γδTCR immunoglobulin constant region domain exchange in human αβTCRs improves TCR pairing without altering TCR gene-modified T cell function

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Abstract. The adoptive genetic transfer of T cell receptors (TCRs) has been shown to be overall feasible and offer clinical potential as a treatment for different types of cancer. However, this promising clinical approach is limited by the serious potential consequence that exogenous TCR mispairing with endogenous TCR chains may lead to the risk of self-reactivity. In the present study, domain-exchange and three-dimensional modeling strategies were used to create a set of chimeric TCR variants, which were used to exchange the partial or complete constant region of αβTCR with corresponding γδTCR domains. The expression, assembly and function of the chimeric TCR variants were examined in Jurkat T cells and peripheral mononuclear blood cells (PBMCs). Genetically-encoded chimeras were fused with a pair of fluorescent proteins (ECFP/EYFP) to monitor expression and the pairing between chimeric TCRα chains and TCRβ chains. The fluorescence energy transfer based on confocal laser scanning microscopy showed that the introduction of γδTCR constant sequences into the αβTCR did not result in a global reduction of mispairing with endogenous TCR. However, the TCR harboring the immunoglobulin-like domain of the γδTCR constant region (i.e., TCRΔIgC), showed a higher expression and preferential pairing, compared with wild-type (wt)TCR. The function analysis showed that TCRΔIgC exhibited the same levels of interferon-γ production and cytotoxic activity, compared with wtTCR. Furthermore, these modified TCR-transduced T cells retained the classic human leukocyte antigen restriction of the original TCR. The other two chimeric TCRs, had either exchange of the cp-tm+ic domain or exchange of the whole C domain (Fig. 1). Ultimately, exchange of these domains demonstrated defective function in the transduced T cells. Taken together, these findings may provide further understanding of the γδTCR constant domain with implications for the improvement of TCR gene transfer therapy.

Introduction

In previous years, the potency of genetically modified T cell-mediated immunity against viruses and certain malignancies has been well established. TCR gene adoptive therapy is a clinically promising approach for the treatment of malignant tumors and viral diseases. Using ex vivo gene transfer, T cells isolated from patients can be genetically engineered to express a novel TCR, and the engineered T cells are re-infused back into the patient to specifically recognize a tumor-associated antigen and thereby selectively lyse tumor cells (1). However toxicity has been observed in clinical trials using genetically modified TCR therapies (2). An important toxic effect is on-target off-tumor activity, which occurs if the peptide target sequence of the TCR is also expressed on non-target proteins (3). This toxic effect may also result from the mixture of TCRs generated by the introduced TCR-α- and β-chains mispairing with the endogenous TCR β- and α-chains. The mispaired TCR increases the risk of unknown specificity causing autoreactivity (4). No formal observations of toxicities mediated by TCR mispairing have been observed in clinical trials to date, however, preclinical studies have demonstrated that mispaired TCRs have the potential to induce the harmful recognition of self-antigens, resulting in graft, vs. host disease (5). These findings indicate the requirement to prevent or reduce TCR mispairing, to...
improve T cell avidity and reduce potential off-target toxicity, including the genetic modification of TCR transgenes (10-13), disruption of endogenous TCR chains via short hairpin RNA or zinc finger nucleases (14,15), αβ TCR transfer to γδT cells or γδ2TCR transduction of αβT cells (16,17).

