Mechanisms and therapeutic implications of hypermutation in gliomas

A high tumour mutational burden (hypermutation) is observed in some gliomas1–5; however, the mechanisms by which hypermutation develops and whether it predicts the response to immunotherapy are poorly understood. Here we comprehensively analyse the molecular determinants of mutational burden and signatures in 10,294 gliomas. We delineate two main pathways to hypermutation: a de novo pathway associated with constitutional defects in DNA polymerase and mismatch repair (MMR) genes, and a more common post-treatment pathway, associated with acquired resistance driven by MMR defects in chemotherapy-sensitive gliomas that recur after treatment with the chemotherapy drug temozolomide. Experimentally, the mutational signature of post-treatment hypermutated gliomas was recapitulated by temozolomide-induced damage in cells with MMR deficiency. MMR-deficient gliomas were characterized by a lack of prominent T cell infiltrates, extensive intratumoral heterogeneity, poor patient survival and a low rate of response to PD-1 blockade. Moreover, although bulk analyses did not detect microsatellite instability in MMR-deficient gliomas, single-cell whole-genome sequencing analysis of post-treatment hypermutated glioma cells identified microsatellite mutations. These results show that chemotherapy can drive the acquisition of hypermutated populations without promoting a response to PD-1 blockade and supports the diagnostic use of mutational burden and signatures in cancer.

Identifying genomic markers of response to immune checkpoint blockade (for example, PD-1 blockade) may benefit cancer patients by providing predictive biomarkers for patient stratification and identifying resistance mechanisms for therapeutic targeting. Gliomas typically have a low tumour mutational burden (TMB) and a highly immunosuppressive microenvironment—two features associated with immunotherapy resistance. Nevertheless, recent work has suggested that a subset of patients with high-TMB (hypermutated) gliomas might benefit from PD-1 blockade6. Although consistent with data from other cancers7–9, these initial observations were derived from uniplediseasercontextsuch as constitutional DNA mismatch-repair (MMR) deficiency syndrome6. Therefore, the extent to which glioma patients at large will benefit from this approach is unknown. While large amounts of genomic data on gliomas exist1,4,5,10,11,12, our understanding of the clinical landscape of hypermutation and the mechanisms that underlie its development remain unclear. Hypermutation is rare in newly-diagnosed gliomas (de novo hypermutation), but common in tumours that have recurred after the use of alkylating agents.

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agents (post-treatment hypermutation)\textsuperscript{4,6,10,11}. Given that gliomas exhibit substantial inter-patient and intra-tumoral genomic variation\textsuperscript{10,12,13}, it remains to be determined whether molecular biomarkers (for example, IDH1 or IDH2 (hereafter IDH1/2) mutations) reliably predict the development of hypermutation or response to immunotherapy.

An association between hypermutation and MMR mutations has been observed in gliomas\textsuperscript{4–11}, but most of the reported MMR mutations were not functionally characterized, and their role in causing hypermutation is unclear. Other studies have suggested that alkylating agents such as temozolomide are the direct cause of hypermutation\textsuperscript{4}. This was supported by the discovery of a mutational signature (single base substitution (SBS) signature 11) characterized by the accumulation of C:C→A:T transitions at non-CpG sites in hypermutated gliomas after exposure to alkylating agents\textsuperscript{46}. However, the fact that hypermutation is undetectable in most gliomas that recur after temozolomide treatment challenges this notion\textsuperscript{14}. Furthermore, it remains unclear whether this mutational pattern enhances tumour immunogenicity and renders gliomas responsive to PD-1 blockade. Not all hypermutated cancers respond to such treatments\textsuperscript{19}; a more accurate characterization of the phenotypic and molecular features of hypermutated gliomas therefore would help clinicians to manage such patients more effectively.

Mutational burden and signatures in gliomas

Previous studies included too few hypermutated gliomas to characterize the landscape of hypermutation in gliomas\textsuperscript{11}. We therefore created a cohort of sufficient scale ($n = 10,294$ and subtype diversity by leveraging large datasets generated from clinical sequencing panels (DFCI-Profile, MSKCC-IMPACT and FMI)\textsuperscript{17,18}. All samples from patients with a histopathological diagnosis of glioma were included and classified into molecular subgroups according to histopathology, mutational status of IDH1/2, and whole-arm co-deletion of chromosomes 1p and 19q (1p/19q co-deletion) (Extended Data Fig. 1, Supplementary Tables 1, 2). We quantified the TMB of all samples (median 2.6 mutations (mut.) per Mb (range 0.0–781.3)), established thresholds for hypermutation by examining the distribution of TMB (Extended Data Fig. 2)\textsuperscript{17,18}, and identified 558 (5.4%) hypermutated gliomas (median TMB 50.8 mut. per Mb (8.8–781.3)) for further analysis.

Using samples with detailed clinical annotation (DFCI-Profile), we found that the prevalence of hypermutation varied between and within subgroups (Fig. 1a, b, Extended Data Fig. 3a, b, Supplementary Table 3). Hypermutation was detected almost exclusively in diffuse gliomas (99.1% of hypermutated samples) with high-grade histology (95.6%) and was more prevalent in recurrent tumours (16.6% versus 2.0% in newly diagnosed tumours; Fisher’s exact test, $P < 10^{-15}$) (Fig. 1b). In samples of recurrent tumours, hypermutation was associated with markers of response to alkylating agents, including IDH1/2 mutation (hypermutation in 1.4% of newly diagnosed versus 25.4% of post-treatment IDH1/2-mutant tumours; Fisher’s exact test, $P = 2.0 \times 10^{-13}$), 1p/19q co-deletion (0.0% versus 33.8%, $P = 7.3 \times 10^{-20}$), and MGMT promoter methylation (2.4% versus 24.2%, $P = 9.0 \times 10^{-15}$). The effect of IDH1/2 mutation was confirmed only in MGMT-methylated tumours (Extended Data Fig. 3c). These findings suggest that selective pressure from therapy may elicit progression towards hypermutation.

The standard treatment for gliomas includes surgery, radiation and chemotherapy with alkylating agents\textsuperscript{10,20}. To assess the role of each of these in the development of hypermutation, we analysed associations between TMB and detailed patterns of treatment in 356 recurrent gliomas. Hypermutation was associated with prior treatment with temozolomide (Fisher’s exact test, $P < 10^{-15}$) in a dose-dependent manner (Fig. 1b, Extended Data Fig. 3d, e), but not with radiation ($P = 0.88$) or nitrosoureas ($P = 0.78$). Among recurrent tumours from patients who had received only one adjuvant treatment modality, TMB was increased only in temozolomide-treated samples (median 16.32 (inter-quartile range (IQR) 6.95–70.32) versus 6.08 (3.80–7.97) with surgery only, $P = 4.0 \times 10^{-7}$; Extended Data Fig. 3f). Of note, the prevalence of hypermutation in post-temozolomide samples correlated with the chemosensitivity of the primary, molecularly defined tumour type (1p/19q co-deleted oligodendrogliomas (59.5%) > IDH1/2-mutant astrocytomas (30.2%) > MGMT-methylated IDH1/2 wild-type glioblastomas (23.1%) > MGMT-unmethylated IDH1/2 wild-type glioblastomas (5.6%); $P = 3.8 \times 10^{-7}$; Fig. 1b). We observed a similar pattern in the FMI validation dataset (Extended Data Fig. 3g-i).

The systematic analysis of somatic mutation patterns by genome sequencing has identified a variety of mutation signatures in human cancer which are driven by known and unknown DNA damage and repair processes\textsuperscript{14}. We examined the contributions of 30 previously reported signatures (COSMIC signatures v2) within our cohort to investigate the biological processes that cause hypermutation in gliomas. We first validated that mutational signatures can be predicted using large targeted panel sequencing in hypermutated samples (Extended Data Figs. 4, 5a–c). The majority of de novo hypermutated gliomas harboured mutational signatures associated with defects in the MMR pathway (COSMIC signatures 6, 15, 26 and 14) or the DNA polymerase POLE (10 and 14)\textsuperscript{16} (69% and 33% of samples, respectively; Extended Data Fig. 5d, e), implying that constitutional deficiency in MMR or POLE was likely to be the underlying genetic cause of hypermutation. By contrast, 98% of post-treatment hypermutated gliomas showed a mutational signature that has been previously associated with temozolomide exposure (signature II). We also identified two distinct mutational signatures that were highly correlated with mutational signature II (Extended Data Fig. 5b, c) including a previously undescribed signature (S2) associated with 1p/19q co-deletion and lack of prior radiation therapy. Finally, half of the samples with a dominant signature II showed a co-existing minor MMR- or POLE-deficiency signature component (Extended Data Fig. 5e), suggesting that defective DNA repair and mutagen exposure cooperate to drive hypermutation in recurrent gliomas.

Molecular drivers of hypermutation

Only a subset of temozolomide-treated samples (58 of 225, 25.8%) showed evidence of hypermutation, suggesting that additional factors are required for its development. Although MMR defects have been consistently observed in hypermutated gliomas\textsuperscript{11,13}, their co-occurrence with high TMB did not enable prior studies to determine the degree to which MMR mutations represent passenger versus hypermutation-causing driver events. We systematically characterized mutations and copy number variants (CNVs; Supplementary Figs. 1, 2) to identify hypermutation drivers using an unbiased approach that controlled for the increased incidence of passenger mutations associated with hypermutation\textsuperscript{11}. In the merged DFCI-Profile/MSKCC-IMPACT dataset, 36 genes were significantly enriched ($q$-value $< 0.01$) in hypermutated tumours (Fig. 2a). Collectively, MMR mutations stood out among the most enriched (91.2% versus 4.9% in non-hypermutated samples, $q < 1.6 \times 10^{-15}$), and mutations in MSH6 showed the highest enrichment (43.0% versus 1.2%, $q = 3.3 \times 10^{-13}$) (Extended Data Figs. 3j–l, 6a, b). MMR-variant allele frequencies (VAFs) and cancer cell fractions (CCFs) in gliomas were most similar to those in MMR-deficient colorectal (CRC) or endometrial cancers and were higher than in MMR-proficient hypermutated cancers (Extended Data Fig. 6c, d). Some MMR variants in post-treatment hypermutated samples matched the canonical signature 11 sequence context (Extended Data Fig. 5f), suggesting that a subset of these variants are likely to have been caused by temozolomide treatment.

As most MMR variants lacked functional annotation, we next integrated sequencing data with immunohistochemistry for protein loss (Extended Data Fig. 6e). Overall, results from both assays were concordant, consistent with MMR mutations leading to loss of function. In rare samples that lacked MMR variants, signature analysis and MMR immunohistochemistry revealed evidence for MMR deficiency.
Mismatch repair deficiency and signature 11

Our analyses indicated that MMR deficiency together with temozolomide exposure might cause signature 11, as opposed to it being a 'pure' temozolomide signature. To test this idea, we exposed isogenic models of MMR deficiency to temozolomide (Extended Data Fig. 9a, b). After treatment with temozolomide, MMR-deficient PDCLs developed hypermutation with signature 11 whereas MMR-proficient controls (expressing sgGFP) did not (Fig. 2d). We then chronically treated temozolomide-sensitive glioblastoma xenografts (PDXs) with temozolomide until resistance was acquired (Fig. 2e, Extended Data Fig. 9c, d). These tumours developed hypermutation with signature 11 (Fig. 2f, Extended Data Fig. 9e) and shared four unique variants; the dominant-negative MSH6 hotspot mutation (p.T1219I) and three non-coding variants (Fig. 2g), consistent with the theory that the MSH6 mutation drives both hypermutation and acquired temozolomide resistance (Extended Data Fig. 9f).

