Identification of neoantigen-reactive T lymphocytes in the peripheral blood of a patient with glioblastoma

Vid Leko 1, Gal Cafri 2, Rami Yossef 1, Biman Paria 3, Victoria Hill 1, Devikala Gurusamy 1, Zhili Zheng 1, Jared J Gartner 1, Todd D Prickett 1, Stephanie L Goff 1, Paul Robbins 1, Yong-Chen Lu 1, Steven A Rosenberg 1

ABSTRACT

The adoptive transfer of naturally occurring T cells that recognize cancer neoantigens has led to durable tumor regressions in select patients with cancer. However, it remains unknown whether such T cells can be isolated from and used to treat patients with glioblastoma, a cancer that is refractory to currently available therapies. To answer this question, we stimulated patient blood-derived memory T cells in vitro using peptides and minigenes that represented point mutations unique to patients’ tumors (ie, candidate neoantigens) and then tested their ability to specifically recognize these mutations. In a cohort of five patients with glioblastoma, we found that circulating CD4+ memory T cells from one patient recognized a cancer neoantigen harboring a mutation in the EED gene (EEDH189N) that was unique to that patient’s tumor. This finding suggests that neoantigen-reactive T cells could indeed be isolated from patients with glioblastoma, thereby providing a rationale for further efforts to develop neoantigen-directed adoptive T cell therapy for this disease.

INTRODUCTION

Glioblastoma, the most common primary brain malignancy in adults, remains essentially incurable with the standard therapy, which includes surgery, radiotherapy and chemotherapy. Given that only 7.2% of patients survive 5 years postdiagnosis, new therapeutic approaches are urgently needed.1

In recent clinical studies, immunotherapy with checkpoint inhibitors,2 cancer vaccines3–7 and CAR T cells8 has largely failed to induce tumor regressions in patients with glioblastoma. A single case of a transient complete response following the intracranial administration of anti-IL13Ra2 CAR T cells was a notable exception.9 Such a low response rate to immunotherapy has been attributed to the paucity of targetable tumor antigens, as well as to the presence of a strongly immunosuppressive tumor microenvironment, neuroanatomical constraints (ie, the blood-brain barrier) and marked tumor heterogeneity.10

Despite these discouraging results, several reports have indicated that glioblastoma may still be susceptible to an immune attack by endogenous T cells. For instance, in two patients whose tumors exhibited excessively high mutational burden due to a mismatch repair deficiency, immune checkpoint inhibition resulted in durable tumor regressions.11

In other studies, checkpoint inhibitors administered before brain surgery were found to promote tumor infiltration by T cells.12 13

Endogenous T cells can mediate an anti-tumor immune attack by recognizing cancer neoantigens, the proteins that are predominantly encoded by genes harboring point mutations unique to patients’ tumors. Neoantigen-reactive T cells can be isolated from patients with various cancer types and, when expanded and used for adoptive cell therapy, can lead to durable tumor regressions.14 However, it remains unknown whether this form of immunotherapy can be used to treat glioblastoma, because naturally occurring neoantigen-reactive T cells have not yet been isolated from patients with this disease.

In this study, we tested whether T cells from patients with glioblastoma could recognize neoantigens unique to patients’ autologous tumors. To this end, five patients with refractory or relapsed glioblastoma, who had been enrolled on an anti-EGFRvIII CAR T protocol,15 were selected based on the availability of blood and archived tumor specimens. Next, their tumors were subjected to whole exome sequencing (WES), and the resulting data were used to construct screening libraries that represented tumor-specific point mutations (ie, candidate neoantigens). Finally, memory T cells were isolated from patients’ blood and, following a stimulation in vitro, tested for recognition of these candidates.
MATERIALS AND METHODS

WES and RNA-seq

Genomic DNA was extracted from formalin-fixed, paraffin-embedded (FFPE) tumors using a column-based truXTRAC FFPE total NA kit (Covaris, Woburn, Massachusetts, USA). DNA and RNA from matched normal blood, as well as from the homogenized fresh tumor (Patient 1), were extracted using the Allprep DNA/RNA Mini-prep extraction kit (QIAGEN, Germantown, Maryland, USA). Following WES and RNA-seq library preparation and sequencing, tumor-specific single nucleotide variants (SNVs) and short insertions and deletions (INDELs) were identified as described previously.16 Where applicable, mutation clonality analysis was performed on WES data using PyClone,17 based on tumor purity and copy number estimates generated using Sequenza.18