Although it has been reported that the transfer of γδ2TCR into αβT cells can prevent the formation of mixed TCR dimers and efficiently kill cancer cell lines in vitro (17), the role of the Vγ9Vδ2 TCR in antigen recognition remains to be fully elucidated, as does the biology of γδTCR, compared with αβTCR. Thus, the present study aimed to examine whether the domains of the γδTCR constant exchanged in αβTCR can improve the pairing and function of αβTCR. Three chimeric TCR variants were constructed, and domain-exchange and three-dimensional (3D) modeling strategies were applied, in which the αβTCR constant was replaced with partial or complete constant regions of γδTCR, leaving the variable domains intact. Subsequently, genetically-encoded reporters coupled with a pair of fluorescent proteins were constructed to monitor the expression and pairing between chimeric TCRα chains and TCRβ chains using a confocal laser scanning microscope (CLSM) in living cells. The data showed that swapping of the αβTCR constant region of immunoglobulin-like (Ig) domain for the corresponding γδTCR domain enhanced expression and reduced mispairing on the cell surface. The other two chimeric TCRs harboring the connecting peptide, transmembrane and intracellular (cp+tm+ic) domains or complete constant (C) domain of γδTCR did not show improved expression, however, the level of mispairing decreased. Finally the function of the chimeric TCR variants were examined in peripheral blood mononuclear cells (PBMCs), which revealed that the introduction of γδTCR constant region of Ig domains in the αβTCR was able to mediate the same levels of interferon (IFN)-γ secretion and cytotoxic activity as the wild-type (wt)TCR when co-cultured with human leukocyte antigen (HLA)αβ human hepatocellular carcinoma cells. However, the other two TCRs containing the γδTCR cp+tm+ic domains or C domains did not trigger the lymphocytes to produce IFN-γ or activate cytotoxic T cells when co-cultured with HLA-A2 or HLA-A2 target cells. Taken together, these findings demonstrated that exchange of the constant region of the Ig domain of γδTCR in αβTCR decreased mispairing without compromising T cell function, however, this was not the case in those containing the γδTCR cp+tm+ic or C domains.

Materials and methods

Cells. PBMCs of HLA-A2 were isolated from the blood of healthy donors (Table I), following the provision of informed consent, using Ficoll gradient centrifugation at 600 x g for 20 min at room temperature, followed by washing in PBS, re-suspension at a concentration of 1x10^6 cells/ml and activated by soluble anti-CD3ε mAb (OKT3, 30 ng/ml, R&D Systems, Inc., Minneapolis, MN, USA) and soluble anti-CD28 mAb (1 ng/ml, R&D Systems, Inc.) and 300 IU/ml recombinant human IL-2 (R&D Systems, Inc.) at 37°C for 48 h. Human PBMCs and Jurkat/E6-1 cells (cat. no. TIB-152; American Type Culture Collection, Manassas, VA, USA) were cultured in RPMI-1640 medium (Gibco; Thermo Fisher Scientific, Inc., Waltham, MA, USA) supplemented with 10% FBS (Gibco; Thermo Fisher Scientific, Inc.) and 100 U/ml penicillin-streptomycin at 37°C and 5% CO₂.

The HEK293 human embryonic kidney cell line, and the HepG2 (HLA-A2), Huh-1 (HLA-A2) and BEL-7402 (HLA-A2) human hepatocellular carcinoma cell lines (Guangdong Province Key Laboratory for Biotechnology Drug Candidates, Guangzhou, China) were cultured in DMEM (Gibco; Thermo Fisher Scientific, Inc.) supplemented with 10% FBS (Gibco; Thermo Fisher Scientific, Inc.) and 100 U/ml penicillin-streptomycin at 37°C and 5% CO₂.

Vector construction. The αβTCR was isolated from tumor-infiltrating lymphocytes of a patient (HLA-A2, α-fetoprotein+) with hepatocellular carcinoma, which was preserved in our laboratory, as described previously (Table II) (18). The γδ2TCR was isolated from healthy human PBMCs and was composed of TRGV9/12/C1 and TRDV2/D3/J1/C1. TCR-V(D)J gene nomenclature was according to http://www.imgt.org. Unmodified wtTCR was used as a control TCR.

Three TCR variants were constructed using a domain-exchange strategy in which the IgC, cp+tm+ic and C regions of αβTCR were exchanged for corresponding regions of γδTCR. These three chimeras were termed TCRαIgC; TCRαCp+tm+ic; and TCRαC, respectively. ∆ indicates a lack of αβTCR domain/s and replacement by corresponding γδTCR domain/s. The exact boundaries of the IgC, cp+tm+ic and C domains of TCRα, TCRβ, TCRγ and TCRδ are described in the legend of Fig. 1. To measure the pairing between the TCR α- and β-chains, the three modified TCRs were coupled to a pair of fluorescent proteins, ECFP and EYFP, and adenoviral particles without fluorescent proteins were constructed and produced, as described previously (18). Primer sequences used for cloning the TCRα-s and β- fusion genes and TCR are provided in Tables III, IV and V. All TCR constructs were sequence verified.

Cell transfection. The TCR variants fused to the pair of ECFP and EYFP fluorescent proteins were transduced into Jurkat cells and BEL-7402 cells at a density of 1x10⁶ cells/ml, using Lipofectamine LTX/PLUS (Invitrogen; Thermo Fisher Scientific, Inc.) according to the manufacturer’s protocol.

