

Endogenous Neoantigen-Specific CD8 T Cells Identified in Two Glioblastoma Models Using a Cancer Immunogenomics Approach

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Abstract

The "cancer immunogenomics" paradigm has facilitated the search for tumor-specific antigens over the last 4 years by applying comprehensive cancer genomics to tumor antigen discovery. We applied this methodology to identify tumor-specific "neoantigens" in the C57BL/6-derived GL261 and VM/Dk-derived SMA-560 tumor models. Following DNA whole-exome and RNA sequencing, high-affinity candidate neopeptides were predicted and screened for immunogenicity by ELISPOT and tetramer analyses. GL261 and SMA-560 harbored 4,932 and 2,171 non-synonymous exome mutations, respectively, of which less than half were expressed. To establish the immunogenicities of H-2K^b and H-2D^b candidate neoantigens, we assessed the ability of the epitopes predicted *in silico* to be the highest affinity binders to activate tumor-infiltrating T cells harvested from GL261 and

SMA-560 tumors. Using IFN γ ELISPOT, we confirmed H-2D^b-restricted Imp3_{D81N} (GL261) and Odc1_{Q129L} (SMA-560) along with H-2K^b-restricted E2f8_{K272R} (SMA-560) as endogenous tumor-specific neoantigens that are functionally immunogenic. Furthermore, neoantigen-specific T cells to Imp3_{D81N} and Odc1_{Q129L} were detected within intracranial tumors as well as cervical draining lymph nodes by tetramer analysis. By establishing the immunogenicities of predicted high-affinity neopeptides in these models, we extend the immunogenomics-based neoantigen discovery pipeline to glioblastoma models and provide a tractable system to further study the mechanism of action of T cell-activating immunotherapeutic approaches in preclinical models of glioblastoma. *Cancer Immunol Res*; 4(12); 1007–15. ©2016 AACR.

Introduction

Glioblastoma is the most common and lethal malignancy of the central nervous system (CNS) in adults. Despite multimodality standard-of-care treatment involving surgery followed by concurrent chemoradiation, patients will eventually relapse or progress with a median overall survival of 12–15 months (1). The unmet need for new treatments has been a strong stimulus to better understand the molecular basis of

glioblastoma in order to identify new therapeutic targets. However, although we now have deep insights into the genomic landscapes of these tumors (2, 3), the "mutation-to-targeted drug" paradigm based on genomically guided precision medicine has not emerged as an effective treatment option as it has for other solid tumors (4). Due to the successes of immunotherapies in other cancer types (5), combined with the realization that the CNS is not an hermetically immunoprivileged site (6), enthusiasm is growing regarding the use of immune-based treatments for patients with glioblastoma. Indeed, vaccine approaches targeting EGFRvIII (7) and other shared epitopes (8), autologous dendritic cell (9) and heat shock protein (10) vaccines, and other modalities including the checkpoint blockade agents are in clinical trials for glioblastoma. However, our understanding of how these immune-based strategies, designed to enhance tumor-specific T-cell recognition and effector function, control glioblastoma progression has been limited by our ability to identify and monitor tumor-specific T-cell responses.

To this end, the "cancer immunogenomics" concept has greatly enabled the search for tumor mutation-specific antigens over the last 4 years by applying cancer genomics information in a new way (11–13). The aim of this approach is to identify expressed tumor-specific, missense mutations that are predicted to bind with high affinity to an individual patient's MHC molecules for presentation as "neoantigens" to host effector T cells. Thus, this view prioritizes the antigenic potential of a somatic mutation in cancer cells over the more traditional "driver" versus "passenger" hierarchy. Since

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its initial conception and application to neoantigen discovery in murine sarcomas (13, 14), this approach has been employed to identify antigenic targets of tumor-specific T cells arising spontaneously or as the result of checkpoint blockade in other preclinical (15–17) and human (18–21) settings.

To better understand the endogenous T-cell immune response to brain tumor antigens, we applied the cancer immunogenomics methodology to the study of two mouse models of glioblastoma. The GL261 model was derived from an intracranially induced methylcholanthrene tumor in C57BL/6 mice (22), whereas the SMA-560 model was derived from a spontaneous astrocytoma that developed in VM/Dk mice (23). Although both models grow progressively when transplanted into syngeneic hosts, both GL261 and SMA-560 are responsive to checkpoint blockade immunotherapy (24–28). To identify candidate targets of the host antitumor CD8 T-cell response, we performed whole-exome DNA and RNA sequencing of both GL261 and SMA-560 gliomas to characterize expressed tumor-specific mutations. We then applied multiple *in silico* MHC class I binding prediction algorithms to identify putative high-affinity H-2D^b and H-2K^b-restricted neoepitopes and then assessed the immunogenicities of these predicted neoantigens. Using IFN γ Enzyme-Linked Immunospot (ELISPOT) assays and tetramer analysis, we confirmed the presence of an endogenous CD8 T-cell response specific to the H-2D^b-restricted neoantigens, Imp3_{D81N} (GL261), and Odc1_{Q129L} (SMA-560). In addition, we identified reactivity to the H-2K^b-restricted SMA-560-derived neoantigen, E2f8_{K272R}. Furthermore, endogenous neoantigen-specific T-cell populations in the brain and draining lymph nodes to Imp3_{D81N} and Odc1_{Q129L} were detected. By characterizing the neoepitope profile and using this information to identify and monitor neoantigen-specific host antiglioma T-cell responses in these preclinical models, we extend the cancer immunogenomics approach to glioblastoma and provide a genomics-based system to further explore the mechanisms of action of immunotherapeutics in glioblastoma.

