺ flux intensity between carCTLs and trCTLs following antigen-specific activation (Fig. 2C), indicating that a threshold level of activation had occurred and that this is not enhanced in carCTL. Taken together, these data indicate that signaling is stronger, more rapid, and of shorter duration in carCTL than in trCTL. Given the fact that a sustained trCTL signal is required for cytokine production, we next examined whether there was any

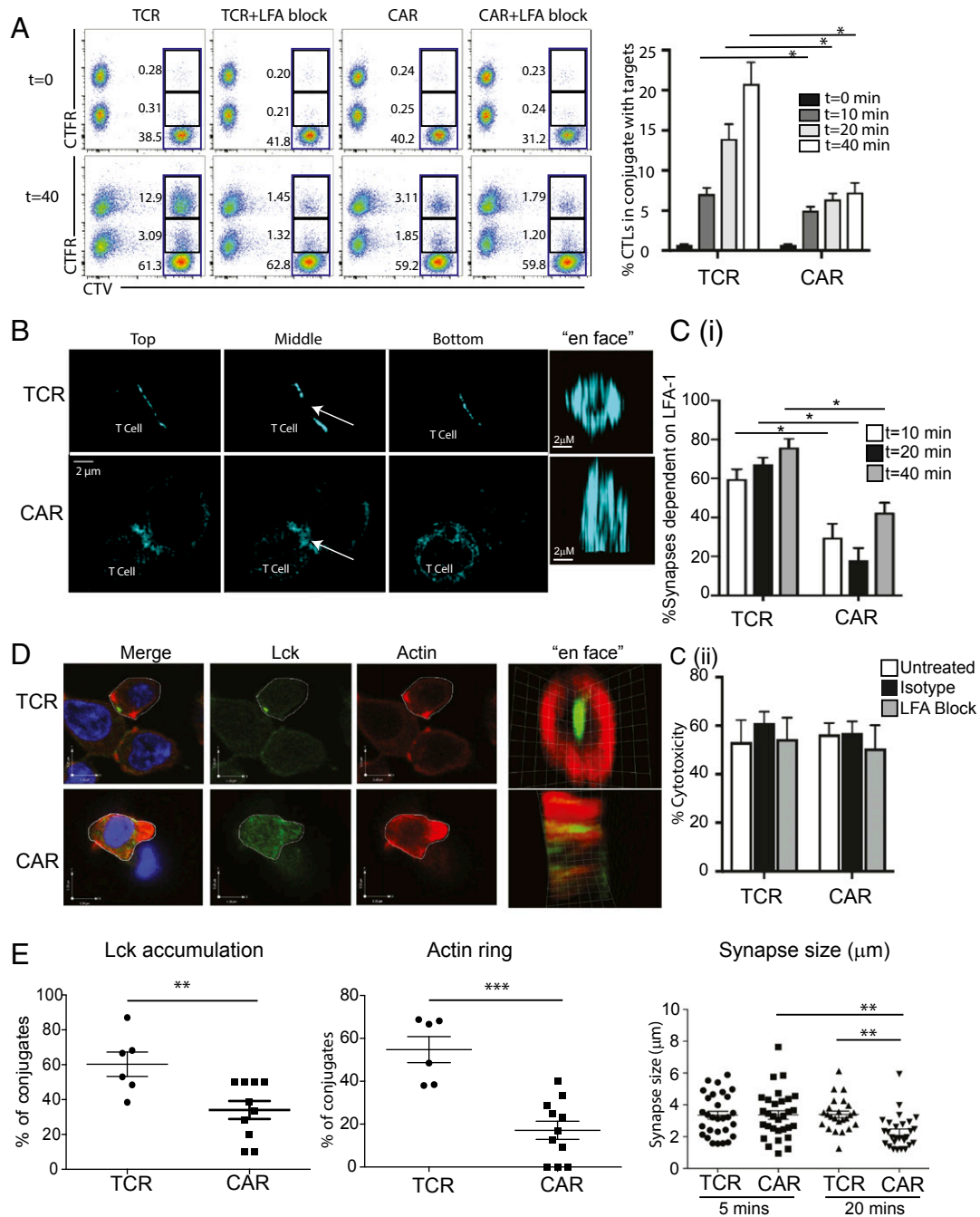


Fig. 1. carCTL synapses are smaller and lack a defined Lck patch. (A) Flow cytometry plots (A, Left) of CTV-labeled effector CAR.OTI CTLs cocultured with EL4 target cells and EL4HER2 (CAR) or EL4OVA₂₅₇ (TCR) each labeled with CTFR^{hi} or parental EL4 (CTFR^{lo}). This conjugate assay was also performed in the absence or presence of LFA-1 blocking antibody. Cells forming conjugates were analyzed over 40 min. The graph (A, Right) shows percentage of CD8⁺ T cells forming conjugates with targets (mean \pm SD) where each condition was performed in triplicate. The figure shows pooled data from four independent experiments. $*P < 0.05$ as determined by a Student's *t* test. (B) Confocal microscopy of CAR.OTI cells in conjugation with MC57 target cells via TCR or CAR, labeled with anti-LFA-1. Montage of confocal planes are shown across the top, middle, and bottom of the T cell. (C, i) Conjugate assay showing the percentage of synapses dependent on LFA-1 binding relative to normal kinetics. CTV-labeled CAR.OTI CTLs were incubated with CTFR-labeled EL4-OVA₂₅₇ (TCR) or EL4-Her2 (CAR) before being analyzed