Image acquisition and fluorescence resonance energy transfer (FRET) analysis. Confocal images of cells were captured using an Olympus Fluoview1000 CLSM with FV10-ASW 1.7 software (Olympus, Tokyo, Japan), and apparent FRET efficiency was calculated, as described previously (18). Briefly, the Jurkat cells transduced with chimeric TCR constructs were immobilized onto a glass-bottomed dish, and the TCR construct-transduced BEL-7402 cells grown on the glass-bottomed dish were washed twice with PBS solution. The ECFP channel was excited with a 458 nm argon laser as a donor and the EYFP channel was excited with a 515 nm argon laser as an acceptor. Subsequently, seven images were captured to calculate FRET efficiency. This was calculated using the sensitized acceptor emission method using the following equation: Efficiency = 1 - I₃₄₀/(I₃₄₀+pFRET(I₃₄₀)(ψ₁₃₄₀/x)(Q₃₄₀/Qₒ)), where pFRET is the processed FRET obtained by removing the donor SBT (DSBT) and the acceptor SBT (ASBT) from the contaminated or uncorrected FRET; I₃₄₀ is the intensity of the donor in the presence of the acceptor; ψ₁₃₄₀ and ψ₃₄₀ are the...
collection efficiencies in the donor and acceptor channel; and $Q_d$ and $Q_a$ are the quantum yield of the donor and acceptor, respectively. Statistical analysis of the mean FRET efficiency were calculated from multiple ($n=4$) cell images in each group and five randomly selected regions of interest (ROI) in each cell image.

**TCR adenovirus construction and transduction of T cells.**

For production of the adenovirus, HEK-293 cells were transfected with the respective TCR-encoding shuttle plasmid and second-generation Ad5F35 adenoviral packaging plasmid (pBHGIoxdelE13Cre; Biovector Science Lab, Inc., Beijing, China). The HEK-293 cells were seeded in 6-well plates at a density of $1\times10^6$ per well. After 24 h, the cells were co-transfected with an equimolar ratio of the two plasmids (2.5 µg total DNA per well) and 6 µl/well Lipofactamine 2000 transfection reagent (Invitrogen; Thermo Fisher Scientific, Inc.) according to the manufacturer’s protocol. The adenovirus supernatants were harvested ~12 days following transfection, cellular debris was removed by centrifugation at 14,000 x $g$ for 10 min at room temperature. Adenovirus particle titers were determined using the TCID50 method and the supernatants were directly used for transduction of the Jurkat T cells and PBMCs, respectively, as previously described (18).

**Flow cytometry and analysis.** The surface expression of the transgenic TCRs on the cells were assessed by fluorescein isothiocyanate (FITC)-conjugated anti-TCRα12.1 mAb (cat. no. TCR2764; dilution, 1:500; Invitrogen; Thermo Fisher Scientific, Inc.) and PE-conjugated anti-TCRβ7.1 mAb (cat. no. IM2287; dilution, 1:500; Beckman Coulter, Inc., Brea, CA, USA). The transduced Jurkat T cells and PBMCs ($5\times10^5$) were stained with the mAbs on ice for 30 min. Following washing with PBS, the cells were fixed in 2% PFA prior to measurements on an Epics-XL flow cytometer (Beckman Coulter, Inc.). Non-transduced Jurkat T cells and PBMCs were used as controls.

**Cytokine release assays.** The unmodified PBMCs and TCR-modified PBMCs were assessed for reactivity in IFN-γ release assays using commercially available ELISA kits (Boster Systems, Inc., Pleasanton, CA, USA). The HepG2 (HLA-A*), Huh-1 (HLA-A*) and BEL-7402 (HLA-A*) target human hepatocellular carcinoma cell lines were cultured in medium at 37°C, followed washing with PBS prior to the initiation of co-cultures. For these assays, $3\times10^5$ responder cells (PBMCs) and $1\times10^5$ stimulator cells were incubated in a 0.2 ml RPMI-1640 medium (Gibco; Thermo Fisher Scientific, Inc.) supplemented with 10% FBS and 100 U/ml

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Table I. Details of blood donors.

