



ZIC1 is a context-dependent medulloblastoma driver in the rhombic lip

Received: 22 March 2023

Accepted: 23 October 2024

Published online: 3 January 2025

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Transcription factors are frequent cancer driver genes, exhibiting noted specificity based on the precise cell of origin. We demonstrate that *ZIC1* exhibits loss-of-function (LOF) somatic events in group 4 (G4) medulloblastoma through recurrent point mutations, subchromosomal deletions and mono-allelic epigenetic repression (60% of G4 medulloblastoma). In contrast, highly similar SHH medulloblastoma exhibits distinct and diametrically opposed gain-of-function mutations and copy number gains (20% of SHH medulloblastoma). Overexpression of *ZIC1* suppresses the growth of group 3 medulloblastoma models, whereas it promotes the proliferation of SHH medulloblastoma precursor cells. SHH medulloblastoma *ZIC1* mutants show increased activity versus wild-type *ZIC1*, whereas G4 medulloblastoma *ZIC1* mutants exhibit LOF phenotypes. Distinct *ZIC1* mutations affect cells of the rhombic lip in diametrically opposed ways, suggesting that *ZIC1* is a critical developmental transcriptional regulator in both the normal and transformed rhombic lip and identifying *ZIC1* as an exquisitely context-dependent driver gene in medulloblastoma.

Malignant transformation of the human rhombic lip results in medulloblastoma, with group 3 (G3), group 4 (G4) and sonic hedgehog (SHH) tumors arising from the upper rhombic lip, and wingless/integrated (WNT) medulloblastoma arising from the lower rhombic lip^{1–13}. There are a number of well-known driver genes for medulloblastoma, particularly SHH pathway genes in SHH medulloblastoma. However, G4 medulloblastoma is less well understood, with mutations of histone modifier genes, members of the *CBFA* complex and amplifications of *MYCN* and *OTX2* (refs. 3,14). A tail of less well understood but recurrent somatically altered genes has been observed across medulloblastoma subgroups¹⁴.

The zinc finger protein in the cerebellum (ZIC) family of transcription factors (TFs) has crucial roles in the development of the central nervous system (CNS), including hindbrain development^{15–17}. There are five human ZIC family genes (*ZIC1*–*ZIC5*), all of which contain conserved tandem C2H2 zinc finger motif repeats that can interact with DNA or other proteins^{15–18}. While ZICs exhibit some overlapping expression patterns throughout the CNS, different mutations are associated with distinct congenital disorders^{15,16,19}. Somatic mutations of *ZIC1* have been identified in distinct medulloblastoma subgroups, and although *ZIC1* is a pan-medulloblastoma master TF associated with an active super-enhancer (SE)²⁰, the specific role of ZIC TFs in the etiology of medulloblastoma is obscure.

ZIC1 and *ZIC4* have multiple critical roles in cerebellar development^{15,16,21}. Heterozygous deletion of the *ZIC1*/*ZIC4* locus in humans²² is a rare cause of Dandy–Walker malformation (DWM), which includes cerebellar hypoplasia¹⁶. Gain-of-function (GOF) mutations at the carboxy terminus of *ZIC1* have been identified in children with craniosynostosis and learning disabilities²³. We now demonstrate that *ZIC1* mutations in medulloblastoma are context dependent, with loss-of-function (LOF) mutations and epigenetic alterations in G4 medulloblastoma, contrasted with GOF mutations in SHH medulloblastoma. Concordantly, expression of *ZIC1* represses malignant phenotypes in G3/G4 medulloblastoma while enhancing malignant phenotypes in SHH medulloblastoma in model systems. *ZIC1* is therefore a stark example of how the same gene can have distinct driver mechanisms in highly similar cancers depending on their specific lineage of origin.

Results

The subgroup-specific H3K27ac/H3K27me3 landscape of medulloblastoma

Due to the high prevalence and recurrence of somatic mutations in genes associated with chromatin modulation in medulloblastoma (~30% of medulloblastomas)¹⁴, we hypothesized that some medulloblastomas

might acquire somatic histone modification alterations (chromatin variants^{24,25}) for driver genes. To test this hypothesis, we profiled H3K27ac and H3K27me3 landscapes across the four medulloblastoma subgroups (including 123 matching samples for H3K27ac and 63 matching samples for H3K27me3) and integrated the data with matching RNA sequencing (RNA-seq), as well as an independent cohort of tumors characterized by H3K27ac ChIP (Fig. 1a, Extended Data Fig. 1a and Supplementary Tables 1 and 2). Hierarchical clustering using either H3K27ac or H3K27me3 chromatin immunoprecipitation followed by sequencing (ChIP-seq) data recapitulated the four subgroups (Fig. 1b). We categorized subgroup-specific H3K27 modification as either subgroup-enriched peaks (signal enrichment) or subgroup-recurrent peaks (peak called recurrently for one subgroup; Fig. 1c–e). A subset of the identified peaks was shared by either SHH/WNT (enriched in SHH versus G3 or G4, but not WNT) or G3/G4 (enriched in G3 versus SHH or WNT, but not G4; Fig. 1d, e) and were documented as such.

The average number of peaks and the proportion of genome coverage for H3K27ac did not significantly differ between subgroups (Fig. 1f). However, H3K27me3 deposition was markedly increased in G3 medulloblastoma (Fig. 1f). Additionally, G3/G4 medulloblastoma-enriched H3K27me3 peaks exhibited a strong preference for gene promoters as compared to WNT/SHH (Extended Data Fig. 1b). Core regulatory circuit analysis of H3K27ac ChIP-seq data identified known and new medulloblastoma subgroup-specific master TFs, including the pan-subgroup master TFs *ZIC1* and *ZIC4* as we reported previously (Extended Data Fig. 1c–e)²⁰. Additionally, H3K27ac ChIP was used to define the enhancer–promoter interactome across medulloblastoma subgroups (Fig. 1g). Integration of H3K27ac ChIP, H3K27ac ChIP-seq and RNA-seq allowed the identification of loops connecting enhancers and promoters of protein-coding genes. Among the enhancer–promoter interacting loops, those with enhancer H3K27ac read counts exhibiting significant positive correlations with the expression of target genes were also identified (adjusted $P < 0.1$) and defined as significantly correlated loops (SCL; Fig. 1g,h). Many SCL-associated enhancers target more than one gene (Extended Data Fig. 1f), and notably, enhancers frequently target genes that are not the most proximal gene (Extended Data Fig. 1g).

We conclude that post-translational modification of H3K27 in medulloblastoma varies by subgroup.

Recurrent single-nucleotide variations (SNVs) and hemizygous H3K27me3 affect *ZIC1* in G4 medulloblastoma

We hypothesized that a subset of medulloblastoma LOF driver genes somatically altered by SNVs, small insertions/deletions (InDels) or copy number aberrations (CNAs) might also be targeted through somatic H3K27me3-mediated repression to achieve the common endpoint of tumor suppressor gene LOF. We determined the intersection between genes affected by genetic mutations and those overlapping either ‘enriched’ or ‘recurrent’ subgroup-specific H3K27me3 peaks (Fig. 2a and Extended Data Fig. 2a)¹⁴. While no overlapping genes were identified for WNT or G3, *BCOR* for SHH, and both *ZIC1* and *FLG* in G4 are affected by both mutation and H3K27me3-modified chromatin. H3K27me3 peaks on the *BCOR* promoter (chromosome Xp11.4) were found predominantly in female SHH tumors, suggesting a link to X chromosome inactivation (Extended Data Fig. 2b,c). Broadening the analysis to genes encompassed by focal deletions identified from our published Affymetrix SNP6 array data^{26,27} identified genes targeted by both deletions and H3K27me3, including the MIR4786 locus in G3 and G4 medulloblastoma (Extended Data Fig. 2d,e and Supplementary Tables 3–13).

The *ZIC1* and *ZIC4* genomic loci are separated by an interposed, shared, bidirectional promoter (Extended Data Fig. 2g). They are coregulated by a SE that is highly active across all four subgroups (Fig. 2b and Extended Data Fig. 2f,g). Both genes are highly expressed across all medulloblastoma subgroups as previously described²⁰, particularly in the G4 (Fig. 2c and Extended Data Fig. 2h). We now

describe a subset of G3 and G4 tumors that exhibit atypical hemizygous H3K27me3 deposition across the *ZIC1/ZIC4* SE locus while showing a robust H3K27ac mark in *trans* on the other allele (Fig. 2d,e). This pattern was associated with reduced *ZIC1/ZIC4* transcript levels (Fig. 2f) and was not recurrently observed in either SHH or WNT medulloblastoma (Fig. 2e). These two functionally opposing marks are usually mutually exclusive at the vast majority of loci, with the ‘H3K27ac–H3K27me3 hemizygous state’ being exceedingly rare (Fig. 2g). We hypothesized therefore that somatic repression of *ZIC1* through acquisition of the ‘H3K27ac–H3K27me3 hemizygous state’ is a chromatin-based driver event in G4 medulloblastoma.

To determine if the H3K27ac and H3K27me3 are indeed found in *trans* on separate alleles within the same cells, allelic frequencies for dbSNP151 annotated heterozygous single-nucleotide polymorphisms (SNPs) were examined in our H3K27ac and H3K27me3 libraries for samples harboring the H3K27ac–H3K27me3 hemizygous state at the *ZIC1/ZIC4* locus (Fig. 2h). While the G3 samples lacked heterozygous SNPs, all SNPs within the examined G4 samples exhibited a strong bias for distinct alleles in the H3K27ac versus H3K27me3 libraries (Fig. 2i), suggesting that the two chromatin marks occur in *trans* within single cells. Inferred SNPs were verified with matching whole-genome sequencing (WGS) data when possible (Extended Data Fig. 2i). While a plurality of G4 medulloblastomas alter activity of *ZIC1* through genetic mutation, an additional nonoverlapping cohort (Supplementary Table 1) of G4 tumors reduce *ZIC1/ZIC4* expression through uni-allelic chromatin variant repression mediated by H3K27me3 deposition, suggesting a convergence of mechanisms underlying *ZIC1* alteration and that *ZIC1* might be a LOF driver gene in G4 medulloblastoma.

