Detection of neoantigen-specific T cells following a personalized vaccine in a patient with glioblastoma

Tanner M. Johanns, Christopher A. Miller, Connor J. Liu, Richard J. Perrin, Diane Bender, Dale K. Kobayashi, Jian L. Campian, Michael R. Chicoine, Ralph G. Dacey, Jiayi Huang, Edward F. Fritsch, William E. Gillanders, Maxim N. Artyomov, Elaine R. Mardis, Robert D. Schreiber, and Gavin P. Dunn

ABSTRACT

Neoantigens represent promising targets for personalized cancer vaccine strategies. However, the feasibility of this approach in lower mutational burden tumors like glioblastoma (GBM) remains unknown. We have previously reported the use of an immunogenomics pipeline to identify candidate neoantigens in preclinical models of GBM. Here, we report the application of the same immunogenomics pipeline to identify candidate neoantigens and guide screening for neoantigen-specific T cell responses in a patient with GBM treated with a personalized synthetic long peptide vaccine following autologous tumor lysate DC vaccination. Following vaccination, reactivity to three HLA class I- and five HLA class II-restricted candidate neoantigens were detected by IFN-γ ELISPOT in peripheral blood. A similar pattern of reactivity was observed among isolated post-treatment tumor-infiltrating lymphocytes. Genomic analysis of pre- and post-treatment GBM reflected clonal remodeling. These data demonstrate the feasibility and translational potential of a therapeutic neoantigen-based vaccine approach in patients with primary CNS tumors.

Introduction

Cancer immunogenomics incorporates next-generation sequencing and bioinformatics to predict which tumor-specific somatic alterations may produce candidate neoantigens that could bind with high affinity to host MHC molecules and potentially be recognized by the anti-tumor immune response. Due to their tumor-restricted expression, neoantigens represent compelling targets for personalized cancer vaccines. In preclinical tumor models, polyvalent neoantigen vaccines can be therapeutically effective. Furthermore, neoantigen-based vaccines can elicit immune responses in patients, although these studies have been restricted to melanoma, which generally possesses high mutational burdens, and therefore, potentially contains a high number of candidate neoantigens. Thus, it remains unclear whether targeting neoantigens is feasible in cancers with lower mutational burdens.

We have previously reported the successful use of an immunogenomics pipeline to identify candidate neoantigens in two preclinical mouse models of glioblastoma. We used IFN-γ ELISPOT and neoantigen-specific tetramer assays to identify the presence of neoantigen-specific T cell responses in two orthotopic transplantable models. These data support the hypothesis that CNS immunosurveillance is operational and capable of inducing an endogenous neoantigen-specific T cell response against intracranial tumors. However, given that these preclinical models harbor a carcinogen-induced high mutational burden not typically seen in patients with glioblastoma, it is unclear if a similar immunogenomics-based neoantigen discovery platform will translate clinically. Notably, a recent report demonstrated the generation of neoantigen-specific T cell responses following an autologous tumor lysate-dendritic cell vaccine in patients with ovarian carcinoma, a tumor with comparable tumor mutational load as glioblastoma.

Here, we report a patient with glioblastoma (GBM) treated with a heterologous personalized vaccine, including an autologous tumor lysate-dendritic cell vaccine (DCVax) followed by a neoantigen-based synthetic long peptide vaccine (GBM.PVax). Analysis of post-treatment peripheral blood demonstrated detectable neoantigen-specific CD8+ and CD4+ T cell responses after peptide vaccination. Similar responses were observed in post-treatment tumor-infiltrating lymphocytes (TIL). Furthermore, genomic and transcriptomic characterization of the mutational

Contact

Gavin P. Dunn gpdunn@wustl.edu Department of Neurological Surgery, Washington University School of Medicine, 660 South Euclid, Box 8057, St. Louis, MO 63110, USA

Color versions of one or more of the figures in the article can be found online at www.tandfonline.com/koni.

© 2019 The Author(s). Published by Taylor & Francis Group, LLC. This is an Open Access article distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivatives License (http://creativecommons.org/licenses/by-nc-nd/4.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited, and is not altered, transformed, or built upon in any way.
landscape and tumor microenvironment pre- and post-treatment demonstrated tumor clonal evolution and evidence of potential immune evasion. Together, these results support the evaluation of immunogenomics-based vaccine strategies in glioblastoma and other tumors exhibiting moderate mutational loads.

**Results**

**Case report**

A 66-year-old previously healthy female presented with new-onset seizures, and a diagnostic brain MRI revealed a right parietal contrast-enhancing lesion. A near gross total resection was performed (>99% extent), and pathology revealed GBM with an unmethylated O6-methylguanine-DNA methyltransferase (MGMT) promoter, presence of mutated IDH1/2 promoter (C228T), and absence of IDH1/2 mutations – molecular features associated with chemotherapy resistance and decreased overall survival.13-15

Prior to surgery, the patient had consented to an autologous tumor lysate-dendritic cell vaccine study (DCVax-L) (NCT00045968) and an institutional personalized neoantigen-based peptide vaccine study (GBM.PVax) (NCT02510950). Therefore, following 6 weeks of standard adjuvant radiation therapy and concurrent temozolomide chemotherapy (CCRT), the patient received DCVax-L plus temozolomide during GBM.PVax preparation (Figure 1). DCVax-L was administered off-trial due to radiographic pseudoprogression noted following CCRT (data not shown). After cycle 4 of DCVax-L, GBM.PVax manufacturing was complete so DCVax-L was discontinued and GBM.PVax was initiated.