Materials and Methods

Animals and cells

Animal studies were approved by the Animal Studies Committee at Washington University. C57BL/6 were purchased from Taconic Biosciences, and VM/Dk mice were obtained from Dr. John Sampson (Duke University). Mice were housed in specific pathogen-free conditions. GL261 was obtained from the NCI Tumor Repository in 2014 and fully characterized by DNA whole-exome and RNA sequencing. SMA-560 was obtained from Drs. John Sampson and Peter Fecci (Duke University) and fully characterized by DNA whole-exome and RNA sequencing. Either 1×10^6 (subcutaneous) or 5×10^4 (intracranial) GL261 or SMA-560 cells were implanted into 6- to 10-week-old naïve syngeneic C57BL/6 mice or VM/Dk, respectively. For intracranial experiments, tumor cells were resuspended in 5 μ L PBS and injected into the right striatum of anesthetized syngeneic mice in a stereotaxic frame. Subcutaneously implanted tumors were harvested when approximately 10 mm in greatest diameter (approximately 2 weeks for SMA-560 and 3 weeks for GL261). Intracranially implanted tumors were harvested when mice became moribund.

Lymphocyte isolation

Subcutaneous or intracranial tumors were minced into 1–2 mm chunks, plated in 12-well plates, and incubated at 37°C in

culture media (RPMI-1640, L-glutamine, penicillin/streptomycin, β -mercaptoethanol, MEM, 10% FBS) with 100 U/mL recombinant human IL2. After 2 to 5 days, tumor-infiltrating lymphocytes (TILs) were harvested and passed through a 70- μ m cell strainer. Draining lymph nodes and spleens were mechanically dissociated and filtered through a 70- μ m cell strainer. Lymphocytes were purified using the Dead Cell Removal Kit (Miltenyi Biotec). For ELISPOT assays, mononuclear cells were isolated from splenocytes by Ficoll-Paque PLUS density gradient (GE Healthcare Life Sciences).

DNA whole-exome and RNA sequencing

Libraries were captured using the Agilent Mouse Exome reagent. Sequencing was performed on an Illumina HiSeq2000 (Illumina Inc.). Sequence coverage was as follows: C57BL/6 normal (92.1X), GL261 tumor (76.7X), VM/Dk normal (86.0X), and SMA-560 tumor (82.4X). Data were aligned to reference sequence using bwa (29) version 0.5.9 and then merged and deduplicated using picard version 1.46 (<https://broadinstitute.github.io/picard/>). Single-nucleotide variants (SNV) were detected using the union of three callers: (1) samtools (30) version r963 intersected with Somatic Sniper (31) version 1.0.2 and processed through false-positive filter v1, (2) VarScan (32) version 2.2.6 filtered by varscan-high-confidence filter version v1 and processed through false-positive filter v1, and (3) Strelka (33) version 0.4.6.2. Indels were detected using the union of three callers: (1) GATK (34) somatic-indel version 5336 pindel version 0.5 filtered with pindel false-positive and VAF filters (params: $-\text{variant-freq-cutoff} = 0.08$), (2) VarScan (32) version 2.2.6 filtered by varscan-high-confidence-indel version v1, and (3) Strelka (33) version 0.4.6.2. SNVs and Indels were further filtered using a Bayesian classifier (<https://github.com/genome/genome/blob/master/lib/perl/Genome/Model/Tools/Validation/IdentifyOutliers.pm>), retaining variants classified as somatic with a binomial log-likelihood of at least 3. Results were filtered to require expression of the mutant allele (FPKM >1 and at least one variant-supporting read in the RNA) and variant allele frequency (VAF) >1%.

MHC class I binding prediction

The potential for GL261 and SMA-560 missense mutations to bind to H-2D^b or H-2K^b molecules was predicted using multiple pipelines: Stabilized Matrix Method (SMM) algorithm, the SMM with a Peptide:MHC Binding Energy Covariance algorithm (SMMPMBEC), Artificial Neural Network (ANN) algorithm, and NetMHCpan algorithm provided by the Immune Epitope Database and Analysis Resource (<http://www.immuneepitope.org>) as well as the Position Specific Scoring Matrices (PSSM). Results were expressed as affinity values ($1/IC_{50} \times 100$; where IC_{50} is the half-maximum inhibitory concentration, nmol/L). The mean affinity value was calculated from predicted binding affinities of all five algorithms. A binding cut-off of $IC_{50} < 500$ nmol/L was empirically applied as an additional filter.