Screening libraries

Mutation data were used to generate tandem minigene (TMG) and peptide pool (PP) screening libraries. TMGs were designed by linking up to 12 minigenes, each encoding either a non-synonymous point mutation (with each mutated amino acid flanked bilaterally by 12 wild type (WT) amino acids) or a frameshift INDEL (with 12 WT amino acids preceding the newly created reading frame, which terminated at the next stop codon). Their sequences were cloned into pcRNA2SL plasmids and transcribed into RNA, as described previously.16

PPs were composed of up to 12 individual crude-grade 25-mer peptides, the sequence of which corresponded to the minigenes. Peptides purified with high-performance liquid chromatography (HPLC) were used in validation experiments. All peptides were obtained from GenScript (Piscataway, New Jersey, USA).

Antigen presenting cells (APCs)

Autologous dendritic cells (DCs) and B cells were used as APCs. They were generated by purifying CD14+ and CD19+ peripheral blood mononuclear cells (PBMCs) using magnetic microbeads (BD Biosciences, San Jose, California, USA) and culturing them as described in Refs. 14 and 19, respectively.

In vitro stimulation (IVS) of memory T cells

IVS was performed following a previously described approach.16,18 Briefly, 0.5–1×10⁶ memory T cells, isolated based on CD62L and CD45RO expression (figure 1A), were cocultured (ie, stimulated) in 4:1 ratio with DCs. TMG-electroporated DCs were used to stimulate memory CD8+ cells, while PP-pulsed DCs were used to stimulate memory CD4+ and—if sample size permitted—memory CD8+ cells. This approach was supported by a recent publication that demonstrated the superior ability of TMGs and PPs to elicit detectable antigen recognition from CD8+ and CD4+ cells, respectively.20

Initial cocultures were performed in 48-well plates, using media supplemented with IL-21 (30 ng/mL) and IL-2 (6000 IU/mL) and added to 1:1 V/V ratio. On day 12, expanded T cells were cocultured with DCs or B cells loaded with the TMGs or PPs used for initial stimulation. The next day, T cells with the highest expression of 4-1BB and OX-40 were sorted and subjected to a rapid expansion protocol using irradiated allogeneic feeders and anti-CD3 antibody (clone OKT3; Affymetrix, San Diego, California, USA). Expanded cells were then tested for the recognition of patient’s TMGs or PPs (see below). Concurrently, IVS with CEFX (JPT Peptide Technologies, Berlin, Germany), a mix of virally derived peptides, was performed as a control.

Assessment of neoantigen recognition

Neoantigen recognition was evaluated by preforming overnight T cell cocultures with APCs, as described previously.16 Prior to coculturing, the APCs were either pulsed for 2 hours with individual peptides or PPs (final concentration of 10 μM) or were electroporated with TMG or full-gene RNA (0.5–1 μg RNA/1×10⁶ cells) using a Neon Transfection System (Life Technologies, Carlsbad, California, USA). T cell activation was assessed by measuring the production of IFN-γ or of multiple cytokines, or by measuring upregulation of 4-1BB and OX40 on T cells using flow cytometry.

IFN-γ production was measured by either ELISA, which was performed using a designated kit (ThermoFisher Scientific, Waltham, Massachusetts, USA), or by ELISPOT, which was performed using appropriate capture and detection antibodies (both from Mabtech, Cincinnati, Ohio, USA). Multicytokine analysis was performed using a customized U-PLEX kit (Meso Scale Diagnostics, Rockville, Maryland, USA).

Flow cytometry

For all experiments, cells were stained with antibodies diluted in PBS/0.5% FBS in 1:50 format wells, based on their growth rate. During each expansion, fresh media containing IL-21 (60 ng/mL) and IL-2 (6000 IU/mL) was added in 1:1 V/V ratio.

On day 12, expanded T cells were cocultured with DCs or B cells loaded with the TMGs or PPs used for initial stimulation. The next day, T cells with the highest expression of 4-1BB and OX-40 were sorted and subjected to a rapid expansion protocol using irradiated allogeneic feeders and anti-CD3 antibody (clone OKT3; Affymetrix, San Diego, California, USA). Expanded cells were then tested for the recognition of patient’s TMGs or PPs (see below). Concurrently, IVS with CEFX (JPT Peptide Technologies, Berlin, Germany), a mix of virally derived peptides, was performed as a control.