by flow cytometry. CD8⁺ T cells (CTV) still in synapse with target (CTV^{hi}/CTFR^{hi}) are compared with proportion of cells in synapse in the presence of LFA-1 blocking antibody and shown is the proportion of total synapses which are lost when LFA-1 is blocked. Data are an average of three independent experiments, pooled, showing mean \pm SEM and statistical differences between TCR and CAR at each time point determined by Student's *t* test, $*P < 0.05$. (C, ii) Chromium release killing assay. Activated CAR.OTI cells were incubated with ⁵¹Cr-labeled MC57-OVA (TCR) or MC57-HER2 (CAR) cells, in the presence of media alone (Untreated, white), isotype control antibody (Isotype, black), or anti-CD11a blocking antibody (LFA Block, Gray) for 5 h at 1:10 E:T ratio, before measuring ⁵¹Cr-release. Data represent the percentage of tumor cell lysis (mean \pm SD) from two independent experiments. (D) Confocal microscopy of CAR.OTI cells in conjugation with MC57 target cells via TCR or CAR, labeled with anti-Lck (green), anti-actin (red), and Hoechst (blue). (E) ISs as shown in D were blindly quantitated using an unbiased approach for the percentage of conjugates displaying a complete discrete Lck patch at the IS, a cleared actin ring, and the size of the actin ring in conjugates after 5 or 20 min of conjugation. Each data point represents a field of view ($n > 6$) with at least 8–10 conjugates from each (therefore >48 cells), from more than three biological replicate experiments. Student's *t* test showed statistical differences $*P < 0.05$, $**P < 0.01$, $***P < 0.001$ as indicated.