To design GBM.PVax, DNA whole exome sequencing of the resected tumor revealed 52 somatic, missense mutations (Figure 2(a), Supplementary File). High-affinity (IC50 <500 nM) candidate neoantigens were identified using the neoantigen discovery pipeline, pVAC-Seq, as well as the NetMHCIIpan 3.2 and NetMHCII 2.3 algorithms,10,16 revealing 2 HLA class I-restricted candidates, 32 HLA class II-restricted candidates, and 14 candidates with high affinity to both HLA class I and II alleles (Figure 2(a)). Eight synthetic long peptides (SLPs) encompassing seven neoantigens were successfully synthesized for inclusion into GBM.PVax (Figure 2(b)).

After cycle 1 of GBM.PVax, the patient developed confusion, and a brain MRI revealed increased T2/FLAIR indicative of edema/inflammation along with increased size of the T1 contrast-enhancing lesion most consistent with pseudoprogression (Figure 2(c)). Symptoms resolved completely with a short steroid taper, and a repeat brain MRI 4 weeks later showed a reduction in T2/FLAIR with stabilization of the T1 contrast-enhancing lesion (Figure 2(c)) coinciding with a significantly improved functional status. The patient continued to do well clinically through cycles 3 and 4 of GBM.PVax and noted interval development of axillary lymphadenopathy. Unfortunately, the subsequent brain MRI was concerning for disease progression (Figure 2(c)), and a repeat craniotomy with subtotal resection was performed. Histopathology demonstrated extensive treatment effect with only focal areas of residual glioma, and no overt tumor recurrence indicating the changes on the MRI was again due to pseudoprogression (Figure 2(d)). The post-surgical course was complicated by a saddle pulmonary embolus treated with embolectomy; intraparenchymal hemorrhage plus gastrointestinal bleeding from antiocoagulation therapy resulting in a declining functional status that precluded additional GBM-related therapy. The patient passed away 21 months after initial diagnosis (Figure 1).

**Immunogenicity of personalized vaccine**

To assess for the presence of neoantigen-specific T cells post-vaccination, we first evaluated the immunogenicity of the known peptides contained within GBM.PVax as the antigens present in DCVax-L were undefined. First, reactivity to the predicted immunodominant minimal epitopes of the GBM.PVax peptides (Table 1) was determined in PBMC obtained 4 months after initiation of GBM.PVax using IFN-γ ELISPOT. Isolated post-vaccination CD8+ T cells reacted to the HLA-B*44:02-restricted neoantigen, GPR133,G126E (mGPR133), while CD4+ T cells were reactive to both IL22,V72I (mIL22) and PTEN,G157S (mPTEN) (Figure 3(a)). An expanded screen of alternative mutation-containing minimal epitopes did not reveal additional reactivity to any of the GBM.PVax-containing neoantigens (data not shown). Reactivity to mGPR133, mIL22, and mPTEN appeared to be mutation-specific as no T cell reactivity was observed to the corresponding wild-type epitopes (Figure 3(b)). In order to determine whether these neoantigen-specific responses were induced or augmented by vaccination rather than as a result of prior therapies, reactivity was also measured in a PBMC sample obtained immediately prior to GBM.PVax but after the completion of DCVax-L administration. A pre-existing CD8+ T cell response to mGPR133 was present above background prior to GBM.PVax but increased after peptide vaccination (Figure 3(c)). As a control for normal variation over time, CD8+ T cell reactivity to HLA-A*24:02 and B*44:02-restricted CMV antigens was observed prior to GBM.PVax and was noted to decrease between

---

**Figure 1.** Schematic representation of treatment course. Note: cycle 3 of temozolomide was delayed due to thrombocytopenia, and cycle 4 was given at a reduced dose (100 mg/m2 down from 150 mg/m2) due to intolerance. Abbreviations: STR = subtotal resection; CCRT = concurrent chemoradiation therapy; TMZ = temozolomide; DCVax = autologous tumor lysate-dendritic cell vaccine.
the pre- and post-GBM.PVax time points (Figure 3(c)), suggesting that mGPR133 reactivity was likely boosted as a result of GBM.PVax immunization. Neither mIL22-specific nor mPTEN-specific CD4⁺ T cell responses were detectable above background prior to GBM.PVax initiation, suggesting both were increased to detectable levels after immunization with GBM.PVax (Figure 3(c)).