Identification, synthesis and transduction of T cell receptors (TCRs)

PP2-reactive T cells from Patient 1 were cocultured for 4 hours with B cells pulsed with dimethyl sulfoxide (DMSO) or PP2 and then subjected to an IFN-γ capture assay (Milenyi, Auburn, California, USA). Next, single IFN-γ+ T cells were sorted into a 96-well microplate, which was subjected to a previously outlined TCR sequencing protocol.21 TCR synthesis and transduction into allogeneic PBMCs was performed as described previously.16
TCR deep sequencing

TCR-β deep sequencing was performed on genomic DNA extracted from the sorted memory T cells by immunoSEQ (Adaptive Technologies, Seattle, Washington, USA). The data were analyzed using the immunoSEQ Analyzer 3.0.

Determination of MHC restriction element and assessment of EED<sup>H189N</sup> gene recognition

The major histocompatibility complex (MHC)-II alleles expressed by Patient 1 were synthesized and cloned into pcDNA3.1 plasmids (GeneOracle, Santa Clara, California, USA). Next, COS7 cells were cotransfected with combinations of individual plasmids and then pulsed with WT or mutant 25-mer EED<sup>H189N</sup> peptide for 2 hours. Following an overnight coculture with TCR-transduced T cells, IFN-γ ELISA was performed. To assess EED gene recognition, cotransfection with plasmids encoding the WT or mutant EED genes was performed instead of peptide pulsing.

Statistical analyses

Statistical analyses were performed on GraphPad Prism 7.0 software (GraphPad Software, La Jolla, California, USA). When applicable, data were expressed as mean±SD.

RESULTS

Study patients

The five patients in this study were referred to the NIH for management of EGFRvIII-positive glioblastoma (online supplemental table 1). The PBMCs used for neoantigen screening were obtained via leukapheresis, which was...
performed before the administration of anti-EGFRvIII CAR T cells.

Concurrently, FFPE tumor biopsies were obtained from the referring institutions and were subjected to WES. For Patient 1, a freshly resected tumor sample, obtained after CART cell administration, was added to the analysis. In the entire cohort, WES revealed between 71 and 115 nonsynonymous mutations per tumor (online supplemental table 1), in accordance with a previous report. These mutations, which consisted predominantly of SNVs, with a minor fraction of INDELs, were incorporated into TMG and PP screening libraries (online supplemental table 2).

**Memory T cells from Patient 1 screened positive for recognition of peptide pools representing candidate neoantigens**

To explore whether neoantigen-reactive T cells can be isolated from patients with glioblastoma, we used an IVS-based method that previously resulted in enrichment of such cells from the blood of patients with gastrointestinal malignancies. To this end, memory CD8+ and CD4+ T cells were first sorted from patients' PBMCs based on CD62L and CD45RO expression (figure 1A, online supplemental table 3). The naïve T cells were excluded from this procedure in an attempt to assess only the physiologically relevant anticancer T cell responses. Next, sorted memory T cells were stimulated with TMGs or PPs (figure 1B), with the goal to expand the reactive cells and thereby facilitate their detection. Finally, the stimulated T cells were screened for recognition of the cognate TMGs and PPs.

As indicated in figure 1C, stimulated memory CD4+ (mCD4+) T cells from Patient 1 exhibited increased IFN-γ production in response to PP1 and PP2. However, only the PP2 response could be reproduced with HPLC-purified peptides (not shown). This response was also associated with increased production of granulocyte-macrophage colony-stimulating factor (GM-CSF) and granzyme B (figure 1D). Concurrently, no significant 4-IBB upregulation was detected (not shown), suggesting that the frequency of PP2-reactive cells in the tested population was very low.

Following the IVS, memory T cells from Patients 2, 3 and 4 did not recognize any of the tumor-specific mutations (online supplemental figure 1). Cells from Patient 5 demonstrated poor expansion after the initial stimulation and thus could not be tested.

**A TCR expressed by memory CD4+ T cells from Patient 1 specifically recognized a neoantigen derived from mutated EED gene**

After the initial screen, PP2-reactive mCD4+ cells from Patient 1 were cocultured with autologous B cells pulsed with either DMSO or PP2 and then subjected to an IFN-γ capture assay. As indicated in figure 2A, 0.44% of CD4+ cells produced IFN-γ in response to PP2, while 0.03% did so in response to DMSO.

Next, captured IFN-γ+CD4+ cells were sorted and subjected to single-cell TCR sequencing. The analysis of productive TCR sequences revealed two predominant T cell clones, bearing either TCR1 or TCR2 (figure 2B). Only the TCR2 was detected exclusively within the PP2-stimulated IFN-γ+CD4+ population, although at a low frequency (3/32 cells).