Mono-allelic SEs regulate *ZIC1/ZIC4* expression in G3/G4 medulloblastoma

Our observation that the *ZIC1/ZIC4* locus undergoes recurrent repression in G4 medulloblastoma through hemizygous deposition of H3K27me3 on its SE prompted us to look for additional mono-allelic SEs in a cohort of 51 medulloblastoma tumors with matching H3K27ac ChIP-seq and WGS data (Fig. 3a). Mono-allelic SEs were rare in SHH medulloblastoma, although a number of further examples were identified for G3 and G4 medulloblastoma, including the known example of *PRDM6* enhancer hijacking in G4 (Fig. 3a)¹⁴. Of the 19 G4 medulloblastoma samples harboring heterozygous SNPs at the *ZIC1/ZIC4* SE locus (to allow assessment of heterozygosity), 9/19 tumors (47% of cases) exhibited a mono-allelic SE in keeping with the H3K27ac–H3K27me3 hemizygous state. A similar, albeit less frequent pattern, was observed in G3 medulloblastoma, but only very rarely in SHH medulloblastoma. Notably, samples with mono-allelic *ZIC1/ZIC4* SE exhibit expression of *ZIC1/ZIC4* mRNA predominantly from the H3K27ac allele (Extended Data Fig. 3a), in keeping with a bona fide repression effect of H3K27me3 deposition. Aside from the SE directly overlapping the *ZIC1/ZIC4* locus, several other genetically proximate SEs that target *ZIC1/ZIC4* were also identified to be recurrently mono-allelic (Extended Data Fig. 3b, c).

We determined the mono-allelic expression pattern of *ZIC1/ZIC4* in a validation cohort of 251 medulloblastomas with matching RNA-seq and WGS data, assembled by combining publicly available and newly generated datasets^{3,4,14,27,28}. We found frequent mono-allelic expression in G3 and G4, but neither SHH nor WNT medulloblastomas (Fig. 3b). Indeed, 55% of G4 tumors (36/65) and 24% of G3 tumors (7/29) exhibit mono-allelic expression of *ZIC1*, and 48.5% (33/68) of G4 tumors and 18.9% (7/37) of G3 tumors have mono-allelic expression of *ZIC4* (Fig. 3b and Extended Data Fig. 3d). In both G3 and G4, mono-allelic expression is associated with reduced expression of *ZIC1/ZIC4*, consistent with chromatin-based suppression (Fig. 3c). The importance of diminished, mono-allelic expression of the *ZIC1/ZIC4* locus in medulloblastomas arising from the rhombic lip is underscored by humans who have hypoplastic cerebella (DWM) secondary to germline hemizygous deletions of *ZIC1/ZIC4* (ref. 16). We conclude that haploinsufficiency of *ZIC1*

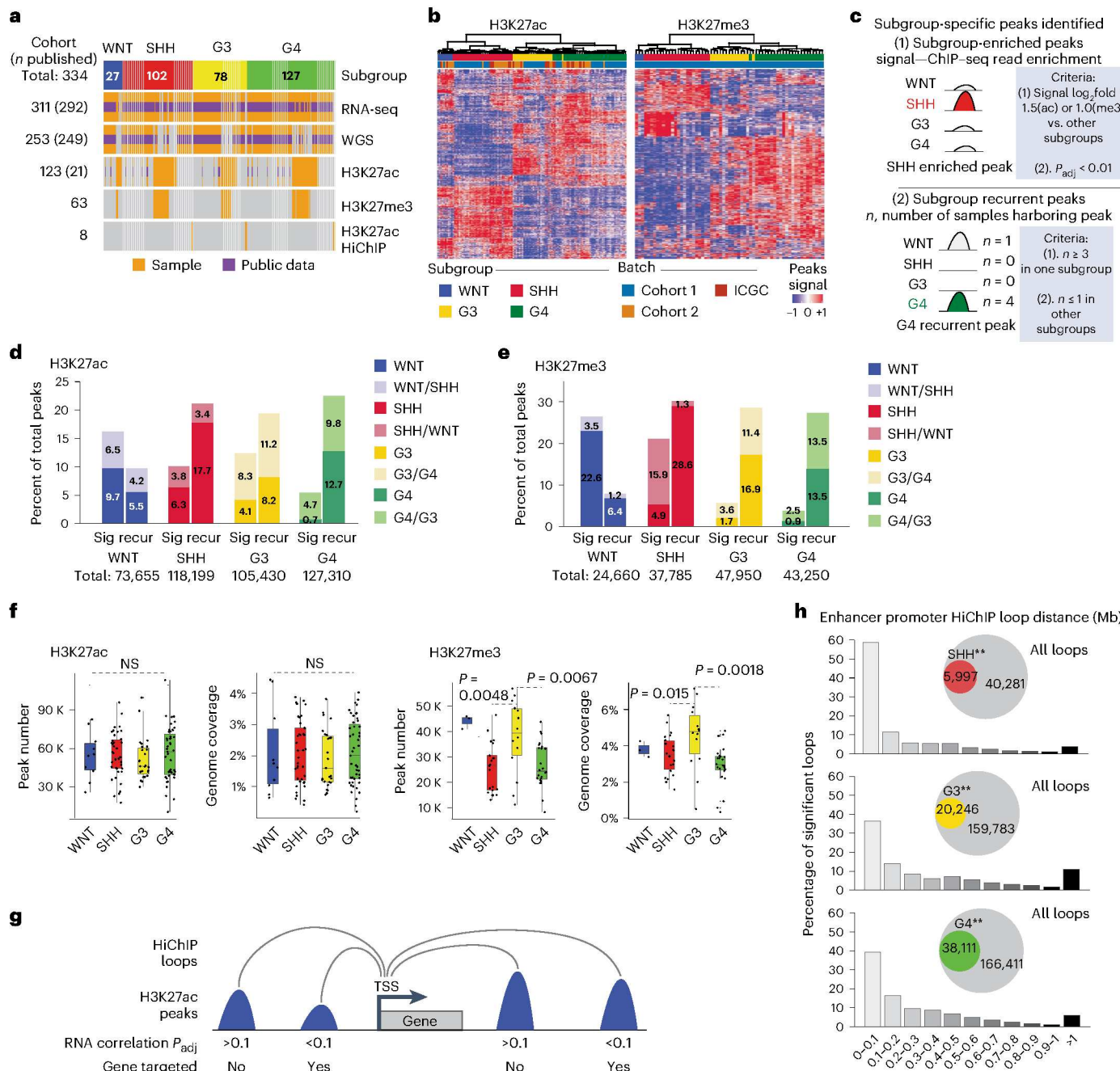


Fig. 1 | Characterization of subgroup-specific chromatin landscape of medulloblastoma. **a**, Summary of the newly generated and public datasets. Number within the bracket indicates the number of tumors with previously published data. **b**, Hierarchical clustering plots generated using the top 10,000 variable H3K27ac and H3K27me3 ChIP-seq peaks. **c**, Schematic representation summarizing different types of ChIP-seq peaks used in downstream analysis. Subgroup-specific peaks were defined by identifying peaks that (1) exhibit subgroup enrichment in ChIP-seq read counts or (2) are recurrently present only for specific subgroups even if the average ChIP-seq read count is not strongly subgroup enriched on average. **d**, Number of subgroup-specific peaks for each subgroup in the H3K27ac cohort. After batch correction, peaks annotated as subgroup enriched for ChIP-seq reads or subgroup recurrent were characterized separately. **e**, Number of subgroup-specific H3K27me3 peaks using the same annotations/criteria as **d**. **f**, Number of peaks and proportion of genome covered

by H3K27ac and H3K27me3 peaks across the medulloblastoma subgroups. P values were calculated by the tailed Mann–Whitney U test. Biological sample size for H3K27ac—G3/G4/SHH/WNT = 27/47/39/10 and H3K27me3—G3/G4/SHH/WNT = 14/24/22/3. Center of box, median. Bounds of box, 25% and 75% percentile. Whiskers show minimum and maximum values within the 1.5 \times interquartile range. **g**, Schematic representation summarizing how high-confidence enhancer–promoter interactions were identified from HiChIP and ChIP-seq data. Adjusted P values were calculated using Pearson correlation between target gene transcript and enhancer H3K27ac read levels, which was corrected for multiple testing. **h**, Summary of distance distribution for high-confidence enhancer–promoter interactions. Proportion of SCLs (**g**; Methods) over a total number of loops is depicted as overlapping Venn diagrams. Double asterisk (**) indicates a significant correlation ($P_{adj} < 0.1$).