Because immunity to neoantigens could also develop secondary to prior therapies (i.e., surgery, radiation, temozolomide and/or DCVax-L) or epitope spreading following peptide vaccination, we evaluated reactivity to predicted high-affinity HLA class I and/or class II candidates. Top candidates were selected for peptide synthesis as long peptides (SLPs). Soluble SLPs are incorporated into GBM.PVax. (b) Table listing the 8 SLPs encompassing 7 neoantigens included in GBM.PVax. The location of the mutated amino acid is enlarged and bolded. (c) Representative brain MRI axial images at indicated time points during treatment with GBM.PVax. (d) Representative H&E sections from initial tumor resection (day 0) and post-treatment tumor resection (day 347). Black line denotes 200 microns.

Presence of neoantigen-specific TIL following treatment
Next, the presence of neoantigen-specific responses in TIL isolated from the second surgical specimen was assessed. Consistent with PBMC, reactivity to mGPR133, mWDR63, mIL22, mPTEN, and mNVL was observed above background among CD8⁺ or CD4⁺ TIL, however, only responses to
mGPR133, mIL22, and mNVL were significant (Figure 4(a)). Spatially, immunohistochemistry demonstrated that T cells were predominantly restricted to the perivascular space, though infiltration into the pauci-cellular intraparenchymal region was also observed (Figure 4(b)).

### Treatment-related clonal evolution following treatment

To explore the effect of treatment on tumor evolution, the mutational landscape and clonal architecture of the pre- and post-treatment tumor specimens were characterized by whole exome DNA sequencing. SciClone analysis revealed loss of two pre-treatment subclonal populations (clusters 2 and 4) but emergence of two additional sub-populations (clusters 3 and 5) post-treatment (Figure 4(c)). T cell reactivity was observed to neoantigens both private to the lost pre-treatment subclones (mGPR133, mMEGF8, and mNUP107) and shared within the founder clone (mWDR63, mPDIA3, mIL22, mPTEN, and mNVL) (Figure 4(c)). Although transcriptional repression of neoantigens has been suggested to facilitate immune escape, no changes were observed in transcriptional expression levels among the founder clone neoantigens between pre- and post-treatment specimens (Supplementary File). Furthermore, there was no observed

### Table 1. Table of minimal epitope sequences of HLA class I and II-restricted GBM.PVax and non-GBM.PVax neoantigens screened by IFN-γ ELISPOT for T cell reactivity including corresponding predicted binding affinity (IC₅₀). The location of the mutated amino acid is enlarged and underlined.

| Neoantigen | Core sequence | IC₅₀ (nM) | GBM.PVax |
|------------|---------------|----------|----------|
| KIAA1958 R115W | HLA-A*01:01 | WTRDSCDFSY | 368 | Yes |
| ZMYND12 R192H | HLA-A*24:02 | HLYHDLNDY | 171 | No |
| GPR133G126E | HLA-B*44:02 | YEYQVSGNF | 260 | Yes |
| VP513C I1804T | HLA-C*05:01 | VTDKMNIEL | 55 | Yes |
| PTEN R475 | HLA-C*06:02 | YSNNDIVDV | 112 | Yes |
| IL22 V72I | HLA-DRB*01:01 | LADNNMTDR | 138 | Yes |
| DLGAP4 G55 | HLA-DRB*01:01 | LSDSRPRHL | 239 | Yes |
| PDA3 T219S | HLA-DRB*01:01 | FEDKSVAYT | 380 | No |
| MEGF8 P238S | HLA-DRB*01:01 | LSSPGGLAV | 383 | No |
| WDR3 T690M | HLA-DRB*01:01 | FYNDIILMV | 109 | No |
| CHD2 N123T | HLA-DRB*01:01 | SRQEPRSFT | 230 | Yes |
| MEGF8 P238S | HLA-DRB*03:01 | AGAFLSPGALPFV | 5 | No |
| SIRPB1 T18M | HLA-DRB1:01 | LLMMLLLGTV | 8 | No |
| NVL I644V | HLA-DRB1:01 | GLNVSVKGPEL | 9 | No |
| NUP107 R555H | HLA-DRB1:01 | ULEFHTLGQK | 10 | No |
| IL22 V72I | HLA-DRB1:01 | NNTDIRLIGEK | 10 | No |
| PTEN R475 | HLA-DRB1:01 | GPAERLEGVSN | 192 | Yes |
| VP513C I1804T | HLA-DRB1:01 | PMHEYLSPVDT | 196 | Yes |
| GPR133 G126E | HLA-DRB1:01 | PSAYEGQVSG | 282 | Yes |
| CHD2 N123T | HLA-DRB1:01 | SMRSQEPSRFT | 325 | Yes |

**Figure 3.** Immunogenicity of GBM.PVax in peripheral blood. (a-d) Bar graphs from IFN-γ ELISPOT data of positively selected CD8⁺ (left column) and CD4⁺ (right column) PBMC stimulated with indicated peptide. Media = negative control (no peptide). (a) Reactivity screen to GBM.PVax neoantigens. (b) Comparison of reactivity between mutated and wild-type GBM.PVax neoantigens that demonstrated positive reactivity in Figure 3(a). (c) Comparison of reactivity between pre- and post-GBM.PVax PBMC. (d) Reactivity screen of non-GBM.PVax neoantigens from Table 1. A, b, and d represent a minimum of two independent experiments done in duplicate. (c) represents a single experiment done in duplicate due to limited sample. (*) represents p < 0.05 compared to negative control unless otherwise indicated.
transcriptional repression or loss of heterozygosity of HLA loci (Supplementary File and data not shown).