Collectively, these findings show that temozolomide exerts a previously underappreciated selective pressure in favour of MMR-deficient cells, which are resistant to temozolomide. Exposing MMR-deficient cells to temozolomide induces hypermutation with signature 11 by causing DNA damage in the absence of functional MMR. Therefore, hypermutation with signature 11 represents a potential biomarker for MMR deficiency and temozolomide resistance in gliomas (Extended Data Fig. 9g).
The current hypothesis behind the response of MMR-deficient gliomas to PD-1 blockade is based on their increased neoantigen burden and immune infiltration. We therefore assessed the association between MMR deficiency and T-cell infiltration in gliomas (n = 43) and CRCs (n = 19). As expected, MMR-deficient CRCs exhibited significantly more infiltrating T-cells than their MMR-proficient counterparts (Fig. 3b). By contrast, both MMR-deficient and MMR-proficient glioma samples lacked significant T-cell infiltrates (Fig. 3c).

We next assessed whether the neoantigen burden was lower in MMR-deficient gliomas than in other hypermutated cancers using samples from the GENIE and TCGA datasets (n = 1,748 and 699 hypermutated cancers, respectively). As neoantigen prediction was not feasible using panel sequencing data, we used the nonsynonymous mutational burden as a surrogate measure. This showed that both de novo and post-treatment MMR-deficient gliomas had an increase in their nonsynonymous mutational burden, when compared to non-hypermutated gliomas, and the glioma nonsynonymous mutational burden was similar to other hypermutated cancers (Fig. 3d, Extended Data Fig. 11a, b, Supplementary Table 6). This finding suggested that the total number of neoantigens is unlikely to explain the differences in immune response between gliomas and other hypermutated cancers.

Recent data suggest that, among mutations associated with MMR deficiency, small insertions and deletions (indels) at homopolymer (microsatellites)—which accumulate in MMR-deficient cells—and can cause frameshift mutations—are crucial for producing ‘high-quality’ neoantigens that are recognized by immune cells. Unexpectedly, although the high TMB was associated with an increased homopolymer indel burden in MMR-deficient CRCs, this association was not found in MMR-deficient gliomas (de novo hypermutated gliomas showed a modest increase; Fig. 3d, Extended Data Fig. 11c). This was validated using testing for microsatellite instability (MSI), a clinical biomarker for MMR.
homopolymer indels are subclonal and below the detection limits for PCR. We hypothesized that, in hypermutated gliomas, more of the tumour-infiltrating lymphocytes are recruited (Fig. 3g, Extended Data Fig. 11l). Strikingly, whereas this tumour harboured only a minor increase in its homopolymer indel burden (Fig. 3e, Extended Data Fig. 11e–h). We therefore performed single-cell whole-genome DNA sequencing (scWGS) of 28 cells from a hypermutated, post-temozolomide glioblastoma with an MSH6(T1219I) mutation, and compared these to 35 non-hypermutated cells from the matched pre-treatment sample (Fig. 3f, Extended Data Fig. III–k). In the post-temozolomide sample, 13 of 28 cells (46.4%) were hypermutated with signature 11 (Fig. 3g, Extended Data Fig. 11l). Strikingly, whereas this tumour harboured only a minor increase in its homopolymer indel burden at the bulk level (0.49 versus 0.0 per Mb), the scWGS analysis showed a ninefold increase in microsatellite mutations in all hypermutated cells (Fig. 3h). This suggested that glioma cells with an MSH6(T1219I) variant harbour a subtle MSI phenotype that is not revealed by standard bulk sequencing or clinical MSI assays (Extended Data Fig. 11m).

**Fig. 3 |** Hypermutated and MMR-deficient gliomas harbour unique phenotypic and molecular characteristics including poor outcome and lack of MSI in bulk sequencing. a. Survival of patients with recurrent high-grade glioma from the time of sample collection according to histomolecular group and TMB status (n = 333 recurrent samples; 238 from DFCI-Profile, 95 from MSKCC-IMPACT). Two-sided log-rank test. b. Quantification of tumour-infiltrating CD3-positive T-cells in CRC samples (n = 19). Left, representative low- and high-magnification images of CD3 immunolabelling (brown; intraepithelial lymphocytes, black arrowheads; stromal lymphocytes, black arrows) and nuclear counterstaining (blue). Dashed lines, border between tumour and stroma. Only intraepithelial lymphocytes were quantified. Scale bars: 100 µm (100×), 50 µm (200×). Right: boxes, quartiles; centre lines, median ratio for each group; whiskers, absolute range. Two-sided Wilcoxon rank-sum test. c. Quantification of tumour-infiltrating CD3-positive T-cells in gliomas according to their MMR status (n = 43). For each group, three areas with the maximal CD3 infiltration were selected for quantification (representative images, left). Scale bars: 500 µm (20×), 50 µm (200×). Right: boxes, quartiles; centre lines, median ratio for each group; whiskers, absolute range. Kruskal–Wallis test and Dunn’s multiple comparison test. d. TMB (top) and homopolymer indel burden (bottom) in hypermutated gliomas compared with other hypermutated cancers from the GENIE dataset. Tukey’s boxplots are shown. Two-sided Wilcoxon rank-sum test with Bonferroni correction. e. Pan-cancer analysis of cancer cell fractions in hypermutated gliomas (post-treatment) compared with other hypermutated cancers from the TCGA and ref. * exome datasets (n = 798). One hundred non-hypermutated samples from the TCGA were randomly selected as controls. Boxes, quartiles; centre lines, median ratio for each group; whiskers, absolute range excluding outliers. Two-sided Wilcoxon rank-sum test with Bonferroni correction. f. Workflow for scWGS and bulk tumour DNA sequencing. g. Single-cell sequencing estimate of the number of G:C→A:T transitions at NCC and NCT trinucleotide contexts in 63 cells from a glioblastoma patient with post-temozolomide hypermutation using 10× scWGS sequencing. Error bars show 95% CI. The absolute computed purity was 0.66 for the primary tumour sample and 0.47 for the recurrent tumour sample in the bulk sequencing. h. Single-cell sequencing estimate of microsatellite mutation rate in eight cells from a patient with glioblastoma with post-temozolomide hypermutation. Eight cells were analysed for the presence of MSI using 10× scWGS sequencing. WGA, whole genome amplification; QC, quality control; nucl, nuclei; seq, sequencing.

deficiency. Whereas MSI was identified in all MMR-deficient CRCs, all tested gliomas with MMR protein loss (n = 15) were microsatellite-stable (MSS) (Extended Data Figs. 7d–f, 11d). We hypothesized that, in hypermutated gliomas, more of the homopolymer indels are subclonal and below the detection limits of bulk sequencing, relative to other MMR-deficient cancers. Indeed, analysis of CCFs indicated that hypermutated gliomas contained a greater burden of subclonal variants than did other hypermutated cancers (Fig. 3e, Extended Data Fig. 11e–h). We therefore performed single-cell whole-genome DNA sequencing (scWGS) of 28 cells from a hypermutated, post-temozolomide glioblastoma with an MSH6(T1219I) mutation, and compared these to 35 non-hypermutated cells from the matched pre-treatment sample (Fig. 3f, Extended Data Fig. III–k). In the post-temozolomide sample, 13 of 28 cells (46.4%) were hypermutated with signature 11 (Fig. 3g, Extended Data Fig. IIIl). Strikingly, whereas this tumour harboured only a minor increase in its homopolymer indel burden at the bulk level (0.49 versus 0.0 per Mb), the scWGS analysis showed a ninefold increase in microsatellite mutations in all hypermutated cells (Fig. 3h). This suggested that glioma cells with an MSH6(T1219I) variant harbour a subtle MSI phenotype that is not revealed by standard bulk sequencing or clinical MSI assays (Extended Data Fig. 11m).
PD-1 blockade in MMR-deficient gliomas

As hypermutation in gliomas that acquire MMR deficiency tends to be subclonal and does not generate optimal antitumour T-cell responses, we hypothesized that these tumours might not have high response rates to PD-1 blockade. We performed a retrospective institutional review of patients treated with PD-1 pathway blockade for which the TMB at treatment initiation was available \( n = 210 \). This identified 11 patients with MMR-deficient glioma (5 de novo, 6 post-treatment) who were treated with PD-1 blockade for a median of 42 days (range 13–145; Supplementary Table 7). Nine (81.8%) had disease progression as their best response (Fig. 4a), and the median progression-free survival (PFS) and OS were 1.38 months (95% CI 0.95–2.69) and 8.7 months (95% CI 2.79–15.08), which were not significantly different from the data for matched patients with non-hypermutated glioma (PFS 1.87 months (95% CI 1.28–2.92), OS 9.96 months (95% CI 7.56–15.08); Fig. 4b, Extended Data Fig. 10d).

Because our prior analyses indicated that patients with hypermutated gliomas might have reduced survival, we used a second set of historical controls to compare the outcome of hypermutated gliomas treated with PD-1 blockade versus other systemic agents (Supplementary Table 7). Unexpectedly, we observed a longer median OS for patients treated with other systemic agents when compared to those treated with PD-1 blockade (16.10 months (95% CI 3.98–22.21) versus 8.07 (95% CI 2.79–15.08); \( P = 0.02 \), two-sided log-rank test; Extended Data Fig. 10e, f, Supplementary Table 8). In one patient with hypermutated glioma that showed rapid imaging changes, histopathologic analysis of samples taken before and after treatment with PD-1 blockade showed highly proliferative tumour in both samples, with no significant evidence of pathologic response or increase in immune infiltrates after PD-1 blockade (Extended Data Fig. 10g).

**DISCUSSION**

Collectively, these results support a model in which differences in the mutation landscape and antigen clonality of hypermutated gliomas relative to other hypermutated cancers markedly affect the strong immune infiltrates are seen at diagnosis. Treatment with anti-PD-1 results in expansion of T cells that recognize these clonal neoantigens and substantial antitumour responses. In gliomas (bottom), few mutations are acquired early during tumorigenesis in the majority of tumours. Temozolomide drives the expansion of cells with MMR deficiency and late accumulation of random temozolomide-induced mutations. Ineffective antitumour responses may result from poor neoantigen quality (high burden of missense mutations versus frameshift-producing indels) and high subclonality associated with an immunosuppressive microenvironment. In some tumours, MMR-proficient subclones that have acquired therapy resistance through other pathways can co-exist with MMR-deficient subclones, giving rise to a mixed phenotype.
resistance to temozolomide that are not associated with hypermutation will need to be addressed.

Finally, our data indicate that the absence of an immune response in gliomas is likely to result from several aspects of immunosuppression in the brain that require further characterization. Approaches that increase infiltration by cytotoxic lymphocytes into the glioma microenvironment will probably be required to improve immunotherapy response. Our data also suggest a change in practice whereby repeated biopsies and sequencing to identify progression and hypermutation could inform prognosis and guide therapeutic management.