| Number | Gender | Age | Hospital admitted to | Dates of blood donation |
|--------|--------|-----|----------------------|------------------------|
| 1      | Female | 24  | The First Affiliated Hospital/School of Clinical Medicine of Guangdong Pharmaceutical University | 2013.9.15, 2013.9.21, 2013.9.30, 2013.10.9, 2013.10.20 |
| 2      | Male   | 25  |                       | 2013.9.15              |
| 3      | Male   | 30  |                       | 2013.9.15, 2013.9.21, 2013.9.30, 2013.10.9, 2013.10.20 |
| 4      | Female | 28  |                       | 2013.9.15              |
| 5      | Male   | 21  |                       | 2013.9.15, 2013.9.21, 2013.9.30, 2013.10.9, 2013.10.20 |

Table II. Patient details of patient with hepatocellular carcinoma.

| Number | Gender | Age | Hospital admitted to | Dates of blood donation |
|--------|--------|-----|----------------------|------------------------|
| 012    | Male   | 54  | The First Affiliated Hospital/School of Clinical Medicine of Guangdong Pharmaceutical University | 2010.5.9               |
penicillin/streptomycin in individual wells of 96-well plates and were co-cultured for 24 h at 37°C and 5% CO₂. The secretion of IFN-γ was measured in the culture supernatants diluted to be in the linear range of the assay.

**CTL assay.** The abilities of the transduced PBMCs to lyse the HLA-A²⁺/HLA-A²⁻ human hepatocellular carcinoma targets were measured using a calcein AM (CAM) release assay (Dojindo Molecular Technologies, Inc., Kumamoto, Japan), as described previously (19). Briefly, 1x10⁸ tumor cells were labeled with 2 μM CAM, which was diluted from a 1 mM stock solution of CAM in dimethyl sulfoxide (Sigma-Aldrich; Merck Millipore, Darmstadt, Germany) for 30 min at 37°C. The labeled target cells were washed three times with PBS and re-suspended at a concentration of 1x10⁷ cells/ml in complete medium. The labeled target cells (1x10⁴ in a volume of 100 μl) were plated in 96-well V-bottomed plates with effector cells in 200 μl of complete medium at effector to target cell ratios of 1:10.

| Gene name | Primer name | Direction | Primer sequence |
|-----------|-------------|-----------|-----------------|
| TRBVC     | P1          | Forward   | ATAGCTAGGCCACCACCAGGCTGCGAGGCTGCTCTG |
|           | P2          | Reverse   | ACATCTGCATCAAGTGTTTCTCCAGTACGTCCTG |
| TRGCm     | O3          | Forward   | CGAGGCTGAGGAGGCTGACCATATCTGCTGCACCCAGGCTGCTG |
|           | O4          | Reverse   | GCTGGAGGCTGAGGAGGCTGACCATATCTGCTGCACCCAGGCTGCTG |
| EYFP      | O5          | Forward   | CTCGAATGAAGAGAAATCAATGGTGAGCAAGGGCGAG |
|           | O8          | Reverse   | CGGCCTCGACATTACTGTACAGCTGCTG |
| TRACV     | X1          | Forward   | ACCGACACACCTGGCCACCACCAGGCTGCGAGGCTGCTCTG |
|           | X2          | Reverse   | GATCCATTGTAGTGACATCTGCTCTACCCAGGCTGCTG |
| TRDCm     | X3          | Forward   | CGAGGCTGAGGAGGCTGACCATATCTGCTGCACCCAGGCTGCTG |
|           | X4          | Reverse   | GCTGGAGGCTGAGGAGGCTGACCATATCTGCTGCACCCAGGCTGCTG |
| ECFP      | X5          | Forward   | GCTGGAGGCTGAGGAGGCTGACCATATCTGCTGCACCCAGGCTGCTG |
|           | X6          | Reverse   | CTCCTGCCCTTGCTCACCACATGATCTGCTGCTG |
|           | X7          | Reverse   | AGTGCGGCCGCTTTACTGTACAGCTGCTG |

Restriction enzyme sites are underlined. SOE-PCR, splicing by overlap extension-polymerase chain reaction.
target (E:T) ratios between 1:1 and 30:1. Following incubation for 4 h at 37˚C with 5% CO₂, the supernatants was harvested and the quantities of calcein released were measured using a Varioskan Flash multimode reader (Thermo Fisher Scientific, Inc.). Spontaneous release was determined by incubating the target cells in medium alone, and maximum release was determined by suspending the cells with 1% Triton X-100. Each data point was an average of four wells. The percentage of PBMC-specific lysis was calculated as follows: Specific lysis (maximum release - spontaneous release) x 100.

