



Cancer-specific loss of *TERT* activation sensitizes glioblastoma to DNA damage

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Most glioblastomas (GBMs) achieve cellular immortality by acquiring a mutation in the telomerase reverse transcriptase (*TERT*) promoter. *TERT* promoter mutations create a binding site for a GA binding protein (GABP) transcription factor complex, whose assembly at the promoter is associated with *TERT* reactivation and telomere maintenance. Here, we demonstrate increased binding of a specific GABPB1L-isoform-containing complex to the mutant *TERT* promoter. Furthermore, we find that *TERT* promoter mutant GBM cells, unlike wild-type cells, exhibit a critical near-term dependence on GABPB1L for proliferation, notably also posttumor establishment *in vivo*. Up-regulation of the protein paralogue GABPB2, which is normally expressed at very low levels, can rescue this dependence. More importantly, when combined with frontline temozolomide (TMZ) chemotherapy, inducible GABPB1L knock-down and the associated *TERT* reduction led to an impaired DNA damage response that resulted in profoundly reduced growth of intracranial GBM tumors. Together, these findings provide insights into the mechanism of cancer-specific *TERT* regulation, uncover rapid effects of GABPB1L-mediated *TERT* suppression in GBM maintenance, and establish GABPB1L inhibition in combination with chemotherapy as a therapeutic strategy for *TERT* promoter mutant GBM.

TERT | cancer | CRISPR | glioblastoma | temozolomide

PPrimary glioblastoma (GBM) is the most common and lethal form of malignant brain cancer in adults. Current treatment strategies are limited, with GBM progression leading to death within 2 y of diagnosis in 90% of cases (1–3). In GBM, as well as the vast majority of other cancers, transcriptional activation of the telomerase reverse transcriptase (*TERT*) gene, which is normally silenced in somatic cells, is a key step in tumorigenesis (4, 5). *TERT* encodes the catalytic subunit of telomerase, and its reactivation in cancer is thought to contribute to cell survival and immortalization (6–8). *TERT* ablation thus has the potential to directly affect both short- and long-term cell viability through telomere-length-dependent and independent pathways (9–15). Prior research has demonstrated that inhibition of *TERT* expression enhances sensitivity of cells to DNA damage by radiation and chemotherapy, suggestive of a possible combination therapy for cancer treatment (16, 17). However, telomerase inhibition is toxic to normal telomerase-dependent stem and germline cells, which has led to the failure of such approaches clinically (18–20). Therefore, understanding genetic contributors to aberrant *TERT* expression, as well as consequences of cancer-specific *TERT* ablation, are critical to develop novel therapeutic avenues for GBM.

A major mechanism by which *TERT* is reactivated in cancer involves the acquisition of somatic mutations in its promoter, which represent the most common noncoding mutations, and the third most common mutations overall, in cancer (21–26). In particular, ~80% of primary GBMs contain one of two common

single-nucleotide mutations that are associated with re-expression of *TERT* messenger RNA (mRNA), referred to as G228A and G250A (22, 24, 27). Both G-to-A transitions generate an identical 11-base-pair sequence (plus strand CCCGGAAGGGG) that creates a binding site for GA binding protein A (GABPA), an E26 transformation-specific (ETS)-family transcription factor (28, 29). Interestingly, these *de novo* ETS binding sites occur within three (G228A) or five (G250A) complete helical turns of two overlapping native ETS binding sites (ETS 195 and ETS 200) in the *TERT*

Significance

Glioblastoma is a highly lethal form of brain cancer with no current treatment options that substantially improve patient outcomes. A key therapeutic challenge is the identification of methods that reduce tumor burden while leaving normal cells unaffected. We show that *TERT*-promoter mutations, common in glioblastoma, lead to *TERT* reactivation through increased binding of GABPB1L-isoform-containing transcription factor complexes. In turn, we find that cancer-cell-specific inhibition of *TERT* through GABPB1L reduction results in near-term anti-growth effects and an impaired DNA damage response that profoundly increase the sensitivity of glioblastoma tumors to frontline chemotherapy. Our results thus provide rationale for GABPB1L inhibition combined with temozolomide chemotherapy treatment as a promising therapeutic strategy for glioblastoma.

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promoter (*TERT*), one or the other of which are also required for *TERT* reactivation (28). GABP transcription factors are obligate multimers that consist of the DNA-binding GABPA subunit (GeneID: 2551) and a transactivating GABPB subunit (30). Humans have two paralogues encoding different beta subunits, GABPB1 (GeneID: 2553) and GABPB2 (GeneID: 126626). Reduced function of the long protein isoform of GABPB1 (GABPB1L) via indel mutations has previously been linked to down-regulation of *TERT* mRNA and long-term telomere attrition (9). This could provide a cancer-specific way to target *TERT*, particularly given that GABPB1L is dispensable for normal murine development while GABPA and total GABPB1 are not (31, 32). However, the extended time period required to induce cell death via progressive telomere shortening limits the therapeutic potential of this approach for high-grade GBM (33). To further our understanding of the GABPB1L-*TERT* axis and its possible therapeutic benefit, here we examined the specificity and binding affinity of GABPB1L for the mutant *TERT* promoter, as well as functional effects of GABPB1L loss in a near-term, clinically relevant timeframe. Notably, we observed a dramatic synergistic effect between GABPB1L reduction and standard-of-care temozolomide chemotherapy, mediated through *TERT* down-regulation and a resulting attenuation of the DNA damage response (DDR) in *TERT* mutant cells. These rapid effects of targeting the cancer-specific GABPB1L-*TERT* axis in combination with frontline chemotherapy in established intracranial GBM xenografts led to substantially increased survival, promising exciting avenues for *TERT*-mutant GBM therapy.

Results

GABPB1L-Containing Complexes Bind and Regulate the Mutant *TERT* Promoter. *GABPB1* encodes two main transcript variants, a short isoform (GABPB1S) and a long isoform (GABPB1L). GABPB1S functions as a heterodimer with GABPA (GABPA₁B₁), while GABPB1L forms a heterotetramer (GABPA₂B₂) due to its unique terminal exon that contains a leucine zipper-like domain (30, 34) (*SI Appendix, Fig. S1A*). Previous work demonstrated that *TERT* mutations create a binding site for the GABPA-B transcription factor complex and linked reactivation of *TERT* in this context to GABP complexes containing GABPB1L (9, 28). Indeed, the fact that two GABP binding sites distanced at complete helical turns from one another are required to enable *TERT* reactivation is suggestive of the recruitment of a GABPB1L-containing heterotetramer (9, 28, 35) (*SI Appendix, Fig. S1A*). However, GABPB1 isoform specificity in mutant *TERT* promoter regulation has not been examined. In order to test this, we made use of a doxycycline-inducible, microRNA-embedded short hairpin RNA (shRNA) system with a green fluorescent protein (GFP) marker reporting shRNA induction (36–38). We used a machine learning algorithm (38) to design shRNAs targeting either GABPB1L or GABPB1S specifically (*SI Appendix, Fig. S1B*). Doxycycline-mediated expression of GABPB1L or GABPB1S shRNAs led to reduction of the respective mRNA and protein levels in *TERT* mutant (U-251) and *TERT* wild-type (WT) (LN-18) GBM cells (Fig. 1A and *SI Appendix, Fig. S1C and D*). Interestingly, GABPB1L—but not GABPB1S—knockdown led to reduced *TERT* mRNA and telomerase activity exclusively in *TERT* mutant cells (Fig. 1B and C and *SI Appendix, Fig. S1E*), suggesting that *TERT* mutant regulation of *TERT* expression is dependent on GABPB1L-containing complexes specifically. Of note, residual *TERT* expression and telomerase activity are observed in *TERT* mutant cells following GABPB1L knockdown when compared with a cancer cell line that is negative for *TERT* expression (U2OS) (Fig. 1B and C and *SI Appendix, Fig. S1E*). This may be due to incomplete shRNA-mediated knockdown (Fig. 1A and *SI Appendix, Fig. S1C*) or to regulation of *TERT* by other factors in conjunction with, or in the absence of, GABPB1L (39–41).