Online content
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Methods

Datasets

For the DFCI-Profile dataset, clinical data and tumour variant calls identified through targeted next-generation sequencing (NGS) panels of 1,628 gliomas sequenced between June 2013 and November 2018 as part of a large institutional prospective profiling program (DFCI-Profile) were included (Extended Data Fig. I). The distinction between photon and proton radiotherapy was not systematically captured; the vast majority of patients underwent photon radiotherapy. For the MSKCC-IMPACT and FMI datasets, clinical data and tumour variant calls from a total of 545 and 8,121 samples, respectively, that could be assigned to a molecular subgroup (see below) were included. For pan-cancer analyses in targeted panel sets, clinical data and tumour variant calls from the GENIE project (a repository of genomic data obtained during routine clinical care at international institutions) were downloaded from Synapse (public data, release v6.1). For pan-cancer analyses in whole-exome sequencing sets, clinical data and tumour variant calls from 17 hypermutated glioblastomas and from the pan-cancer TCGA dataset were downloaded from the NCI Genomic Data Commons. In addition, 247 gliomas collected at one site between 2009 and 2017 were analysed for protein expression of four MMP proteins (MMP2, MMP6, MLH1, and PMS2) using immunohistochemistry. Written informed consent or IRB waiver of consent was obtained from all participants. Patients of the FMI dataset were not consented for release of raw sequencing data. The study, including the consent procedure, was approved by the institutional ethics committees (10-417/11-104/17-000; Western Institutional Review Board (WIRB), Puyallup, WA).

Tumour genotyping and diagnosis

For the majority of samples, genomic testing was ordered by the pathologist or treating physician as part of routine clinical care to identify relevant genomic alterations that could potentially inform diagnosis and treatment decisions. Patients who underwent DFCI-Profile testing signed a clinical consent form, permitting the return of results from clinical sequencing. In total, 1,628 gliomas were sequenced as part of a cohort of 21,992 tumours prospectively profiled between June 2013 and November 2018. Research tumour diagnoses were reviewed and annotated according to histopathology, mutational status of IDH1 and IDH2 genes, and whole-arm co-deletion of chromosomes 1p and 19q (1p/19q co-deletion), according to WHO 2016 criteria. All samples were assigned to one of four molecular subgroups: IDH1/2-mutant and 1p/19q co-deleted oligodendrogliomas (high- and low-grade), IDH1/2-mutated astrocytomas (high- and low-grade), IDH1/2 wild-type glioblastomas (high-grade only), and IDH1/2 wild-type gliomas of other histologies (high- and low-grade), the latter including grade I pilocytic astrocytomas, glioneuronal tumours and other unclassifiable gliomas. For simplification, IDH1/2 wild-type grade III anaplastic astrocytomas and grade IV diffuse intrinsic pontine gliomas were assigned to the group of IDH1/2 wild-type glioblastomas in all analyses. Samples for which the clinical diagnosis of glioma could not be confirmed (other histology or possible non-tumour sample) and five samples with missing minimal clinical annotation were excluded from all analyses. For the MSKCC-IMPACT and FMI datasets, patients also signed a consent form, and samples were classified using the same procedure. MGMT promoter methylation status was determined as part of routine clinical care using chemical (bisulfite) modification of unmethylated, but not methylated, cytosines to uracil and subsequent PCR using primers specific for either methylated or the modified unmethylated DNA in the Cpg island of the MGMT gene (GenBank accession number AL355331 nt46931-47011).

Targeted panel next-generation sequencing (DFCI-Profile) was performed using the previously validated OncoPanel assay at the Center for Cancer Genome Discovery (Dana-Farber Cancer Institute) for 277 (POPv1), 302 (POPv2), or 447 (POPv3) cancer-associated genes. In brief, between 50 and 200 ng tumour DNA was prepared as previously described, hybridized to custom RNA bait sets (Agilent SureSelect TM, San Diego, CA) and sequenced using Illumina HiSeq 2500 with 2 × 100 paired-end reads. Sequence reads were aligned to reference sequence b37 edition from the Human Genome Reference Consortium using bwa, and further processed using Picard (version 1.90, http://broadinstitute.github.io/picard/) to remove duplicates and Genome Analysis Toolkit (GATK, version 1.6-5-g557da77) to perform localized realignment around indel sites. Single-nucleotide variants were called using MuTect v1.1.4,9, insertions and deletions were called using GATK Indelocator, and variants were annotated using Oncotator. Copy number variants and structural variants were called using the internally developed algorithms RobustCNV40 and Breakmer41 followed by manual review. To filter out potential germline variants, the standard pipeline removes SNPs present at >0.1% in Exome Variant Server, NHLBI GO Exome Sequencing Project (ESP), Seattle, WA (http://evs.gs.washington.edu/EVS/, accessed May 30, 2013), present in dbSNP, or present in an in-house panel of normal tissue, but rescues those also present in the COSMIC database. For this study, variants were further filtered by removing variants present at >0.1% in the gnomAD v2.1.1 database or annotated as benign or likely benign in the ClinVar database.42,43 Arm-level copy number changes were generated using an in-house algorithm specific for panel copy number segment files followed by manual expert review. We set a copy number segment mean log, ratio threshold at which we could accurately call arm amplification and deletion based on the average observed noise in copy number segments. Chromosome arms were classified as amplified or deleted if more than 70% of the arm was altered. A sample was considered co-deleted if more than 70% of both 1p and 19q were deleted.

Sequencing data from MSKCC-IMPACT were generated at the Memorial Sloan Kettering Cancer Center using a custom targeted panel capture to examine the exons of 341 (IMPACT341) or 398 (IMPACT410) cancer-associated genes as previously described. The FMI dataset comprised specimens sequenced as a part of clinical care using a targeted next-generation sequencing assay as previously described (FoundationOne or FoundationOne CDx, Cambridge, MA).12,13 Germline variants without clinical significance were further filtered by applying an algorithm to determine somatic or germline status. Results were analysed for genomic alterations, TMB, MSI and mutational signatures. TMB was assessed by counting all mutations and then excluding germline and known driver mutations. The remaining count was divided by the total covered exonic regions. MSI status was determined as previously described. A log-ratio profile for each sample was obtained by normalizing the sequence coverage at all exons and ~3,500 genome-wide SNPs against a process-matched normal control. This profile was corrected for GC-bias, segmented and interpreted using allele frequencies of sequenced SNPs to estimate tumour purity and copy number at each segment. Loss of heterozygosity (LOH) was defined as local copy number loss or, if local copy number was 2, with an estimated tumour minor allele frequency of 0%. To assess 1p/19q co-deletion, we calculated the percentage of each chromosome arm that was monoallelic (under LOH). A sample was considered 1p/19q co-deleted if both 1p and 19q were >50% monoallelic.

For the DFCI-Profile and FMI datasets, the appropriate cutoffs for hypermutation (17.0 and 8.7 mut/Mb, respectively) were determined by examining the distribution of TMB in all samples and further confirmed using segmented linear regression analysis (Extended Data Fig. 2). For the MSKCC-IMPACT datasets, a threshold previously validated in this dataset was used. In all analyses, the homopolymer indel burden was calculated by computing the number of single base insertions or deletions in homopolymer regions of at least 4 bases in length and dividing the count by the total genomic indel burden as previously established. Somatic variants were annotated as previously described. In addition, for the DFCI-Profile and MSKCC-IMPACT datasets, variants in a selected list of glioma- and DNA-repair associated genes (IDH1, IDH2,
TERT, ATRX, CIC, H3F3A, HIST1H3B, EGF, PDGFA, FGFR1, FGFR2, FGFR3, MET, KRAS, NRAS, HRA5, BRF1, PTPN11, PTEN, PIK3CA, PIK3CB2, PIK3R1, CDKN2A, CDKN2B, CDKN2C, CDK4, CDK6, CCND2, RBL1, TP53, MDM2, MDM4, TP53BP1, PPM1D, CHEK1, CHEK2, RAD51, BRCA1, BRCA2, ATM, ATR, MLI1, MLH3, PMS1, PMS2, MSH2, MSH6, EPCAM, SETD2, POLE, POLD1, MUTYH, WRN) were manually reviewed for oncosignature using several clinical databases for variant annotation (OncoKB, ClinVar, COSMIC, ExAC, and ARUP).

Mutational signature analyses

All variants detected by the sequencing pipeline covered by at least 30× read depth were stringently filtered for germline origin using the gnomAD (population allele frequency greater than 0.1%), and ClinVar (benign or likely benign annotation) databases as well as manual review of VAF distributions and variants with VAFs consistent with possible germline origin (45–55% or over 95%). The mutational spectrum of variants filtered during these steps was similar to a previously published germline mutation spectrum. Signature analysis was performed for hypermutated samples in a two-step approach starting with the Somatic Signatures package in R for de novo signature extraction within each group. To account for the inherent heuristic quality of the NMF approach, the NMF clustering step was repeated 100 times and chosen result was selected based on identifying signatures with the strongest Pearson’s correlation coefficients when compared to the 30 well-established COSMIC signatures v2 (https://cancer.sanger.ac.uk/cosmic/signatures_v2) (Extended Data Fig. 5a–c). We then used the DeconstructSigs package in R to estimate the contribution of identified signatures using a regression model. To account for the potential overfitting of a regression approach owing to either lack of important signatures in the model, or inclusion of uninvolved signatures—we used only the signatures identified by the decomposition approach in step one, supplemented by any strong signature predictions identified through a first pass run of DeconstructSigs with the 30 COSMIC signatures to check for samples that may show strong correlation to an outlier signature. For the FMI dataset, mutational signatures were called as previously described. All point mutations were included in the analysis except known oncogenic driver mutations and predicted germline mutations. A sample was deemed to have a dominant signature if a mutational signature had a score of 0.4 or greater.

To assess the ability of this method to detect hypermutation-associated signatures in targeted panel sequencing data, we compared the signature calls of exome-sequenced samples using all variants (previously published DeconstructSigs signature predictions) versus using only variants that overlapped with the panel-targeted regions. Somatic variant calls for bladder cancer, colon adenocarcinoma, rectal adenocarcinoma, skin cutaneous melanoma, and lung adenocarcinoma (study abbreviations BLCA, COAD, READ, SKCM, LUAD) from the TCGA MC3 dataset were used to assess the detection of COSMIC mutational signatures associated with APOBEC (signatures 2 and 13), mismatch repair (signature 6), ultraviolet light (signature 7), POLE (signature 10), and tobacco (signature 4). Variant calls for 17 hypermutated and 12 non-hypermutated glioma exome-sequenced samples were used for assessing temozolomide (signature 11) detection. There were two COAD samples with known POLE exonuclease domain oncogenic mutations and a POLE signature predicted by DeconstructSigs; these were used for assessing POLE signature detection. For a given threshold number of variants (X), we considered how many samples had at least X variants, and what percentage of these samples could correctly predict the exome-based signature using panel-restricted variants (with a predicted signature fraction greater than 0.1 − 0.2). This analysis showed that panel-based signature calls for the APOBEC, mismatch repair, tobacco, and ultraviolet light signatures reached 90% sensitivity with at least 20 somatic variants. Owing to the low number of samples with POLE-associated and temozolomide-associated hypermutation, we did not assess the sensitivity of signature detection at each variant count threshold; we instead downsampled the number of variants in positive control samples to find the minimum number of variants necessary to reproducibly predict the known signature, which was also determined to be 20 somatic variants (Extended Data Fig. 4).