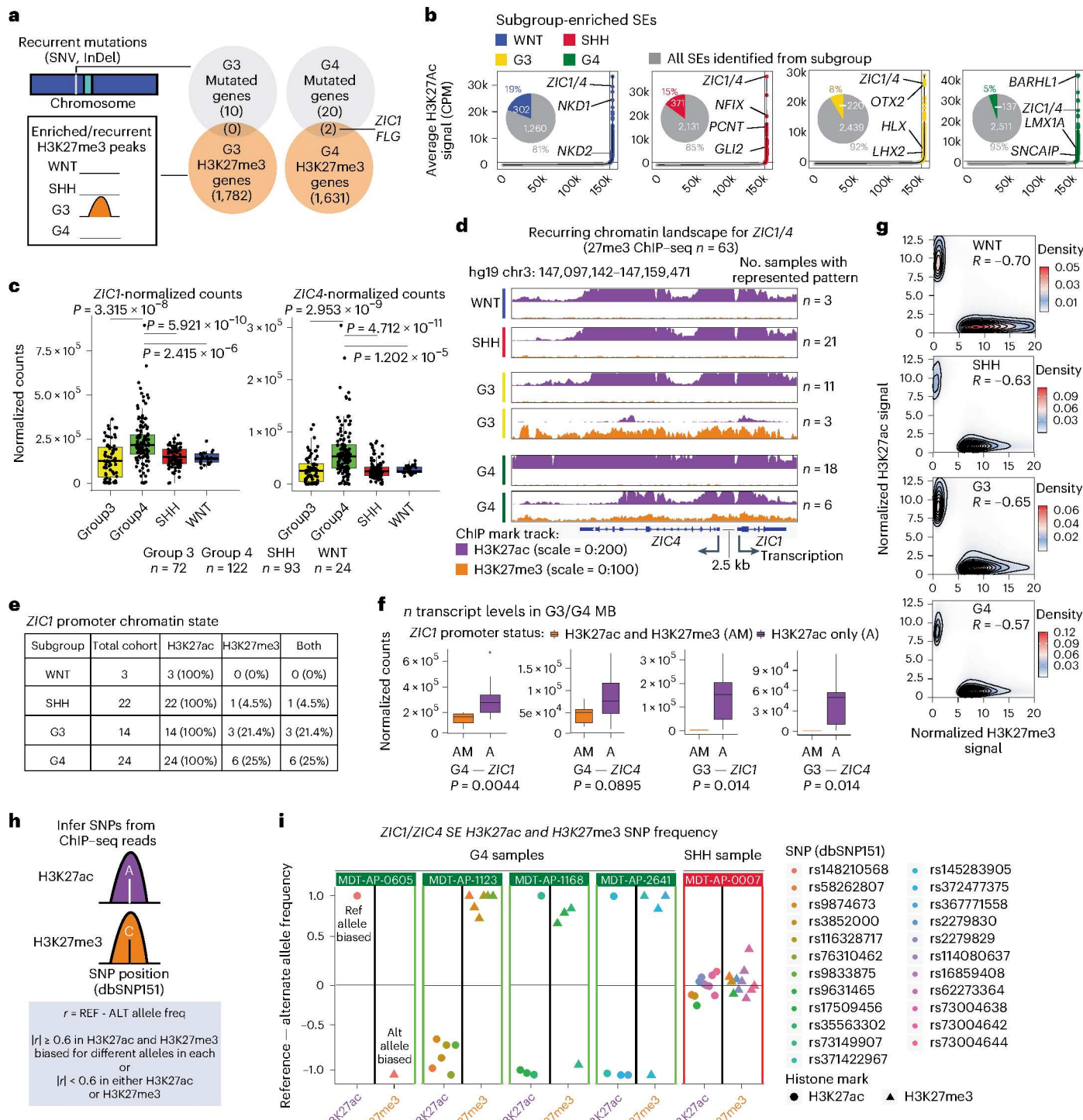


Fig. 2 | ZIC1 is recurrently mutated and repressed by H3K27me3 in G4 medulloblastoma. **a**, Overlap between genes regulated by subgroup-specific H3K27me3 peaks in G3, G4 medulloblastoma and genes recurrently mutated in each subgroup. **b**, Ranking of SEs across medulloblastoma subgroups, showcasing the number of total SEs identified (in gray) as well as the proportion of subgroup-enriched SEs in pie charts. **c**, ZIC1 and ZIC4 expression patterns across medulloblastoma subgroups. Biological sample size—G3/G4/SHH/WNT = 72/122/93/24. *P* values from two-tailed Mann–Whitney *U* test. Center of box, median. Bounds of box, 25% and 75% percentile. Whiskers show minimum and maximum values within 1.5× interquartile range. **d**, Sequencing depth normalized bigwig tracks showcasing recurrent ($n \geq 3$ per subgroup) ZIC1 and ZIC4 chromatin states across four subgroups. **e**, Summary of chromatin states observed at the ZIC1 promoter across all samples in the ChIP-seq libraries with

both H3K27ac and H3K27me3 modifications. **f**, Expression levels of ZIC1 and ZIC4 in G3/G4 medulloblastoma samples that harbor both H3K27ac and H3K27me3 (AM) or just H3K27ac (A) peaks on the ZIC1 promoter. Biological sample size for G4—AM/A = 6/18 (24 total) and G3—AM/A = 3/11 (14 total). *P* values from two-tailed Mann–Whitney *U* test. Same whisker box plot parameters as **c**. **g**, Density plot summarizing H3K27ac versus H3K27me3 signal at H3K27ac and H3K27me3 peaks. Correlations between H3K27ac and H3K27me3 were calculated by Pearson correlation on merged peak coordinates. **h**, Method for inferring heterozygous SNPs using H3K27ac and H3K27me3; two mutually exclusive histone modification marks. **i**, Distribution of inferred heterozygous SNPs across H3K27ac and H3K27me3 libraries of four G4 samples and one SHH sample with H3K27ac and H3K27me3 peaks on the ZIC1 promoter.

due to either germline or somatic events, with consequent diminished transcription, has critical effects on the biology of the rhombic lip, either *in toto* (DWM) or possibly in distinct somatic subclones (medulloblastoma).

ZIC1 is a presumed medulloblastoma driver gene that recurrently harbors SNVs in G4 and SHH medulloblastoma¹⁴. We now demonstrate that *ZIC1* mutations in G4 medulloblastoma are found in the DNA-binding zinc finger domain, whereas SHH medulloblastoma SNVs are found in the 3' end of the gene, encoding a carboxy-terminal intrinsically disordered region (IDR) of currently unknown function (Fig. 3d)¹⁴. Intriguingly, SHH medulloblastoma *ZIC1* somatic mutations are found in the same 3' region of the *ZIC1* gene as previously reported germline GOF *ZIC1* mutations in humans with craniostenosis²³. Within our 251 medulloblastoma validation cohort, three G4 tumors and two SHH tumors with *ZIC1* mutations were identified. In all three G4 tumors, the variant allele frequency (VAF) of mutants comprised nearly 100% of all *ZIC1* reads from RNA-seq, whereas they were below 50% in the matching WGS libraries (Fig. 3e). Conversely, SHH medulloblastoma mutants exhibited VAF near 50% in both WGS and RNA-seq reads. Examination of *ZIC1* VAF from our published medulloblastoma RNA-seq cohort^{3,27} produced similar results (Fig. 3f). These data are consistent with a model in which G4 medulloblastomas acquire LOF genetic and chromatin variants, while SHH medulloblastomas acquire GOF variants.

Mono-allelic *ZIC1* expression occurs in a subset of G4 medulloblastoma

PRDM6 overexpression secondary to a tandem duplication of the *SNCAIP* locus is a suspected G4 medulloblastoma driver gene¹⁴, and in our dataset it is found only in G4 tumors with mono-allelic expression of *ZIC1* or *ZIC4* (Fig. 4a and Extended Data Fig. 4a). G4 *ZIC1/ZIC4* mono-allelic samples were significantly enriched ($P = 0.0196$) for mutations in chromatin modifiers including *KDM6A*, *KMT2C* and *KMT2D* (Fig. 4b). In G3, *KMT2D* mutation was significantly enriched ($P = 0.0215$) in *ZIC1/ZIC4* mono-allelic samples (Fig. 4c,d). Conversely, *KBTBD4* InDel mutations were enriched ($P = 0.0041$) in G3/4 *ZIC1/ZIC4* bi-allelic samples (Fig. 4b,c). SHH tumors with *ZIC1* mutations always co-occurred with mutations of the *U1* splicing factor (Extended Data Fig. 4b), consistent with our previous publication in which *ZIC1* mutations were found in SHH α and SHH β tumors where *U1* mutations occur²⁷. Notably, we observe cases of G4 medulloblastoma with mono-allelic *ZIC1/ZIC4* expression but without H3K27me3 deposition, suggesting that additional cryptic genetic/epigenetic routes to allelic silencing of *ZIC1/ZIC4* exist (Fig. 4e-h). G3/G4 medulloblastoma tumors exhibit a spectrum of *ZIC1* expression levels as well as differentiation signatures (Supplementary Table 14), with G4 medulloblastoma exhibiting higher levels of both (Extended Data Fig. 4c,d), potentially rehighlighting the known role of *ZIC1* in cerebellar development²⁹.

One possible explanation for the H3K27ac–H3K27me3 hemizygous state is that it occurs naturally during the differentiation of the rhombic lip subventricular zone (RL-SVZ), where G4 medulloblastoma is thought to arise^{2,3}. However, hierarchical clustering of G3 and G4 medulloblastoma by both overall transcriptome or neuronal gene expression does not segregate tumors by *ZIC1/ZIC4* expression status, suggesting that the observed repression of the *ZIC1/ZIC4* locus from chromatin variants is not purely secondary to a transient developmental state in the RL-SVZ (Extended Data Fig. 4e,f). Mono-allelic *ZIC1/ZIC4* expression may also arise from local or distal mutations/structural variations affecting *ZIC1/ZIC4* transcription. However, mutational mining of the region surrounding the *ZIC1/ZIC4* locus for the presence of noncoding mutations that could account for the observed epigenetic repression failed to yield any likely candidates (Extended Data Fig. 4g,h). Taken together, we hypothesize that the acquisition of somatic mutations and/or aberrant activity of histone-modifying complexes may result in unusual regulation of the *ZIC1/ZIC4* locus, although this concept remains largely speculative.

Opposing *ZIC1/ZIC4* CNAs in G3/G4 versus SHH medulloblastoma

Previous studies have reported recurrent copy loss of chromosome 3q (chr3q), which contains the *ZIC1/ZIC4* locus, in G4 medulloblastoma^{26,30}. Examining CNAs at the *ZIC1/ZIC4* locus using published SNP6 array data²⁶ validates this finding and further showcases an intriguing pattern—the *ZIC1/ZIC4* locus was recurrently deleted in G3/G4; however, the same locus exhibits recurrent genomic gains in SHH (Fig. 5a), as determined by GISTIC³¹, and pairwise comparison of CNAs across subgroups (Fig. 5b,c). Frequencies of chr3q deletions and focal deletions harboring the *ZIC1/ZIC4* locus within G4 medulloblastoma were examined at the subtype level as we annotated previously³⁰. These deletions exhibited subtype specificity, being notably depleted in G4 β (Fig. 5d), whereas chromatin-based repression of the locus is very frequent in G4 β (Fig. 5e). Tumors that target *ZIC1* through either a genetic or a chromatin route show loss of heterozygosity at the level of mRNA (Fig. 5f,g). SHH samples affected by copy number gains exhibited concomitant increased expression of both *ZIC1* and *ZIC4* (Fig. 5h). SNP6 and expression array data^{26,30} demonstrate that G4 γ samples with focal and broad deletions of the *ZIC1/ZIC4* locus exhibit diminished expression of *ZIC1* and *ZIC4* transcripts as compared to balanced controls (Fig. 5i). Because the *ZIC1/ZIC4* locus can be targeted by both genetic- and chromatin-based mechanisms, we examined the overall proportion of samples within the validation cohort medulloblastomas (251 tumors with RNA-seq and WGS) affected by either chromatin or genetic variants. We identified the copy number status for the *ZIC1/ZIC4* locus within these samples using control-FREEC on the WGS data³². Annotating samples by *ZIC1/ZIC4* allelic expression status, copy gain within SHH, copy loss within G3/G4 medulloblastoma and *ZIC1* SNV status revealed that close to 20% of SHH samples harbor genetic variants promoting *ZIC1/ZIC4* expression (Fig. 5j). Conversely, approximately 33% of G3 and 60% of G4 samples harbored genetic/epigenetic variants associated with repression of *ZIC1/ZIC4* expression (Fig. 5j). These results are consistent with a model in which *ZIC1*, and possibly *ZIC4*, are LOF drivers in G4 medulloblastoma and GOF drivers in SHH medulloblastoma.