Though binding of GABP to the *TERT* promoter has previously been demonstrated (28), possible GABPA₂B₁L₂ heterotetramer formation at the mutant promoter has not directly been examined. To determine the nature of the GABPA-B1L interaction with the *TERT* promoter, we titrated purified heterodimer (*SI Appendix, Fig. S1F and G*) against a fixed concentration of radiolabeled wild-type *TERT* DNA, mutant *TERT* DNA (G228A), and control probes with single (G201T) or double (A197T and G201T) mutations in the native ETS sites contained within the WT promoter. We observed that GABPB1L-containing complexes bind with ~twofold increased affinity to the mutant *TERT* relative to wild type. However, calculating K_d values that accurately reflect GABP binding was challenging due to the observation of multiple intermediate states, which may be reflective of heterodimer binding, as has been previously suggested (42, 43) (Fig. 1D and *SI Appendix, Fig. S1H*). Thus, we segregated observed states into unbound, intermediate bound, and fully bound states. The comparison of signal ratios in each state for each target DNA tested across multiple GABP concentrations showed that the *TERT* mutation (G228A) leads to increased formation of the fully bound state, which we attribute to a likely tetrameric complex (42, 43) (Fig. 1D and E). We saw that GABP has the ability to bind all *TERT* promoter sequences tested in this study. However, binding to WT and native ETS site mutants (G201T or A197T and G201T) showed an increased number of intermediate states when compared with the G228A-bearing promoters and a decrease in the fully bound state (Fig. 1D and E and *SI Appendix, Fig. S1H*). Together, these results demonstrate that *TERT* mutations result in increased binding affinity for probable GABPB1L-containing heterotetramer complexes, which help drive *TERT* reactivation.

GABPB1L Knockout Impairs Near-Term Growth of *TERT* Promoter Mutant GBM Cells. For GBM therapy, short-term effects of *TERT* reduction, rather than long-term consequences of canonical telomere attrition, would increase the clinical relevance of targeting the cancer-specific GABP-*TERT* axis (33). Given that GABPB1L regulates *TERT* in the context of *TERT* mutations, our data raise the question of possible early functional consequences of GABPB1L inhibition on *TERT*-mutant cell growth and viability. CRISPR-Cas methods enable the precise dissection of genetic dependencies through the generation and analysis of specific heterozygous and homozygous genetic deletions in isogenic cell lines (44). Therefore, to assess near-term consequences of complete GABPB1L deletion, we used CRISPR-Cas9 and single-guide RNAs (sgRNAs) targeting the intron and 3' untranslated region (UTR) flanking the unique exon 9 of *GABPB1L* (Fig. 2A and *SI Appendix, Fig. S2A and B and Table S1*). We compared a set of three *TERT* mutant GBM cell lines (U-251, LN-229, and T98G) and four *TERT* WT cell lines (HEK293T, HAP1, NHA-S2, and LN-18) by generating monoclonal isogenic derivatives. The *TERT* WT lines include an immortalized human astrocyte line (NHA-S2) as well as a GBM line (LN-18). Sequencing of the *TERT* promoter confirmed *TERT* mutation status (*SI Appendix, Fig. S2C and D*).

To identify ideal guide RNAs, all possible combinations of three intronic and three 3' UTR targeting guides were compared in HEK293T and U-251 cells to assess editing efficiency (Fig. 2B and *SI Appendix, Fig. S2E*). Next, the two most efficient pairs of sgRNAs were used to edit all seven cell lines, and 169 monoclonal cell line derivatives were established, genotyped, and subjected to further analysis (Fig. 2C and D and *SI Appendix, Figs. S3A–H and S4A–C*). Strikingly, we observed that while full homozygous knockout of GABPB1L occurred at an appreciable rate across all *TERT* WT cell lines (13 to 36%), we were able to generate only one full GABPB1L knockout line, LN229-C19, among the 76 *TERT* mutant clones that were screened (Fig. 2E and *SI Appendix, Fig. S3A–H*). Importantly, the discrepancy between *TERT* WT and mutant cell lines did not appear to result