Enrichment analysis

Mutation enrichment was statistically determined through a permutation test to control for confounders including variable mutability of different genes as well as sample mutation rates, which is of particular importance when assessing enrichment in hypermutated samples. First, we generated a list of every mutation in each of our samples. We calculated the difference in the mutation counts (Δ) between the group of interest and the reference group. We then randomly permutated the mutations 100,000 times, preserving sample and gene mutation counts, and computed the Δ for each gene in each permutation. The P value for a given gene was determined by the fraction of permutations 1–n (in our case, n = 100,000) for which Δ ≥ Δ'. Storey q values were generated using the qvalue package in R to adjust for multiple comparisons. The analysis was first performed in the merged DFCI-Profile and MSKCC-IMPACT dataset, and further revalidated in the FMI dataset in an independent analysis.

Single-cell whole-genome sequencing

Frozen glioma samples were mechanically dissociated into pools of single nuclei as previously described, following which single nuclei were isolated by flow cytometry, using a DAPI-based stain. Nuclei were subjected to whole-genome multiple displacement (MDA) amplification (Qiagen, REPLI-g) followed by next-generation sequencing library construction for Illumina Sequencing (Qiagen QIAseq FX DNA library kit). Libraries were sequenced on the Illumina HiSeq platform in paired end mode. Single cells were sequenced to 0.1–1× coverage. Bulk pooled nuclei were sequenced to 60× coverage while matched germline DNA (extracted from blood) was sequenced to 30× coverage.

Reads were aligned to hg38 using bwa mem, and variants were jointly called across bulk normal tissue, primary tumour single cells, and recurrent tumour single cells using the GATK best practices pipeline without variant quality score recalibration. Somatic mutations in single cells were called if they were monomorphic, had a homoyzygous reference genotype call but no alternate-allele support in bulk normal tissue, and had at least three supporting reads in a single cell. Germline heterozygous mutations (gHets) were called if they were monomorphic, were found in dbSNP (version 138, http://www.ncbi.nlm.nih.gov/snp), and had at least one supporting read and a heterozygous genotype call in bulk normal tissue. To assess sensitivity in each single cell, we computed the fraction of gHets detected with at least three supporting reads, analogous to our procedure for calling somatic mutations. To estimate the total number of somatic mutations present in each cell, we divided the total number of somatic mutations detected by sensitivity. To obtain 95% confidence intervals on the total mutational burden, we modelled the measurement of sensitivity using a beta distribution with Jeffrey’s prior, in which the beta parameters (α, β) are equal to the number of detected gHets + 0.5 and the number of undetected gHets + 0.5, respectively. We identified recurrent tumour single cells as hypermutated if their mutational burden was at least 1.5 times the highest mutational burden detected among primary tumour cells.

The method to detect microsatellite mutations was based on read-based phasing and was previously validated using scWGS data from neurons (I.C.-C. et al., manuscript in preparation). First, the human genome was scanned to define a reference set of microsatellite repeats that can be captured using short reads (that is, between 6 and 60 bp) as previously described. Heterozygous SNPs were then detected in the bulk normal sample using the variant caller GATK. Next, the reads in a given cell mapping to each heterozygous SNP allele detected in the bulk sample and their mates were extracted. If any of the microsatellites in the reference set were covered by these reads, the distribution
of the allelic repeat lengths supported by the data was obtained by collecting the lengths of all intra-read microsatellite repeats mapped to the microsatellite locus under consideration. To discount truncated microsatellite repeats, we required 10-bp flanking sequences (both 5′ and 3′) of the intra-read microsatellite repeats to be identical to the reference genome. The same procedure was applied to the bulk sample. Finally, the distributions of microsatellite lengths from the single cell and the bulk sample were compared using the Kolmogorov–Smirnov test. The rates of microsatellite instability for each cell were computed as the number of sites mutated divided by the total number of microsatellites for which a call could be made. We applied FDR correction using 0.05 as a threshold for statistical significance, with a minimum of 8 single cell and 10 normal reads required to make a call. All the code is publicly available (https://github.com/parklab/MSIprofiler).

Immunohistochemistry
For the revalidation of MMR defects in an independent set, all prospectively collected surgical samples representing consecutive relapses of diffuse glioma following treatment with alkylating agents in adult patients (surgery between 2009 and 2015) were included. An expert neuropathologist reviewed histological samples from the IHC Pitie Salpetriere cohort (Supplementary Table 2) in order to assess the WHO 2016 integrated diagnosis and to select the tumour areas for immunohistochemistry and for DNA extraction when molecular testing from formalin-fixed paraffin-embedded (FFPE) tissue material was required. Diffuse gliomas harbouring unambiguous positive IDH1(R132H) immunostaining were classified as IDH1/2-mutant. IDH1/2 status was tested by targeted sequencing in all diffuse gliomas harbouring negative or ambiguous IDH1(R132H) immunostaining. IDH1/2-mutant diffuse gliomas with loss of ATRX expression in tumour cells were classified as non IDP1/19q co-deleted. 1p/19q co-deletion was tested in all IDH1/2-mutant diffuse gliomas with maintained ATRX expression. MGMT status was assessed in IDH1/2 wild-type gliomas. FFPE sections (3 µm thick) were deparaffinized and immunolabelled with a Ventana Benchmark XT autostainer (Roche, Basel, Switzerland). The secondary antibodies were coupled to peroxidase with dianaminobenzidine as brown chromogen. For immunohistochemistry performed at Pitie-Salpetriere (PSL) Hospital, the following antibodies were used: mouse monoclonal anti-ATRX (Bio SB, clone BSB-108, BSB3296, 1:100), mouse monoclonal anti-IDH1 (Dianova, clone H09, DIA-H09, 1:100), rabbit monoclonal anti-CD3 (Roche, clone 2G6, 790-4341, prediluted), rabbit polyclonal anti-IBAI (Wako, WI0199-19741, 1:500), mouse monoclonal anti-MLH1 (Roche, clone ML, 790-4355, prediluted), mouse monoclonal anti-MSH2 (Roche, clone G219-1129, 760-4265, prediluted), mouse monoclonal anti-MSH6 (Roche, clone 44, 760-4455, prediluted), rabbit monoclonal anti-PMS2 (Roche, clone EPR3947, 760-4331, prediluted). For immunohistochemistry performed at BWH, the following antibodies were used: mouse monoclonal anti-MLH1 (Leica, clone ES05, MLHL1-C, 1:75), mouse monoclonal anti-MSH2 (Merck Millipore, clone Ab-2-FE1, NA27, 1:200), mouse monoclonal anti-MSH6 (Leica, clone PU29, MS6-H1-C, 1:50), mouse monoclonal anti-PMS2 (Cell Marque, MRQ-28, 285M-14-AR, 1:100). An expert neuropathologist blinded to the molecular status of MMR deficiency analysed the immunostaining. If loss of expression of one or several MMR proteins was observed in tumour cells, this result was confirmed in an independent laboratory by a second expert pathologist with separate stainer and reagents: FFPE sections were immunolabelled with a BOND stainer (Leica, Wetzlar, Germany). Primary antibodies were as follows: mouse monoclonal anti-MLH1 (clone G168-728, BD Pharmingen), mouse monoclonal anti-MSH2 (clone 25D12, Diagnostic BioSystems), mouse monoclonal anti-MSH6 (clone 44, Diagnostic BioSystems), mouse monoclonal anti-PMS2 (clone A16-4, BD Pharmingen). The loss of expression of MMR proteins was defined as the total absence of nuclear labelling in tumour cells associated with a maintained expression in normal cells (as a positive internal control in the same tissue area). The density of the immune infiltrate was studied after immunolabelling of T lymphocytes by CD3 and of macrophage/microglial cells by IBA1. The number of immunopositive cells was quantified by visual counting in the three areas (one square millimetre) of tumour tissue containing the highest density of immunopositive cells and a mean density was calculated.

Patient-derived cell lines
All PDCLs with a name starting with BT were established from tumours resected at Brigham and Women’s Hospital and Boston Children’s Hospital (Boston, MA) and were maintained in neurosphere growth conditions using the NeuroCult NS-A Proliferation Kit (StemCell Technologies) supplemented with 0.0002% heparin (StemCell Technologies), EGF (20 ng/ml), and FGF (10 ng/ml; Miltenyi) in a humidified atmosphere of 5% CO2 at 37 °C. The N16-1162 PDCL was established by the Glio Tex team (Glioblastoma and Experimental Therapeutics) at the Institut du Cerveau et de la Moelle épinière (ICM) laboratory and maintained as described above. SU-DIPC-XIII (DIPC13) cells were provided by Dr. Michelle Monje at Stanford University and were maintained in neurosphere growth conditions in a humidified atmosphere of 5% CO2 at 37 °C in tumour stem medium (TSM) consisting of Dulbecco’s modified Eagle’s medium: nutrient mixture F12 (DMEM/F12), neurobasal-A medium, HEPEES buffer solution 1×, sodium pyruvate solution 100 mM, non-essential amino acids solution 10 mM, Glutamax-1 supplement and antibiotic-antimycotic solution (Thermo Fisher). The medium was supplemented with B-27 supplement minus vitamin A (Thermo Fisher), 20 ng/ml human-EGF (Miltenyi), 20 ng/ml human-FGF-basic (Miltenyi), 20 ng/ml human-PDGF-AA, 20 ng/ml human-PDGF-BB (Shenandoah Biotech) and 2 µg/ml heparin solution (0.2%, Stem Cell Technologies). The identity of all cell lines established was confirmed by short tandem repeat assay or sequencing. All cell lines were tested for the absence of mycoplasma. Cell lines, xenographs, and model data available from the DFCI Center for Patient Derived Models.

Viability assays
For short-term viability assays, cells were plated in 96-well plates and treated the following day with temozolomide (Selleckchem) or CCNU (Selleckchem) for 7–9 days incubation. Fresh medium was added after four days of incubation. Cell viability was assessed using the luminescent Cell Titer-Glo reagent (Promega) according to the manufacturer’s protocol. Luminescence was measured using the Modulus Microplate Reader (Promega). The surviving fraction (SF) for each [X] concentration was calculated as SF = mean viability in treated sample at concentration [X]/mean viability of untreated samples (vehicle). Dose–response curves and IC50 were generated using Prism 8 (GraphPad Software, San Diego, USA) after log transformation of the concentrations. Curves were extrapolated using nonlinear regression with four-parameter logistic regression fitting on triplicates from survival fractions of three independent replicates, following the model: y = Bottom + (Top – Bottom) / (1 + 10( logIC50 – x ) × Hillslope).

Generation of Isogenic MMR-deficient cell lines
Oligos of the form 5′-CAGCC[N20] (where [N20] is the 20-nucleotide target sequence; sgGFp, GAGCTGGACGGCGACGTAA; sgMSH2, ATTCTGTTCTTATCCATGA; sgMSH6, TTATTTGAGTCTAGTACGTC; sgMLH1, ACTACCCAAATGCTCATAC; sgPMS2, TCATCTGACGAGGATGT) and 5′-AAC[rc20]C (where [rc20] is the reverse complement of [N20]) were purchased from Integrated DNA Technologies (IDT). For DIPC13 cells, oligos containing the sgRNA target sequence were annealed with their respective reverse complement and cloned into the lentivector plasmid pXPR_BRD001; now available as pXPR_BRD023 lentICRiSPRv2 from the Broad Institute Gene Perturbation Platform (GPP). For BT145 cells, oligos containing the sgRNA target sequence were annealed with their respective reverse complement and cloned into the pXPR_BRD051
CRISPRko all-in-one sgRNA/Cas9-delivery lentiviral expression vector (available from the Broad Institute Genetics Perturbation Platform, GPP). Successful cloning of each sgRNA target sequence was confirmed via Sanger Sequencing. To generate lentiviruses from these vectors, HEK293T cells were transfected with 10 μg of each expression plasmid with packaging plasmids encoding PSPAX2 and VSVG using lipofectamine. Lentivirus-containing supernatant was collected 48 and 72 h after transfection. DIPG3 and BT145 cells were seeded in a 12-well plate at 1 × 104 cells/well in 3 ml medium and spin-infected (2,000 rpm for 2 h at 30 °C with no polybrene) with pLX311-Cas9 (DIPG13) or pXPR_BRD051 (BT145) lentiviral vectors and selected with blastidicin (10 μg/ml, DIPG13) or hygromycin (300 μg/ml, BT145) to generate Cas9-expressing or knockout cells. DIPG13-Cas9 cells underwent a subsequent lentiviral spin-infection with the lentCRISPR sgGFP, sgMsh2, or sgMsh6 vectors described above. Puromycin selection (0.4 μg/ml for DIPG13 cells) commenced 48 h post-infection.