In the post-treatment tumor, 7 of the 25 private mutations were predicted to generate at least one high-affinity HLA class I-restricted neoantigen, a similar frequency (28%) to the shared founder clone (33.3%) and private pre-treatment subclones (23%) (Figure 4(d)) suggesting equivalent somatic mutation and neoantigen rates within the sub-populations. However, while RNA expression was detected for 2 of 3 and 8 of 13 neoantigens in the private pre-treatment subclones and founder clone, respectively, no private post-treatment neoantigen RNA expression was detectable (Figure 4(d) and Supplementary File), suggestive of possible immunoediting.

**Immunologic evolution during treatment**

To characterize the effects of treatment on the broader immune microenvironment, we compared pre- and post-treatment transcriptomes. Analyses revealed increased relative expression levels for genes associated with both T cell (CD4, CD8A) subsets (Figure 5(a)). An increase in cytotoxic CD8 T cell effector function genes (PRF1, GZMA, GZMB) was also noted to be upregulated (Figure 5(a)). Among the CD4 T helper lineage, increased expression of the Treg, Th2 and Th17 lineage-defining transcription factors FOXP3, GATA3, and RORC, respectively, were upregulated post-treatment, while the Th1 master regulator transcription factor, T-bet (TBX21), was downregulated (Figure 5(a)). Consistent with the decreased expression of TBX21 and increased expression of FOXP3, IFNG was not found to be expressed in the post-treatment specimen (data not shown), while IL10 was noted to be upregulated (Figure 5(a)). Among the Th17 effector cytokine, IL-17, was upregulated (IL17B and IL17D) though other isoforms (IL17A, IL17C, IL17E, IL17F) were not detected (Figure 5(a) and Supplementary File and data not shown).

**Figure 4.** Clonal evolution of neoantigens after treatment. (a) Bar graphs from IFN-γ ELISPOT data of positively selected CD8+ (left column) and CD4+ (right column) ex vivo expanded TIL isolated from the post-GBM.PVax specimen and stimulated with indicated peptide. Media = negative control (no peptide). Graphs represent a single experiment done in duplicate due to limited sample. (*) represents p < 0.05 compared to negative control. (b) Representative immunohistochemistry sections of anti-CD3 and anti-CD8 staining in the post-treatment resection specimen. Black arrow identifies perivascular lymphocytes. Black line denotes 200 microns. (c) SciClone plot of clonal and subclonal populations present in pre- and post-treatment tumor specimens. VAF = variant allele fraction. Cluster 1 = founder clone mutations. Bolded neoantigens represent those with observed reactivity by ELISPOT. (d) Venn diagrams of missense mutations (left) and predicted high-affinity HLA class I neoantigens (right) from the pre- and post-treatment tumor specimens. Numbers indicate respective private or shared missense mutations or neoantigens. Parentheses represent the number of neoantigens with confirmed expression by RNA-seq.
Expression of the Th2 effector cytokine, IL-4 (IL4), was also not detected in either the pre- or post-treatment specimens (data not shown). Together, these data demonstrate that treatment with radiation, chemotherapy, and vaccination resulted in an overall increase in the relative gene expression profile of cytotoxic CD8 T cells and non-Th1 CD4 T cell subsets, including regulatory T cells, within the tumor microenvironment.

A similar mixed response was seen among the myeloid compartment following treatment with a noted increase in relative expression levels of both M1 (CD68, STAT1) and M2 (CD163, STAT3) macrophages/microglia compared to pre-treatment. However, there appeared to be an overall skewing toward a predominant M2 environment with the noted upregulation of M2-associated genes CSF1R, CD206, ARG1, and TGFB1, whereas M1-associated genes NOS2 and IL12A were relatively downregulated. Consistently, a similar increase of M2 macrophages has been reported in GBM patients treated with DCVax following standard chemoradiation.

Next, we compared expression patterns of select co-stimulatory and co-inhibitory immune checkpoint molecules. Genes associated with the co-stimulatory molecules OX40, 4–1BB, CD27, and CD70, as well as the co-inhibitory molecules PD-1, PD-L1, TIM-3, GITR, VISTA, TIGIT, and LAG3 were broadly upregulated (Figure 5(c)). Of note, other clinically actionable immune checkpoint targets (i.e. CTLA4, IDO1, ICOS, and PDL2) were unchanged or downregulated after treatment (Figure 5(c)) suggesting there was not a global upregulation of these molecules between specimens.

To provide an integrated overview of treatment-related changes in the immune landscape, we applied a previously described gene set enrichment analysis (GSEA) approach in which non-overlapping metagene sets corresponding to functional immune modules – antigen processing/presentation (MHC), checkpoints (CP), effector cells (EC), and suppressor cells (SC) – were normalized to 166 TCGA GBM samples. We then compared GSEA analysis of transcriptomic data before and after treatment to this reference dataset (Figure 6). This
analysis demonstrated that all four modules were comparatively downregulated in the pre-treatment sample compared to the averaged TCGA GBM samples. In contrast, after treatment, all four modules exhibited increases over the averaged TCGA GBM z-scores, supporting the observation that both pro-inflammatory and immunoregulatory mechanisms were upregulated following treatment with radiation, chemotherapy, DCVax-L, and GBM.PVax.