Statistical analysis. Differences among the TCRs in various assays were examined using Student’s t-test (unpaired; two-tailed) and two-way analysis of variance with a Bonferroni’s multiple comparisons test using GraphPad Prism 6 software (GraphPad Software Inc., La Jolla, CA, USA). P<0.05 was considered to indicate a statistically significant difference. Data are expressed as the mean ± standard deviation.

Results

Expression of wtTCR and chimeric TCR constructs. In the present study, three chimeric TCR variants were generated, in which the IgC, cp+tm+ic and C regions of 8βTCR were replaced with corresponding regions of γδTCR (Fig. 1). The wtTCR and chimeric TCR genes were cloned separately into the pDC315 shuttle plasmid to produce the Ad5F35 adenovirus, and transduction of Jurkat T cells was performed with subsequent fluorescence-activated cell sorting analysis. Double immunofluorescent staining with anti-TCRV12 mAb<sup>FITC</sup> and anti-VC<sub>7mAb</sub><sup>PE</sup> showed that Jurkat cells transduced with TCRAlγC exhibited higher surface coexpression (17.1%), compared with wtTCR (10.7%), and TCRACP+tm+ic exhibited a marginally higher surface expression (13.4%), compared with wtTCR. By contrast, TCRAC exhibited a lower level of surface expression, compared with wtTCR (Fig. 2). To further examine the expression of the chimeras in living cells, the C terminus of chimeric TCR<sub>γδ</sub> chains and TCRβ chains were fused to a pair of cyan and yellow fluorescent proteins, ECFP and EYFP respectively (Fig. 3A and B). In Jurkat cells, the fluorescence observation using a CLSM showed that TCRAlγC was expressed more markedly on the cell surface, which was in accordance with the flow cytometry results. By

### Table V. Primers for amplifying TCRβΔC-EYFP and TCRαΔC-ECFP fusion genes (regular polymerase chain reaction (PCR) and SOE-PCR).  

| Gene     | Primer name | Direction | Primer sequence                  |
|----------|-------------|-----------|----------------------------------|
| TRBV     | P1          | Forward   | ATAGCTAGGCCACCATGGGCTGAGCTTCTTG  |
|          | P2          | Reverse   | ACATCTGCACTAAGTGGTTTCTCCAGTACGGTCAGCT  |
| TRGC+GCm | P3          | Forward   | AGGCTGACCGTACTGGAGAAACACCTTGAGCAAGCTGT  |
|          | O4          | Reverse   | CCTCGCCCTTTGCTCACATTGATTTTCTCTCCATTTGCAG  |
| EYFP     | O5          | Forward   | CTGCAATGGAGAGAAAATCAATGGTAGCAAGGGGC  |
|          | P8          | Reverse   | CGGCCGTGCAGTTACTTGTACAGCTGCAG  |
| TRAV     | X1          | Forward   | ACGCCACAAACTTGGCACCATGATATCCCTTAGAGATT  |
|          | X2          | Reverse   | GGTTTTGATGAGGCTGACATTTGTGTTTTACGTCACTGCTG  |
| TRDC+DCm | X3          | Forward   | CCAGACTGACATGAAACACATTGCCTCATACCAAACC  |
|          | H4          | Reverse   | CCTGCCCTTTGCTCACATCAAGAAATACCTGGACGAT  |
| ECFP     | H5          | Forward   | ACTGCCAAAGTTATTTTCTTGATGGTAGCAAGGGCGAG  |
|          | X8          | Reverse   | AGTCGCCAGCTATTTCTTGACTACAGCTGCAGCAT  |

Restriction enzyme sites are underlined. SOE-PCR, splicing by overlap extension-polymerase chain reaction.