Chronic treatment and sequencing of isogenic MMR-deficient cell lines
DIPG13-sgGFP, -sgMsh2, and -sgMsh6 cells were seeded at 8 × 104 cells/well in 4 ml medium in a 6-well ULA plate. Each line was grown for 3 months under 3 conditions: no treatment, temozolomide (100 μM, Selleckchem), or DMSO vehicle. Cells were grown under these conditions in the absence of both blastidicin and puromycin. Cells were re-dosed with temozolomide or DMSO every 3–5 days, splitting over-confluent cells 1:2 or 1:4 as needed. After 3 months, genomic DNA was extracted using the QiAmp DNA Mini Kit. DNA was subjected to whole-exome Illumina sequencing. Reads were aligned to the Human Genome Reference Consortium build 38 (GRCh38). WES data were analysed using the Gertz Lab CGA whole-exome sequencing characterization pipeline (https://docs.google.com/document/d/1V02Kx_ggUd03mB59nljULWGZu794WbTep6B3c9g08/edit#heading=h.yby87l2ztbcj) developed at the Broad Institute which uses the following tools for quality control, calling, filtering and annotation of somatic mutations and copy number variation: Picard Tools (http://broadinstitute.github.io/picard/), ContEst38, MuTect39, Strelka39, Orientation Bias Filter39, DeTN40, AllelicCapSeg41, MAFPoNFilter41, RealignmentFilter, ABSOLUTE44, GATK45, Variant Effect Predictor46, and Oncotator47.

Subcutaneous xenografts and drug treatment
BT145 cells (2 × 105) were reseeded in equal parts Hank’s buffered salt solution (Life Technologies) and Matrigel (BD Biosciences) and then injected into both flanks of eight-week-old NU/NU male mice (Charles River). Tumour-bearing mice (n = 8) were randomly assigned to the treatment or vehicle arm when tumours measured a volume of 100 mm3. Animals received 12 mg/kg/day temozolomide or vehicle (Ora-Plus oral suspension solution, Perrigo, Balcatta, Australia) by oral gavage for 5 consecutive days per 28-day cycle. An additional 4 weeks resting period without treatment was observed before the second cycle. Tumour volumes were calculated using the formula: 0.5 × length × width2. Body weights were monitored twice weekly. The investigators were not blinded to allocation during experiments and outcome assessment. Mice were euthanized when they showed signs of tumour-related illness or before reaching the maximum tumour burden. Tumours were subsequently removed, and a subset were submitted to Oncopanel sequencing for analyses of exonic mutations (POPv3, 447 genes) and mutational signature as defined above. To separate human and mouse sequenced reads in the DNA sequencing data generated for the PDX models, the ‘raw’ data were mapped to both the hg19 human and mm10 mouse reference genomes using BWA-MEM-0.7.17. The output of the alignment was name sorted by Samtools. We then used the software package Disambiguate (ngs_disambiguate-1.0) to assign each read to the human or mouse genome and to produce final alignment files in BAM format. Final hg19 BAM files were coordinate sorted by Samtools-1.7. Duplicate reads were marked and removed from the BAM files using Picard-2.0.1. GATK4.1.0.0 base recalibration was performed using BestRecalibration and Applying Recalibration followed by CollectFIR2Counts and LearnReadOrientationModel to create a model for read orientation bias. Variant calling was performed using GATK-4.1.0.0/Mutect2 pipeline with the default parameters and filters except for the following modifications: (i) ‘af-of-alleles-not-in-resource’ was set to 0; (ii) ‘MateOnSameContigOrNoMappedMateFilter’ was disabled; (iii) the output of Step8 was used for the read orientation model; and (iv) a germline resource from the gnomAD database was included (https://console.cloud.google.com/storage/browser/details/gatk-best-practices/somatic-b37/af-only-gnomad.raw.sites.vcf). The capture targets intervals used for Mutect2 were POPv3. The generated variant calls were further filtered using the FilterMuteCtCalls module of GATK4.1.0.0 and the final output in VCF format was annotated with Ensemble Variant Effect Predictor (ensembl-vep-96.0) using vcf2maf-1.6.16. The calls were additionally annotated with the OncoKB dataset using oncokb-annotator and sorted as MAF files.

All in vivo studies were performed in accordance with Dana-Farber Cancer Institute animal facility regulations and policies under protocol number 09-016.

Immunoblotting
Proteins were extracted in lysis RIPPA buffer (50 mM Tris, 150 mM NaCl, 5 mM EDTA, 0.5% sodium deoxycholic acid, 0.5% NP-40, 0.1% SDS) supplemented with protease inhibitor cocktail (Roche Molecular). Proteins were quantified using the PierceBCA Protein Assay Kit, according to the manufacturer’s protocol. Samples were then prepared with 1× NuPAGE (Invitrogen) LDS sample buffer, and NuPAGE (Invitrogen) sample reducing agent followed by heating to 95 °C for 5 min. The samples were then loaded onto NuPAGE 4–12% Bis-Tris Gel (Invitrogen) with NuPAGE MOPS SDS (Invitrogen) buffer and run through electrophoresis. The transfer onto membrane was then done at 40 V overnight at 4 °C in NuPAGE transfer buffer (Invitrogen) with 10% methanol. Membranes were blocked with 5% skim milk in TBST for 1 h, then incubated with the following primary antibodies added to 5% BSA and incubated overnight at 4 °C on a shaker: mouse monoclonal anti-MGMT (Millipore, MT3.1, MAB6200, 1:500), mouse monoclonal anti-MSH2 (Calbiochem, FE11, NA27, 1:1,000), mouse monoclonal anti-MSH6 (Biosciences, 44, 610918, 1:500), mouse monoclonal anti-MLH1 (Cell Signaling, 4C9C7, 3515, 1:500), mouse monoclonal anti-PMS2 (BD Biosciences, A16-4, 556415, 1:1,000), mouse monoclonal anti-beta-actin (Sigma, AC-74, A2228, 1:10,000). After several cycles of washing and incubation with secondary goat anti-mouse antibody (Invitrogen 31430, LI:10,000), membranes were imaged by chemiluminescence using the Biorad ChemidocTM MP imaging system.

Microsatellite instability analysis
PCR amplification of the five mononucleotide markers (BAT25, BAT26, NR21, NR24, MONO27) was performed with the MSI Analysis System kit (Version 1.2, Promega). PCR products were analysed by an electrophoretic separation on the polymer POP7 50cm in an Applied Biosystems 3130XL sequencer and using Genemapper Software 5.

Outcome of patients treated with PD-1 blockade
For comparison of PFS and OS in patients treated with PD-1 pathway blockade according to TMB and MMR statuses, we retrospectively identified patients with glioma who had been treated with PD-1 blockade (alone or in combination with bevacizumab) for recurrent disease at our institutions. Patients for whom sequencing was not performed at the time of recurrence were excluded. Magnetic resonance imaging (MRI) tumour assessments were reviewed using the Response Assessment in Neuro-Oncology (RANO) criteria by three independent reviewers (M.J.L.-F., S.A., and R.Y.H.) who were blinded to the groups. PFS and OS duration were calculated from cycle 1 day 1 of PD-1 blockade therapy.
Statistical analyses

Data were summarized as frequencies and proportions for categorical variables and as median and range for continuous variables. Continuous variables were compared using Mann–Whitney or Kruskal–Wallis tests; categorical variables were compared using Fisher’s exact or Chi-squared tests. Survival and PFS were estimated using the Kaplan–Meier method, and differences in survival or PFS between groups were evaluated by the log-rank test. Survival for subjects who were alive or lost to follow-up at the time of last contact or on data cutoff was censored at the date of the last contact. Patient matching in a A-to-k fashion was conducted using coarsened exact matching according to diagnosis, primary versus recurrent status, and prior treatments. For evaluation of response to PD-1 blockade, patients with glioma from the DFCI-Profile cohort who were treated with anti-PD(L)-1 antibodies or other treatments (total n = 210) as part of their management were included in the analysis. For multivariable analysis, Cox proportional hazard regression was used to investigate the variables that affect survival. P-values were considered statistically significant when <0.05. Statistical analyses were performed using STATA (v14.2, StataCorp LLC, College Station, USA), Prism 8 (GraphPad Software, San Diego, USA), and MedCalc Statistical Software, version 19.0.3 (MedCalc Software bvba, Ostend, Belgium). For enrichment analyses, mutated genes were considered significant when Q < 0.01. Where applicable, the means of population averages from multiple independent experiments (s.d. or s.e.m.) are indicated. No statistical methods were used to pre-
determine sample size.

Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this paper.

Data availability

Clinical and sequencing data from 1,493 samples from the DFCI-Profile and 544 samples from the MSKCC-IMPACT datasets are publicly available (GENIE v6.1://https://genie.cbiportal.org or https://www.synapse.org/). All data for samples from the GENIE v6.1 and TCGA pan-cancer datasets are publicly available. Data for samples from the FMI dataset are not publicly available, but de-identified, aggregated data can be accessed on request. dbGaP Study Accession: phs001967.v1.p1. All other data are available on request.

Code availability

The code for the detection of microsatellite mutations in single-cell DNA sequencing is publicly available (https://github.com/parklab/MSIprofi ler).
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**Additional information**

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**Extended Data Fig. 1** Overview of the clinical characteristics of the patients in the study and analyses performed. 

**a** | Clinical datasets analysed and main demographics including age, histomolecular subtype and disease stage. 1,628 glioma samples from adult and paediatric patients were sequenced as part of a large institutional prospective sequencing program of consented patients (DFCI-Profile) and subsequently clinically annotated. We identified 545 and 8,121 gliomas with sequencing from the MSKCC-IMPACT and FMI datasets, respectively, and used them as a replication set (total set of 10,294 sequenced samples). In addition, 314 tumours—including 247 consecutive recurrent gliomas—were analysed for protein expression of four MMR proteins (MSH2, MSH6, MLH1, and PMS2) using immunohistochemistry. 