Discussion

Here, we describe the successful use of an immunogenomics pipeline to identify candidate neoantigens for incorporation into a personalized vaccine to treat GBM. Furthermore, the same immunogenomics pipeline was used to guide screening for neoantigen-specific T cell reactivity that may have been generated endogenously, as a result of prior treatment (surgery, radiation, chemotherapy or DCVax-L), or through epitope spreading. In total, we were able to detect reactivity to three HLA class I- and five HLA class II-restricted neoantigens from PBMC obtained post-treatment. Moreover, a similar pattern of reactivity was seen in post-treatment TIL, suggesting that systemically generated responses are capable of homing to and infiltrating CNS tumors. Lack of adequate pre-vaccination samples restricted our analysis regarding the kinetics of these responses, but we were able to demonstrate an augmentation of neoantigen-specific T cell responses following vaccination. Regardless, these results support the rationale for targeting neoantigens even in lower mutational burden tumors like GBM, where the number of potentially immunogenic candidates is perhaps more limited than higher mutational tumors, such as melanoma. These results are further supported by the recent observation that an autologous tumor lysate-dendritic cell vaccine was shown to induce neoantigen-specific responses in ovarian cancer, another lower mutational burden tumor.¹¹,¹²

These results combined with previously published reports²,⁶,⁷ support the use of an immunogenomics pipeline to identify candidate neoantigens for personalized vaccines across multiple tumor types. It will now be necessary to determine the therapeutic value of neoantigen vaccine on improving clinical outcomes in patients, particularly in the setting of the recent phase 3 study showing no difference in overall survival with the addition of rindopepimut, a peptide vaccine targeting the neoantigen, EGFRvIII, to standard of care therapy for newly diagnosed GBM.²² To this end, ongoing work by our group and others (NCT02149225, NCT02287428, NCT03223103, NCT03412877) is directed at evaluating the clinical efficacy of neoantigen-based vaccines in GBM.

Figure 6. Modified immunophenoscore diagrams derived from Charoentong et al.²¹ for pre-treatment (top) and post-treatment (bottom) specimens. The gene or gene set comprising each point around the circle is labeled and grouped in corresponding quadrants. Red to blue color scale represent averaged z-score generated from 166 TCGA GBM samples. Grayscale represents weighted z-score associated with the composite score of each quadrant corresponding to the genes or gene sets within the quadrant. MHC = Major histocompatibility complex-associated and antigen presentation genes; CP = checkpoint molecules; EC = effector cells; SC = suppressor cells.
Although the Rindopepimut vaccine targeting the juncional sequence created by the EGFRvIII variant also represented a neoantigen targeting approach, there are several differences between this approach and the cancer immunogenomics approach in this report. First, the EGFRvIII target is heterogeneously expressed, which could limit the efficacy of a vaccine directed at a single target due to therapy-induced immunoediting. However, by using a multi-valent vaccine platform such as the one described here, one can incorporate clonal neoantigens as well as neoantigens that may be expressed by different subclones to enhance the potential of targeting a greater proportion of tumor cells. Second, while rindopepimut efficiently generated robust EGFRvIII-specific humoral responses, it has not been shown to induce a cytotoxic T cell response. Thus, a vaccine approach that can generate neoantigen-specific cell-mediated immune responses may be more effective. Third, patients were not selected for rindopepimut vaccine by MHC haplotype. Although predictions of neoantigen binding to HLA molecules are imperfect, the consideration of target affinity may increase the likelihood of effective T cell priming and target recognition in patients following vaccine treatment.

It is likely that combination immunotherapies will be necessary to increase therapeutic efficacy. Specifically, many personalized neoantigen vaccine approaches are combining tailored vaccines with immune modulating agents, such as checkpoint blockade therapy. Since neither vaccines nor immune checkpoint blockade therapy have been shown to improve survival of patients with GBM as monotherapies,22-24 the combination of several complementary immune-based strategies may potentially be more effective. For example, the addition of a vaccine may improve checkpoint blockade therapy by inducing or increasing the number of tumor-specific T cells present within the tumor microenvironment.8,17,18 Conversely, checkpoint blockade therapy may improve the effector function of vaccine-induced T cells once they home to the immunosuppressive tumor microenvironment.25,26 Thus, while immunotherapies have yet to translate into improved survival for patients with GBM, a combinatorial immune-based approach may lead to better outcomes in the future.

It will also be important to determine the optimal methods of vaccination to stimulate the most robust and functional neoantigen-specific responses to the largest number of candidates per patient. For example, in this case, a heterologous prime-boost personalized vaccine approach of DCVax-L followed by synthetic long peptides was administered. Additional vaccine platforms – including nucleic acids and viral/bacterial vectors – warrant study. Moreover, identifying the ideal adjuvant including TLR agonists, immune checkpoint agonists/antagonists, or cytokines is an additional variable that has yet to be fully addressed.