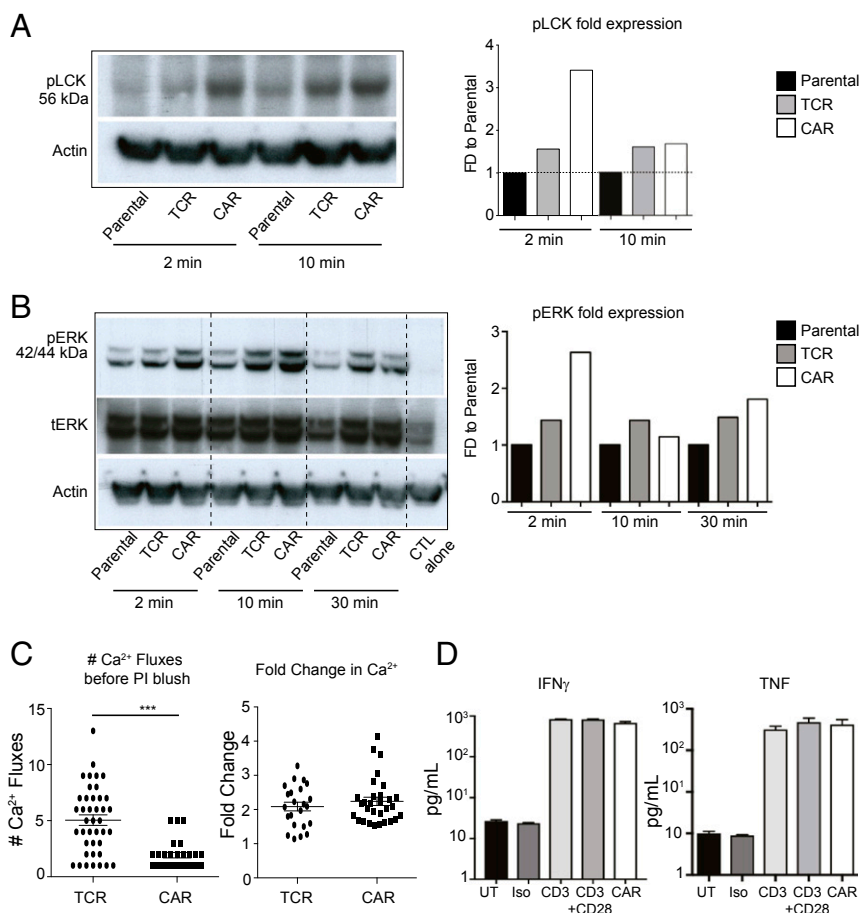


Fig. 2. CAR ligation initiates faster proximal and distal T cell signaling than TCR ligation. CAR.OTI CTLs cocultured with parental, OVA₂₅₇, or HER2-expressing MC57 targets and Western blotting performed for (A) pLck; (B) pErk, total Erk; and actin. Quantitation utilized Fiji imaging software and was normalized to actin loading. Data represented in graphs as mean fold change from parental expression. Western blots are representative of two (A) and three (B) experiments. FD, fold difference. (C) Effector CAR.OTI CTLs were labeled with Fluo4-AM and added to adherent MC57 cells expressing either the TCR antigen (OVA₂₅₇) or the CAR antigen (HER2) and imaged using live-cell microscopy. Individual conjugates were quantitated for the time taken after T cell calcium flux until PI entering the target cell cytosol and the fold change in Ca²⁺ intensity, as measured by Fluo-4-AM MFI. Each point represents one conjugate and data pooled from three independent experiments. Student's *t* test, ****P* < 0.001. (D) CAR.OTI CTLs were added to plates precoated with either PBS (UT), isotype control (Iso), anti-CD3, anti-CD3 + anti-CD28, or anti-myc-tag (CAR). Supernatant was collected after 5 h and analyzed for IFN- γ and TNF levels. Cytokine expression levels (mean \pm SEM) in picograms per milliliter of three independent pooled experiments are shown. Statistical significance was determined using Student's *t* test between groups, **P* < 0.05.

difference in IFN- γ and TNF α secretion from carCTL and tcrCTL following activation. The second-generation CAR used in this study contains the CD28 and CD3 ζ endodomains, and therefore we activated CAR.OTI CTLs via either the TCR alone, TCR and CD28, or the CAR (via the c-myc tag domain). However, regardless of stimulation conditions, similar levels of both IFN- γ and TNF were produced (Fig. 2D).

To further explore differences between carCTL and tcrCTL signaling we employed a large-scale reverse-phase 308 protein array to identify any signaling pathways that were differentially activated. We found no significant difference in signaling protein expression at 2 min; however, at 30 min, 43 of 308 proteins in tcrCTL versus carCTL were significantly changed (Fig. 3A). Out of the 43 differentially expressed proteins we identified signature proteins that are involved in TCR signaling pathways, protein trafficking, and cell survival and proliferation (Fig. 3B). Interestingly, the proximal signaling protein PKC δ , which has previously been shown to localize to the secretory lysosomes in CD8 T cell killing (25), was down-regulated in carCTLs compared with tcrCTLs by 30 min, supported by a similar reduction in downstream phosphorylation of JNK and ERK pathways. This suggested that carCTL signaling is more rapid and of shorter

duration compared with tcrCTLs. Despite this shorter signaling duration carCTLs maintained a higher expression of both Bcl-2 and PEA-15 over 30 min (Fig. 3B). The cellular functions of PEA-15 are yet to be fully elucidated, but it has been previously shown that PEA-15 can negatively regulate ERK1/2, the Fas-associated death domain, and phospholipase D1 proteins, indicating that it may have antiapoptotic, antiproliferative, and antiinflammatory properties (26). Overexpression of BCL2 has also been shown to enhance the therapeutic efficacy of tumor-specific T cells (27). How CAR signaling affects levels of Bcl-2 or the long-term survival of T cells has yet to be determined. cMET is a receptor tyrosine kinase which activates a large range of cellular pathways including PLC γ 1 and PI3K and was down-regulated in carCTLs compared with tcrCTLs. We validated these differences in distal ERK signaling using Western blot analysis, which showed a rapid decrease in phospho-ERK levels by 30 min in carCTLs (Fig. 3C). Taken together, these data confirm our observations in the tumor coculture system that CAR-initiated signaling is more rapid and resolves faster than signaling through the TCR. This may have functional sequelae in carCTL, including altered kinetics of cytotoxic granule movement compared with that in tcrCTL.