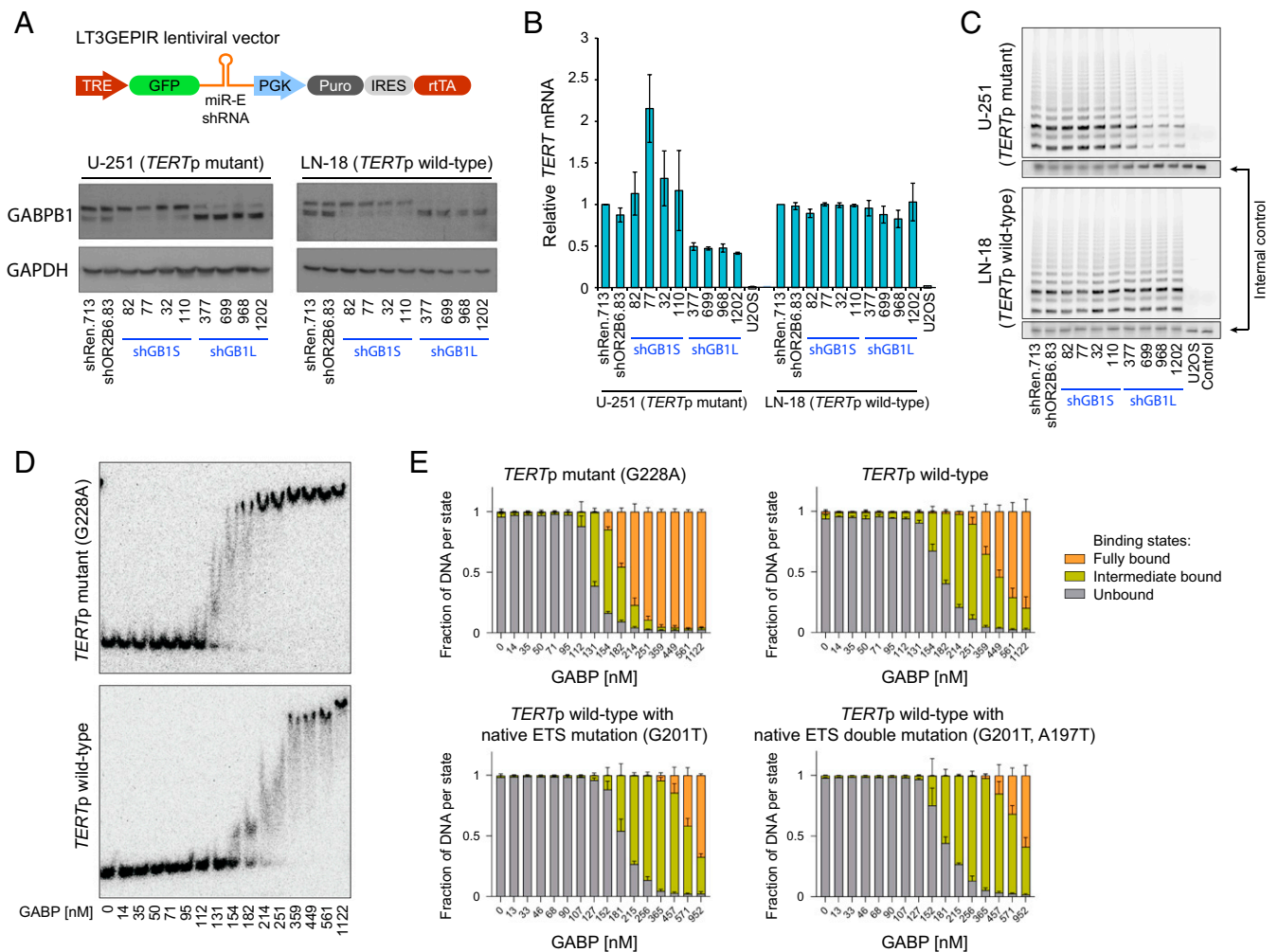


Fig. 1. GABPB1L-containing complexes bind and regulate the mutant *TERT* promoter. (A) Representative immunoblots of GABPB1 in U-251 and LN-18 cells expressing doxycycline-induced shRNAs targeting GABPB15 (shGB15.82, shGB15.77, shGB15.32, and shGB15.110) and GABPB1L (shGB1L.377, shGB1L.699, shGB1L.968, and shGB1L.1202) compared with negative control (olfactory receptor OR2B6, shOR2B6.83) and nontargeting (renilla luciferase, shRen.713) shRNAs. Cells were incubated with doxycycline for 6 d prior to harvest. The lower bands represent GABPB15, and upper bands represent GABPB1L. (B) *TERT* mRNA expression measured via qRT-PCR in cell lines from A compared with a control cell line (U2OS) lacking *TERT* expression. (C) Representative gels of telomerase activity measured via TRAP assay in cell lines from A compared with a control cell line (U2OS) lacking *TERT* expression. The control condition reflects no lysate. (D and E) Representative gels (D) and quantification (E) of electrophoretic mobility shift assays (EMSA) comparing binding affinity of GABPB1L heterodimers to the mutant (G228A) *TERT* promoter (additional ETS binding site), WT *TERT* promoter (native ETS binding sites), and control WT *TERT* promoter sequences lacking native ETS binding sites (G201T: native ETS single mutant; G201T and A197T: native ETS double mutant).

from differences in overall editing efficacy between cell lines, as equal numbers of heterozygous knockouts were observed (Fig. 2E and SI Appendix, Fig. S3H). Thus, loss of GABPB1L may have near-term effects on growth and survival of *TERTp* mutant cells specifically that prevent the generation of full knockout clones. Prior studies from our laboratory demonstrated that impaired GABPB1L protein function through indel-based editing led to long-term telomere attrition over a more than 90 d timeframe (9). However, the present data suggest that full knockout of GABPB1L causes *TERTp* mutant cells to become either slow growing or nonviable over the ~30 d editing and selection process, a timeframe that is not consistent with gradual telomere shortening (9, 15, 45).

GABPB1L Knockdown Slows Growth of Established *TERTp* Mutant Tumors. To more directly assess the therapeutic potential of GABPB1L reduction, we next examined its effect on growth of established GBM tumors in vivo, using the above-described doxycycline-inducible shRNA system (Fig. 1A and SI Appendix,

Fig. S1B). We first tested the system for in vivo efficacy using a bona fide essential gene, Replication Protein A1 (RPA1; GeneID: 6117), along with controls. Addition of doxycycline to U-251 cells stably transduced with an inducible vector encoding RPA1-targeting shRNAs resulted in reduced *RPA1* mRNA in vitro (SI Appendix, Fig. S5A) and prolonged animal survival in vivo (SI Appendix, Fig. S5B and C). Because the doxycycline-inducible vectors encode an shRNA within the 3' UTR of a GFP construct (Fig. 1A), the percentage of GFP-positive cells was used as a proxy readout for shRNA expression. Postmortem tumor analysis showed that in vivo expression of the RPA1-targeting shRNA construct was reduced compared with controls (SI Appendix, Fig. S5D), consistent with a survival advantage for loss of RPA1 knockdown (37).

We then made use of our GABPB1L-targeting shRNAs to determine the effect of GABPB1L knockdown on growth of existing GBM tumors. Two separate GABPB1L-targeting shRNAs tested in orthotopic intracranial xenografts of U-251 cells slowed tumor growth within 12 d of shRNA induction when induced post tumor implantation (Fig. 2F and SI Appendix, Fig. S5E) and