Splicing by overlap extension (SOE)-PCR. This method was used to generate the fusion genes of three chimeric TCR variants. Variable fragments for generating TCRαΔC-EYFP variants (including TCRAV, TRDC, TRACm and ECFP) and TCRβΔC-EYFP variants (including TRBV, TRGC, TRBCm and EYFP) were amplified using a set of forward primers and reverse primers (Table III). The first step of SOE-PCR reactions were performed with 100 ng of the template without primers, 10X Buffer, 2 mM dNTPs, 25 mM MgSO<sub>4</sub>, 0.5 U KOD-Plus-Neo Polymerase (Toyobo Co., Ltd., Osaka, Japan) in 25 µl reaction volume. The PCR cycling conditions were as follows: Denaturation at 94˚C for 2 min, followed by 5 cycles at 94˚C for 30 sec, at 57˚C for 30 sec, and at 68˚C for 90 sec, and completed with a final extension at 68˚C for 7 min. The PCR generated overlapping gene segments that are then used as template DNA for the second step of SOE-PCR to create a full-length product. Therefore, another 25 µl reaction mixture (contained 10X Buffer, 2 mM dNTPs, 25 mM MgSO<sub>4</sub>, 0.5 KOD-Plus-Neo Polymerase and 1 µM forward primer X1/P1 and reverse primer X8/P8; Invitrogen; Thermo Fisher Scientific, Inc.) were added to the first reaction mixture for the second step of SOE-PCR. The reaction conditions were initial denaturation at 94˚C for 2 min, followed by 30 cycles at 94˚C for 30 sec, at 62˚C for 30 sec, and at 68˚C for 90 sec and a final extension at 68˚C for 7 min. The other two TCR (TCRACP+tm+ic and TCRAC) fusion genes were generate using the same methods, as the primer sequences for the amplification of variable regions and fusion chains are given in Tables IV and V.
contrast, the other two modified TCRs, TCRΔcp+tm+ic and TCRΔC, exhibited no apparent difference in expression levels, compared with wtTCR. These data suggested that the modified αβ TCRs harboring the γδ TCR constant region were expressed on the Jurkat cell surface and that the FRET reporters were able to used for monitoring the expression and interaction of TCR variants.

**Analysis of TCR mispairing.** The Jurkat cell line (cloneE6-1) was used as a recipient T cell model for TCR gene transfer, to determine whether the chimeric TCRα- and β-chains assembled preferentially when there was a pair of endogenous TCRs. It was hypothesized that the introduced TCR α- and β-chains comprising heterodimers on the cell surface results in FRET efficiency between the donor (ECFP) and acceptor (EYFP) fluorescent proteins. When the introduced TCR α- and β-chains and the endogenous TCR β- and α-chains mispair, FRET is not detected between the mispairing TCRs (Fig. 3C). As shown in Fig. 3A, seven images were used to remove the DSBT and ASBT from the contaminated FRET to obtain the FRET efficiency images. The corrected FRET efficiency images showed that TCRΔIgC exhibited a higher FRET efficiency, compared with wtTCR. The images of TCRΔcp+tm+ic and TCRΔC exhibited no differences in FRET efficiencies, compared with wtTCR. A total of four independent cell images and five ROIs in each cell image were selected for FRET efficiency analysis, respectively, in each group. The statistical results showed that the average FRET efficiency between the TCRΔIgC α- and β-chains (10.69±0.76%) was significantly higher, compared with the average FRET efficiency between the wtTCR α- and β-chains (7.92±1.32%; P<0.01). No significant differences were found between the FRET efficiencies of the other two chimeric TCRs (9.07±0.61 and 8.95±0.89%), compared with wtTCR (P>0.05; Fig. 3D).

To further investigate the extent of mispairing of the modified TCR with the endogenous TCR, BEL-7402 cells deficient in TCR and CD3 molecules were selected as the next recipient cell model. It was hypothesized that as there was no endogenous TCR in the BEL-7402 cells, the pairing of the introduced TCR α- and β-chains will not be interfered with, allowing the extent of mispairing to be measured by comparing the FRET efficiencies of BEL-7402 and Jurkat cells. The images showed that the TCRΔIgC exhibited a higher FRET efficiency, compared with the wtTCR in the BEL-7402 cells (Fig. 3B). The statistical results showed that the average FRET efficiency of wtTCR in the Jurkat cells (7.92±1.32%) was lower, compared with that in the BEL-7402 cells (10.59±1.02%; P<0.05; Fig. 3D), suggesting that the wtTCR was mispaired with the endogenous TCR in Jurkat cells, and the level of mispairing with endogenous TCR was 25%. Statistical analysis also showed that the FRET efficiencies of TCRΔIgC were decreased in the Jurkat cells (10.69±0.76%), compared with the BEL-7402 cells (13.34±0.40%), indicating a 20% mismatch rate in the Jurkat cells (Fig. 3D). TCRΔcp+tm+ic and TCRΔC also showed mispairing with the endogenous TCR (19 and 14%, respectively). Together, these data showed that replacement with various constant domains of γδ TCR in αβ TCR reduced mispairing to the same extent, but were unable to prevent mispairing.