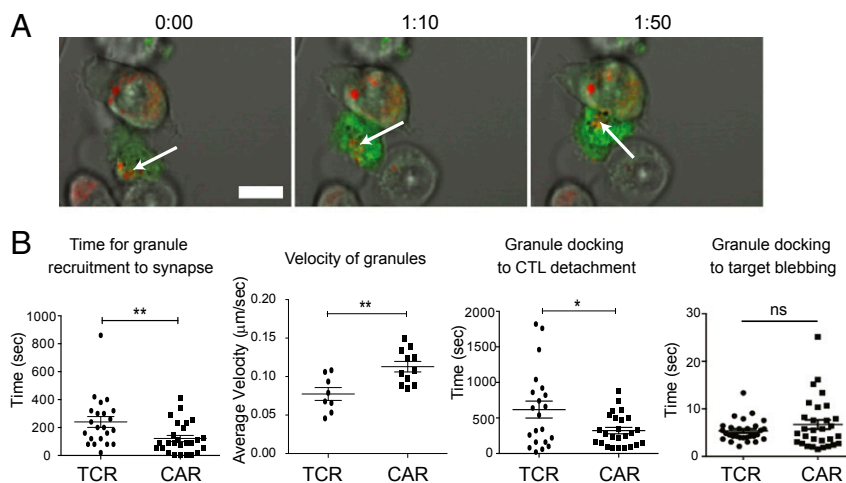


Fig. 4. CAR ligation results in faster granule recruitment to the IS. Effector CAR.OTI CTLs were labeled with Fluo4-AM and lysotracker red, added to adherent MC57 cells expressing either the TCR antigen (OVA₂₅₇) or the CAR antigen (HER2), and imaged using live-cell microscopy. (A) Representative montage of a carCTL conjugate polarizing granules toward the IS. White arrow highlights granules. Time in minutes:seconds. (Scale bar: 10 μm.) (B) Individual conjugates were quantitated for the time taken after calcium flux for granule mass to be recruited to the IS, the velocity of the granule mass, and the time from granule recruitment to CTL detachment. Each point represents one conjugate and data pooled from three independent experiments. Student's *t* test, ***P* < 0.001, **P* < 0.01. See [Movie S2](#).

insufficient antigen present for high-avidity T cells to initiate cytotoxicity (31). Whether or not CARs with different affinities and avidities will affect the kinetics of cytotoxicity is yet to be determined. However, this may be an essential feature to consider for designing better CAR-T responses.

Concluding Remarks

Our study compares CAR and TCR ligation events in the same T cell. Taken together, our dual-receptor CAR-T cell model revealed a number of distinct features of CAR-T cell IS structure, signaling, and cytotoxic functional responses which were different depending upon which receptor encountered antigen. This is summarized in Fig. 5. carCTL detection of cognate antigen initiated an alternate IS characterized by a disorganized multifocal pattern of Lck arrangement, confirming previous reports of a convoluted synaptic structure (16), a small actin ring, and diffuse LFA-1 distribution (Fig. 5B). This was in stark contrast to the IS structure observed in trCTL that comprised a classic bull's-eye structure with a peripheral LFA-1 adhesion ring (Fig. 5A). Work over the past decade has highlighted the importance of endosomal trafficking of Lck as a critical mediator of T cell signaling (32–34). Alcover and coworkers (35) have suggested that Rab11 interacting with family members such as FIP3 (Rab11 family interacting protein-3) controls Lck trafficking to the IS through its interaction with dynein and microtubules. Therefore, when assessing the disrupted Lck labeling at the IS in our studies we cannot rule out Lck localization at an intracellular pool less than 1 μm from the plasma membrane.