ZIC1/ZIC4 represses G3 medulloblastoma model growth *in vitro* and *in vivo*

Due to the lack of accurate, robust G4 medulloblastoma cell lines, we examined the functional importance of *ZIC1/ZIC4* by overexpressing blue fluorescence protein (BFP) empty vector, *ZIC1*, *ZIC4* or *ZIC1* and *ZIC4* together in D425 and D283 G3 medulloblastoma cell lines. Because G3 and G4 medulloblastomas are (1) molecularly similar and (2) exhibit highly similar genetic and epigenetic dysregulation of the *ZIC1/ZIC4* locus, G3 medulloblastoma cell lines were considered relevant for these experiments. Overexpression of *ZIC1* led to a significant reduction in the proliferative potential of D425 with evidence for some additive activity with *ZIC4* (Fig. 6a). Similar results were observed for D283 in a cell proliferation assay (Fig. 6b,c). Overexpression of *ZIC1/ZIC4* in G3 medulloblastoma lines followed by transcriptional profiling revealed increased expression of genes involved in neuronal differentiation, consistent with a model in which LOF of *ZIC1/ZIC4* might hinder differentiation (Fig. 6d). Cerebellar xenografting of NOD SCID γ (NSG) mice with D425 cells overexpressing *ZIC1/ZIC4* or BFP empty vector demonstrated a significant difference in both bioluminescence imaging (BLI) signal and survival (Fig. 6e-g). The patient-derived G3 xenograft, MB051, harbors single allele chromatin-based suppression of the *ZIC1/ZIC4* locus (Fig. 6h,i and Supplementary Table 15). Restoring *ZIC1/ZIC4* expression in MB051 significantly reduces BLI signal, as well as prolonging survival *in vivo* (Fig. 6j-m) in a setting with pre-existing *ZIC1/ZIC4* chromatin repression. Upon endpoint, *ZIC1* expression was minimal with the *ZIC1/ZIC4* overexpression construct (but higher than an empty vector), suggesting a possible negative selection for

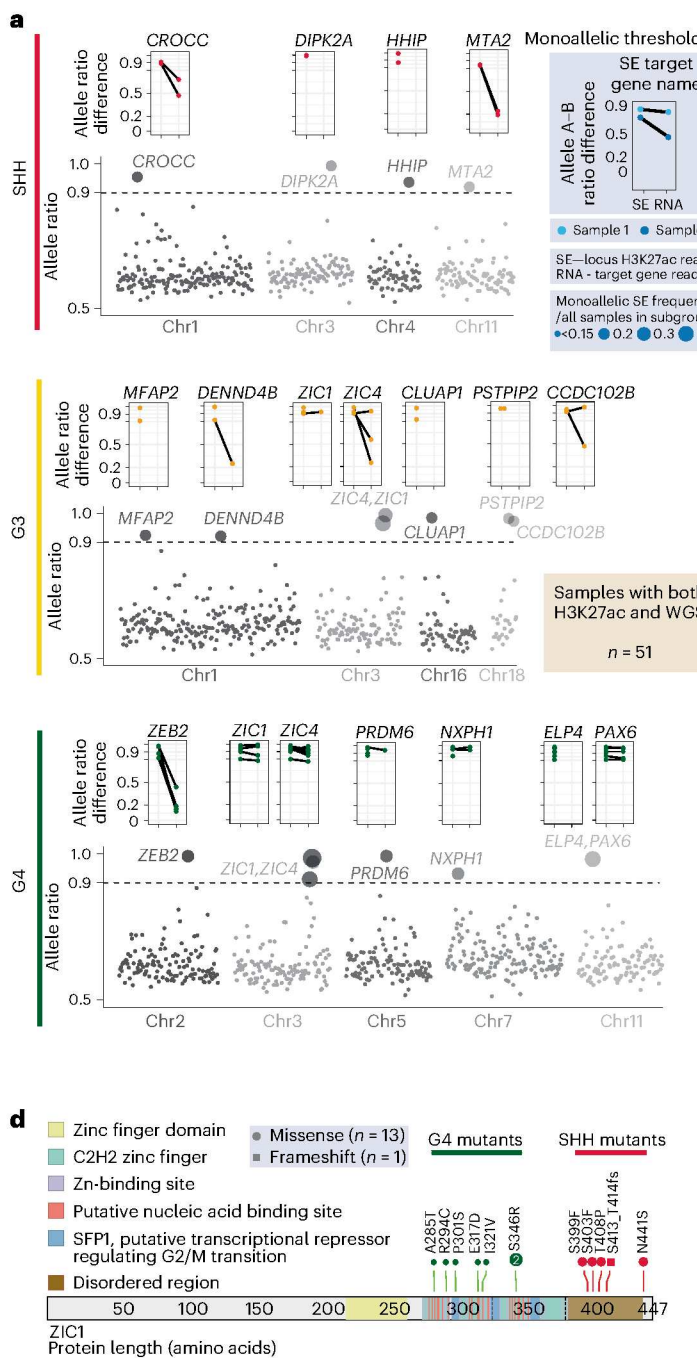
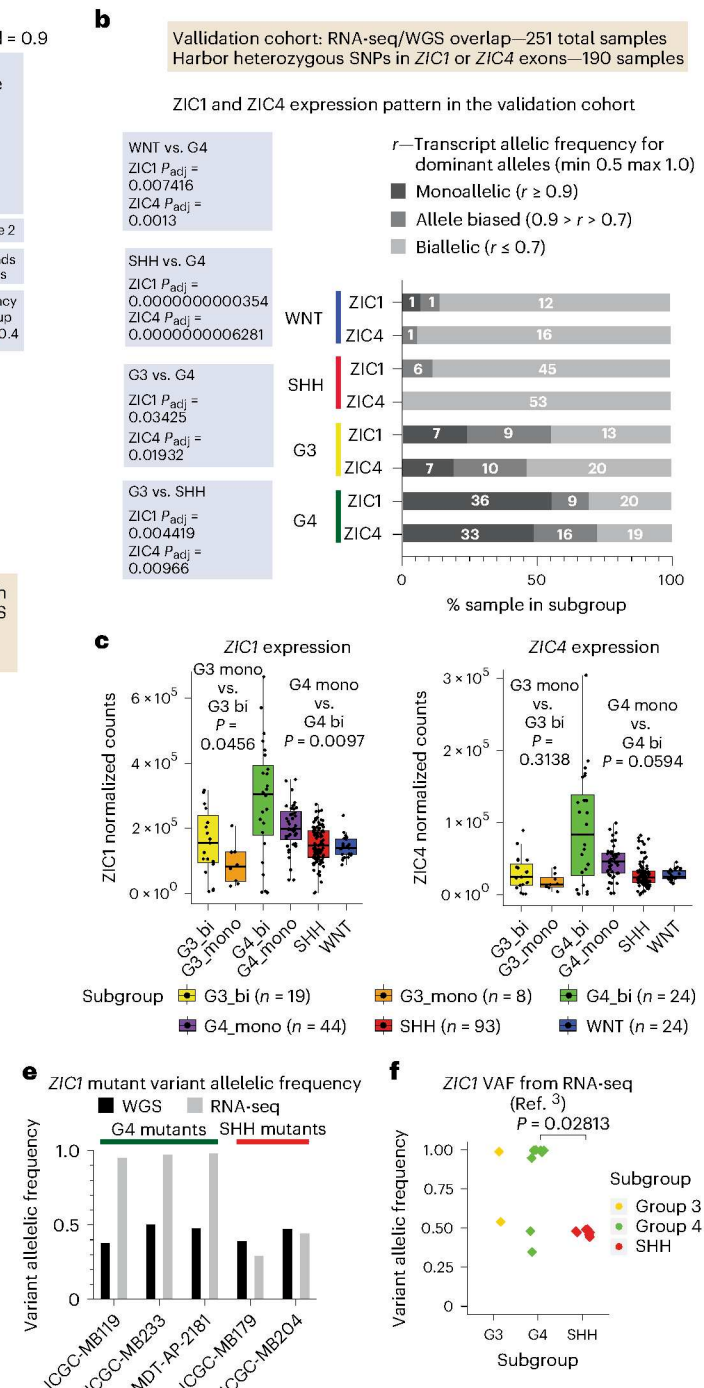


Fig. 3 | ZIC1/ZIC4 exhibit mono-allelic expression patterns in G3 and G4.
a, SEs that are recurrently ($n \geq 3$ for G4 and $n \geq 2$ for others) mono-allelic across different medulloblastoma subgroups. SEs that harbor SNPs (phased and pooled for each allele) that are heterozygous in WGS but homozygous (normalized allelic frequency ≥ 0.9) in H3K27ac ChIP-seq reads (same SNPs) from the same sample were defined as mono-allelic. Dot plots above each SE show differences in pooled allelic frequencies for heterozygous SNPs (allele A-B) in (1) H3K27ac reads from the SE (left) and (2) RNA-seq reads from the SE target gene (right). Matching samples are connected by lines between SE and RNA. **b**, Allelic frequency summary for heterozygous germline SNPs for ZIC1 and ZIC4 transcripts in RNA-seq within the validation cohort (251 samples with both WGS and RNA-seq data). Adjusted P values from two-tailed pairwise Fisher's exact test. **c**, Whisker box plots summarizing ZIC1 and ZIC4 expression across the medulloblastoma subgroups, but G3 and G4 are divided according to mono-allelic (mono) versus bi-allelic (bi) expression of ZIC1 or ZIC4. Biological sample size: G3.bi/G3_mono = 19/8, G4.bi/G4_mono = 24/44 and SHH/WNT = 93/24. P values from two-tailed Mann-Whitney U test. Center of box, median. Bounds of box, 25% and 75% percentile. Whiskers show minimum and maximum values within the $1.5 \times$ interquartile range. **d**, Mutational landscape of ZIC1 in G4 and SHH. **e**, Allelic frequency distribution for ZIC1 mutations in G4 ($n = 3$) and SHH ($n = 2$) samples from the assembled validation cohort. **f**, ZIC1 VAF obtained from published medulloblastoma RNA-seq data. P value from two-tailed Mann-Whitney U test.



exact test. **c**, Whisker box plots summarizing ZIC1 and ZIC4 expression across the medulloblastoma subgroups, but G3 and G4 are divided according to mono-allelic (mono) versus bi-allelic (bi) expression of ZIC1 or ZIC4. Biological sample size: G3.bi/G3_mono = 19/8, G4.bi/G4_mono = 24/44 and SHH/WNT = 93/24. P values from two-tailed Mann-Whitney U test. Center of box, median. Bounds of box, 25% and 75% percentile. Whiskers show minimum and maximum values within the $1.5 \times$ interquartile range. **d**, Mutational landscape of ZIC1 in G4 and SHH. **e**, Allelic frequency distribution for ZIC1 mutations in G4 ($n = 3$) and SHH ($n = 2$) samples from the assembled validation cohort. **f**, ZIC1 VAF obtained from published medulloblastoma RNA-seq data. P value from two-tailed Mann-Whitney U test.