**b** | Analyses performed and key clinical questions that were addressed in the study.
Extended Data Fig. 2 | Distributions of TMB, homopolymer indels, and SNV mutation spectra in the datasets used. a, DFCI-Profile (de novo gliomas only); b, MSKCC-IMPACT; c, FMI (total n = 9,938). After examining the distribution of TMB in each dataset for breakpoints, thresholds for hypermutation were further confirmed using segmented linear regression analysis (analysis restricted to de novo gliomas for DFCI-Profile). This method showed the presence of a breakpoint at 17.0 and 8.7 mutations per Mb for the DFCI-Profile and FMI datasets, respectively. For the MSKCC-IMPACT dataset, the cutoff used for hypermutation (13.8 mutations per Mb) was previously determined17. The frequency of hypermutation was similar in the three datasets (85 (5.2%) in DFCI-Profile; 29 (5.3%) in MSKCC-IMPACT; 444 (5.5%) in FMI). The median tumour mutation burden (TMB) in the combined datasets was 2.6 mutations per Mb (range, 0.0–781.3). Compared with non-hypermutated gliomas, hypermutated tumours showed atypical patterns of SNVs, consistent with abnormal mutational processes operating in these samples. Bars represent median and interquartile range for each dataset (right). HPI, homopolymer indels.
Extended Data Fig. 3 | See next page for caption.
Extended Data Fig. 3 | Integrated analysis of tumour mutation burden in hypermutated gliomas in the DFCI-Profile, MSKCC-IMPACT and FMI datasets. a, Distribution of TMB, homopolymer indels, MMR mutations, and SNV mutational spectrum according to molecular status of IDH1/2, 1p/19q co-deletion (1p/19q), gain of chromosome 7 and/or deletion of chromosome 10 (7gain/10del), and MGMT promoter methylation, histological grade, age at initial diagnosis, and history of prior treatment with alkylating agents or radiation therapy (the distinction between photon and proton therapy was not systematically captured) in the DFCI-Profile dataset (n = 84, data not shown for the single sample from other gliomas, IDH1/2-wt subgroup). b, Top, distribution of histomolecular groups in non-hypermutated and hypermutated gliomas from the combined sequencing dataset (n = 2,173). Bottom, distribution of molecular groups in de novo and post-treatment hypermutated gliomas from the DFCI-Profile dataset (n = 85) (annotation not available for the MSKCC-IMPACT set). c, Prevalence of hypermutation according to MGMT promoter methylation and IDH1/2 mutation status in post-temozolomide gliomas from the DFCI-Profile dataset (n = 150). Two-sided Fisher’s exact test. d, Number of temozolomide cycles according to IDH1/2 mutation status in post-temozolomide diffuse gliomas from the DFCI-Profile dataset (n = 211 gliomas). Patients who received combined chemoradiation but no adjuvant temozolomide were included. Two-sided Wilcoxon rank-sum test. e, Boxplots of TMB in post-treatment hypermutated gliomas according to the number of temozolomide cycles received before surgery. Kruskal–Wallis test and Dunn’s multiple comparison test. F, TMB in recurrent gliomas according to treatments received before surgery. Patients who received multiple treatment modalities were excluded. Kruskal–Wallis test and Dunn’s multiple comparison test. Boxes, quartiles; centre lines, median ratio for each group; whiskers, absolute range (d–f). g, Integrated analysis of the FMI dataset (n = 8,121 gliomas) depicting tumour mutation burden, the number of indels at homopolymer regions, and the SNV mutation spectrum detected in each tumour according to molecular status of IDH1/2 and 1p/19q co-deletion (1p/19q), MSI status, and age at initial diagnosis. Dominant mutational signatures detected in hypermutated samples are depicted. The dotted line indicates the threshold for samples with a high mutation burden (8.7 mutations per Mb). h, Prevalence of hypermutation among molecularly defined subgroups in the FMI dataset (n = 8,121 gliomas). Chi-squared test. i, Dominant mutational signatures detected in hypermutated samples in the FMI dataset (n = 8,121 gliomas). Chi-squared test. j, Mutated genes and pathways enriched in hypermutated gliomas in the FMI dataset (n = 8,121). Enrichment was assessed using a permutation test to control for random effects of hypermutability in tumours with high TMB. k, l, Proportion of TMB-high versus TMB-low samples with mutations in selected DNA repair genes and glioma drivers (e) and in the MMR pathway (MSH2, MSH6, MLH1 and PMS2; f). Permutation test; ****P < 0.0001, ***P < 0.001, **P < 0.01; ns, not significant.
Extended Data Fig. 4 | See next page for caption.
Extended Data Fig. 4 | Validation of known hypermutation-associated signatures using TCGA datasets. Mutational signatures were predicted using exome-sequencing variants that overlapped with the panel-targeted regions, and then compared to previously published DeconstructSigs signature predictions based on all exome variants. The TCGA MC3 dataset was used to assess the detection of COSMIC mutational signatures associated with APOBEC (signatures 2 and 13), mismatch repair (signature 6), ultraviolet light (signature 7), POLE (signature 10), and tobacco (signature 4). Variant calls for 17 hypermutated and 12 non-hypermutated glioma exome-sequenced samples were used to assess temozolomide (signature 11) detection.

**a**, Detection of APOBEC-associated mutational signature in TCGA BLCA samples (n = 129 out of 411 samples).

**b**, Detection of ultraviolet-associated mutational signature in TCGA SKCM samples (n = 237 out of 466 samples).

**c**, Detection of tobacco smoking-associated mutational signature in TCGA LUAD samples (n = 250 out of 513 samples).

**d**, Detection of MMR-associated mutational signature in TCGA COAD (n = 188 out of 380 samples).

**e**, Detection of POLE-associated mutational signature in TCGA COAD and READ samples (n = 277 out of 380 samples).

**f**, Detection of temozolomide-associated mutational signature in ref. 4 (n = 29).

**g**, Unsupervised clustering of hypermutated samples. A total of 865 hypermutated tumour samples from exomes (pan-TCGA and Wang et al.4) and targeted panels (DFCI-Profile and MSK-IMPACT) were analysed for known hypermutation signatures (tobacco, UV, MMRD, POLE, TMZ, APOBEC). Samples and signatures underwent 2D hierarchical clustering based on Euclidean distance.

**h**, Performance of cancer panel versus other genesets in mutational signature calling. We analysed 622 hypermutated tumour exomes (pan-TCGA and Wang et al.4, black) for their mutational signature contributions when restricted to variants from i) DFCI-Profile OncoPanel cancer panel genes (red), or ii) 9 randomly selected gene sets (grey) of similar total capture size to the cancer panel. For each sample, we assessed known hypermutation signatures for cancer panels and gene sets for which at least 20 single base substitutions were retained in the sample after restriction. Samples and signatures underwent 2D hierarchical clustering based on Euclidean distance.

**i**,The violin plots represent the number of variants (top) and the cosine similarity of signature contributions (bottom) when using all exonic variants versus restriction to cancer panel or the 49 random gene sets. Boxes, quartiles; centre lines, median ratio for each group; whiskers, absolute range. Two-sided Welch’s t-test.
Extended Data Fig. 5 | See next page for caption.
Extended Data Fig. 5 | Mutational signature analysis of primary and secondary hypermutated cohort (n = 111). a, Mutational signature analysis of newly diagnosed hypermutated gliomas in the DFCI-Profile dataset (n = 24). b, Mutational signature analysis of secondary hypermutated gliomas (samples in which hypermutation was detected in the recurrent tumour) in the DFCI-Profile dataset (n = 58). The novel COSMIC Signature 11-related signature (S2) was associated with 1p/19q co-deletion and lack of prior radiation therapy (66.7% of samples with high S2 versus 26.2% of samples with high S1 signature, Fisher P = 0.016). c, Mutational signature analysis of hypermutated gliomas from the MSKCC-IMPACT dataset (n = 29). d, Mutational signature analysis in de novo (hypermutated at first diagnosis, n = 26, left) and post-treatment hypermutated gliomas (hypermutation in a recurrent tumour, n = 59, right). Percentage of samples exhibiting the most common mutational signatures and their hypothesized causes are displayed. MMR, C6, C14, C15, C26; age-related, C1; POLE, C10, C14. Chi-squared test. e, Mutational signatures identified in individual de novo hypermutated gliomas (hypermutated at first diagnosis, n = 26, left) and post-treatment hypermutated gliomas (hypermutation in a recurrent tumour, n = 59, right). f, Mutational signature analysis of MMR variants in hypermutated gliomas from the DFCI-Profile and MSKCC-IMPACT datasets (n = 114). Ninety variants of the MMR genes MSH2, MSH6, MLH1, and PMS2 were merged into two groups (de novo, n = 18; post-treatment, n = 72) according to the type of sample in which they were found and analysed for mutational signatures using a regression model (Rosenthal et al.52). In each sample, only the MMR variant with the highest VAF was included, to limit the inclusion of possible passenger variants. For signature discovery in both cohorts (a–e), variants were analysed using the non-negative matrix factorization (NMF) method and correlated with known COSMIC mutational signatures using Pearson correlation.
Extended Data Fig. 6 | See next page for caption.
Extended Data Fig. 6 | Characteristics of MMR molecular variants in hypermutated gliomas. a, Proportion of TMB$^{\text{H}}$ versus TMB$^{\text{L}}$ samples with mutations in selected DNA repair genes and glioma drivers (a) and in the MMR pathway (MSH2, MSH6, MLH1 and PMS2) (b) in the merged DFCI-Profile/MSKCC-IMPACT dataset ($n = 2,173$). Permutation test; $^{****}P < 10^{-5}$, $^{**}P < 10^{-2}$, $^{*}P < 0.05$. c, CCFs of MMR gene mutations in post-treatment hypermutated gliomas versus other hypermutated cancers in the FMI dataset. Horizontal line, median. Two-sided Wilcoxon rank-sum test with Benjamini–Hochberg correction. d, VAF distribution of mutations in post-treatment hypermutated gliomas, non-glioma MMR-deficient cancers (diverse histologies) and other non-glioma hypermutated samples (diverse histologies) from the TCGA and MSKCC-IMPACT datasets. Each dot represents a mutation found in an individual sample (represented vertically). MMR mutations are depicted in red. e, Integrated view of mutational signatures and MMR gene mutations and protein expression in hypermutated gliomas ($n = 114$). Tumours with the mutational hotspot MSH6(T1219I) (11.9% of post-treatment hypermutated gliomas) are highlighted. f, Mutation diagram of MSH2, MSH6, MLH1, and PMS2 mutations found in hypermutated gliomas from the DFCI-Profile and MSKCC-IMPACT datasets ($n = 114$). The hotspot MSH6 missense variant p.T1219I was found in nine samples. g, Hotspot MSH6 p.T1219I variant mapped to the bacterial MutS 3D structure (PDB 5YK4). h, Representative immunohistochemistry (IHC) images of the MMR proteins MSH2, MSH6, MLH1 and PMS2 in a hypermutated glioblastoma with MSH6(T1219I) mutation. Three independent samples were stained. Scale bar, 100 µm.
Extended Data Fig. 7 | See next page for caption.
Extended Data Fig. 7 | Results of MMR IHC screening in 213 consecutive recurrent gliomas and patterns of MMR protein expression loss in three de novo or post-treatment MMR-deficient gliomas. a, Recurrent patterns of MMR protein loss identified by IHC in gliomas. Scale bar, 50 µm. b, Summary of MMR IHC screening results for 213 consecutive recurrent gliomas. All monocentric consecutive relapses of diffuse gliomas in adult patients following treatment with post-alkylating agents (surgery between 2009 and 2015) were included in the immunohistochemistry analysis. Further sequencing of samples in which MMR protein loss was identified showed hypermutation with MMR molecular defects in 18/19 (94.7%) samples. c, Percentage of tumour MMR protein loss in glioma samples with de novo (n = 16) or post-treatment (n = 46) MMR deficiency. Samples were scored by two pathologists in blinded fashion. Regional heterogeneity of MMR protein loss for the four MMR proteins MSH2, MSH6, MLH1, and PMS2 was scored as to the maximal percentage of protein loss among tumour cells for each sample (5% increments). Boxes, quartiles; centre lines, median ratio for each group; whiskers, absolute range, excluding outliers. Two-sided Wilcoxon rank-sum test. d, Clonal MMR deficiency in a de novo high-grade glioma. Top left, low magnification of haematoxylin and eosin (H&E) staining of the large surgical tumour pieces obtained from surgical resection. Right, high magnification images of H&E staining, showing highly cellular tumour and an astrocytic phenotype, and PMS2 IHC, showing loss of expression of PMS2 in all tumour cells (open arrowheads). Normal cells have maintained PMS2 expression (internal control, solid arrowheads). Bottom left, Microsatellite testing via PCR amplification of five mononucleotide markers (BAT25, BAT26, NR21, NR24, and MONO27) showed the tumour to be MSS. NGS showed a TMB of 120.1 per Mb and homopolymer indel burden of 3.8 per Mb, with contributions from temozolomide (90%) and MMR-deficiency (10%) mutational signatures. A missense (p.P648L) hotspot MLH1 mutation known to be pathogenic from patients with Lynch syndrome with a VAF of 0.73 and loss of heterozygosity was present in this case. Scale bars, top left, 5 mm; right 100 µm. e, Clonal MMR deficiency in a hypermutated post-treatment IDH1-mutant glioblastoma. Top left, low magnification image of H&E staining of tissue obtained from surgical resection, with three areas of tumour selected for images. Red dashed line delimits normal brain. Right, high-magnification images of H&E staining, showing highly cellular tumour and an astrocytic phenotype, and PMS2 IHC, showing loss of expression of PMS2 in all tumour cells (open arrowheads). Normal cells have maintained PMS2 expression (internal control, solid arrowheads). Bottom left, Microsatellite testing via PCR amplification of five mononucleotide markers (BAT25, BAT26, NR21, NR24, and MONO27) showed the tumour to be MSS. NGS showed a TMB of 236.5 per Mb and homopolymer indel burden of 2.3 per Mb, with 95% contribution of temozolomide mutational signature. Scale bars, top left 5 mm; right 100 µm.
Extended Data Fig. 8 | See next page for caption.
Extended Data Fig. 8 | Characterization of high-grade glioma PDCLs and their sensitivity to temozolomide and CCNU.