In our previous publication we showed that carCTLs displayed a faster time to detach from dying target cells compared with trCTLs. Here we show that the carCTL IS-initiated signaling was more rapid but of shorter duration than that observed for trCTL (Fig. 5C). This rapid carCTL signaling induced an accelerated delivery of cytotoxic granules to the IS, leading to faster killing of tumor targets. The carCTL displayed an exquisite killing function, induced by a signal of shorter duration (Fig. 5C), and a lack of LFA-1 clustering. Together this enabled the carCTL to detach from the dying tumor cell faster and move on to the next tumor target (serial killing). In contrast, the trCTL classic IS structure drove a slower, more protracted signaling resulting in slower but more extended killing.

CAR-T cell functional responses are heavily influenced by CAR design, including the scFv affinity for antigen, spacer length, and signaling endodomains (reviewed in ref. 36). Future studies exploring the effect of lower-affinity HER2 scFv in the second-generation CAR, coupled with differing signaling endodomains (e.g., CD137), may reveal a CAR-T cell with slower, more protracted signaling and cytotoxic responses, similar to that generated in trCTL. The endodomain design of CARs will ultimately influence the programming of the T cell. In recent studies, swapping CD28 with the CD137 costimulatory domain resulted in increased persistence of CAR-T cells (37). Whether or not the initial synapse structure contributes to this remains to be determined. Future studies focusing on the CAR IS in live imaged carCTL interactions or high-resolution structured illumination microscopy would be helpful to elucidate the exact nature of the CAR IS as well as the recruitment and retention of important synaptic molecules. Additionally, based on our findings, the dual-antigen receptor concept in human CAR-T cells could be used to “tune” the CAR-T cytotoxic response to a desired response depending on the disease status resulting in (i) a more rapid serial killer (via the CAR) where the disease burden is initially high or (ii) a slower but more protracted cytotoxic response (via the TCR), particularly in the case of tumor relapse. In addition, our prior data (8) showed that carCTL responses initiated loss of CAR expression after 20 h and loss of cytotoxic response to tumor. Similar data were also observed with TCR-engineered T cells; however, reexpression of the engineered TCR was observed following stimulation of the endogenous TCR (38). Thus, stimulating carCTL cells via the TCR (e.g., via a vaccine) may induce reexpression of the CAR and recovery of tumor cytotoxic function. Recent reviews have highlighted technologies that could help to quantify the carCTL IS, similar to that of the trCTL (23, 39). These technologies include high-speed, live-cell microscopy techniques recently used in TCR cell biology published by Griffiths and coworkers (29).

In conclusion, this study defined the combination of unique factors that drive CAR-T cells to be efficient killers. In doing so, we provide potential mechanisms which contribute to a CAR-T cell dose eradicating large tumor burden or controlling minimal residual disease.