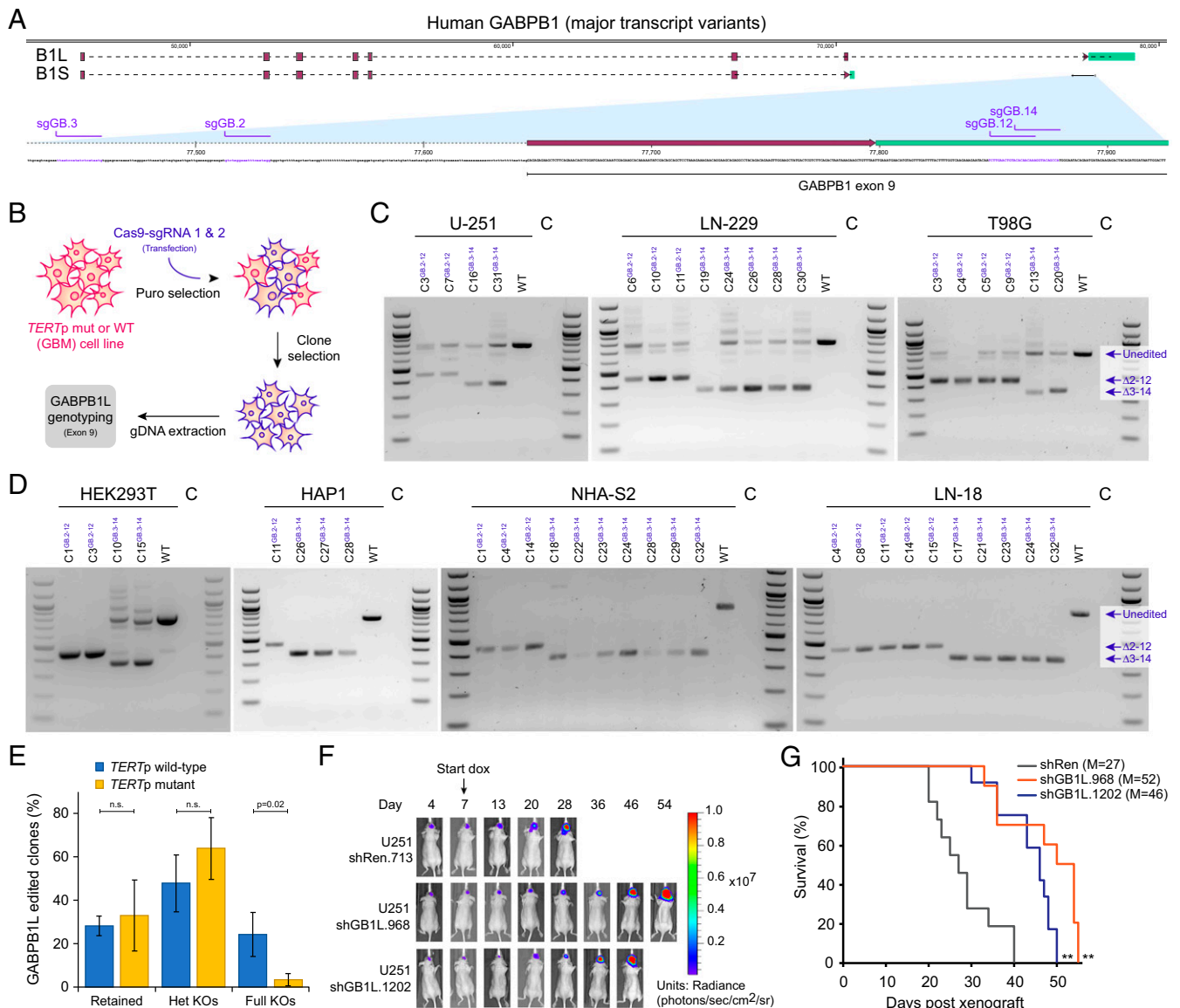


Fig. 2. Inhibition of GABPB1L rapidly prevents growth of *TERT* promoter mutant cells. (A) *GABPB1L* exon 9 excision strategy. A diagram of the major human *GABPB1* transcript variants, encoding either a short (*GABPB1S*) or long (*GABPB1L*) isoform. Red rectangles represent the coding sequence, and green rectangles represent the 3' UTR. For clarity, the first exon containing the 5' UTR is not shown. *GABPB1L* sgRNA targeting sites are highlighted in purple. (B) The editing strategy for scarless knockout of the long isoform of GABPB1 (*GABPB1L*). In each case, a pair of sgRNAs were used targeting *GABPB1L* intron number 8 and the 3' UTR of *GABPB1L*, respectively. (C and D) Genotyping analysis of *GABPB1L* edited monoclonal *TERT* promoter mutant U-251, LN-229, and T98G GBM cell lines (C) and *TERT* promoter WT HEK293T, HAP1, NHA-S2, and LN-18 cell lines (D). The superscript characters indicate the sgRNA pair used for editing. WT: parental WT cell line. C, PCR no template control. Unedited, full-length genotyping band; $\Delta 2$ -12, band after excision with sgGB2 and sgGB12; $\Delta 3$ -14, band after excision with sgGB3 and sgGB14. (E) A comparison of *GABPB1L* editing outcomes between *TERTp* WT (blue) and *TERTp* mutant (yellow) lines. The bars represent the percentage of each editing outcome averaged over three *TERTp* mutant and four *TERTp* WT lines. Retained refers to clones where the *GABPB1L* exon 9 was overall retained, even though they may contain indels at either or both of the sgRNA target sites flanking exon 9. *P* value (unpaired, two-tailed Student's *t* test). n.s., not significant (alpha level = 0.05). (F and G) Representative bioluminescence images (F) and Kaplan–Meier survival curve (G) of mice injected with U-251 cells expressing a control shRNA (shRen.713) or *GABPB1L*-targeting shRNAs (shGB1L.968, shGB1L.1202). Mice were fed doxycycline chow to induce shRNA expression starting at day 7 postinjection. *n* = 10 to 12 mice per condition. ***P* < 0.01 compared with shRen.713 (Kaplan–Meier log-rank test). M, median survival.

increased median survival by 70 to 90% (Fig. 2G). *GABPB1L* shRNA-mediated target knockdown and linked GFP expression were maintained in vivo via analysis of ex vivo dissociated tumor cells postmortem (SI Appendix, Fig. S5 F and G). A third tested *GABPB1L*-targeting shRNA (shGB1L.699) did not impact survival (SI Appendix, Fig. S5H). Correspondingly, postmortem analysis demonstrated minimal *GABPB1L* protein reduction in tumors with this shRNA and showed that only ~30% of tumor cells retained GFP expression (SI Appendix, Fig. S5 F and G), indicating that sustained *GABPB1L* knockdown is necessary to slow tumor growth.

A short-term cell-culture-based competitive proliferation assay with the same shRNAs did not show any toxicity (SI Appendix, Fig. S5I), pointing toward a specific in vivo effect of sustained *GABPB1L* knockdown. Together, these data demonstrate that shRNA-mediated *GABPB1L* suppression in established tumors significantly prolongs animal survival, highlighting its therapeutic value.

GABPB2 Promotes Resistance to *GABPB1L* Loss in *TERTp* Mutant GBM Cells. Given the potential therapeutic benefit of *GABPB1L* inhibition for *TERTp* mutant GBM, we chose to analyze heterozygous

knockout clones generated from *TERT*_p mutant lines compared with full and heterozygous knockouts from *TERT*_p WT lines in order to better understand the molecular impact of GABPB1L loss (Fig. 2 C–E). Full knockout of GABPB1L led to a complete loss of GABPB1L mRNA and protein in most cell lines, whereas heterozygous knockout led to various degrees of GABPB1L reduction, consistent with the expected effects of total versus partial removal of GABPB1L loci (SI Appendix, Fig. S6 A and B). We also observed unchanged or up-regulated total *GABPB1* mRNA (all transcript variants) and substantially up-regulated *GABPB1S* mRNA and protein across all cell lines (SI Appendix, Fig. S6 A–F), supporting the possibility that all *GABPB1* transcripts from a *GABPB1L* knockout locus are now *GABPB1S*. Notably, GABPB1L reduction was associated with a decrease in *TERT* mRNA expression and reduced telomerase activity in *TERT*_p mutant cells only (SI Appendix, Fig. S6 G–J), confirming regulation of *TERT* by GABPB1L in a *TERT*_p mutant-specific manner.

The single *TERT*_p mutant GABPB1L full knockout cell line (LN229-C19) that we obtained did not exhibit reduction of either *TERT* mRNA or telomerase activity (SI Appendix, Fig. S5 G, I, and J), suggesting a compensatory mechanism to regulate *TERT* independent of GABPB1L. *GABPB1* and *GABPB2* are distinct genes with high sequence conservation that perform similar but

separate functions, though GABPB2 is thought to function exclusively as a heterotetramer with GABPA (30, 34), mediated by the presence of a leucine zipper-like domain (Fig. 3A and SI Appendix, Fig. S7A). Since GABPB2 is expressed at lower levels than GABPB1 in GBM based on data from the TCGA Research Network (<https://www.cancer.gov/tcga>) (SI Appendix, Fig. S7B), we probed for possible GABPB2 up-regulation in our GABPB1L knockout clones. While deletion of GABPB1L did not affect *GABPB2* mRNA in any other line, the LN229-C19 clone showed a 4.5-fold elevation in *GABPB2* transcript levels (Fig. 3B and SI Appendix, Fig. S7C). In support of *GABPB2* up-regulation as a potential compensatory mechanism, stable GABPB2 over-expression in U-251 cells rescued cell dropout following CRISPR editing of the *GABPB1* locus (Fig. 3C). In addition, small interfering RNA-mediated inhibition of *GABPB2* showed that *TERT* mRNA levels are responsive to GABPB2 knockdown in LN229-C19 but not in WT LN-229 cells or heterozygous knockout clones (Fig. 3D). These data suggest that GABPB2 up-regulation can compensate functionally for *GABPB1L* deletion in *TERT*_p mutant cells and indicate that *TERT*_p mutant GBM can tolerate total loss of GABPB1L in the context of alternative mechanisms rescuing *TERT* expression. This finding reveals a possible resistance mechanism for GABPB1L-targeting therapies.