ELISPOT assay

Naïve splenocytes were plated at a concentration of 100,000 to 200,000 cells/well in 100 μ L serum-free C.T.L. (Cellular Technology, Ltd.) media on precoated murine IFN γ ELISPOT plates (Cellular Technology, Ltd.). Harvested TILs were added at a final concentration of 25,000 cells/well in 200 μ L with peptide (10 μ mol/L; Peptide 2.0 Inc.). Conavalin A (1 μ g/well) was a positive

control. Plates were incubated overnight at 37°C and analyzed using the C.T.L. ImmunoSpot Kit (Cellular Technology, Ltd.).

Tetramers and flow cytometry

Recombinant H-2D^b heavy chains and human β2-microglobulin were produced in BL21-CodonPlus (DE3)-RIPL *Escherichia coli* (Agilent) and purified from inclusion bodies. Purification of MHC class I heavy and light chain by size-exclusion FPLC was performed as described (35). Peptide-specific monomers were generated by UV-mediated exchange as described (36). MHC class I multimers were generated using streptavidin-conjugated PE or APC (Invitrogen). For TIL and lymph nodes, 1 × 10⁶ cells were dual stained with PE- and APC-peptide:MHC class I tetramers for 15 minutes at 37°C. Cells were then stained with CD8α-FITC, Thy1.2-PE/Cy7, and Zombie NIR Fixable Viability Kit (BioLegend) and analyzed on a BD Fortessa flow cytometer.

Statistical analysis

The intergroup difference in mean number of spots on ELISPOT was evaluated using a Student *t* test with *P* < 0.05 as statistically significant.

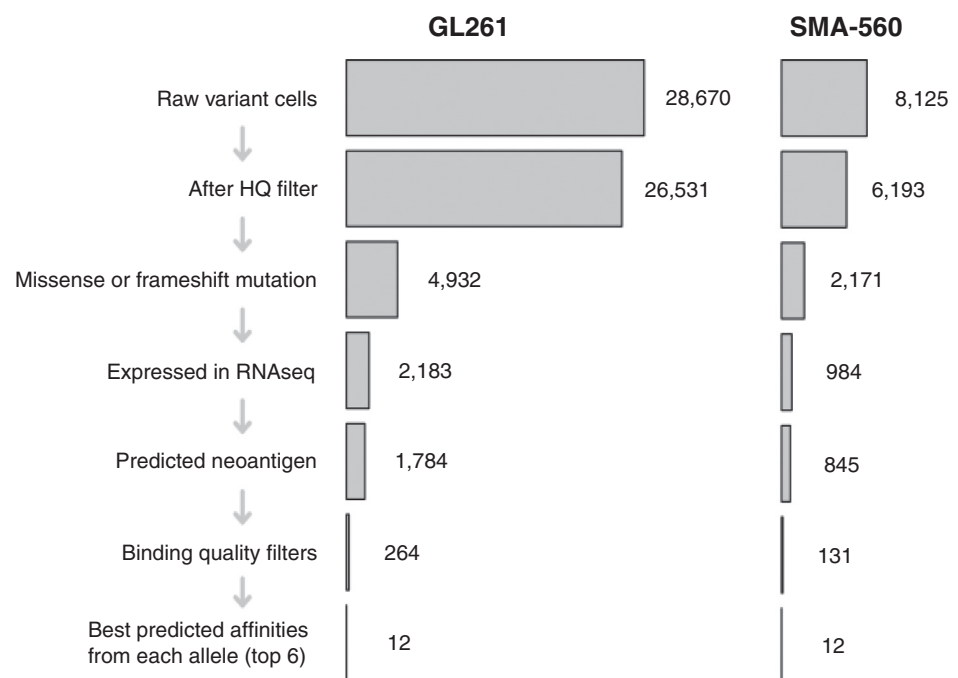
Results

To identify candidate neoantigens within GL261 and SMA-560, we applied a cancer immunogenomics discovery pipeline similar to the approach that has been described previously (11, 14, 15). Briefly, unbiased genomic characterization is performed to detect expressed tumor-specific variants, and existing *in silico* algorithms are used to determine the predicted affinity strength with which translated peptide sequences bind to particular MHC class I molecules. DNA whole-exome sequencing was employed to identify tumor-specific, somatic missense SNVs. GL261 harbored a total of 26,531 somatic mutations compared with the syngeneic C57BL/6 genome, of which 4,932 were missense or frameshift