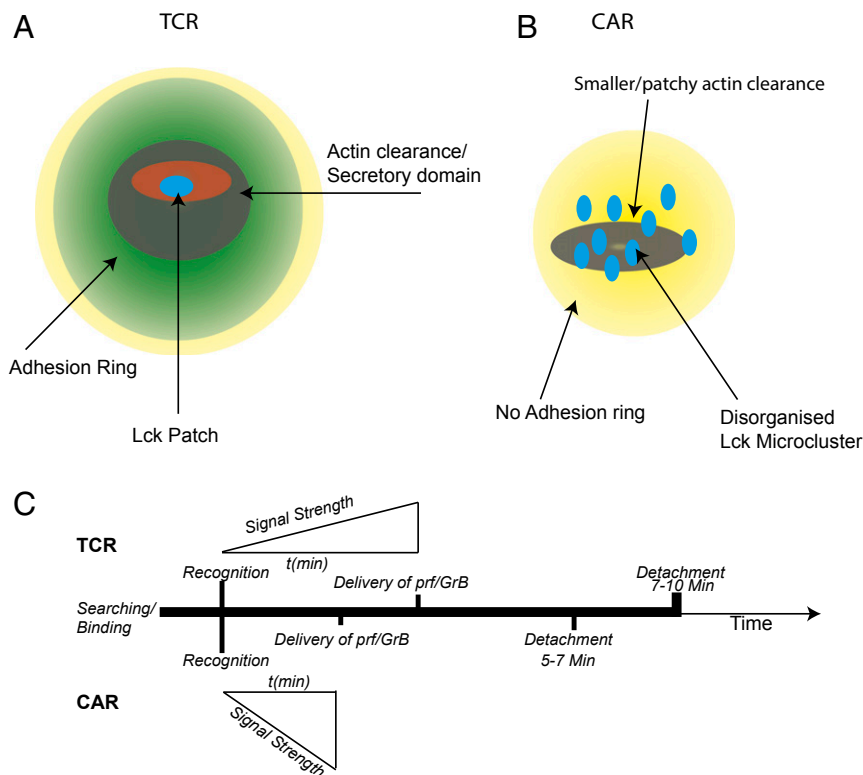


Fig. 5. Schematic model of the CAR and TCR IS and temporal killing event differences. Schematic of the IS in CTLs when activated via the CAR or TCR. (A) TCR-mediated conjugates display a classical bull's-eye structure with well-characterized SMAC domains; however, signaling via CAR (B) results in disorganized Lck patterns and lack of a pSMAC domain, as labeled by LFA-1 with a smaller actin clearance. (C) Timeline depicting temporal differences in killing events by CAR. OTI CTLs when activated by the TCR or CAR.

Materials and Methods

Cell Culture. MC57 (mouse fibrosarcoma cell line), EL4 (mouse T cell thymoma cell line), and their retrovirally transduced derivatives (OVA and Her2) (8) were cultured in complete media: RPMI-1640 comprising 10% FCS, 1 mM sodium pyruvate, 2 mM L-glutamine, 0.1 mM nonessential amino acids, 100 U/mL penicillin/streptomycin (Invitrogen, Life Technologies), and 5 μ M 2ME (called cRPMI). All T cell cultures were maintained in cRPMI with the addition of 100 U/mL of recombinant human IL-2 (T cell media).

Antibodies and Reagents. Antibodies for confocal immunofluorescence included the rat anti-mouse LFA-1 (clone C71/10; Biobynt), anti-mouse Lck (clone 3A5; Abcam), rabbit anti-mouse actin (clone 13E5; Cell Signaling), and rabbit anti-mouse γ -tubulin (clone T5192; Sigma). We used Hoechst 33342 from Sigma-Aldrich to counterstain the nucleus. Secondary antibodies included polyclonal goat anti-mouse-488 (A11001), goat anti-rabbit-546 (A11035), and goat anti-rat-647 (Invitrogen). Antibody for the LFA-1 blocking assay was purified rat anti-mouse CD11a (LFA-1 α subunit) (clone M17/4; Biolegend) and the corresponding rat IgG2a isotype control (clone GL117; WEHI-Bundoora Antibody Production Facility).

Antibodies for Western blotting included the rabbit anti-pLck (pTyr394) (GTX38561; GeneTex Inc.), rabbit anti-Lck (clone GTX24896; GeneTex Inc.), rabbit anti-PhosphoERK p44/42 (clone D13.14.4E; Cell Signaling Technology), rabbit anti-ERK1/2 p44/42 (clone 137F5; Cell Signaling), and mouse anti-actin (clone 13E5; Cell Signaling). Secondary HRP-conjugated antibodies included anti-rabbit Ig (E0432) and anti-mouse Ig (F0232) (Dako). OVA₂₅₇₋₂₅₄ (SIINFEKL) peptide was obtained from GenScript. CellTrace Violet (CTV), and CellTrace Far Red (CTFR) were obtained from Invitrogen. Reagents used for live-cell imaging studies included Fluo-4-AM (1 μ M) and Pluronic F-127 (0.02% wt/vol) (Thermo Fischer) and lysotracker red (DND99 RED, used at 1:1,000; Invitrogen). Protein kinase C ζ pseudosubstrate inhibitor (P1614) (used at 10 μ M) was obtained from Sigma-Aldrich.

Mouse Model and Generation of CAR.OTI CTLs. The dual-receptor T cells used in this study have been described previously (8). In brief, the recently described "CAR" mouse model, in which a second-generation CAR (anti-HER2-CD28-

CD3 ζ) is expressed in all immune system cells under the control of the *vav* promoter (40), was crossed with the OTI mouse to create the CAR.OTI mouse. F1-generation CAR.OTI mouse splenocytes were activated via their TCR as previously reported (41). Briefly, naive splenocytes were cultured in cRPMI media with 10 nM SIINFEKL peptide and 100 U/mL IL-2 for 3 d. Viable cells were recovered by density gradient centrifugation and cultured for a further 4 d, before being utilized in experiments as activated effector CD8⁺ T cells (termed CAR.OTI CTL).