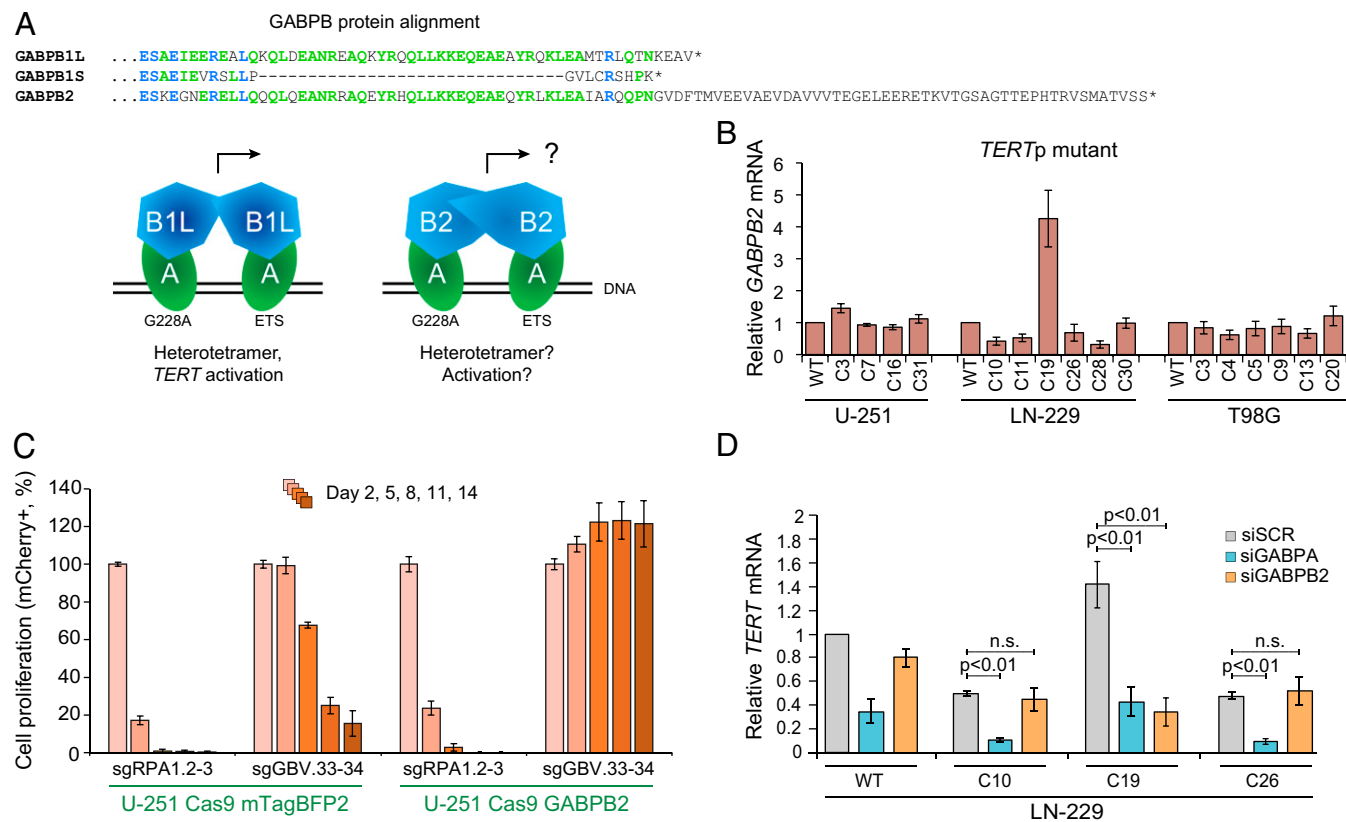


Fig. 3. GABPB2 up-regulation can compensate for loss of GABPB1L. (A) Similarity between GABP proteins. Partial protein alignment (MUSCLE) of GABPB1L, GABPB1S, and GABPB2. GABPB1L is the long isoform and GABPB1S the short isoform of the *GABPB1* gene. *GABPB2* is a distinct gene. Amino acids matching across all three proteins are highlighted in blue, and those matching across two proteins are highlighted in green. GABPA (alpha) subunits of the GABP complex bind to both the native ETS site (ETS) and the mutation-derived ETS sites (G228A or G250A) at the *TERT* promoter locus. The GABPB1 short (B1S) or long (B1L) isoform subunits bind to the alpha subunits to form either heterodimers (GABPA₁B1S₁) or heterotetramers (GABPA₂B1L₂). Heterotetramer formation is presumably mediated through the leucine zipper-like domain of GABPB1L, which is also present in GABPB2. (B) *GABPB2* mRNA expression measured via qRT-PCR in GABPB1L knockout clones, plotted relative to control (WT) cells, in *TERT*_p mutant cell lines. The data represent mean ± SEM. (C) Competitive proliferation assay in U-251 cells using pairs of sgRNAs targeting a positive control locus (sgRPA1.2 and 3) and total GABPB1 (targeting exon 3, sgGBV.33 and 34) in the presence of a lentiviral vector expressing either GABPB2 or mTagBFP2 (control). The data represent mean ± SD of triplicates. (D) *TERT* mRNA expression measured via qRT-PCR following siRNA-mediated knockdown of GABPA or GABPB2 in LN-229 WT, full knockout, or heterozygous knockout cells. p, P value (unpaired, two-tailed Student's t test). n.s., not significant (alpha level = 0.05). The data represent mean ± SEM.

Increased Response of GABPB1L-Reduced GBM Tumors to Chemotherapy. Currently, most patients diagnosed with primary GBM are treated with a regimen that includes temozolomide (TMZ) chemotherapy, a DNA-alkylating chemotherapeutic agent that is most effective at eliminating cells that lack O-6-methylguanine-DNA methyltransferase (MGMT) through generation of single- and double-strand DNA breaks (46, 47). Prior studies have indicated that reduction of *TERT* in *TERT*-expressing cells may sensitize them to DNA damage from ionizing radiation and chemotherapy (16, 17). Reduction of *TERT* mRNA by RNA interference has been shown to prevent the repair of DNA breaks in vitro by inhibiting central components of the DDR, such as histone H2AX phosphorylation (γH2AX), through an incompletely understood mechanism (16). Loss of DNA damage signaling increases DNA damage-related senescence or cell death, possibly due to a reduction in normal cell cycle arrest (16, 48, 49). Therefore, we wondered whether a GABPB1L-mediated down-regulation of *TERT* mRNA in *TERT*^p mutant GBM might promote TMZ response, enabling a possible path to GBM-cell-specific chemotherapy sensitization (Fig. 4A).

U-251 cells expressing doxycycline-inducible shRNAs targeting either *TERT* or GABPB1L were treated with a dose titration of TMZ, and the DDR was assessed through analysis of γH2AX levels. Compared with control, shRNA-mediated *TERT* knockdown significantly reduced the usual increase in γH2AX 20 h after treatment with TMZ (Fig. 4B and *SI Appendix, Fig. S8 A and B*). Strikingly, GABPB1L knockdown led to an equally blunted γH2AX increase following TMZ treatment (Fig. 4B and *SI Appendix, Fig. S8B*). Similarly, TMZ-mediated γH2AX induction was reduced in both U-251 and LN-229 clones with heterozygous *GABPB1L* knockout, compared with WT cells (*SI Appendix, Fig. S8 C and D*). However, the full *GABPB1L* knockout clone LN229-C19 with GABPB2 up-regulation and maintained *TERT* expression exhibited a TMZ-induced γH2AX increase that mirrored WT, consistent with the impairment of the DDR being mediated by *TERT* down-regulation (*SI Appendix, Fig. S8D*).

To directly assess whether GABPB1L knockdown leads to a reduced γH2AX increase through reduced *TERT*, we generated U-251 cells that stably express either blue fluorescent protein (BFP; control) or *TERT* (rescue) constructs (*SI Appendix, Fig. S8E*). While the BFP construct showed no effects, overexpression of *TERT* completely rescued the loss of γH2AX induction following shRNA-mediated knockdown of both *TERT* and GABPB1L (Fig. 4C). In the control shRNA condition (shRen.713), *TERT* overexpression significantly increased γH2AX levels following TMZ treatment when compared with WT *TERT* levels, providing further evidence for regulation of the DDR by *TERT* (Fig. 4C and *SI Appendix, Fig. S8E*).

Ablated DDR signaling can lead to a blunting of the G2 cell cycle arrest that is commonly observed following treatment with DNA damaging agents such as TMZ, causing an increased loss of viability as cells continue to divide (48, 50, 51). In concordance, here we also observe reduced G2 arrest in both U-251 and LN-229 GABPB1L heterozygous knockout clones treated with TMZ, as compared with both WT cells and the LN-229 full GABPB1L knockout clone C19 (Fig. 4D and *SI Appendix, Fig. S8 F and G*). Lack of MGMT expression was confirmed in all cell lines and corresponding clones (*SI Appendix, Fig. S8H*).

Both partial and full loss of H2AX, or inhibition of its phosphorylation, have been shown to significantly impact cell survival following exposure to DNA damage (52–54). Similarly, inhibition of G2 arrest following TMZ administration is associated with decreased cell viability and growth, resulting in TMZ sensitization (50). Therefore, we hypothesized that *TERT*^p mutant GBM cells with reduced GABPB1L would be more sensitive to TMZ chemotherapy. We first assessed the therapeutic implications of this finding in vivo through administration of a TMZ regimen following orthotopic intracranial xenograft of either U-

251 or LN-229 WT and *GABPB1L* knockout cells. Initial analysis—in the absence of TMZ therapy—of *GABPB1L* heterozygous knockout cell tumors in vivo demonstrated that GABPB1L reduction slowed growth of both U-251 and LN-229 tumors and increased animal survival (*SI Appendix, Fig. S8 I–L*), while the full knockout LN229-C19 clone with maintained *TERT* expression formed tumors at a similar rate to WT cells. Strikingly, in vivo administration of chemotherapy in combination with *GABPB1L* heterozygous status radically ablated tumor growth of U-251 clones (*SI Appendix, Fig. S9 A–D*). A similar potentiation of treatment efficacy was also observed, though to a lesser extent, in GABPB1L-reduced LN-229 tumors, again with the expected exception of the full knockout LN229-C19 clone that maintained WT *TERT* mRNA levels (*SI Appendix, Fig. S9 E–H*).