variants (Fig. 1; Supplementary Table S1). Conversely, SMA-560 harbored 6,193 somatic mutations compared with the syngeneic VM/Dk exome, of which 2,171 were missense or frameshift variants (Fig. 1; Supplementary Table S2). To determine which of the identified SNVs were expressed and could be translated into candidate neoepitopes, RNA sequencing was performed on both tumors. These data showed that fewer than half of somatic SNVs were expressed at the transcript level; GL261 and SMA-560 expressed 2,183 and 984 mutations, respectively (Fig. 1; Supplementary Tables S1 and S2). Because both C57BL/6 and VM/Dk genetic backgrounds possess the H-2^b haplotype, we used *in silico* computational approaches to determine which mutant proteins represented candidate neoepitopes that were predicted to bind with high affinity to the associated class I MHC, H-2D^b and H-2K^b. We calculated the median binding affinities from a combination of five MHC class I binding prediction algorithms, and after applying an affinity cutoff of IC₅₀ < 500 nmol/L, we identified 181 H-2D^b and 1,599 H-2K^b predicted high-affinity neoepitopes in GL261 (Fig. 2A and B; Supplementary Table S1). Similarly, there were 77 H-2D^b and 647 H-2K^b predicted high-affinity neoepitopes in SMA-560 (Fig. 2C and D; Supplementary Table S2). The top six predicted binding affinities for each allele were prioritized for further evaluation (Tables 1 and 2). These data showed that both GL261 and SMA-560 harbored a high mutational load and expressed potentially high-affinity candidate neoantigens.

To determine the immunogenicities of the top predicted neoantigen candidates, we isolated and cultured TIL from established subcutaneously implanted tumors and performed IFNγ ELISPOT assays to assess the presence of neoantigen-specific T cells within intratumoral TIL populations. GL261-derived TIL demonstrated increased activation following stimulation with the H-2D^b-restricted peptide, Imp3_{D81N} (referred to as mImp3; Fig. 3A and B). SMA-560 TIL stimulated with the H-2D^b-restricted candidate neoantigen, Odc1_{Q129L} (referred to as

Figure 1. Mutational burden of GL261 and SMA-560. Schematic of workflow for GL261 and SMA-560 neoantigen discovery. Numbers to right of bar graph indicate number of mutations identified following application of indicated filter on left side of bar graph.



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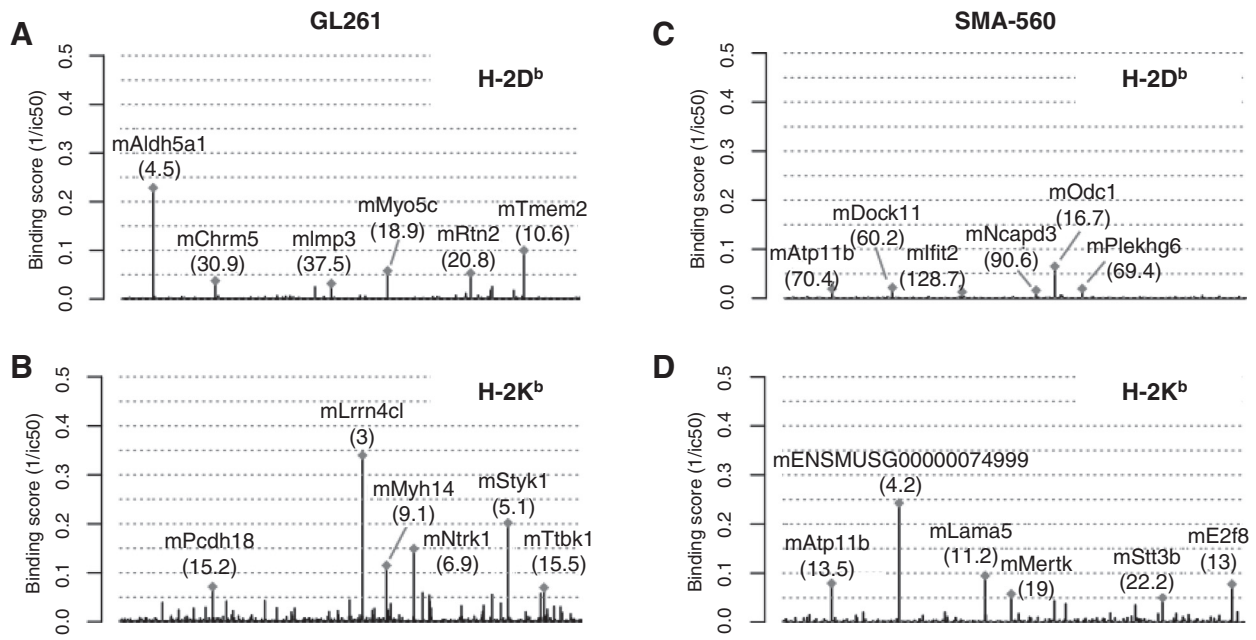


Figure 2. Neoantigen landscape of GL261 and SMA-560. Manhattan plot of mean binding affinity (1/IC₅₀) of putative candidate GL261-derived neoantigens for H-2D^b (A) and H-2K^b (B). Manhattan plot of mean binding affinity (1/IC₅₀) of putative candidate SMA-560-derived neoantigens for H-2D^b (C) and H-2K^b (D). Labeled are the six highest predicted binding affinity candidate neoantigens. Numbers in parentheses represent calculated IC₅₀ (nmol/L) for each candidate neoantigen.

mOdc1), as well as the H-2K^b-restricted epitope, E2f8_{K272R} (referred to as mE2f8), had significantly increased IFN γ (Fig. 3C–F). Of note, none of the top predicted H-2K^b-restricted GL261 neoantigens induced increased IFN γ production from GL261-derived TIL (data not shown).