In order to determine the function of these TCRs, allogeneic PBMCs were transduced with either TCR1 or TCR2 and then tested for recognition of individual peptides from PP2. As indicated in figure 2C, TCR1 did not recognize any of the candidate peptides, suggesting that the cells harboring this receptor were activated in a non-specific fashion. However, TCR2 recognized a 25-mer peptide (PITMQCICHVNGNGNAELKFP) derived from Embryonic Ectoderm Development gene that contained a histidine-to-asparagine mutation (EEDH189N). TCR-V8 deep sequencing of unstimulated PBMCs from Patient 1 confirmed that cells harboring this receptor were indeed derived from memory but not the naïve T cell subsets in the peripheral blood (online supplemental figure 2A).

Next, several analyses were performed to verify that EEDH189N truly exhibited properties of a cancer neoantigen. First, EED gene expression was confirmed in Patient 1’s freshly resected tumor sample by RNA-seq (online supplemental figure 2B). Its expression pattern could not be assessed due to the lack of simultaneous multiregion tumor sampling. However, in a WES-based clonality analysis of a single tumor biopsy, the fraction of cancer cells predicted to harbor the EEDH189N mutation was 0.99, indicating that this mutation was clonal in at least one tumor region.

Second, specificity of EEDH189N recognition was verified after coculturing TCR2-transduced T cells from two unrelated donors with Patient 1’s B cells that were pulsed with EEDH189N or its WT counterpart (PITMQCICHVNGNGNAELKFP). As indicated in figure 2D, TCR2-transduced cells from both donors recognized only the EEDH189N in a dose-dependent manner.

Third, MHC-restricted recognition of EEDH189N was confirmed by coculturing TCR2-transduced T cells with COS7 cells that were first transfected with pairs of plasmids encompassing all MHC-II molecules identified in Patient 1 and were then pulsed with WT or EEDH189N 25-mer peptides. As indicated in figure 2E, IFN-γ production was detected only when the COS7 cells were pulsed with EEDH189N in presence of both HLA-DRA1*01:01 and HLA-DRB3*02:02:01.

Finally, the ability of EEDH189N to be processed and presented on the cell surface MHC-II molecules was confirmed by coculturing TCR2-transduced T cells with COS7 transfected with plasmids encoding the HLA-DRA1*01:01 and HLA-DRB3*02:02:01, as well as the full-length WT or MUT EED genes. As indicated in figure 2F, IFN-γ production was detected only in presence of both HLA and EEDH189N genes.
DISCUSSION

In this study, we isolated memory T cells from the blood of five patients with refractory glioblastoma and, following an IVS, tested them for recognition of tumor-specific mutations. (SNVs and short INDELs). This led to the discovery of a rare mCD4+ T cell clone from Patient 1 that expressed an HLA-DRB3*02:02-restricted TCR, which specifically recognized a neoantigen derived from mutated EED gene (EEDH189N).

EED is a ubiquitously expressed protein that serves as an essential component of polycomb repressive complex 2 (PRC2), which mediates gene silencing through histone methylation. EED deletions and inactivating EED mutations, which may hinder the PRC2 function, have been previously reported in several cancers, including glioblastoma. However, the oncogenic repercussions of the EEDH189N mutation remain unknown.

Among 393 patients in the TCGA glioblastoma dataset (TCGA-GBM), EED mutations were reported in only two cases (https://www.cancer.gov/tcga). Neither of them had the EEDH189N mutation, which thus appears unique to Patient 1 in our study. This is consistent with our previous findings in other cancer types, where the vast majority of immunogenic mutations were found to be unique and not shared among the patients. T cell-mediated targeting of such mutations, which can be discovered only by performing personalized neoantigen screens, has been shown to provide durable clinical benefits in patients with select metastatic solid tumors.
The discovery of EED^{H180N}-reactive memory T cells provides evidence that the immune system can mount a natural T cell response against a glioblastoma-specific neoantigen. Due to the limited scope of our study, the role of this response in cancer immunosurveillance remains unclear, as does the mechanism that led to its induction. It may have occurred following the in situ phagocytosis of tumor cells by the specialized APCs (eg, microglia), or following the phagocytosis of tumor cells that have emigrated from the tumor site by the APCs in the deep cervical lymph nodes.