Although we recognize the limitations from a single patient study, further consideration is needed with regard to monitoring the anti-tumor effects of vaccines in the CNS. Specifically, it is challenging to determine the efficacy of immune-based treatments in a tissue site, in this case, the CNS, in which it is more difficult to sample tissue iteratively to understand in situ immune activity. As a proxy for immune activity at the tumor site, we identified neoantigen-specific T cell responses in the blood, and we were able to culture neoantigen-specific TIL from the redo surgery sample. However, we did not observe a robust T cell infiltrate at the time of the patient’s second surgery, a finding that was supported by our gene expression analysis. These findings raise the possibility that further study is needed to understand the kinetics of T cell homing to the CNS following vaccination. Additionally, these data underscore the influence of the large myeloid compartment within the GBM microenvironment. Further study is necessary to understand how to effectively enhance neoantigen-specific T cell activity, the effects of vaccines on the microenvironment, and how these parameters change with combination treatment.

The presence of pseudoprogression on the post-vaccination MRI represents another important consideration for vaccine trials in GBM as well as other novel immunotherapeutic strategies targeting CNS malignancies such as CAR T cells.27 Radiographically assessing treatment response may be difficult as there are currently no criteria that can accurately and reliably delineate immune reactivity or inflammation from true tumor progression. This has begun to be addressed in the setting of immune checkpoint blockade therapy and brain metastases with the development of the Immunotherapy Response Assessment in Neuro-Oncology (iRANO).28 However, in order to spare patients unnecessary surgical procedures or changes in otherwise effective therapies, novel imaging modalities or adjunct non-invasive monitoring assays will be needed.

Lastly, further work is needed to understand the emergence of resistant subclones post-treatment despite the presence of antigen-specific T cells. Neoantigenic and molecular profiling of the post-treatment tumor and microenvironment, respectively, may provide insight into potential immune evasion mechanisms. Specifically, the lack of expressed immunogenic neoantigens private to the resistant subclones suggests escape of active immunosurveillance. Conversely, while not observed in this case, it is plausible to postulate that treatment-induced changes from radiation and alkylating chemotherapies may lead to the development of new candidate neoantigens that could be targeted iteratively. Furthermore, the concomitant upregulation of immunoregulatory factors, such as inhibitory checkpoint molecules and suppressive immune cell subsets, supports the rationale for combining neoantigen-based vaccines with additional immunomodulatory strategies such as immune checkpoint blockade and microenvironment modulators5,7 to augment the potency of these individual strategies.

In summary, the results presented herein provide proof-of-principle evidence supporting the feasibility of an immunogenomics-guided neoantigen discovery platform to design personalized vaccines in both CNS and lower mutational burden tumors.

Materials and methods

Informed consent was signed prior to enrollment. All procedures and experiments were performed in accordance with the Declaration of Helsinki and an Institutional Review Board-
approved protocol at the Washington University School of Medicine. This study is NCT02510950 on clinicaltrials.gov.

**Whole exome DNA and RNA sequencing**

Library construction, sequencing, and variant detection were performed as described.7 Briefly, normal and tumor exome libraries were captured using the Nimblegen VCRome kit and cDNA was prepared with the TruSeq Stranded Total RNA kit. Each library was sequenced on an Illumina HiSeq2500. Somatic variants were detected using an ensemble approach incorporating statistical and heuristic filters. RNA-sequencing (RNA-seq) data were processed with Tophat v2.0.8 and Cufflinks v2.1.1 to generate gene expression values.29

**HLA class I and II neoantigen predictions, prioritization, and selection**

Clinical HLA typing was performed. HLA alleles, annotated somatic variants, and FPKM-based variant expression values were used as pVAC-Seq inputs (www.pvactools.org),16 which queries the IEDB algorithm collection to predict HLA class I neoantigen binding. Averaged binding affinities of NetMHCIIpan 3.2 and NetMHCII 2.3 were used for predicting HLA class II antigens. Candidate neoantigens for GBM.PVax were based on multiple factors including predicted HLA class I and/or II binding affinity, RNA-seq expression level, clonality, predicted peptide solubility, and targeted HLA allele diversification.

**Long peptide synthesis and GBM.PVax administration**

Fifteen neoantigens were selected for synthetic long peptide production. Ultimately, eight peptides targeting seven candidate neoantigens were soluble and incorporated into GBM.PVax (Figure 1(b)). GMP grade peptides were produced by CS Bio (Menlo Park, CA). Three peptide pools containing 2–3 peptides/pool were dissolved in DMSO and diluted to 1 mg/peptide in D5W containing succinate. For vaccination, each peptide pool was co-administered subcutaneously with 1.5 mg poly-ICLC (Hiltonol; Oncovir, Washington, DC) into one of four rotating injection sites (right arm, left arm, right groin, left groin). Vaccinations were given on days 1, 3, 5, 8, 15, and 22 (±1 day) of cycle one and then on day 22 of every subsequent 28-day cycle (Figure 1(a)).