a, Clinico-molecular characteristics of four native newly diagnosed or recurrent glioma PDCL models harbouring hypermutation and MMR deficiency. B, Thirty glioma PDCLs, including four PDCLs derived from patients with de novo (BT1160, N16-1162, both established from patients with Lynch syndrome) or post-treatment (BT237, BT559) MMR deficiency were molecularly characterized using whole-exome sequencing. The panels show the tumour mutational burden (left) and homopolymer indel burden (right) in each model. Boxes, quartiles; centre lines, median ratio for each group; whiskers, absolute range. Two-sided Wilcoxon rank-sum test.

b, A panel of 12 glioma PDCL models representing the different MGMT and MMR classes was selected and assessed for sensitivity to temozolomide in a short-term viability assay (e; dots represent means). The temozolomide AUC was compared between the three groups using a Kruskal–Wallis test and Dunn’s multiple comparison test (f; mean ± s.d.).

c, Mutational signature analysis was performed in the PDCL models of constitutional and post-treatment MMR deficiency using the R package DeconstructSigs to estimate the contributions of mutational signatures using a regression model (Rosenthal et al. 52). For each PDCL, the contribution of the main COSMIC mutational signatures identified is expressed as decimal. d, Boxplots of temozolomide AUC in non-hypermutated versus hypermutated PDCLs. Boxes, quartiles; centre lines, median ratio for each group; whiskers, absolute range. Two-sided Wilcoxon rank-sum test.

e, f, A panel of 12 glioma PDCL models representing the different MGMT and MMR classes was selected and assessed for sensitivity to temozolomide in a short-term viability assay (e; dots represent means). The temozolomide AUC was compared between the three groups using a Kruskal–Wallis test and Dunn’s multiple comparison test (f; mean ± s.d.).

f, Western blot of the glioblastoma patient-derived cell line (BT145) in which the genes MSH2, MSH6, MLH1 or PMS2 have been knocked-out using the CRISPR–Cas9 system. h, i, A panel of 11 glioma PDCL models representing the different MGMT and MMR classes was selected and assessed for sensitivity to CCNU in a short-term viability assay (h; dots represent means). No CCNU data was available for the model BT172. The CCNU AUC was compared between the three groups using a Kruskal–Wallis test and Dunn’s multiple comparison test (i; mean ± s.d.).
Extended Data Fig. 9 | See next page for caption.
Extended Data Fig. 9 | MMR-deficient models of glioma, continued.

a, b, CRISPR–Cas9 MSH2 and MSH6 gene knockout in DIPG13 high-grade glioma cell line. a, Integrated genomics viewer (IGV) plots depicting MSH2 reads in between the guide RNAs in the MSH2 unedited line (sgGFP, left) and the MSH2 CRISPR knockout line (right) confirming knockout in the MSH2 edited line. b, IGV plots depicting MSH6 reads in between the guide RNAs in the MSH6 unedited line (sgGFP, left) and the MSH6 CRISPR knockout line (right) confirming knockout in the MSH6 edited line.

c, Overview of in vivo temozolomide resistance study. Treatment of subcutaneous BT145 PDX-bearing animals was initiated at a volume of 100 mm³ and eight nude mice per group were randomized to 12 mg/kg/day temozolomide or vehicle for five consecutive days per 28-day cycle. Mice were treated until tumours reached a volume of 1,500 mm³, and tumours were sequenced to identify mutations and mutational signature.

d, Survival of mice with BT145 xenografts (n = 8 mice per group) during treatment with vehicle (blue) or temozolomide (red). Two-sided log-rank test.

e, Unique variants found in three sequenced BT145 tumours (two temozolomide-treated, and one vehicle-treated) were analysed for correlation with known mutational signatures. COSMIC Signature 11 was found in the two temozolomide-treated tumours. Mutational signatures could not be called in the vehicle-treated tumour (too few variants). After filtering of truncal variants common to all tumours, the two temozolomide-treated tumours shared only four variants, including an MSH6(T1219I) mutation and three noncoding variants.

f, BT145 xenografts chronically treated with vehicle (n = 1) or temozolomide (n = 2) were removed, dissociated and cultured in serum-free medium to establish cell lines. After three passages in culture, sensitivity to temozolomide was assessed. The results of the short-term viability assays (mean ± s.e.m.) and temozolomide AUC of each cell line are depicted.

g, Model of acquired hypermutation with mutational signature 11 in gliomas. Top, MMR-proficient cells repair TMZ damage and do not develop signature 11. Resistance in these cells is mediated by non-MMR pathways (for example, MGMT expression). Bottom, TMZ induces and/or selects resistant subclonal MMR-deficient cells. Further TMZ exposure produces accumulation of mutations at specific trinucleotide contexts, detected as hypermutation with signature 11.
Extended Data Fig. 10 | See next page for caption.
Extended Data Fig. 10 | Extended outcome data. 

**a**–**c**, Survival of patients with recurrent high-grade glioma (WHO grade III or IV) from the time of initial diagnosis according to TMB status (solid curves, TMB<sup>low</sup>; dotted curves, TMB<sup>high</sup>). The curves include 240 recurrent samples from DFCI-Profile with available survival data from initial diagnosis. Two-sided log-rank test. 

**a**, Survival of patients with recurrent high-grade 1p/19q co-deleted oligodendroglioma from the time of initial diagnosis. 

**b**, Survival of patients with recurrent high-grade IDH1/2-mutant astrocytoma from the time of initial diagnosis. 

**c**, Survival of patients with recurrent IDH1/2 wild-type glioblastoma from the time of initial diagnosis. 

**d**, PFS of 11 patients with hypermutated and MMR-deficient glioma who were treated with PD-1 blockade (red curves). A cohort of hypermutated patients treated with other systemic agents is depicted as control (best matches according to diagnosis, primary vs recurrent status, and prior treatments were selected from the cohort of sequenced gliomas, purple curves). Two-sided log-rank test. Clinical and histomolecular characteristics of patients from both cohorts are provided in Supplementary Table 7. 

**e**, **f**, PFS (**e**) and OS (**f**) of 11 patients with hypermutated and MMR-deficient glioblastoma (pembrolizumab) in a patient with post-treatment hypermutated MMR-deficient glioblastoma. A cohort of patients with non-hypermutated glioma who were treated with PD-1 blockade is depicted as control (n = 10, best matches according to diagnosis, primary versus recurrent status, and prior treatments, blue curve). A two-sided log-rank test is used. 

**g**, Lack of immune response following PD1 blockade (pembrolizumab) in a patient with post-treatment hypermutated MMR-deficient glioblastoma. Top, timeline; middle, MRI images; bottom, H&E images and IHC for PMS2 expression and tumour infiltration with CD3-positive T cells and IBA1-positive macrophages in the primary (S1), recurrent pre-pembrolizumab (S3) and recurrent post-pembrolizumab (S4) tumours. The tumour acquired a focal PMS2 two-copy deletion, protein loss, and hypermutation in the post temozolomide recurrent tumour (S3). Scale bar, 50 µm.
Extended Data Fig. 11 | See next page for caption.
Extended Data Fig. 11 | Molecular characteristics of hypermutated gliomas. a, Pan-cancer analysis of TMB and homopolymer indel burden in the GENIE dataset (n = 44,389). Tumour samples from the GENIE dataset (v6.1) were analysed for mutational and homopolymer indel burden. Statistical comparisons between groups are provided in Supplementary Table 6. b, TMB in hypermutated gliomas (post-treatment) versus MMR-deficient cancers and other hypermutated cancers from the TCGA and Wang et al. exome datasets (n = 798). Two-sided Wilcoxon rank-sum test with Bonferroni correction. c, Pan-cancer analysis of the homopolymer indel burden in hypermutated gliomas (post-treatment) versus MMR-deficient cancers and other hypermutated cancers from the TCGA and Wang et al. exome datasets (n = 798). d, Results of MSI analysis using the standard pentaplex assay in glioma (n = 39) and CRC samples (n = 19) according to MMR status (MMR-d, MMR deficient; MMR-p, MMR-proficient). e, Pan-cancer analysis of cancer cell fractions in hypermutated gliomas (post-treatment) versus MMR-deficient cancers and other hypermutated cancers from the TCGA and Wang et al. exome datasets (n = 798). Two-sided Wilcoxon rank-sum test with Bonferroni correction. f, Weighted TMB in hypermutated gliomas (post-treatment) versus MMR-deficient cancers and other hypermutated cancers from the TCGA and Wang et al. exome datasets (n = 798). The weighted TMB was calculated by weighing each individual mutation to its cancer cell fraction. Two-sided Wilcoxon rank-sum test with Bonferroni correction. g, Distribution of VAFs (left) and mutation spectrum analysis of low-allelic frequency variants (<0.1, right) in TMBhigh gliomas (n = 1,543, top), de novo hypermutated gliomas with MMR deficiency mutational signature (n = 12, middle), and post-treatment hypermutated gliomas (n = 59, bottom) from the DFCI-Profile dataset. h, Distribution of VAFs (left) and mutation signature analysis of low-allelic frequency variants (<0.1, right) in TMBhigh CRCs (n = 1,265, top) and TMBlow CRCs with MMR deficiency mutational signature (n = 110, bottom) from the GENIE dataset. i, Clinical timeline for the patient with hypermutated glioblastoma with an MSH6(T1219I) mutation in whom bulk and single-cell WGS was performed. j, Distribution of VAFs of mutations in the recurrent bulk sample. The median VAF in the recurrent sample was 0.11. The MSH6(T1219I) mutation had the 18th-highest VAF out of 4,350 coding mutations. k, Cancer cell fractions (CCFs) of mutations in the primary and recurrent tumour bulk samples. Each dot represents a coding mutation. The horizontal and vertical axes are estimated clonal frequency for each mutation in the primary and recurrent samples, respectively. Mutations of the four main MMR genes are depicted in red. l, Mutational spectra in 35 cells from the primary tumour (orange) and 28 from the recurrent tumour (green) submitted to scWGS sequencing (1×). Mutational signature analysis showed a strong contribution of mutational signature II in hypermutated cells from the recurrent tumour.
Reporting Summary