In contrast to this endogenous T cell response, previous studies have successfully elicited in vivo T cell responses by vaccinating glioblastoma patients with peptides representing selected mutations unique to their tumors. However, such responses may not represent naturally occurring immune reactions against the tumor, and it remains uncertain whether the T cells that mediate them can be successfully used for therapy. Due to its small size, this study cannot be used to accurately determine the proportion of all patients with glioblastoma who harbor neoantigen-reactive T cells in their blood. The low prevalence reported here (1/5 patients) appears lower than the prevalence (5/8 patients) reported in a study that explored the same IVS method in patients with gastrointestinal malignancies. This discrepancy could be due to strong T cell suppression that characterizes glioblastoma, but other factors could also possibly have affected the results. For instance, the use of limited tumor biopsies for WES may have prevented the capture of all potentially immunogenic mutations due to the marked intratumoral heterogeneity that hallmarks glioblastoma. Furthermore, PBMCs from Patients 3 and 4 were obtained while the patients were receiving dexamethasone, an agent known to adversely affect the T cell function in this disease.

In addition to the established therapeutic benefits of adoptively transferred CD8+ T cells directed against MHC-I neoantigens, targeting MHC-II neoantigens with CD4+ cells can also be efficacious. For instance, administration of neoantigen-reactive CD4+ T cells to a patient with metastatic cholangiocarcinoma, a malignancy characterized by largely negative or heterogeneous MHC-II expression, has led to a durable, near complete tumor destruction. However, the mechanisms that would allow such responses in cancer types without strong MHC-II expression, including the glioblastoma, remain unclear. They may involve therapy-induced APC trafficking into the tumor with subsequent antigen cross-presentation onto MHC-I molecules, or recognition of new antigens, presented by either MHC class I or II molecules (so-called “antigen spreading”), or MHC-II upregulation on the tumor cells induced by in vivo secreted IFN-γ. In support of the latter, IFN-γ was shown to upregulate MHC-II expression and presentation of an unmutated antigen in glioblastoma cell lines.

To establish the true prevalence and characteristics (eg, CD8+ vs CD4+) of neoantigen T cell responses in glioblastoma, peripheral blood of a larger number of patients could be tested using the IVS, preferably in the absence of steroid treatment. Furthermore, tumor infiltrating lymphocytes, which can be successfully grown from glioblastoma specimens obtained by surgery, could also be tested for neoantigen recognition.

In summary, we report a successful isolation of cancer neoantigen-reactive memory T cells from the blood of a patient with glioblastoma, a cancer hallmark by its immunosuppressive properties. This finding provides a rationale for further efforts to investigate neoantigen responses in these patients and to potentially develop neoantigen-directed T cell therapies for their treatment.

Acknowledgements The authors thank Sivasish Sindiri, Noam Levin and Yong Li for suggestions and technical support.

Contributors VL: conceptualized the project, designed and conducted experiments, analyzed data and wrote manuscript. GC, RY, BP, VH, DG, ZZ, JJG, TDP and SLG: provided resources and analyses for the study and edited the manuscript. PR and Y-CL: supervised research, edited the manuscript. SAR: conceptualized the project, supervised research and edited the manuscript.

Funding This study was funded by Intramural Research Program of the National Cancer Institute.

Competing interests None declared.

Patient consent for publication Not required.

Ethics approval Patients were enrolled on a protocol approved by the NCI Institutional Review Board (NCT01454586). They had provided an informed consent for all subsequent analyses of their banked blood and tumor samples, which were performed in the current study.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement All data relevant to the study are included in the article or uploaded as supplementary information.

Supplemental material This content has been supplied by the author(s). It has not been vetted by BMJ Publishing Group Limited (BMJ) and may not have been peer-reviewed. Any opinions or recommendations discussed are solely those of the author(s) and are not endorsed by BMJ. BMJ disclaims all liability and responsibility arising from any reliance placed on the content. Where the content includes any translated material, BMJ does not warrant the accuracy and reliability of the translations (including but not limited to local regulations, clinical guidelines, terminology, drug names and drug dosages), and is not responsible for any error and/or omissions arising from translation and adaptation or otherwise.

Open access This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See http://creativecommons.org/licenses/by-nc/4.0/.

ORCID iDs
VL Leko http://orcid.org/0000-0001-7863-5030
Stephanie L Goff http://orcid.org/0000-0003-3317-9804
Paul Robbins http://orcid.org/0000-0002-1260-8123

REFERENCES
32 Soos JM, Krieger JI, Stüve O, et al. Malignant glioma cells use MHC class II transactivator (CIITA) promoters III and IV to direct IFN-gamma-inducible CIITA expression and can function as nonprofessional antigen presenting cells in endocytic processing and CD4(+) T-cell activation. *JILA* 2001;36:391–405.