Lastly, in order to assess the therapeutic potential of this combination therapy in a more clinically relevant manner, we performed orthotopic xenografts of U-251 cells engineered to express doxycycline-inducible shRNAs targeting either *TERT* (shTERT.3952) or GABPB1L (shGB1L.968) compared with a control (shRen.713). Following tumor engraftment, mice were placed on doxycycline and treated with a regimen of either TMZ or vehicle, then either assessed 30 d postxenograft for immunohistochemistry (Cohort 1) or followed to endpoint for survival analysis (Cohort 2) (Fig. 4E). We verified that doxycycline and TMZ do not interfere with each other to affect either tumor growth or shRNA induction in vivo (*SI Appendix, Fig. S9 I and J*). Interestingly, knockdown of *TERT* and GABPB1L both slowed growth of tumors and prolonged survival to a similar extent, which was accompanied by a significant reduction in the percentage of Ki-67 positive tumor cells in these conditions (Fig. 4 F–H and *SI Appendix, Fig. S9 K–M*). More importantly, both *TERT* and GABPB1L reduction in established intracranial xenografts resulted in a similar potentiation of TMZ efficacy, such that tumor growth was either dramatically slowed or eliminated (Fig. 4H and *SI Appendix, Fig. S9 K and L*). The median animal survival was increased beyond a simple addition of effects (Fig. 4I), showing the signature of a synergistic combination therapy. Ki-67 status was not affected by TMZ treatment, as assessed 14 d following the TMZ regimen (Fig. 4G and *SI Appendix, Fig. S9M*). Together, the dramatic anti-tumor effects of TMZ combined with GABPB1L reduction and the associated cancer-specific down-regulation of *TERT* mRNA are of significant clinical interest, as they provide a path to increased TMZ response for *TERT*^p mutant GBMs. Indeed, our data suggest that MGMT-silenced *TERT*^p mutant GBM could be effectively targeted through a combination of GABPB1L reduction and TMZ treatment.

Discussion

A major challenge in GBM therapy is finding efficient methods to reduce growth of established tumors while leaving normal cells unaffected. Here, our data directly demonstrate increased binding of GABPB1L-isoform-containing complexes to the mutant compared with WT *TERT* promoter. Additionally, we show that GABPB1L knockdown in vivo, and full knockout in vitro, result in near-term anti-growth effects on *TERT*^p mutant cells and that up-regulation of GABPB2 can compensate for the loss of GABPB1L in both contexts. Mechanistically, the early appearance of these anti-growth effects hints toward non-canonical functions of *TERT* rather than telomere attrition or immediate vulnerability of cells with short telomeres, though this warrants further investigation (9–11, 15, 45). Additionally, these effects may be the direct result of *TERT* down-regulation following GABPB1L reduction, a combined effect between reduced *TERT* and other factors associated with GABPB1L loss, or involve other differences in gene expression specific to *TERT*^p mutant cells (55). Regardless, our results provide evidence that