Having demonstrated the functional immunogenicities of candidate neoantigens by ELISPOT, we next asked whether we could directly detect the presence of TIL-derived CD8 T cells specific to the cognate neoantigen:MHC complex using tetramer analysis. For these experiments, we focused solely on the H-2D^b-restricted neoantigens. In TIL derived from established GL261 subcutaneously implanted tumors, approximately 3.4% (range, 2.5%–3.9%) of short-term cultured CD8 T cells were specific for the H-2D^b-mImp3 tetramer, whereas there were no cells that stained

with the negative control H-2D^b-mOdc1 tetramer (Fig. 4, top left graph). Likewise, approximately 1.84% (range, 1.1%–3.1%) of CD8 T cells in TIL derived from established subcutaneously implanted SMA-560 tumors were found to be tetramer positive for mOdc1 (Fig. 4, top left graph). Of note, no detectable tetramer-positive CD8⁺ T-cell populations specific to other predicted GL261 or SMA-560 H-2D^b-restricted putative neoantigens were appreciated (data not shown). Together with the ELISPOT data, these results demonstrate the presence of functional, endogenous neoantigen-specific CD8 T cells in each of these transplantable glioblastoma models.

Finally, due to the distinct features of immune responses in the CNS compared with those in extracranial sites, we asked whether we could detect evidence of endogenous host immune responses to brain tumor neoantigens following orthotopic intracerebral implantation. We isolated TIL from intracranial GL261 and SMA-560 tumors and assessed for the presence of mImp3- and mOdc1-specific CD8 T cells, respectively, by tetramer analysis. Both mImp3- and mOdc1-specific CD8 T cells were detectable at comparable frequencies (GL261, 2.8%; range, 1.7%–4.3%; SMA-560, 1.6%; range, 0.8%–3.2%; Fig. 4, top right graph). Because it has been shown that there may be a physical lymphatic connection between the brain and the cervical lymph nodes (37, 38), we next asked whether we could identify neoantigen-specific T cells within the draining cervical lymph nodes in mice orthotopically transplanted with GL261 or SMA-560 tumors. Surprisingly, a small but discrete population of tetramer-positive, neoantigen-specific CD8 T cells were detectable within the cervical lymph nodes of both C57BL/6 mice harboring intracranial GL261 tumors and VM/Dk mice harboring intracranial SMA-560 mice (Fig. 4, bottom right graph). Taken together, these data show endogenous neoantigen-specific CD8 T-cell responses are

Table 1. Top 6 *in silico* predicted H-2D^b and H-2K^b candidate GL261-derived neoantigens with the corresponding amino acid sequence

H-2D ^b restriction	
Amino acid mutation	Neoantigen sequence
mAldh5a1 V444F	F AIANAAEV
mTmem2 K1042N	V M LENGYTI
mMyo5c L822M	Y M VRNLYQL
mRtn2 L405F	GA I FNGFTL
mChrm5 R503W	YALCN R TFW
mImp3 D81N	AALL N KLYA
H-2K ^b restriction	
Amino acid mutation	Neoantigen sequence
mLrrn4cl V200L	VTLVYAAL
mStyk1 L429I	ISY S FSVI
mNtrk1 H470Q	MSL Q FMTL
mMyh14 G135V	LIY T SYVL
mPcdh18 Q1012R	MSSV F RRL
mTtbk1 C450R	RSL R YRRV

NOTE: The mutated amino acid is denoted by bold and underlining.

Table 2. Top 6 *in silico* predicted H-2D^b and H-2K^b candidate SMA-560–derived neoantigens with the corresponding amino acid sequence

H-2D ^b restriction	
Amino acid mutation	Neoantigen sequence
mOdc1 Q129L	YAASNGVLM
mDock11 G1958V	SVQVNAVPL
mPlekhg6 G10C	FGPPNECP
mAtp11b K884M	FFYMNVCFI
mNcapd3 V510L	NTVLNPSPL
mIfit2 V60I	ATMCNILAYI
H-2K ^b restriction	
Amino acid mutation	Neoantigen sequence
mENSMUSG00000074999 C15F	SSFYIYAI
mLama5 G897V	VRFFVFNPL
mE2f8 K272R	MSQRVFMV
mAtp11b K884M	VQYFFYMNV
mMertk R682P	TFLLYSPL
mStt3b G323V	AAVVFALL

NOTE: The mutated amino acid is denoted by bold and underlining.

detectable in both orthotopically transplanted brain tumors and cervical lymph nodes.