**Isolation of PBMC and Ex Vivo Expansion of TIL**

Heparinized blood was collected by venipuncture prior to GBM.PVax initiation and by leukapheresis after GBM.PVax cycle 4. Peripheral blood mononuclear cells (PBMC) were isolated by Ficoll-Paque (GE Healthcare Life Sciences, Pittsburgh, PA) density gradient centrifugation and cryopreserved in 10% DMSO, 20% FBS, and culture media (RPMI-1640, L-glutamine, penicillin/streptomycin, β-mercaptoethanol, sodium bicarbonate, sodium pyruvate, 10% heat-inactivated FBS).

TIL were isolated by mincing the resected specimen into 1–2 mm chunks and incubating at 37°C in culture media with 2500 U/mL recombinant human IL-2 (EMED Medical Company, Maryland Heights, MO). Media was changed every 3–5 days. After 3 weeks, lymphocytes were harvested by Percoll (GE Healthcare Life Sciences) density gradient centrifugation and cryopreserved. Cells were stored in liquid nitrogen.

**IFN-γ ELISPOT assay**

CD8+ and CD4+ cells were isolated from PBMC and TIL using sequential magnetic bead-based positive selection kits (STEMCELL Technologies, Vancouver, Canada). Approximately 100,000 double-negative autologous PBMC were incubated on pre-coated human IFN-γ ELISPOT plates (Cellular Technology, Ltd., Shaker Heights, OH) with 10 μM of indicated peptide (GenScript, Piscataway, NJ) plus ~400,000 CD8+ or CD4+ PBMC, or ~20,000 CD8+ or CD4+ TIL for 18–20 hours at 37°C. Plates were analyzed using the C.T.L. ImmunoSpot kit (Cellular Technology, Ltd., Shaker Heights, OH).

**Analysis of clonal evolution and transcriptome**

Clonal evolution was analyzed as described previously.29 Briefly, the SciClone algorithm was used to infer clonal hierarchy,30 and clonEvol was used to reconstruct tumor phylogeny.31 For gene set enrichment analysis, gene-specific expression levels from pre- and post-treatment specimens were z-score normalized across 166 TCGA-GBM patients as described previously.21 Immunophenogram figures were generated using the publicly available R code on GitHub (https://github.com/mui-icbi/Immunophenogram).

**Statistical analysis**

Intergroup differences in mean number of spots on ELISPOT were evaluated using a Student’s t-test with $p < 0.05$ as statistically significant (GraphPad Prism, San Diego, CA).

**Disclosure of Potential Conflicts of Interest**

R.D. Schreiber is a co-founder and has ownership interests in Neon Therapeutics and Joune Therapeutics, and is a consultant/advisory board member for BioLegend, Constellation, Merix, NGM Biopharmaceuticals, and Lytix. G.P. Dunn is a co-founder of Immunovuvalent Therapeutics. E.F. Fritsch is a co-founder and an employee of Neon Therapeutics. E.R. Mardis is a member of the scientific advisory boards of PACT Pharma and Moderna LLC. All other authors have no potential conflicts of interest to disclose.

**Funding**

This study was funded by National Institutes of Health grant K08NS092912 (G.P.D.), Cancer Research Institute (G.P.D.), Damon Runyon Cancer Research Foundation (G.P.D.), and the Physician-Scientist Training Program at Washington University School of Medicine (T.M.J.) and NIH grant R01CA190700 (R.D.S.).

**References**

1. Schumacher TN, Schreiber RD. Neoantigens in cancer immunotherapy. Science. 2015;354:689–74. doi:10.1126/science.aaa4971.

2. Carreno BM, Magrini V, Becker-Hapak M, Kaabinejadian S, Hundal J, Petti AA, Ly A, Lie W-R, Hildebrand WH, Mardis ER, et al. Cancer immunotherapy. A dendritic cell vaccine increases the breadth and diversity of melanoma neoantigen-specific T cells. Science. 2015;348:803–808. doi:10.1126/science.aaa3828.

3. Gubin MM, Artyomov MN, Mardis ER, Schreiber RD. Tumor neoantigens: building a framework for personalized cancer immunotherapy. J Clin Invest. 2015;125:3413–3421. doi:10.1172/JCI80008.

4. Johanns TM, Bowman-Kirigin JA, Liu C, Dunn GP. Targeting neoantigens in glioblastoma: an overview of cancer
immunogenomics and translational implications. Neurosurgery. 2017;64:165–176. doi:10.1093/neuros/nyx321.

5. Johanns TM, Dunn GP. Applied cancer immunogenomics: leveraging neoantigen discovery in glioblastoma. Cancer J. 2017;23:125–130. doi:10.1097/PPO.0000000000000277.

6. Ott PA, Hu Z, Kwon DS, Shukla SA, Sun J, Bozym DJ, Zhang W, Luoma A, Giobbie-Hurder A, Peter L, et al. An immunogenic personal neoantigen vaccine for patients with melanoma. Nature. 2017;547:217–221. doi:10.1038/nature22991.