**Analysis of TCR-transduced primary T cells.** The Ad5F35 adenovirus encoding the wtTCR and the chimeric TCRs were used to transfer into PBMCs, and PBMCs expressing the
wtTCR and chimeric TCRs were co-cultured with the HepG2 (HLA-A2+, Huh-1 (HLA-A2+)) and BEL-7402 (HLA-A2+) human hepatocellular carcinoma cell lines. The secretion of IFN-γ into the medium was measured using an ELISA procedure. As shown in Fig. 4A, when co-cultured with HLA-A2+ cell lines, the PBMCs transduced with TCRΔIgC secreted the same quantity of IFN-γ as the PBMCs transduced with wtTCR. Unexpectedly, the PBMCs transduced with TCRΔcp+tm+ic and TCRΔC secreted lower levels of IFN-γ, compared with wtTCR, with levels close to those of the PBMCs containing no TCR transgene. This indicated that TCRΔcp+tm+ic and TCRΔC did not trigger cytokine secretion. The levels of IFN-γ in the PBMCs transduced with the three chimeric TCRs or wtTCR were not above background levels following incubation with the HLA-A2+ cell line.

The cell-mediated cytotoxicity of human PBMCs expressing either wtTCR or the modified TCR variants was also compared in a 4-h CAM release assay. The Ad5F35 adenoviruses encoding the wtTCR and chimeric TCRs were transferred into PBMCs. PBMCs expressing the wtTCR and chimeric TCRs were co-cultured with CAM-labeled human hepatocellular carcinoma cell lines. It was observed that wtTCR and TCRΔIgC were able to mediate specific lysis of the HLA-A2+ hepatocellular carcinoma cell lines, as shown in Fig. 4B. The lymphocytes expressing TCRΔIgC showed equivalent lysis in the HLA-A2+ HepG2 and Huh-1 target cell lines, compared with the wtTCR at an E:T ratio of 30:1 (11.42±0.75 and 12.03±1.10%, respectively for HepG2 target cells; 11.10±0.92 and 12.13±1.17%, respectively for Huh-1 target cells). These results indicated that the lymphocytes expressing TCRΔIgC exhibited similar cytotoxic activity to wtTCR. However, the lymphocytes expressing TCRΔcp+tm+ic and TCRΔC exhibited lower lytic activity, compared with the wtTCR, however, lysis was equivalent to the control PBMCs. In the HLA-A2- target cell line, no significant lysis was observed by any of the TCR transgene PBMCs.

Taken together, TCRΔIgC substitution of the γδTCR IgC domain was functionally equivalent to the wtTCR, whereas TCRΔcp+tm+ic and TCRΔC affected the recognition and cytotoxic abilities.
**Discussion**

The αβ and γδTCRs are two types of antigen receptor expressed on distinct T cell populations. γδTCR is homologous to αβTCR in the variable and constant regions, and the αβ and γδ TCRs are heterodimers linked by disulfide bonds (20). The γδT cells carrying Vγ9Vδ2 TCRs are primarily found in peripheral blood, where they constitute a minor fraction of total T cells and respond to non-peptidic intermediates of the mevalonate pathway, termed phosphoantigens (21). It has been reported that Vγ9Vδ2TCR can be efficiently expressed in αβT cells without mispairing with αβTCR, and mediates the tumor-specific proliferation of αβT cells (17). In the present study, three chimeric TCR variants were generated by swapping the partial or complete constant regions of αβTCR with those of γδTCR (Fig. 1). These constructs were assessed for surface expression, mispairing with endogenous TCR chains, and TCR transgene-mediated functions in Jurkat T cells and the human PBMCs expressing either wtTCR or modified TCR variants were co-cultured with different tumor cell lines for 24 h, and concentrations of IFN-γ secreted into the co-culture supernatant were measured using ELISA. (B) Specific cytotoxicity of tumor cell lines. The human PBMCs expressing the wtTCR, TCRAlgC, TCRtcp+tm+ic or TCRAC transgenes, or without a TCR transgene were co-cultured with CAM-labeled tumor cell lines at the indicated E:T ratios for 4 h, following which specific lysis was calculated. Results represent the average of three independent experiments, performed with three donors. *P<0.05, **P<0.01, ***P<0.001. TCR, T cell receptor; PBMCs, peripheral blood mononuclear cells; wt, wild-type; Ig, immunoglobulin-like; cp+tm+ic, connecting peptide, transmembrane and intracellular; C, complete constant; CAM, calcein AM; IFN-γ, interferon-γ; HLA, human leukocyte antigen; E:T, effector to target cell; ns, non-significant.
primary human T cells. The subsequent observations revealed for these chimeric TCR variants that the introduction of the γδTCR IgC domain in the αβTCR improved surface expression, reduced mispairing and did not compromise the function of the unmodified wtTCR. The other two TCRs containing cp+tm+ic or C domain of γδTCR showed decreased mispairing with endogenous TCR, but impaired function in T cells.