Discussion

We employed a cancer immunogenomics approach to characterize the neoantigen landscape of two distinct, well-studied, murine orthotopic transplant models of glioblastoma. Our work represents the first application of this approach for neoantigen discovery in preclinical brain tumor model systems. The presence of H-2D^b–restricted Imp3_{D81N} (GL261) and Odc1_{Q129L} (SMA-560) as well as H-2K^b–restricted E2f8_{K272R} (SMA-560) neoantigen-reactive TIL was detected by screening candidate neoantigens with the highest predicted binding affinities using a functional IFN γ ELISPOT assay. H-2D^b–restricted tetramer analysis validated the presence of mImp3- and mOdc1-specific CD8 T-cell populations both within intracranial tumors and cervical lymph nodes.

The growing use of T cell–activating immunotherapies in cancer treatment has stimulated further studies into the identities of the antigens recognized by this effector subset. More broadly, we have had a longstanding interest in how the concept of cancer immunoeediting applies to malignant glioma (39). Although there are several types of tumor antigens that T cells may recognize (40), neoantigens derived from somatic mutations are tumor-restricted in most cases and therefore harbor a lower likelihood of inducing tolerance or autoimmunity. Especially in the brain, it is important to limit the cross-reactivity of induced immune responses against normal tissue. The identities of the neoantigens we characterized underscore the emphasis of the cancer immunogenomics perspective on prioritizing somatic mutant proteins as antigens rather than as obvious drivers or passengers. Although Imp3, a small ribonucleoprotein, and Odc1, ornithine decarboxylase 1, have been implicated in cancer in some studies (41, 42), neither gene is recurrently mutated. Moreover, the contributions of mutant E2f8, an atypical E2F family repressor, to oncogenesis remain to be clarified.

Although our results provide evidence that predicted neoantigens can be validated as immunogenic in brain tumor models, the functional validation of only a subset of the highest predicted binding candidates suggests that additional work is needed to address further refinements to the *in silico* approach in order to increase the concordance between neoantigenicity and immuno-

genicity. Furthermore, the identified neoantigen-specific T cells represented a small subset of the total CD8 TIL population suggesting the presence of yet to be identified neoantigens. Although there exists substantial experience in prediction algorithm development (43–45), further refinements are necessary to enhance our ability to predict immunogenicity. Specifically, further work is focusing on the incorporation of other parameters—such as mutant transcript expression levels, IC₅₀ thresholds of neoantigen binding to MHC molecules, comparative binding affinities of mutant peptides to wild-type counterparts, and location of amino acid substitutions within the TCR/peptide:MHC topography—to enhance the predictive potential of the neoantigen discovery pipeline. Moreover, ongoing work is directed at determining the extent of neoantigen screening needed to capture the entire immunogenic landscape, especially in tumors with high mutational burdens such as these preclinical models, and how best to exploit neoantigen identification for therapeutic approaches. Nevertheless, the cancer immunogenomics approach to neoantigen detection has become a powerful method to probe the cancer immunome (11).

Our identification of the H-2D^b–restricted, GL261-derived immunodominant neoantigen, Imp3_{D81N}, supports the findings of a previous study (46), providing further validity to our immunogenomics approach. Specifically, an attenuated oncolytic strain of HSV-1 was injected into an established subcutaneously implanted tumor, and a GL261-specific cytotoxic T-cell line was derived from the isolated splenocytes. Using a cDNA expression cloning approach, an antigen encoded by the *1190002L16Rik* gene was identified and termed "GARC-1." Strikingly, during the preparation of our article, we determined that the amino acid sequence of mutated GARC-1 recognized by this GL261 T-cell clone is identical to the Imp3_{D81N} 9-mer we identified, AALLNKLYA. Thus, these data provide independent validation of the immunogenomics pipeline and further corroboration of the immunogenicity of the Imp3_{D81N} antigen in the GL261 model.

Our observations of neoantigen-specific CD8 T cells within brain tumors and cervical lymph nodes provide further opportunities to study the cellular immunobiology of spontaneously arising immune responses to endogenous glioma antigens. Although it is acknowledged at this point that the brain is not "immunoprivileged" (6), it is clearly immunologically specialized. A number of studies have provided evidence that there may be a connection between the cerebrum and secondary lymphoid tissues, particularly the cervical lymph nodes (38). Recent work identified a dural lymphatic network that may represent the physical conduit by which transit of antigens from the brain to the extracranial secondary lymph nodes takes place (37, 47). Of note, we did not detect tumor cells within the lymph nodes we analyzed (data not shown). Additional work will be necessary to clearly establish the cellular basis for the antiglioma immune response both in naturally occurring immunosurveillance and following vaccination or other therapeutic interventions.