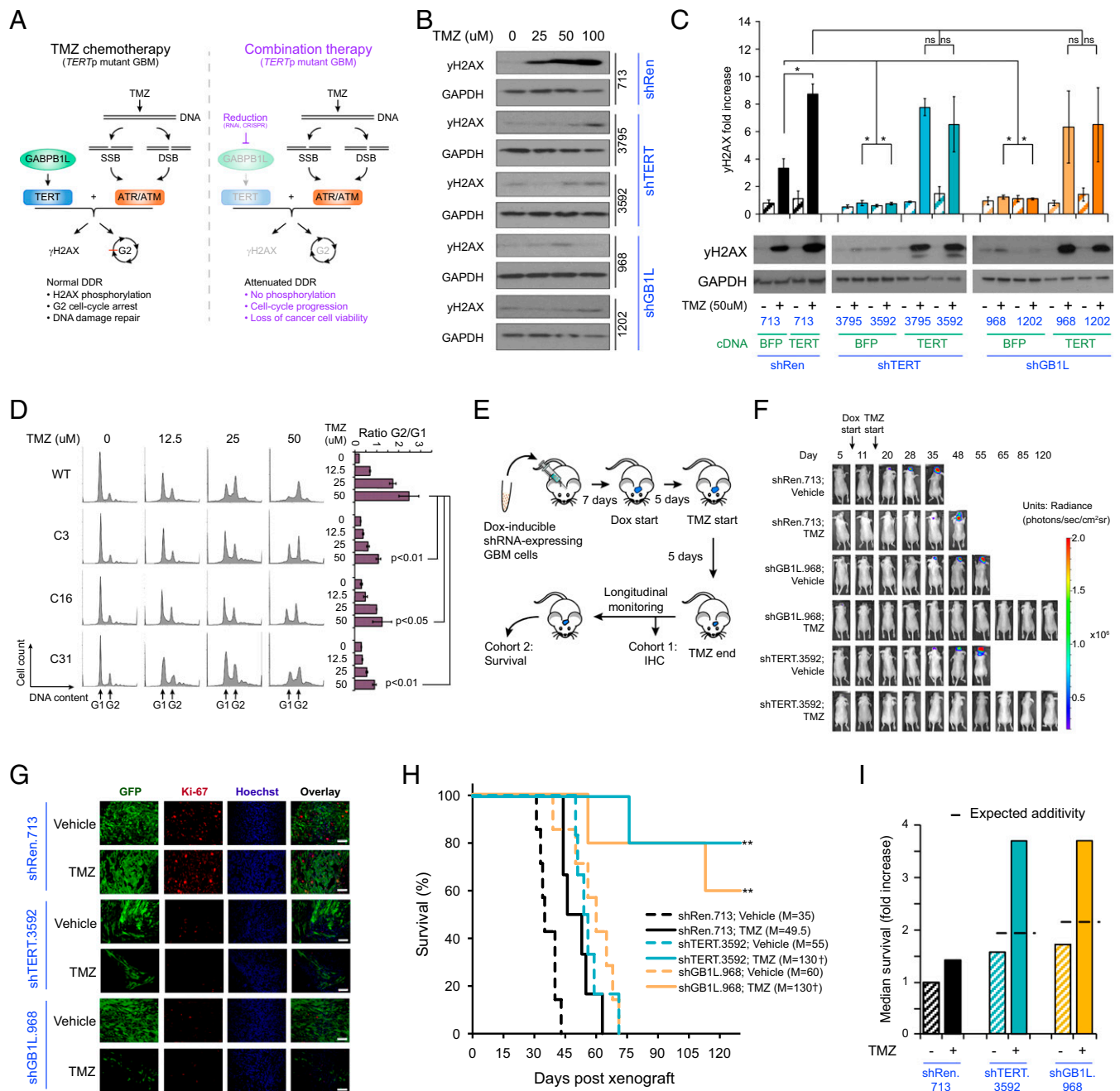


Fig. 4. Reduction of GABPB1L potentiates anti-tumor response of *TERT*p mutant GBM to chemotherapy. (A) A schematic for possible sensitization of *TERT* promoter mutant GBM cells to TMZ through GABPB1L inhibition. Inhibiting GABPB1L leads to reduced *TERT* expression, resulting in a blunted DDR and lack of normal cell cycle arrest, ultimately reducing cancer cell viability following DNA damage. SSB, single-strand break; DSB, double-strand break; DDR, DNA-damage response. (B) Representative immunoblots of yH2AX in U-251 cells expressing doxycycline-induced shRNAs targeting *TERT* (shTERT.3795 and shTERT.3592) or GABPB1L (shGB1L.968 and shGB1L.1202) compared with a nontargeting (renilla luciferase, shRen.713) shRNA. (C) Representative immunoblots (Bottom) and quantification (Top, from triplicate blots) of yH2AX in U-251 cells engineered to stably express either BFP (control) or *TERT* (rescue) in the presence of shRNAs targeting *TERT* (shTERT.3795 and shTERT.3592) or GABPB1L (shGB1L.968 and shGB1L.1202) compared with a nontargeting shRNA (renilla luciferase, shRen.713). (B and C) Cells were incubated with doxycycline for 6 d prior to harvest and treated with specified dose(s) of TMZ 20 h prior to harvest. (D) Representative images (Left) and G2/G1 ratio quantification (Right) from flow cytometry-based cell cycle analysis of U-251 WT and GABPB1L heterozygous knockout cells treated with a dose titration of TMZ 72 h prior to harvest. *p*, *P* value (unpaired two-tailed Student's *t* test). (E) A schematic of in vivo TMZ treatment. Mice were xenografted with U-251 cells expressing doxycycline-inducible shRNAs and placed on doxycycline chow 7 d postorthotopic xenograft. Mice were then treated with TMZ or a vehicle control by oral gavage starting 12 d postxenograft for 5 d. IHC, immunohistochemistry. (F) Bioluminescence imaging of mice injected with U-251 cells engineered to express shRNAs targeting *TERT* (shTERT.3592), GABPB1L (shGB1L.968), or a nontargeting control shRNA (shRen.713). Mice were placed on doxycycline chow and per os (p.o.) dosed with either TMZ or a vehicle control as described in E. (G) Representative images of immunofluorescence staining in tumors from mice described in E, cohort 1. Mice were euthanized 30 d postorthotopic xenograft for analysis. Doxycycline-induced tumor cells are GFP positive. *n* = 5 images per mouse, three mice per condition. (Scale bar, 50 μ m). (H) Kaplan–Meier survival curves for mice described in E, cohort 2. For statistics, survival of mice with no tumor burden was set at the experimental endpoint (130 d). *n* = 5 to 7 mice per condition. ***P* < 0.01, relative to all vehicle conditions (Kaplan–Meier log-rank test). M, median survival. (I) A comparison of the relative fold increase in median survival based on overall survival in H. The dashed line represents a simple addition of effects of TMZ plus GABPB1L or *TERT* knockdown. The survival of mice xenografted with WT cells and treated with vehicle control was used for normalization.

even after tumor initiation, GABPB1L knockdown rapidly impairs growth of GBM tumors *in vivo*, which is of great therapeutic interest.

More strikingly, we also find that GABPB1L reduction combined with TMZ treatment dramatically potentiates anti-tumor effects *in vivo* through cancer-specific loss of TERT, emphasizing the clinical potential of this combination therapy. Specific targeting of TERT through the long isoform of GABPB1 (GABPB1L) is particularly promising given that both GABPA and total GABPB1 (the short and long isoform together) are required for normal murine development, while the GABPB1L isoform is dispensable (9, 31, 32). Previous studies have observed that TERT reduction sensitizes cells to chemotherapy and radiation by inhibiting the normal DDR, measured in part by a loss of γ H2AX increase (16). Here, our data show that reducing TERT through knockdown of GABPB1L prevents both the induction of γ H2AX and the G2 cell cycle arrest normally seen following exposure of *TERT*^p mutant cells to standard-of-care TMZ chemotherapy. The mechanism of action of TMZ is largely through mismatch repair pathway-associated single-strand breaks but also results in DNA double-strand breaks, and γ H2AX signaling may occur in response to single-strand breaks as well as canonically to double-strand breaks (46, 47, 56, 57). Loss of γ H2AX signaling leads to reduced cell viability following DNA damage induced by radiation and chemotherapy (48, 49, 54). Our findings extend this by demonstrating that *TERT*^p mutant GBM tumors with reduced *TERT* mRNA, either through knockdown of GABPB1L or by targeting TERT directly, show a dramatically increased response to TMZ *in vivo*.

Targeted cancer therapies have the potential to be highly impactful, but cancer cells commonly develop resistance to anti-tumor drugs (58). Uncovering mechanisms of resistance often allows development of second-generation therapies that stunt tumor relapse. Our work points to sporadic GABPB2 up-regulation as a possible resistance mechanism to loss of GABPB1L, providing preliminary evidence that GABPB2, which is typically expressed only at very low levels in GBM, could be targeted in conjunction with GABPB1L for increased treatment robustness. Multiple other transcription factors have also been linked to TERT regulation, both in normal and oncogenic contexts (59–62). Such factors could play an additional role in basal expression of TERT from the mutant promoter or compensate for loss of GABPB1L and merit further study. While clinical methods to inhibit GABPB1L do not currently exist, and transcription factors such as the GABP complex are generally considered difficult to target, viable pathways might include approaches such as anti-sense oligo therapy, which has been successful at targeting transcription factors through regulation of splicing (63), or development of small molecules that prevent GABPA₂B₂ heterotetramer formation. The possible clinical benefit of these approaches warrants further study. Together, our data suggest that GABPB1L inhibition combined with TMZ treatment can provide a tumor-specific path to improve disease outcomes for patients with *TERT*^p mutant GBMs and possibly the many other cancers harboring *TERT*^p mutations.

Materials and Methods

See *SI Appendix, Methods* for additional experimental details. See *SI Appendix, Table S1* for sequence information for sgRNAs, shRNAs, oligonucleotides, and vectors.

Protein Purification. GABPB1L and GABPA subunits were cloned into the pET-Duet-1 vector with Gibson assembly. The GABPB1L subunit was inserted after the first ribosome binding site and a plasmid-encoded HisTag. The HisTag and GABPB1L were separated by insertion of a tobacco etch virus (TEV) cleavage sequence. GABPA was inserted after the second ribosome binding site without any affinity purification tags.

Mammalian Cell Culture. HEK293T human kidney cells (293FT; Thermo Fisher Scientific, #R70007; RRID: [CVCL_6911](#)), NHA-PC5 normal human astrocyte cells (a kind gift from Russell Pieper, University of California, San Francisco [UCSF]), and derivatives thereof were grown in Dulbecco's Modified Eagle Medium (DMEM; Corning Cellgro, #10-013-CV) supplemented with 10% fetal bovine serum (FBS; Seradigm #1500-500), and 100 Units/mL penicillin and 100 μ g/mL streptomycin (100-Pen-Strep; Gibco, #15140-122). U-251 human GBM cells (Sigma-Aldrich, #09063001; RRID: [CVCL_0021](#)), LN-229 human GBM cells (American Type Culture Collection [ATCC], #CRL-2611; RRID: [CVCL_0393](#)), T98G human GBM cells (ATCC, #CRL-1690; RRID: [CVCL_0556](#)), LN-18 human GBM cells (ATCC, #CRL-2610; RRID: [CVCL_0392](#)), and derivatives thereof were cultured in DMEM/Nutrient Mixture F-12 (DMEM/F-12; Gibco, #11320-033 or Corning Cellgro, #10-090-CV) supplemented with 10% FBS and 100-Pen-Strep. HAP1 cells (a kind gift from Jan Carette, Stanford University) and derivatives thereof were grown in Iscove's Modified Dulbecco's Medium (IMDM; Gibco, #12440-053 or HyClone, #SH30228.01) supplemented with 10% FBS and 100-Pen-Strep. HAP1 cells had been derived from the near-haploid chronic myeloid leukemia cell line KBM7 (64).

Establishment of a Puromycin-Sensitive Normal Human Astrocyte Cell Line. The normal human astrocyte cell line NHA-PC5 was previously described (45) and is puromycin resistant. For our experiments, we established a puromycin-sensitive monoclonal derivative, termed NHA-S2, through CRISPR editing of the previously inserted puromycin-resistance cassette.

Plasmid and Lentiviral Vectors. To generate monoclonal knockout cell lines, sgRNAs and Cas9 were expressed from either the pCF120 or pCF123 vectors. In brief, the plasmid vector pCF120, expressing a U6-promoter-driven sgRNA and an EF1-alpha short (EFS)-promoter-driven Cas9-P2A-Hygro cassette was derived from pX459 (65) by replacing the chicken beta-actin promoter with an EFS promoter from pCF204 (66), swapping the T2A-Puro cassette with a P2A-Hygro cassette and exchanging the guide RNA scaffold for a more efficient variant (67). The plasmid vector pCF123, expressing a U6-driven sgRNA and an EFS-driven Cas9-T2A-Puro cassette, was generated based on pX459 by replacing the chicken beta-actin promoter with an EFS promoter and exchanging the guide RNA scaffold for a more efficient variant (67).