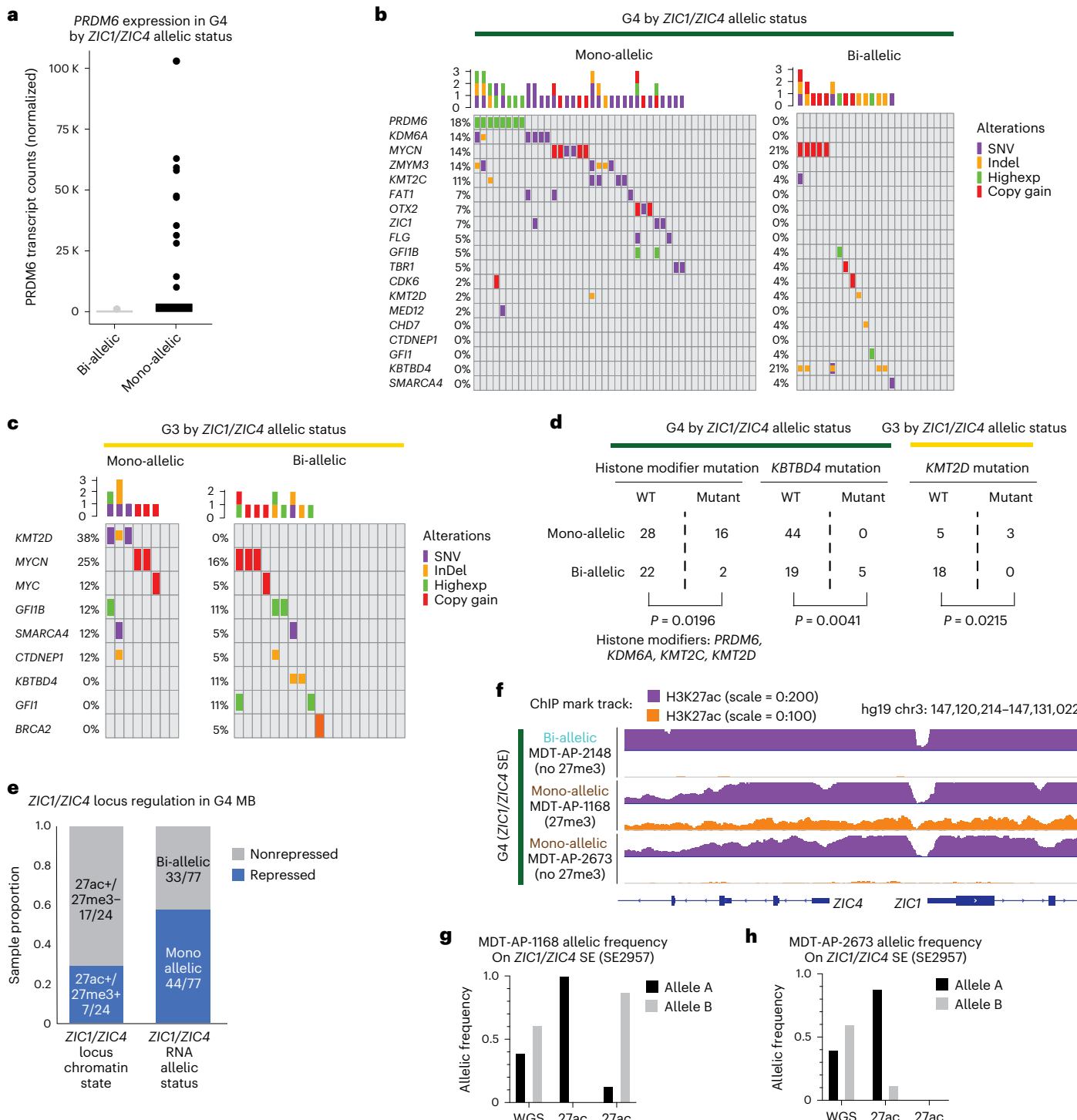


Fig. 4 | ZIC1/ZIC4 mono-allelic and bi-allelic G3/G4 medulloblastomas enrich for distinct mutations. **a**, Whisker box plot of normalized PRDM6 transcript counts in bi-allelic versus mono-allelic ZIC1/ZIC4 G4 samples. PRDM6 transcription occurs exclusively in the context of single allele inactivation of ZIC1/ZIC4. **b**, Oncoplot showcasing mutation status of previously published recurrently mutated genes in mono-allelic and bi-allelic G4 samples. Each column represents different samples. Each row represents different genes that are recurrently mutated in medulloblastoma. Distinct types of mutations for a gene in each patient are depicted with different size/colored bars. **c**, Oncoplot showcasing mutation status of previously published recurrently mutated genes in mono-allelic and bi-allelic G3 samples. **d**, Sample distribution summary for the significance of enrichment for two-tailed Fisher's exact test outputs for the significance of enrichment for

chromatin modifier mutations in ZIC1/ZIC4 mono-allelic G4 and G3, as well as KBTBD4 mutation in ZIC1/ZIC4 bi-allelic G4. **e**, Summary of different proportions of G4 medulloblastoma samples exhibiting transcriptional repression within the chromatin (H3K27me3) data or RNA (mono-allelic expression) data. **f**, Sequencing depth normalized bigwig tracks for H3K27ac and H3K27me3 in one G4 sample with bi-allelic ZIC1/ZIC4 SE and two G4 samples with mono-allelic ZIC1/ZIC4 locus SE. Not all G4 samples with mono-allelic ZIC1/ZIC4 SE harbor H3K27me3 peak on the locus. **g,h**, Allelic frequencies for heterozygous SNPs in WGS, H3K27ac and H3K27me3 ChIP-seq data in the two mono-allelic G4 samples: MDT-AP-1168, where H3K27me3 is observed, and MDT-AP-2673, where H3K27me3 is absent on the ZIC1/ZIC4 locus.

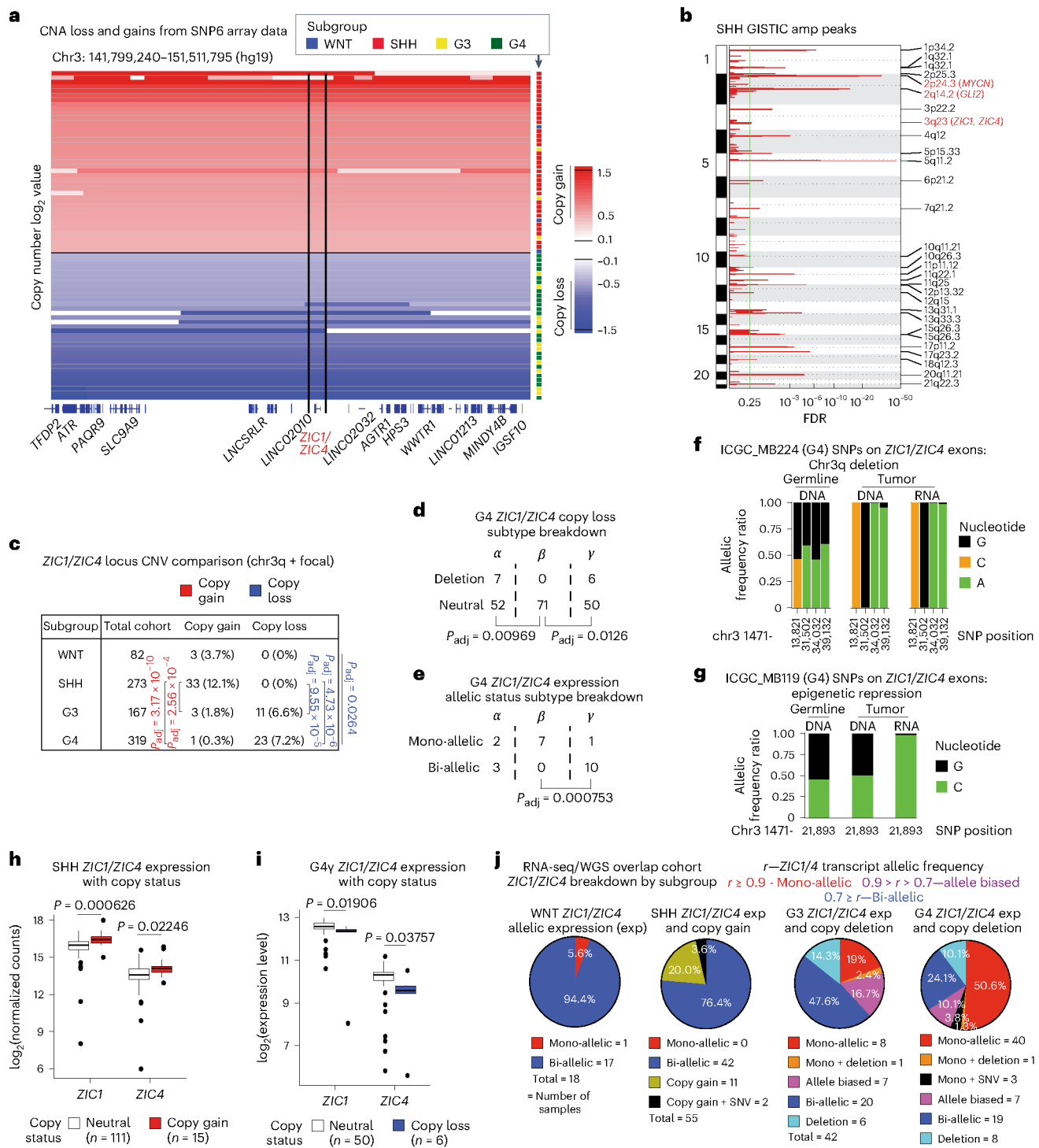


Fig. 5 | ZIC1/ZIC4 locus exhibits distinct genomic rearrangements in G3/G4 and SHH medulloblastoma. **a**, CNA track for medulloblastoma samples exhibiting ZIC1/ZIC4 locus copy gain/loss. **b**, GISTIC output for SHH medulloblastoma, highlighting 2p24.3 (*MYCN*), 2q14.2 (*GLI2*) and 3q23 (ZIC1/ZIC4) gain. FDR, false discovery rate. **c**, CNA summary for the ZIC1/ZIC4 locus per medulloblastoma subgroups. Adjusted *P* values from two-tailed pairwise Fisher's exact test. **d**, Chr3q and ZIC1/ZIC4 focal copy deletion frequency across three subtypes of G4 medulloblastoma. *P* values from two-tailed Fisher's exact test and Hochberg correction. **e**, Breakdown of chromatin repression of a single allele of ZIC1/ZIC4 locus across three subtypes of G4 medulloblastoma. *P* values were calculated by two-tailed Fisher's exact test followed by Hochberg multiple correction. **f,g**, Allelic frequencies for heterozygous germline SNPs across normal tumor DNA and tumor RNA from a representative G4 sample with (f) chr3q deletion

and (g) epigenetic suppression of the ZIC1/ZIC4 locus. **h**, Whisker box plots for ZIC1 and ZIC4 expression in SHH medulloblastoma tumors with chr3q copy gain versus neutral. Expression values from RNA-seq data with matching SNP6 array data. *P* values were calculated from the two-tailed Mann–Whitney *U* test. Center of box, median. Bounds of box, 25% and 75% percentile. Whiskers show minimum and maximum values within the 1.5× interquartile range. **i**, Whisker box plots for ZIC1 and ZIC4 expression in G4y medulloblastoma with chr3q copy loss versus copy neutral. Expression values from expression array data with matching SNP6 array data. Same statistical test and whisker box plot parameters as **h,j**. Breakdown of ZIC1/ZIC4 allelic expression pattern, ZIC1/ZIC4 CNA and ZIC1 SNVs in medulloblastoma samples with both RNA-seq and WGS data available, as well as harboring heterozygous germline SNPs in ZIC1/ZIC4 exons.