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Statistics

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

- The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement
- A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
- The statistical test(s) used AND whether they are one- or two-sided
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- For null hypothesis testing, the test statistic (e.g. F, t, r) with confidence intervals, effect sizes, degrees of freedom and P value noted
  Give P values as exact values whenever suitable.
- For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
- For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
- Estimates of effect sizes (e.g. Cohen’s d, Pearson’s r), indicating how they were calculated

Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection No software used for data collection

Data analysis

For WES analyses, we used the CGA WES Characterization pipeline developed at the Broad Institute to call, filter and annotate somatic mutations and copy number variation. The pipeline employs the following tools: MuTect[1], ContEst[2], Strelka[3], Orientation Bias Filter[4], DeTiN[5], AllelicCapSeg[6], MAPPoNFilter[7], RealignmentFilter, ABSOLUTE[8], GATK[9], PicardTools[10], Variant Effect Predictor[11], Oncotator[12].

1. MuTect1: Cibulskis, K, Lawrence, MS & Carter, SL. Sensitive detection of somatic point mutations in impure and heterogeneous cancer samples. Nature ... (2013). doi:10.1038/nbt.2514
2. ContEst: Cibulskis, K. et al. ContEst: estimating cross-contamination of human samples in next-generation sequencing data. Bioinformatics 27, 2601–2 (2011).
3. Strelka: Saunders, C. T. et al. Strelka: accurate somatic small-variant calling from sequenced tumor-normal sample pairs. Bioinformatics 28, 1811–7 (2012).
4. Orientation Bias Filter (oxoG, FFPE): Costello, M. et al. Discovery and characterization of artifactual mutations in deep coverage targeted capture sequencing data due to oxidative DNA damage during sample preparation. Nucleic Acids Res. 41, e67 (2013).
5. DeTiN: Taylor-Weiner, A. et al. DeTiN: overcoming tumor-in-normal contamination. Nat Methods 15, 531–534 (2018).
6. AllelicCapSeg: Landau, D. A. et al. Evolution and impact of subclonal mutations in chronic lymphocytic leukemia. Cell 152, 714–26 (2013).
7. MAPPoNFilter: Lawrence, M. et al. Discovery and saturation analysis of cancer genes across 21 tumour types. Nature 505, 495 (2014).
8. ABSOLUTE: Carter, S. L. et al. Absolute quantification of somatic DNA alterations in human cancer. Nat. Biotechnol. 30, 413–21 (2012). doi: 10.1038/nbt.2203.
9. GATK (Mutect2, somatic CNV): McKenna, A. et al. The Genome Analysis Toolkit: a MapReduce framework for analyzing next-generation DNA sequencing data.Genome Res. 20, 1297–303 (2010).
10. Picard Tools: https://software.broadinstitute.org/gatk/documentation/tools/docs/4.0.1.0/picard_fingerprint_CrosscheckFingerprints.php
https://software.broadinstitute.org/gatk/documentation/tools/docs/4.0.0.0/picard_analysis_CollectMultipleMetrics.php
Additional tools used for PDX analyses, mutational signature analyses and statistical analyses included:
1. SAMtools (1.7): Li, H. et al A statistical framework for SNP calling, mutation discovery, association mapping and population genetical parameter estimation from sequencing data. Bioinformatics 2011;27:2987-93
2. BWA-MEM (0.7.17): https://github.com/lh3/bwa
3. Disambiguate (ngs_disambiguate-1.0): https://github.com/AstraZeneca-NGS/disambiguate
4. Genemapper Software (5): https://www.thermofisher.com/order/catalog/product/4475073#/4475073
5. SomaticSignature (3.1): https://bioconductor.org/packages/release/bioc/html/SomaticSignatures.html
6. DeconstructSigs (1.8): https://github.com/raerose01/deconstructSigs
7. STATA (v14.2): https://www.stata.com
8. MedCalc (19.0.3): https://www.medcalc.org
9. Graphpad Prism (8): https://www.graphpad.com/scientific-software/prism/

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors/reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.

Data
Policy information about availability of data
All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:
- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

Clinical and sequencing data from 1495 samples from the DFCI-Profile and 545 samples from the MSKCC-IMPACT datasets are publicly available (GENIE v4.1: https://www.synapse.org/). All data for samples from the GENIE v6.1 and TCGA pan-cancer datasets are publicly available. Data for samples from the FMI dataset are not publicly available. Detailed clinical annotation for the DFCI-Profile and MSKCC-IMPACT cohorts is provided in supplementary table 1.

Field-specific reporting
Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

Life sciences

All studies must disclose on these points even when the disclosure is negative.

Sample size
No statistical test was used to determine the sample size. We systematically collected data from 10,294 glioma samples from three large independent datasets. Based on prior literature (prevalence of hypermutation of 2-5% in gliomas), we hypothesized that this sample size would provide enough power (200-500 hypermutated samples) for clinical and molecular association studies.

Data exclusions
47 samples for which the clinical diagnosis of glioma could not be confirmed (other histology or possible non-tumor sample) and 5 samples with missing clinical annotation were excluded from all analyses. Exclusion criteria were pre-established.

Replication
In vivo experiments were performed using groups of 8 mice per group. All mice were included in the analysis. In vitro sensitivity assays were replicated in at least 3 independent experiments. All experiments that were technically valid were included in the analysis.

Randomization
For in vivo experiments, mice were randomized.

Blinding
No blinding was performed. Blinding was not relevant to our study.

Reporting for specific materials, systems and methods
We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.
## Materials & experimental systems

| n/a | Involved in the study |
|-----|-----------------------|

- Antibodies | [ ]
- Eukaryotic cell lines | [ ]
- Palaeontology | [ ]
- Animals and other organisms | [ ]
- Human research participants | [ ]
- Clinical data | [ ]

## Methods

| n/a | Involved in the study |
|-----|-----------------------|

- ChIP-seq | [ ]
- Flow cytometry | [ ]
- MRI-based neuroimaging | [ ]

### Antibodies

**Antibodies used**

For immunohistochemistry, the following antibodies were used: mouse monoclonal anti-ATRX (Bio SB, clone BSB-108, BSB3296, 1:100), mouse monoclonal anti-IDH1 R132H (Dianova, clone H09, DIA-H09, 1:100), rabbit monoclonal anti-CD3 (Roche, clone 2GV6, 790-4341, prediluted), rabbit polyclonal anti-IBA1 (Wako, W1WD19-19741, 1:500), mouse monoclonal anti-MLH1 (Roche, clone M1, 790-4535, prediluted), mouse monoclonal anti-MSH2 (Roche, clone G219-1129, 760-4265, prediluted), mouse monoclonal anti-MSH6 (Roche, clone 44, 760-4455, prediluted), rabbit monoclonal anti-PMS2 (Roche, clone EPR3947, 760-4531, prediluted). For immunohistochemistry performed at BWH, the following antibodies were used: mouse monoclonal anti-MLH1 (Leica, clone E505, MLH1-L-CE, 1:75), mouse monoclonal anti-MSH2 (Merck Millipore, clone Ab-2-FE11, NA27, 1:200), mouse monoclonal anti-MSH6 (Leica, clone PU29, MSH6-L-CE, 1:50), mouse monoclonal anti-PMS2 (Cell Marque, MRQ-28, 288M-14-ASR, 1:100). For immunoblotting, the following antibodies were used: mouse monoclonal anti-MGMT (Millipore, MT3.1, MAB16200, 1:500), mouse monoclonal anti-MSH2 (Calbiochem, FE11, NA27, 1:1000), mouse monoclonal anti-MSH6 (Biosciences, 44, 610918, 1:500), mouse monoclonal anti-MLH1 (Cell Signaling, 4C9C7, 3515, 1:500), mouse monoclonal anti-PMS2 (BD Biosciences, A16-4, 556415, 1:1000), mouse monoclonal anti-beta-actin (Sigma, AC-74, A2228, 1:10000).

**Validation**

All antibodies used are commercially available and were validated by the manufacturer.

### Eukaryotic cell lines

**Policy information about cell lines**

**Cell line source(s)**

All patient-derived cell lines (PDCLs) and xenografts (PDXs) with a name starting with "BT" were established from tumors resected at Brigham and Women’s Hospital and Boston Children’s Hospital (Boston, MA). SU-DIPG-XIII (DIPG13) cells were provided by Dr. Michelle Monje at Stanford University.

**Authentication**

The identity of all cell lines established were confirmed by short tandem repeat assay.

**Mycoplasma contamination**

All cell lines were tested for the absence of mycoplasma.

**Commonly misidentified lines**

No commonly misidentified cell lines were used.

### Animals and other organisms

**Policy information about studies involving animals; ARRIVE guidelines recommended for reporting animal research**

**Laboratory animals**

Eight-week-old NU/NU male mice (Charles River).

**Wild animals**

The study did not involve wild animals.

**Field-collected samples**

The study did not involve samples collected from the field.

**Ethics oversight**

All in vivo studies were performed in accordance with Dana-Farber Cancer Institute animal facility regulations and policies under the protocol #09-016.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

### Human research participants

**Policy information about studies involving human research participants**

**Population characteristics**

We systematically analyzed clinical data and somatic tumor variants identified through targeted next-generation sequencing (NGS) panels of 1,628 gliomas sequenced between June 2013 and November 2018 as part of a large institutional prospective profiling program (DFCI-Profile). All samples were assigned to a molecular subgroup based on histopathology, mutational status of IDH1 and IDH2 genes, and whole-arm co-deletion of chromosomes 1p and 19q (1p/19q co-deletion). A total of 545 independent samples from the GENIE project (a repository of genomic data obtained during routine clinical care at international institutions) were also identified and assigned to molecular subgroups. The combined sequencing set comprised 2,173 glioma samples, which included a broad spectrum of newly-diagnosed and recurrent glioma types including IDH1/2 wild-type.
| Recruitment | Genomic testing was ordered by the pathologist or treating physician as part of routine clinical care to identify relevant genomic alterations that could potentially inform diagnosis and treatment decisions. All patients undergoing genomic testing signed a clinical consent form, permitting to return results from clinical sequencing. No systematic bias likely to impact results were known at the time of data analysis. |
|---|---|
| Ethics oversight | The study, including the consent procedure, was approved by the institutional ethics committees (10-417/11-104/17-000; WIRB, Puyallup WA). Note that full information on the approval of the study protocol must also be provided in the manuscript. |