Our results point to several other clinically relevant topics. First, personalized vaccination targeting mutated antigens is an active area of investigation (11, 12, 48), and clinical trials testing this possibility in the treatment of patients with glioblastoma are presently accruing patients (<http://clinicaltrials.gov>: NCT02287428, NCT02510950). The cancer immunogenomics neoantigen discovery pipeline employed in the current study therefore establishes a preclinical setting in which various

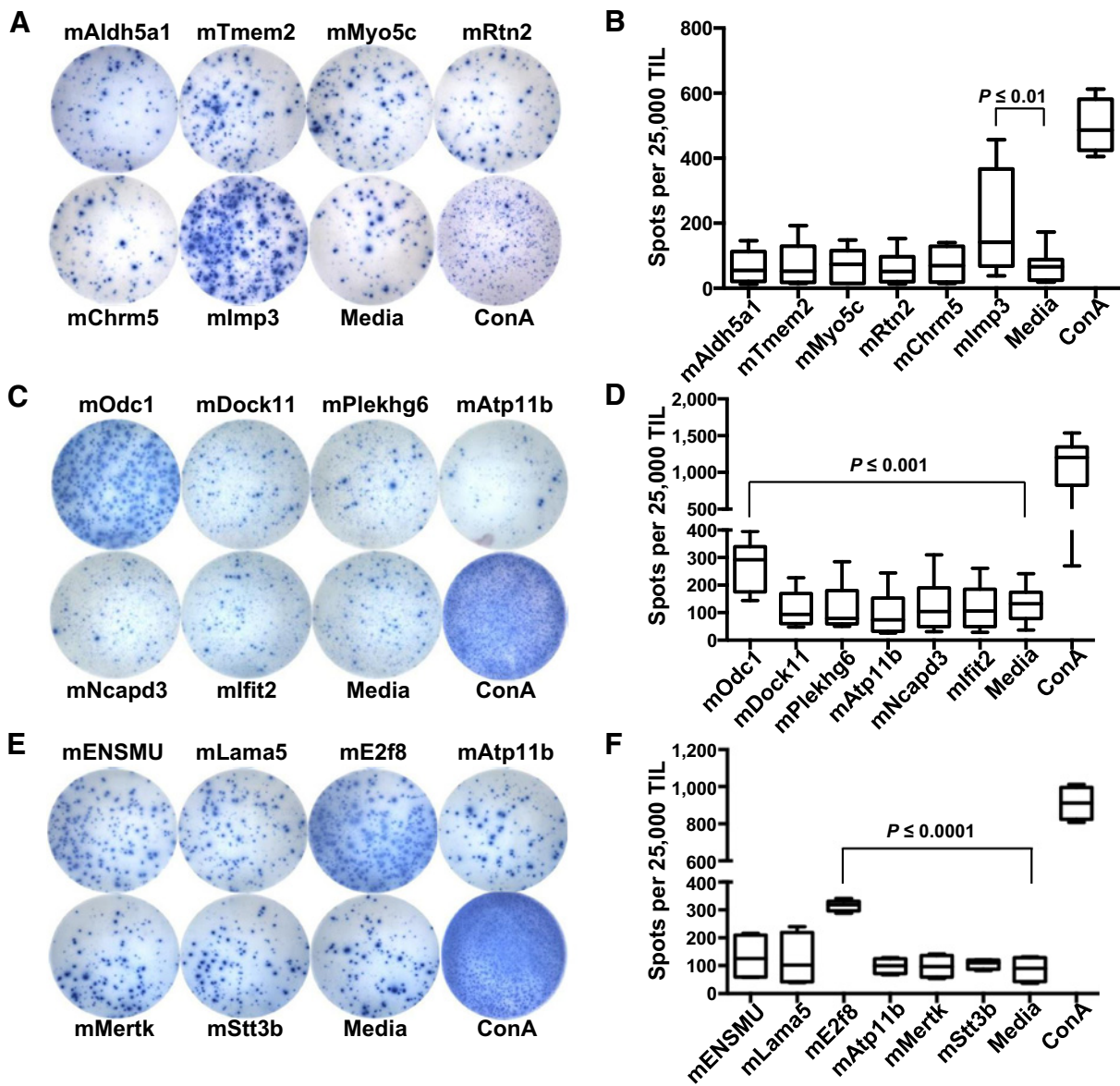


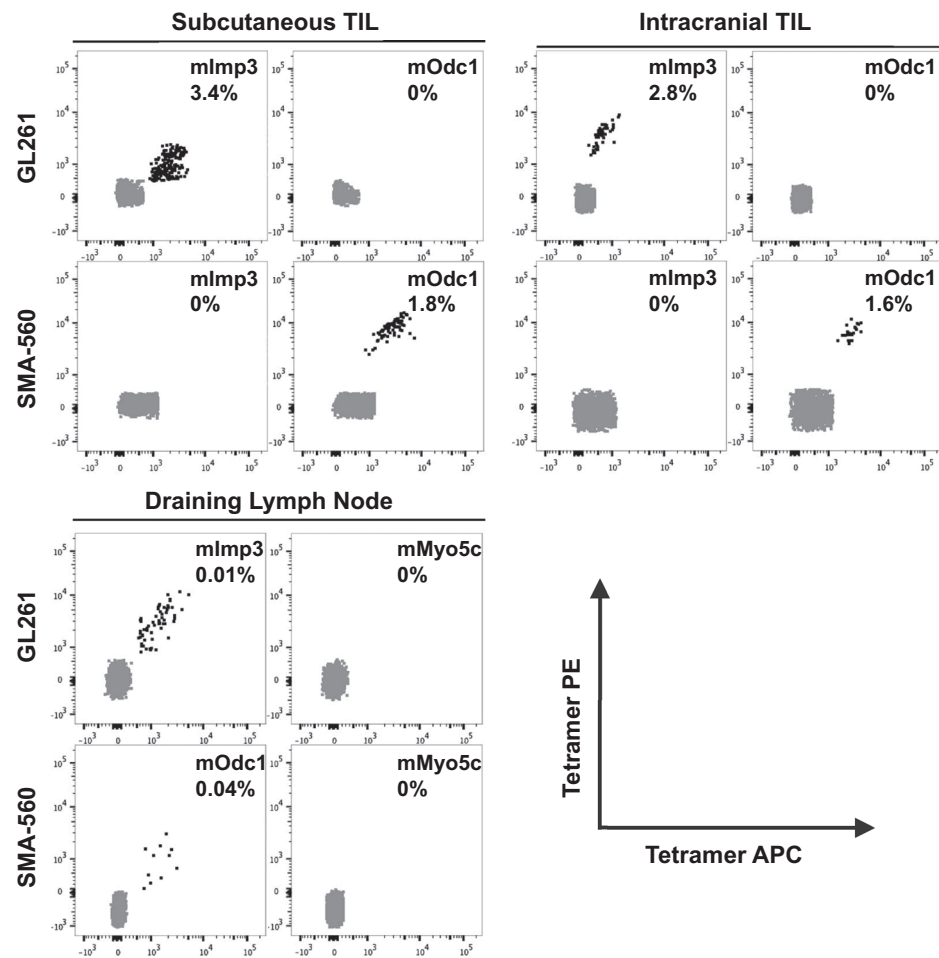
Figure 3. Identification of neoantigen-reactive TIL in GL261 and SMA-560. **A**, Representative images from IFN γ ELISPOT of GL261 TIL stimulated with H-2D^b-restricted candidate neoantigens. TILs were isolated from established subcutaneously implanted GL261 at day 21 and incubated for 4 days in IL2 (50 U/mL). Cultured TILs (25,000 TIL/well) were then incubated overnight with the indicated peptide (10 μ mol/L) and assessed for IFN γ production the following day by ELISPOT. **B**, Bar graph quantifying number of IFN γ spots per well. **C**, Representative wells from IFN γ ELISPOT of SMA-560 TIL stimulated with H-2D^b-restricted candidate neoantigens. **D**, Bar graph quantifying number of IFN γ spots per well. **E**, Representative wells from IFN γ ELISPOT of SMA-560 TIL stimulated with H-2K^b-restricted candidate neoantigens. **F**, Bar graph quantifying number of IFN γ spots per well. Presented data depict pooled results from at least three experiments with 2 to 3 mice per experiment.

methods of vaccination against genomically identified glioma neoantigens can be rigorously tested. Second, the mutational load of both the GL261 and SMA-560 tumors is much higher than is typically observed in primary human glioblastoma in which the tumor exome mutational load is usually less than 100 (2). Thus, the number of potentially targetable neoantigens may be more limited in contexts harboring this mutational burden. However, work over the last 10 years has identified that approximately 20% to 30% of recurrent glioblastomas harbor a "hypermutator" phenotype (49–53) in which the number of