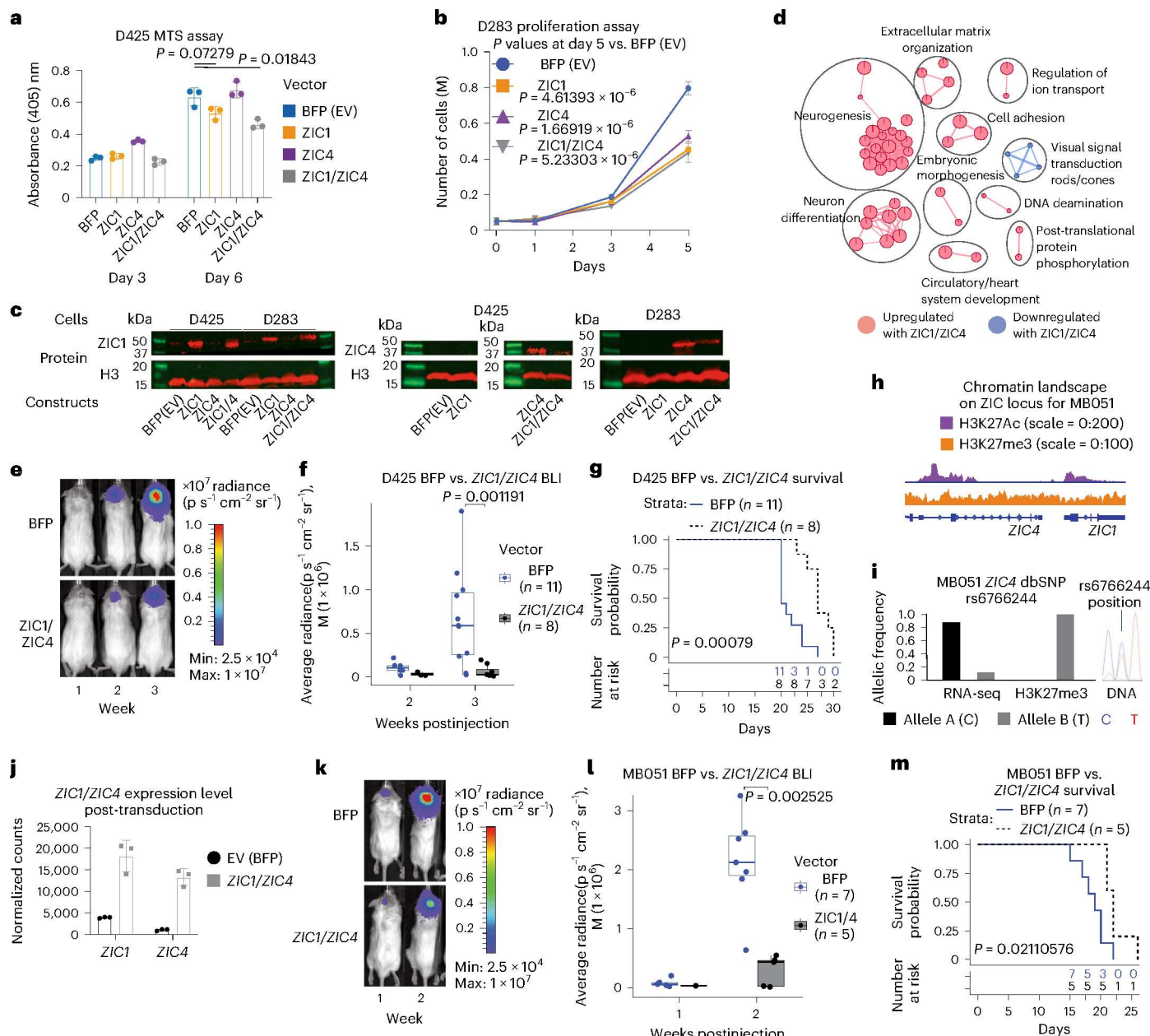


Fig. 6 | *ZIC1/ZIC4* reduces G3 medulloblastoma cell proliferation both in vitro and in vivo. **a**, 3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium (MTS) cell proliferation assay results (mean \pm s.d.) for D425. Three biological replicates. *P* values from two-tailed Welch *t*-test. **b**, Cell proliferation assay results for D283. *P* values from two-tailed Welch *t*-test. Data points show mean \pm s.d. Five biological replicates. Center of box, median. Bounds of box, 25% and 75% percentile. Whiskers show minimum and maximum values within the 1.5 \times interquartile range. **c**, Western blot validation of *ZIC1/ZIC4* overexpression in D283 and D425. **d**, Pathway analysis for *ZIC1/ZIC4* versus EV (BFP) overexpressing D425 (RNA-seq, biological *n* = 3). **e**, **f**, Representative images (**e**) and whisker box plots (**f**) summarizing BLI signals in BFP versus *ZIC1/ZIC4* overexpressing D425-injected mice. *P* values were calculated by two-tailed Welch *t*-test. Same whisker box plot parameters as **b**. **g**, Survival curves for BFP versus

ZIC1/ZIC4-transduced D425-injected mice. *P* values from two-tailed log-rank test. **h**, Normalized bigwig tracks showcasing chromatin state of *ZIC1/ZIC4* locus in patient-derived G3 xenograft line MB051. **i**, Allelic frequency of heterozygous SNP rs6766244 on coding exon of *ZIC4* from MB051 RNA-seq and H3K27me3 ChIP-seq counts, and Sanger sequencing result from the tumor DNA for the same SNP. **j**, *ZIC1/ZIC4*-normalized counts from RNA-seq in MB051 (biological *n* = 3 for EV and *ZIC1/ZIC4* constructs). Mean \pm s.d. **k**, **l**, Representative images (**k**) and whisker box plots (**l**) summarizing BLI signals in BFP versus *ZIC1/ZIC4* overexpressing MB051-injected mice. *P* values were calculated by two-tailed Welch *t*-test. Same whisker box plot parameters as **b**. **m**, Survival curves for BFP versus *ZIC1/ZIC4*-transduced MB051-injected mice. *P* values from two-tailed log-rank test. H3, histone 3; EV, empty vector.

cells highly expressing *ZIC1* over time in vivo (Extended Data Fig. 5a). MB051 also exhibited upregulation of neuronal differentiation-associated genes with *ZIC1/ZIC4* overexpression in vivo (Extended Data Fig. 5b–f), although morphological changes were not evident (Extended Data Fig. 6). Taken together, our results show tumor suppressive roles of genes in the *ZIC1/ZIC4* locus, especially *ZIC1*.

SHH and G4 medulloblastoma *ZIC1* mutants exert opposite phenotypes

As the CNAs in SHH (gain) and G4 (deletion) are diametrically opposed, we hypothesized that the SHH medulloblastoma SNVs would have divergent biological activity compared to G4 medulloblastoma SNVs, consistent with GOF and LOF phenotypes, respectively. To test

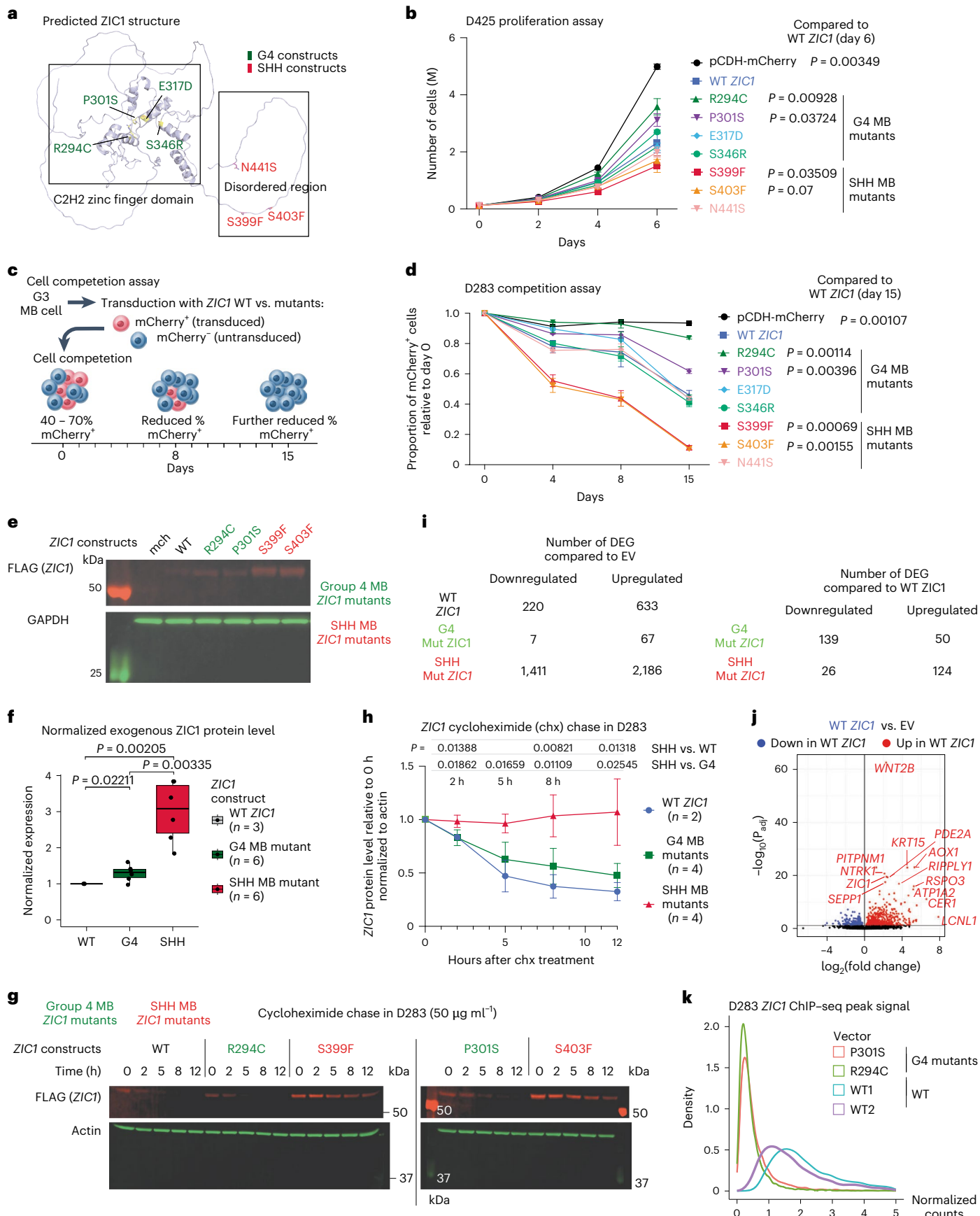


Fig. 7 | *ZIC1* mutations from G4 and SHH medulloblastoma are functionally distinct. **a**, AlphaFold2 predicted structure of *ZIC1*. Mutant constructs generated and used in the study are summarized in the structure. **b**, Proliferation assay for D425 G3 cell line transduced with *ZIC1* mutant constructs and mCherry EV. Three technical replicates for each construct. Mean \pm s.d. *P* values from two-tailed Welch *t*-test. **c**, Schematic representation for the cell competition assay using D283. **d**, Cell competition assay results using D283 transduced with *ZIC1* mutant constructs and mCherry EV. Three technical replicates for each construct. Mean \pm s.d. *P* values from two-tailed Welch *t*-test. **e**, Representative western blot visualization of exogenous *ZIC1* expression in D283 transduced with FLAG-*ZIC1* constructs. **f**, Whisker box plots showing exogenous *ZIC1* expression in D283 transduced with FLAG-*ZIC1* constructs. Signals were normalized by transduction efficiency and *GAPDH* levels. Center of box—median. Bounds of