Generation of CEA-Specific CAR OTI T Cells. To generate CEA-specific second-generation (anti-CEA-CD28-CD3 ζ) dual-specific CAR T cells, OTI mouse splenocytes were activated via their TCR, as described above for 24 h, before being transduced using retrovirus harvested from the GP.E86 viral packaging cell line as previously published (42). Briefly, following activation with 10 nM SIINFEKL peptide, T cells were transduced using spinfection for 1 h at 1,000 \times g, 22 $^{\circ}$ C with Retronectin (32 μ g/mL). Following transduction, CAR T cells were cultured in cRPMI media with 100 U/mL IL-2 for 5 d before being utilized in experiments as activated effector CD8⁺ T cells.

Conjugate Assay with LFA-1 Blocking. Preactivated effector CAR.OTI CTLs were labeled with 0.5 μ M CTV. Target parental EL4 cells were labeled with 12.5 nM CTFR and EL4-OVA and EL4-Her2 were labeled with 100 nM CTFR. The CTLs were incubated in the presence or absence of 20 μ g/mL LFA-1 blocking antibody at 37 $^{\circ}$ C for 30 min. The targets were added in an equal volume without washing, giving a final ratio of 1:1:1 EL4:EL4-OVA:CAR.OTI or 1:1:1 EL4:EL4-Her2:CAR.OTI. Cell mixtures were incubated for 10, 20, or 40 min at 37 $^{\circ}$ C, fixed with paraformaldehyde (2% final concentration), washed and analyzed by flow cytometry.

Cytokine Bead Array. Activated CAR.OTI T cells (10^5) were cultured in 96-well flat-bottom plates in T cell media, pre-coated overnight with PBS, 1 μ g mouse IgG2a Isotype control (Cell Signaling) (clone7H8/50), 1 μ g/mL anti-CD3 (145-2C11) (BD Biosciences), or 1 μ g/mL anti-myc-Tag antibody (9B11; Cell Signaling Technology) for 5 h. Cytokine production was measured from 10 μ L of culture supernatant of CTLs using CBA flex sets

(BD Biosciences) for mouse IFN- γ and TNF, according to the manufacturer's instructions. Samples were analyzed on a FACS VERSE using FACS Array software version 3.0 (Softflow) and the concentration of cytokine was determined to a standard curve and plotted as picograms per milliliter of cytokine.

Confocal Immunofluorescence. Effector CTLs and target cells were resuspended 1:1 in serum-free RPMI at 4×10^6 cells per mL, centrifuged at 200 rpm, and incubated for 5 min at 37 °C. Cells were then added to glass multiwell slides (Hendley Slides) and allowed to adhere to glass for 15 min before fixation on ice for 5 min with -20 °C 100% methanol. Slides were then blocked in PBS 2% BSA for 1 h at room temperature and incubated with primary antibodies for 1 h at room temperature and with secondary antibodies for 1 h at room temperature. Nuclei were stained with Hoechst (1:10,000; Sigma-Aldrich) in PBS for 5 min before mounting with 1.5 coverglass and mowiol-mounting medium (Sigma-Aldrich). Slides were examined using a Nikon C2 confocal microscope or Elyra Zeiss Microscope with SIM (63 \times objective under oil) and lasers exciting at 405 nm (41.5 mW), 488 nm (21.2 mW), and 561 nm (20.2 mW). Image analysis was conducted using MetaMorph Imaging Series 7 software (Universal Imaging), Imaris 7 (Bitplane), or Volocity (PerkinElmer). Images are presented as either single planes or projections, as stated in legends. We used an unbiased blinded approach for the analysis of scoring protein labeling at the IS. For blind scoring of Lck labeling at the IS a 2- μ m slice across the IS was isolated for confocal examination, excluding intracellular pools of Lck from the analysis.