exome mutations are significantly higher than at presentation and similar to the number of mutations seen in these preclinical tumors representing a subpopulation of glioblastoma patients that may benefit from neoantigen-based personalized vaccination. Moreover, due to the growing literature that tumors with hypermutated phenotypes may be more sensitive to checkpoint blockade immunotherapy (54), preclinical models with similar genomic landscapes may represent valuable models to explore the immunologic basis for these observations and whether they are generalizable to the CNS.

Figure 4.

Detection of neoantigen-specific CD8 T cells by tetramer within TIL and draining lymph node. TILs isolated from subcutaneously implanted (top left graph) or intracranially implanted (top right graph) GL261 (top row) and SMA-560 (bottom row) tumors were cultured for 4 days in IL2 (50 U/mL) prior to tetramer staining. Indicated peptides were loaded into H-2D^p-restricted tetramers and dual labeled with PE and APC fluorochromes. Tetramer-positive cells are identified as double-positive populations. Draining cervical lymph nodes were surgically removed at time of sacrifice and stained directly with indicated tetramers (bottom left graph). Representative FACS plots of at least three experiments containing pooled TIL from 2 to 5 mice with similar results are shown. Number in each FACS plot represents average percentage of tetramer-positive cells within CD8⁺ population of cells.



Disclosure of Potential Conflicts of Interest

R.D. Schreiber has ownership interest (including patents) in, and is a consultant/advisory board member for, Igenica, Jounce, and Neon. No potential conflicts of interest were disclosed by the other authors.

Compliance with Ethical Standards

The authors declare they have no conflicts of interest to disclose. All applicable international, national, and/or institutional guidelines for the care and use of animals were followed. All procedures performed in studies involving animals were in accordance with the ethical standards of the institution or practice at which the studies were conducted. This article does not contain any studies with human participants performed by any of the authors.

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