box—25% and 75% percentile. Whiskers show minimum and maximum values within the 1.5 \times interquartile range. *P* values from two-tailed Welch *t*-test.

g, Representative cycloheximide chase results for WT and mutant *ZIC1* constructs in D283. **h**, Comparison of *ZIC1* protein level across varying exposure times to cycloheximide for WT ($n = 2$), G4 medulloblastoma mutant ($n = 4$) and SHH medulloblastoma *ZIC1* mutant ($n = 4$) constructs. *n*, biological replicates. Mean \pm s.d. *P* values from two-tailed Welch *t*-test. **i**, Number of DEG (DESeq2 output) for *ZIC1* constructs when compared against EV or WT *ZIC1*. *Q* value cutoff of 0.05. **j**, Volcano plot summarizing differentially expressed genes between WT *ZIC1* and EV. **k**, Distribution of normalized reads from FLAG ChIP-seq peaks from FLAG-tagged WT versus G4 medulloblastoma mutant *ZIC1*-transduced D283. DEG, differentially expressed genes.

this hypothesis, we generated *ZIC1* expression constructs with mutations from G4 medulloblastoma (G4 medulloblastoma *ZIC1* mutants) in the zinc finger regions or with mutations from SHH medulloblastoma (SHH medulloblastoma *ZIC1* mutants) in the carboxy terminus IDR (Fig. 7a). Consistent with our hypothesis, cell proliferation assays in D425 and cell competition assays in D283 demonstrated a reduced anti-proliferative effect for the G4 medulloblastoma *ZIC1* mutants compared to the wild-type (WT) *ZIC1*, whereas SHH medulloblastoma *ZIC1* mutants exhibited even more profound growth repression (Fig. 7b–d). We noted marked overexpression after Western blotting for SHH medulloblastoma *ZIC1* mutant proteins as compared to WT controls or G4 medulloblastoma *ZIC1* mutant proteins (Fig. 7e,f). Cycloheximide pulse-chase assays demonstrated that SHH medulloblastoma *ZIC1* mutant proteins exhibit significantly higher protein stability, as compared to WT *ZIC1*, or G4 medulloblastoma *ZIC1* mutant proteins, suggesting that the carboxy terminus IDR exerts control over the stability of the *ZIC1* protein (Fig. 7g,h). Overexpression of G4 medulloblastoma *ZIC1* mutant constructs in G3 medulloblastoma cell lines leads to tenfold fewer upregulated genes, as compared to WT *ZIC1*, whereas overexpression of the SHH medulloblastoma *ZIC1* mutant constructs resulted in more differentially expressed genes as compared to WT controls (Fig. 7i,j and Extended Data Fig. 7a–c). WT *ZIC1* overexpression led to activation of pathways involved in development and organogenesis, which was dampened with the G4 medulloblastoma *ZIC1* mutants but further augmented with the SHH medulloblastoma *ZIC1* mutants (Extended Data Fig. 7d–f). ChIP-seq against Flag-*ZIC1* demonstrates reduced DNA-binding affinity of G4 medulloblastoma *ZIC1* mutant proteins, offering a mechanistic insight underlying the reduction of *ZIC1* target gene induction (Fig. 7k and Extended Data Fig. 7g). As the G4 medulloblastoma *ZIC1* point mutations occur in the DNA-binding domain, we conclude therefore that loss of DNA binding is at least partially responsible for the phenotype of G4 medulloblastoma *ZIC1* mutants.

ZIC1 is a GOF driver in SHH medulloblastoma

Contrary to *ZIC1* suppressing the growth of G3 medulloblastoma, we hypothesized that *ZIC1* would promote the growth of SHH medulloblastoma. Indeed, overexpression of *ZIC1* constructs in mouse granule neuron progenitor (GNPs) cells (the cell of origin for SHH medulloblastoma)^{10,12} results in increased cellular proliferation, which was more pronounced with the SHH medulloblastoma *ZIC1* mutants as compared to WT *ZIC1* or G4 medulloblastoma *ZIC1* mutants (Fig. 8a,b). Cycloheximide chase in GNPs transduced with *ZIC1* mutant constructs revealed that SHH medulloblastoma *ZIC1* mutants also increase protein stability in GNPs, demonstrating the conservation of mutant mechanism across different cell types (Fig. 8c,d). *ZIC1* ChIP-seq in GNPs transduced with *ZIC1* mutant constructs also demonstrated reduced DNA-binding affinity for G4 medulloblastoma *ZIC1* mutants similar to results observed in D283 (Extended Data Fig. 8a,b). Transduction of GNPs with *ZIC1* constructs promoted higher expression of cell cycle pathway genes as well as *Gli2*, the main effector of SHH signaling (Fig. 8e–g and Extended Data Fig. 8c,d)^{14,26}. *Gli2* is a known

oncogene for SHH medulloblastoma, which exhibits a highly SHH medulloblastoma-enriched expression pattern as well as *ZIC1*-binding motif enrichment in its promoter (Extended Data Fig. 8e and Supplementary Table 16). Re-analysis of published datasets³³ demonstrates that *Zic1* binds the *Gli2* promoter in the mouse cerebellum and that loss of *Zic1* is associated with diminished expression of *Gli2* (Extended Data Fig. 8f–h). These data are consistent with a model in which *ZIC1* expression represses cell growth in maturing unipolar brush cell (UBC) progenitors of the RL-SVZ (origin of G4 medulloblastoma)^{2,3}, whereas it promotes growth of GNPs (origin of SHH medulloblastoma) in the developing cerebellar external granule layer (EGL). In the mouse, after the generation of eomesodermin (EOMES)+ excitatory deep cerebellar nuclear neuron committed cells at E10.5–E12.5 (refs. 34,35), the RL-SVZ arises as a bipotent progenitor zone capable of producing both GNPs and UBCs from E13.5 (refs. 35,36). Publicly available data on developing human cerebellum^{3,37}, as well as newly generated RNA-scope results, demonstrated that both *ZIC1* and *ZIC4* are highly expressed in UBC progenitors of the RL-SVZ (Extended Data Fig. 9a–g). The genetic and chromatin variants of *ZIC1* and *ZIC4* in G4 and SHH medulloblastoma suggest a model in which the activity of ZIC TFs has context-dependent roles in UBC and granule neuron lineage cells, which cumulatively constitute the majority of the neurons in a human brain (Fig. 8h,i).

Discussion

G3 and G4 medulloblastoma are molecularly distinct medulloblastoma subgroups that are highly related to each other and share many oncogenic drivers³⁸. We report similar *ZIC1* LOF phenotypes manifesting in G3 and G4 (epigenetic suppression, copy deletion and LOF mutation), albeit at different proportions, suggesting that the *ZIC1/ZIC4* locus has similar roles within each subgroup and possibly within their cells of origin. On the other hand, while SHH medulloblastoma shares a direct developmental relationship with G4 medulloblastoma, *ZIC1/ZIC4* events confer a GOF phenotype. These findings suggest that *ZIC1/ZIC4* has opposing roles in G3/G4 medulloblastoma versus SHH medulloblastoma, raising the possibility that these genes may also have distinct roles in the cells of origin for these similar but distinct tumor types.

Our genetic and experimental data provide robust support for a model in which LOF mutations/chromatin variants in the *ZIC1/ZIC4* locus promote G4 medulloblastoma, while GOF mutations promote SHH medulloblastoma within the different lineages of the rhombic lip. *ZIC1* events in the current cohort are found in 20% of SHH medulloblastoma and 60% of G4 medulloblastoma, making *ZIC1* one of the most frequently affected driver genes in medulloblastoma biology. While *ZIC4* is coregulated with *ZIC1* through recurrent epigenetic suppression and copy number changes, the functional role of *ZIC4* in G3 medulloblastoma cell lines is minimal compared to that of *ZIC1*. Furthermore, somatic point mutations have only been identified for *ZIC1* and not for *ZIC4*. As such, we predict that *ZIC1* has a more dominant role in medulloblastoma tumorigenesis, with *ZIC4* potentially providing some additive effects.