For CRISPR-Cas9-mediated competitive proliferation assays, sgRNAs were expressed from the previously established pCF221 lentiviral vector and Cas9 from the pCF226 lentiviral vector (66). For low-level stable overexpression of *GABPB2* (NCBI gene ID: 126626) and mTagBFP2 (negative control), we first modified the pCF525-mTagBFP2 lentiviral vector (68), encoding an EF1a-Hygro-P2A-mTagBFP2 cassette, by replacing the strong EF1-alpha promoter with a weak EFS promoter. This new vector, EFS-Hygro-P2A-mTagBFP2, was named pCF554. Next, we cloned the coding sequence of homo sapiens GA binding protein transcription factor subunit beta 2 (*GABPB2*, transcript variant 1, and mRNA (NCBI reference sequence: NM_144618.2) into pCF554 by replacing mTagBFP2, yielding lentiviral vector pCF553.

For doxycycline-inducible, microRNA-embedded miR-E shRNA expression, both *in cell culture* and *in vivo*, we used the lentiviral vector LT3GEGPIR (36).

Lentiviral Transduction. Lentiviral particles were produced in HEK293T cells using polyethylenimine (Polysciences #23966)-based transfection of plasmids, as previously described (66).

Design of sgRNAs for CRISPR-Cas9 Genome Editing. Standard sgRNA sequences were either designed manually or using GuideScan (69). To generate specific genomic excisions rather than indels, pairs of sgRNAs were designed and used in tandem. This strategy was applied to remove sequences specific to the long isoform of *GABPB1L* while not affecting the short isoform.

CRISPR-Cas9 Competitive Proliferation Assay. CRISPR-Cas9 competitive proliferation assays were used to assess whether Cas9-mediated editing of genes of interest leads to a change in proliferation speed compared with unedited cells. First, cell lines (U-251) were stably transduced with a lentiviral vector expressing Cas9 (pCF226) and selected on puromycin (1.0 to 2.0 μ g/mL) as previously described (66). Subsequently, Cas9-expressing cell lines were further stably transduced with pairs of lentiviral vectors (pCF221) expressing various mCherry-tagged sgRNAs.

Design of shRNAs for Reversible Target Inhibition. MicroRNA-embedded miR-E shRNA sequences were predicted using SplashRNA (38) and cloned into LT3GEGPIR (36), an all-in-one doxycycline-inducible miR-E shRNA expression vector. The shRNA numbers refer to the position of the shRNA, specifically the 3' nucleotide of the guide strand, on the target transcript (38, 70). All

shRNA sequences are shown (SI Appendix, Table S1) and were cloned as previously described (70).

Generation of Isogenic GABPB1 Knockout Cell Lines. To generate isogenic monoclonal knockout cell lines of the long isoform of GABPB1 (NCBI gene ID: 2553), pairs of sgRNAs flanking the long isoform-specific exon 9 of *GABPB1* were used.

GABPB1 locus editing was assessed by genotyping with the primers oCF862_GAL-fw1 (AAAAGTACAGGTGCCAGTTTG) and oCF863_GAL-rev2 (GCCTAACCAACAACGATCAC), yielding an 808-base-pair band in WT cells. Where mentioned, Sanger sequencing of the GABPB1 locus and Tracking of Indels by Decomposition analysis (71) were also carried out using the primers oCF862_GAL-fw1 and oCF863_GAL-rev2.

Telomerase Repeated Amplification Protocol. Cy5-based telomerase repeated amplification protocol (TRAP) assays were performed as previously described (72). Briefly, 150,000 cells per condition were collected via trypsinization and resuspended in Nonidet P-40 lysis buffer (10 mM Tris HCl pH 8.0, 1 mM MgCl₂, 1 mM ethylenediaminetetraacetic acid (EDTA), 1% Nonidet P-40, 0.25 mM sodium deoxycholate, 10% glycerol, 150 mM NaCl, 5 mM 2-mercaptoethanol, and 0.1 mM AEB5F) at a concentration of 2,500 cells/ul to maintain telomerase activity. A total of 1 μL of lysate (or 1 μL of Nonidet P-40 lysis buffer for the negative control) was added to a PCR master mix containing a Cy5-labeled telomeric substrate primer, an internal standard control, and an extension primer mix at the described ratios (72).

Orthotopic Xenografts and Bioluminescence Imaging. Animal procedures conformed to care guidelines approved by the UCSF Institutional Animal Care and Use Committee (protocol numbers: AN179550, AN17599, and AN179565). U-251 and LN-229 cell lines were stably transduced with Firefly Luciferase Lentiviral Purified Lentiviral Particles (Genecopoeia, #LPP-FLUC-Lv105) at a multiplicity of infection (MOI) of 3. All cell lines were verified to be stably expressing luciferase at similar levels in vitro 48 h prior to xenograft. Either 4 × 10⁵ U-251 cells or 1 × 10⁵ LN-229 cells were injected in a 4 μl total volume into the right frontal cortex (mediolateral [ML]: 1.5 mm, anteroposterior [AP]: 1 mm, and dorsoventral [DV]: -3.5 mm, relative to bregma) of 6 to 7 wk old female athymic *nulnu* mice (inducible shRNA TMZ combination experiment: The Jackson Laboratory, RRID: IMSR_JAX:002019; all other experiments: Envigo, Hsd:ATHymic Nude-Foxn1^{nu}, #069).

Imaging and Analysis. For immunofluorescence-based analysis of GFP, Ki-67, and glial fibrillary acidic protein (GFAP) expression in inducible shRNA-expressing U-251 tumors treated with vehicle or TMZ, animals were euthanized 30 d postxenograft to compare across conditions at identical time-points. Animals were perfused and brains fixed in 4% paraformaldehyde (PFA) for 24 h. Brains were then washed in phosphate-buffered saline (PBS) and soaked in 30% sucrose prior to freezing in optimal cutting temperature (OCT) compound blocks for cryosectioning. Slides were washed in PBS to

remove OCT, blocked in 5% bovine serum albumin + 0.3% Triton-X in PBS for 1 h, then incubated with primary antibodies toward Ki-67 (Thermo Fisher, #PA5-19462) or GFAP (Abcam, #ab4674-50) overnight, followed by secondary antibodies for 1 h (Alexa Fluor 647, Abcam #ab150171 or Alexa Fluor 555, and Thermo Fisher #A27039). Slides were incubated with Hoechst stain for 10 min prior to mounting. GFP immunofluorescence was detectable without the need for antibody staining. Imaging was performed using a Keyence digital microscope by a researcher blinded to experimental condition. Then, 2× representative images were taken of the entire tumor field for each of three mice per condition, and 40× images near the tumor edges were obtained for analysis of Ki-67 expression, since Ki-67 expression is highest at the tumor edge and is used as a pathology-based marker for tumor grade. Ki-67 image quantification was carried out using similar methods to previously published work (73–75).

Statistical Analysis. Statistical analyses were performed using GraphPad Prism 9 software. Two-tailed, unpaired Student's *t* tests were used with alpha = 0.01 or alpha = 0.05, as specified. Statistical significance between Kaplan Meier survival curves was performed using a Kaplan Meier log-rank test with alpha = 0.01 or alpha = 0.05, as specified. The Wilcoxon signed-rank test was used for comparison of GABPB1 and GABPB2 expression levels from TCGA RNA sequencing datasets. Statistical analysis for qPCR data were performed among specific samples, rather than relative to WT control, due to normalization on WT control using the deltaCT method. Quantified data represent mean ± SD or mean ± SEM, as referenced in the figure legends. Values of "n" are listed in relevant methods and/or figure legends.

Data Availability. All study data are included in the article and/or supporting information.

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