T Cell-Signaling Protein Detection. CAR.OTI cells were activated via either (i) solid phase antibodies or (ii) antigen-expressing tumor target cells. (i) CAR. OTI CTLs were cultured at 4×10^6 /mL with plate bound antibodies at 1 μ g/mL; Armenian hamster anti-CD3e (clone 145-2C11; BD Bioscience), hamster anti-CD28 (clone 37-51; BD Bioscience), or mouse anti-c-myc Tag (clone 9B11; Cell Signaling). (ii) CAR.OTI CTLs and MC57 or MC57HER2 tumor target cells were incubated in serum-free media ($\pm 1 \mu$ M SIINFEKL peptide) for 1 h before coculturing 1:1 and incubating at 37 °C for various time points, before snap freezing (LN₂) in the presence of RIPA buffer (25 mmol/L Hepes, 0.25 mol/L NaCl, 2.5 mmol/L EDTA, and 0.1% Triton X-100), protease inhibitor mixture, and PhosphoStop (Roche Diagnostics). SDS PAGE followed by Western blot was then performed on the cell lysates using a 4–12% Bis-Tris gel (Invitrogen). PVDF membranes were probed with antibodies for 16 h at 4 °C, washed in PBS-T, and probed for 1 h at room temperature with HRP-conjugated secondary antibodies (anti-rabbit Ig or anti-mouse Ig; Dako), washed in PBS-T (3 \times 5 min at room temperature) before digital imaging.

Reverse-Phase Protein Array and Bioinformatics Analyses. CAR.OTI CTLs (1×10^7 /mL) were transferred to 48-well plates (5 $\times 10^5$ per well) precoated for 16 h (4 °C) with purified activating antibodies in PBS. Cells from two individual mice were incubated at 37 °C in 5% CO₂ for 2 min, 10 min, or 30 min before being snap-frozen (LN₂) and delivered to the MD Anderson reverse-phase protein array (RPPA) core facility on dry ice. Cell pellets were then lysed in RIPA buffer and serially diluted twofold for five dilutions and arrayed using a nitrocellulose collated slide. Protein lysate samples were probed with antibodies using tyramide-based signal amplification and visualized using DAB-colorimetric reaction. All sample slides were then scanned with a high-resolution scanner and spots from samples were

identified and density quantified with Array-Pro Analyzer. Relative protein levels were determined for each dilution curve from a standard curve included on each slide. The standard curve was created in R by the RPPA core at MD Anderson. RPPA staining was performed with 308 different protein antibodies and internally quality-controlled, with a quality control score above 0.8 indicating good antibody labeling. NormLog₂ data from RPPA analysis was then used to plot graphs in GraphPad Prism. NormLog₂ data were converted to their original scale and then quantile-normalized. These normalized values were log₂-transformed and fed into the Bioconductor R package limma for linear modeling (43). Protein level changes were assessed using the empirical Bayes moderated-*t* statistic (44). Empirical Bayes moderated-*t* *P* values were adjusted to control the false discovery rate (FDR). Proteins with significant level changes were called if they achieved a FDR of 0.25 or less. Data shown are a box-and-whiskers plot of *n* = 2 from minimum to maximum.

Time-Lapse Live Microscopy. Interactions between CTLs and tumor cells were assessed by time-lapse live microscopy, using well-established protocols published previously (24, 41, 45). Briefly, adherent MC57- or HER2-expressing MC57 (MC57-HER2) target cells were prepared for live-cell imaging by seeding 3×10^4 cells in each well of an eight-well chamber slide (ibidi) and incubating overnight at 37 °C/10% CO₂. MC57 target cells were then incubated for 1 h with 1 μ M SIINFEKL, OVA₂₅₇ peptide. CAR.OTI CTLs (3×10^4 per well) were labeled with Fluo-4-AM and Pluronic F-127 carrier for 20 min at 37 °C/10% CO₂. LysoTracker red was added to each chamber well to monitor granule movement kinetics. To monitor perforation of the target cell membrane 100 μ M propidium iodide was added (24). Chamber slides were mounted on a heated stage within a temperature-controlled chamber maintained at 37 °C and constant CO₂ concentrations (5% "The Brick"; ibidi). Optical sections were acquired through sequential scans of Fluo-4 (excitation 488 nm), LysoTracker Red (excitation 561 nm), or Brightfield/DIC on a Leica SP5 confocal microscope (Leica Microsystems) using a 403 (NA 0.85) air objective and Leica LAS AF software. Images were acquired every 20 s. Image analysis was performed using Leica LAS AF Lite software and MetaMorph Imaging Series 7 software (Universal Imaging), Imaris 7 (Bitplane), or Velocity (Perkin-Elmer).

Chromium Release Assay. CAR.OTI CTLs were cocultured with HER2-expressing EL4 cells (5:1 E:T ratio) in a 4-h ⁵¹Cr release killing assay (as described previously in ref. 8) in the presence or absence of protein kinase C ζ pseudosubstrate inhibitor (46, 47).

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