FACS analysis in Jurkat cells showed that TCRαlgC exhibited improved surface expression, and the CLSM images also showed TCRαlgC exhibited on the cell surface of the Jurkat cells and the non-T cells (BEL-7402). The fluorescent images of the reporter were subjected to FRET analysis, which showed that TCRαlgC exhibited a higher FRET efficiency, compared with the wtTCR in BEL-7402 cells and Jurkat cells. Detailed statistical analysis of FRET efficiencies between BEL-7402 cells and Jurkat cells showed that TCRαlgC reduced mispairing, but failed to prevent mispairing with endogenous TCR. The function of TCRαlgC in cytotoxic lymphocytes showed equivalent IFN-γ secretion and cytotoxic activity as in wtTCR when targeting HLA-A2- HepG2 and Huh-1 cell lines, however not the HLA-A2- cell line. These results suggested that the γδTCR IgC domain substituted for αβTCR preserved the recognition and lytic abilities of wtTCR, and even classic HLA restriction. The TCR αβ heterodimer has three conserved basic residues (R, K and K) in the transmembrane regions. These residues are considered to drive the associations between TCR and CD3 components by forming pair-wise ionic interactions, however, the association between this charged residue and the αβTCR heterodimer remains to be elucidated (22). The present study hypothesized that TCRαlgC prevents mispairing with endogenous TCR to a certain extent, perhaps due to harboring the γδTCR IgC domain, and the preserved ability of the original TCR may be due to the αβTCR transmembrane region interacting with the CD3 complex.

By contrast, the TCRαcp+tm+ic and TCRαC did not increase the surface expression significantly, compared with the wtTCR when monitoring these TCR variants in living cells. No significant differences in FRET efficiencies were found for the TCRαcp+tm+ic and TCRαC, compared with the wtTCR in BEL-7402 cells and Jurkat cells. However, the statistical analysis of FRET efficiencies between BEL-7402 cells and Jurkat cells showed these two modified TCRs decreased mispairing. At present, the molecular mechanisms determining the efficiency of TCR pairing remain to be elucidated. Studies have shown that the variable region sequences are important in determining the efficiency of the expression of TCRs (23). In the present study, which examined TCR constant region modification (TCRαcp+tm+ic and TCRαC), minimal effect was found in the their expression efficiencies, compared with wtTCR. It was hypothesized that the variable region sequences drive efficient αβ pairing, which can proceed despite modifications in the constant region. Unexpectedly, when their function was assessed in cytotoxic lymphocytes, the present study observed that TCRαcp+tm+ic and TCRαC were unable to trigger the secretion of IFN-γ, and failed to mediate cytotoxicity in either the HLA-A2- nor HLA-A2- hepatocellular carcinoma cell lines. The TCRβ chain contains a conserved transmembrane glutamic acid, which is not found in the γ chain, and this residue may be a key determinant in the differential CD3 composition of the αβ and γδ complexes (24).

The chimeric TCRs in the present study contained γ instead of β residues in the transmembrane domain, which may have affected the composition of the CD3 subunits. Therefore, the present study hypothesized that the differences in CD3 subunit composition between the αβ- and γδTCR/CD3 complexes may have resulted in the chimeric TCRs containing a substituted cp+tm+ic domain of γδTCR failing to transduce a signal through the TCR complex. This may explain why TCRαcp+tm+ic and TCRαC lost the functions of recognition and lysis in primary T cells.

In conclusion, the present study showed that the modified αβTCR, substituted for by the IgC domain of γδTCR, improved expression and pairing on the cell surface, and did not compromise the function of the already present wtTCR.

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