7. Sahin U, Derhovanessian E, Miller M, Bp K, Simon P, Lower M, Bukur V, Tadmor AD, Luxumburger U, Schrörs B, et al. Personalized RNA mutanome vaccines mobilize poly-specific therapeutic immunity against cancer. Nature. 2017;547:222–226. doi:10.1038/nature23003.

8. Gubin MM, Zhang X, Schuster H, Carreno BM, Petti AA, Linette GP, Griffith OL, et al. Predicting immunogenic tumour mutations by combining mass spectrometry and exome sequencing. Nature. 2014;515:572–576. doi:10.1038/nature14001.

9. Johanns TM, Ward JP, Miller CA, Wilson C, Kobayashi DK, Baumgartner P, Stevenson B.J., Iseli C, Dangaj D, et al. Personalized cancer vaccine effectively mobilizes antitumor T cell immunity in ovarian cancer. Sci Transl Med. 2018;10(436):eaao5931. doi:10.1126/scitranslmed.aao5931.

10. Bobisse S, Genolet R, Roberti A, Tanyi JL, Racle J, Stevenson BJ, et al. Sensitive and frequent identification of high avidity neo-epitope specific CD8+ T cells in immunotherapy-naive ovarian cancer. Nat Commun. 2018;9:1092. doi:10.1038/s41467-018-03301-0.

11. Eckel-Passow JE, Lachance DH, Molinaro AM, Walsh KM, Decker PA, Sicotte H, Pekmezci M, Rice T, Kosel ML, Smirnov IV, et al. Glioma groups based on 1p/19q, IDH, and TERT promoter mutations in tumors. N Engl J Med. 2015;372:2499–2508. doi:10.1056/NEJMoa1407279.

12. Bobisse S, Genolet R, Roberti A, Tanyi JL, Racle J, Stevenson BJ, Iseli C, Michel A, Le BitoüX M-A, Guillaume P, et al. Sensitive and frequent identification of high avidity neo-epitope specific CD8+ T cells in immunotherapy-naive ovarian cancer. Nat Commun. 2018;9:1092. doi:10.1038/s41467-018-03301-0.

13. Stupp R, Mason WP, Cloughesy T, Voloschin A, Ramkissoon SH, Ligon KL, Lates K, et al. Nivolumab with or without ipilimumab in patients with recurrent glioblastoma: results from exploratory phase I cohorts of CheckMate 143. Neuro Oncol. 2018;20:674–686. doi:10.1093/neuonc/nox208.

14. Lopes A, Vanvarenberg K, Kos S, Lucas S, Colau D, Van den Eynde B, Prät V, Vandermeulen G. Combination of immune checkpoint blockade with DNA cancer vaccine induces potent antitumor immunity against P815 mastocytosis. Sci Transl Med. 2018;10:15732. doi:10.1126/scitranslmed.aap3933-7.

15. Kinked HL, Hopkins A, Lutz E, Wu A, Zhao W, Van den Eynde B, Prat V, Vandermeulen G. Combining STING-based neoantigen-targeted vaccine with checkpoint modulators enhances antitumor immunity in murine pancreatic cancer. JCI Insight. 2018;3(20):122857. doi:10.1172/jci.insight.122857.

16. O’Rourke DM, Nasrallah MP, Desai A, Melenhorst JJ, Mansfield K, Morrisette JJD, Martinez-Lage M, Brem S, Maloney E, Shen A, et al. A single dose of peripherally infused EGFRVIII-directed CAR T cells mediates antigen loss and induces adaptive resistance in patients with recurrent glioblastoma. Sci Transl Med. 2017;9(399):eaao9884. doi:10.1126/scitranslmed.aao9884.

17. Okada H, Weller M, Huang R, Finocchiaro G, Gilbert MR, Witzik W, Ellingson BM, Hashimoto N, Polack IF, Brandes AA, et al. Immunotherapy response assessment in neuro-ontology: a report of the RANO working group. Lancet Oncol 2015;16: e534–e42. doi:10.1016/S1470-2045(15)00088-1.

18. Johanns TM, Miller CA, Dorward IG, Tissen C, Chang E, Perry A, Upalurali R, Ferguson C, Schmidt RE, Dahiya S, et al. Immunogenomics of hypermutated glioblastoma: a patient with germline POLE deficiency treated with checkpoint blockade. Cancer Discov. 2018;6:1230–1236. doi:10.1158/2326-4482.CD-16-0575.

19. Miller CA, White BS, Dees ND, Griffith M, Welsh JS, Griffith OL, Vij R, Tomasson MH, Graubert TA, Walter MJ, et al. SciClone: inferring clonal architecture and tracking the spatial and temporal patterns of tumor evolution. PLoS Comput Biol. 2014;10: e1003665. doi:10.1371/journal.pcbi.1003665.

20. Ding L, T L, De L, Ca M, Dc K, Is W, Ritchey JK, Young MA, Lamprecht T, McLellan MD, et al. Clonal evolution in relapsed acute myeloid leukaemia revealed by whole-genome sequencing. Nature. 2012;481:506–510. doi:10.1038/nature10738.