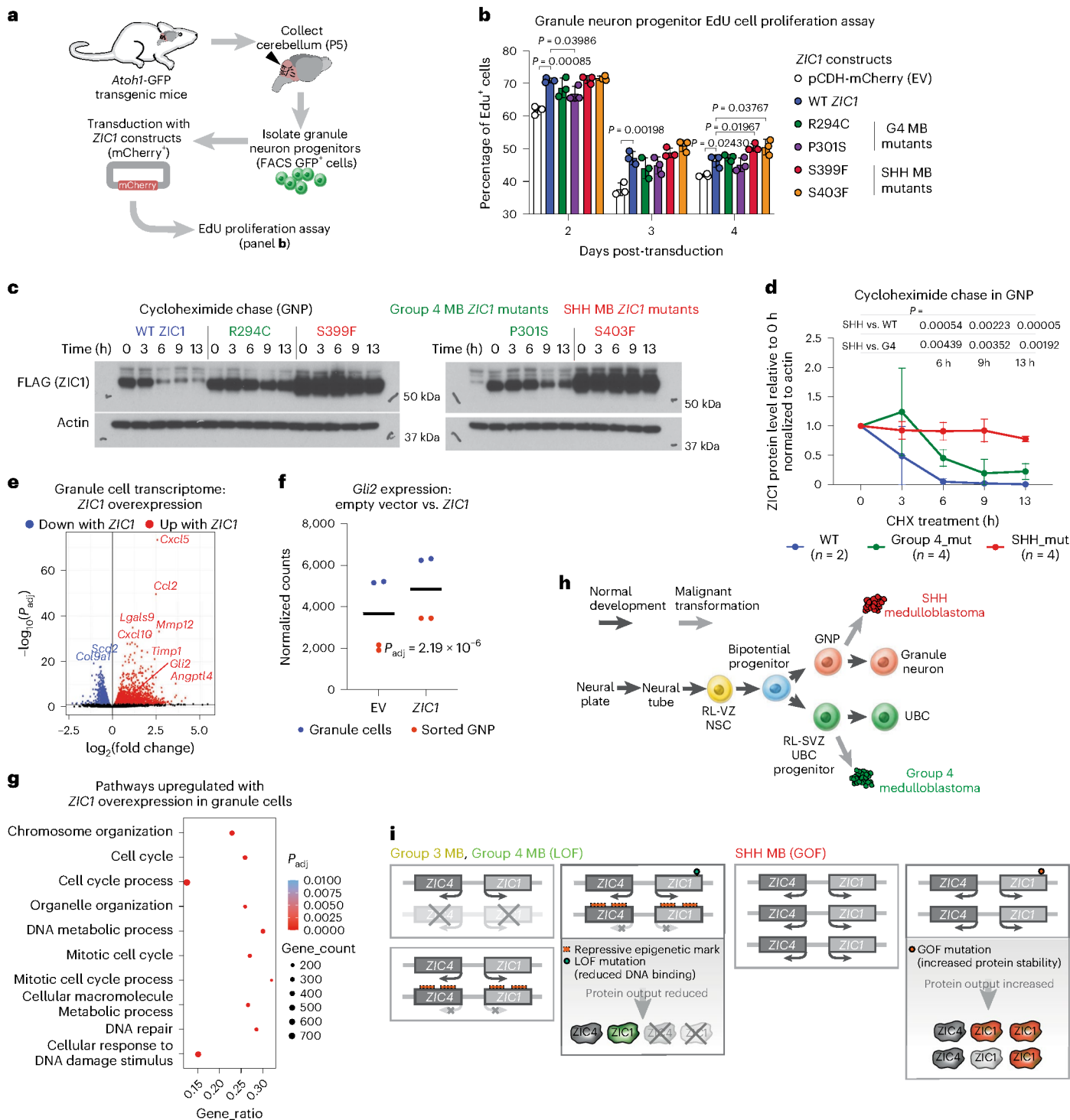


Fig. 8 | ZIC1 is a GOF driver oncogene in SHH medulloblastoma. **a**, Schematic representation summarizing GNP 5-ethynyl-2'-deoxyuridine (EdU) proliferation assay. FACS, fluorescence activate cell sorting. **b**, Summary of EdU proliferation assay for GNP transduced with ZIC1 mutant constructs and mCherry EV. GNPs enriched from multiple mouse cerebellums were used to generate biological triplicates for each construct. Mean \pm s.d. as error bars. *P* values were calculated by two-way ANOVA. **c**, Representative results from two independent replicates from running cycloheximide (CHX) chase on GNP transduced with WT ZIC1 construct, two G4 medulloblastoma ZIC1 mutant constructs and two SHH medulloblastoma ZIC1 mutant constructs. **d**, Comparison of ZIC1 protein level from GNP across varying exposure times to cycloheximide for WT ($n=2$), G4 medulloblastoma mutant ($n=4$) and SHH medulloblastoma ZIC1 mutant ($n=4$) constructs. *n*, biological replicates. Mean \pm s.d. *P* values were calculated by two-tailed Welch *t*-test. **e**, RNA-seq-derived volcano plot summarizing DEG (DESeq2 output) between ZIC1 (mch^+ ZIC1 $^+$) and EV (mch^+) transduced granule cells. Two biological replicates were generated for bulk granule cells and sorted GNPs (biological *n* = 4). *Q* value cutoff of 0.05. **f**, Normalized RNA-seq counts for Gli2 transcript in EV (mch^+) and ZIC1 (mch^+ ZIC1 $^+$) transduced GNPs. Adjusted *P* value from differential expression was calculated from DESeq2 differential expression analysis. **g**, Top ten pathways upregulated with ZIC1 overexpression in bulk granule cells and GNPs. **h,i**, Summary of normal rhombic lip development (**h**) as well as epigenetic and genetic events (**i**) that lead to ZIC1 LOF in G3 and G4 medulloblastoma and ZIC1 GOF in SHH medulloblastoma. ANOVA, analysis of variance; NSC, neural stem cell.

($n=4$) constructs. *n*, biological replicates. Mean \pm s.d. *P* values were calculated by two-tailed Welch *t*-test. **e**, RNA-seq-derived volcano plot summarizing DEG (DESeq2 output) between ZIC1 (mch^+ ZIC1 $^+$) and EV (mch^+) transduced granule cells. Two biological replicates were generated for bulk granule cells and sorted GNPs (biological *n* = 4). *Q* value cutoff of 0.05. **f**, Normalized RNA-seq counts for Gli2 transcript in EV (mch^+) and ZIC1 (mch^+ ZIC1 $^+$) transduced GNPs. Adjusted *P* value from differential expression was calculated from DESeq2 differential expression analysis. **g**, Top ten pathways upregulated with ZIC1 overexpression in bulk granule cells and GNPs. **h,i**, Summary of normal rhombic lip development (**h**) as well as epigenetic and genetic events (**i**) that lead to ZIC1 LOF in G3 and G4 medulloblastoma and ZIC1 GOF in SHH medulloblastoma. ANOVA, analysis of variance; NSC, neural stem cell.

Our discovery of a H3K27me3/H3K27ac heterozygous chromatin state in G4 medulloblastomas at the *ZIC1/ZIC4* locus demonstrates a convincing complementation group in which some tumors achieve repression of *ZIC1* through deletion or somatic mutations of genomic DNA, while other tumors reach the same phenotype through chromatin variants that impose epigenetic repression. This may be through somatic acquisition of chromatin variants, akin to *de novo* allele-specific 'epimutations' that have been described to be associated with oncogenesis^{39,40}. Indeed, this robust complementation group provides strong evidence for the biological importance of somatic chromatin variants in the pathogenesis of cancer. We suggest that the observed chromatin events drive the clonal selection of tumor cells and are not merely passenger events.

We were unable to use current technologies to identify local or distal cryptic noncoding mutations driving the H3K27me3/H3K27ac heterozygous chromatin state, although we acknowledge that these may occur and be currently cryptogenic. It is also possible that there exists a minor unidentified population in the rhombic lip that is temporally or anatomically restricted and passes through a state with the H3K27me3/H3K27ac heterozygous chromatin state, and that these particular cells are at increased risk for transforming to G4 medulloblastoma. An additional possible mechanism is somatic 'epimutation', in which aberrant H3K27me3 marks repress *ZIC1* expression, and this heritable chromatin state results in clonal expansion and eventually G4 medulloblastoma. The consistent co-occurrence of somatic mutations of histone lysine modifier genes in G4 medulloblastomas that also harbor somatic chromatin variants of *ZIC1* is consistent with a model in which aberrant control of the epigenome leads to 'epigenetic instability', with clones that by error contain *ZIC1* silencing chromatin events undergoing clonal selection. Similarly, it has been previously shown that succinate dehydrogenase deficiency can induce aberrant epigenetic remodeling *mono-allelically*⁴¹. Which of the three outlined mechanisms, or mechanisms not currently suspected, is responsible for the H3K27me3/H3K27ac heterozygous chromatin state is, however, not currently known, nor readily determined using current technologies, although we favor the somatic chromatin variant model.

G4 medulloblastoma comprises cells similar to the UBC progenitors within RL-SVZ, while SHH medulloblastoma cells resemble GNPs of the EGL. These highly related cell types likely arise from the same bipotential progenitors. The clear difference between the LOF phenotypes (G4) versus GOF phenotypes (SHH) suggests a model in which *ZIC1* and/or *ZIC4* have context-dependent roles in UBC progenitors and GNP during rhombic lip development. In GNPs, *ZIC1/ZIC4* may work in conjunction with other SHH pathway genes, such as *GLI2*, to promote cell proliferation and granule-cell-like transcriptome. Tight regulation of *ZIC1/ZIC4* activity is likely critical to prevent overexpansion of GNPs during EGL formation. Conversely, UBC progenitors likely require higher levels of *ZIC1/ZIC4* activity for normal differentiation, as shown by the UBC lineage-enriched *ZIC1/ZIC4* expression pattern. Perturbation of *ZIC1/ZIC4* activities in these different contexts likely contributes to improper rhombic lip development and favors oncogenic transformation, where LOF genetic/chromatin variants promote the transformation of the UBC progenitors and GOF variants promote the transformation of the GNPs.

We maintain that LOF/GOF mutations of *ZIC1* are true driver events, as overexpression of *ZIC1* represses malignant phenotypes in G3 medulloblastoma models while promoting malignancy in SHH medulloblastoma precursor cells, both *in vitro* and *in vivo*. Indeed, our data support a model in which *ZIC1* is the paramount example of a context-specific cancer driver gene, as it appears to show diametrically opposing biological activity in these two different cell types that arise from the exact same progenitors and which occur on either side of a very specific cell fate decision during rhombic lip development.

Online content

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41588-024-02014-z>.

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Methods

Research ethics board (REB)

This study obtained full ethics approval from the Hospital for Sick Children (REB 0020020238 and REB 1000055059) as well as McGill University Health Centre (REB MCH003-26). All materials were collected after receiving written informed consent from patients, including consent to publish the generated data. All primary sample collection and experimental procedures (in vitro and in vivo) were done in accordance with guidelines from the REB of Hospital for Sick Children (REB 0020020238 and REB 1000055059), McGill University Health Centre (REB MCH003-26) and the Centre for Phenogenomics (AUP 22-0151H).

Experimental model and subject details

Primary tumor collection. Primary tumors used in the study were obtained from the Medulloblastoma Advanced Genomics International Consortium and International Cancer Genome Consortium. All materials were collected after receiving written informed consents, including consent to publish the generated data, as per guidelines from REB from the following institutes: Agostino Gemelli University Hospital, Children's Hospital of Minnesota, Cooperative Human Tissue Network, David Geffen School of Medicine at University of California Los Angeles, Duke University, Emory University, Erasmus University Medical Centre, German Cancer Research Centre (DKFZ), Hospital Cantonal De Geneve, Hospital Infantil de Mexico Federico Gomez, Hospital Sant Joan de Deu, Ludwig Maximilians University, Masaryk University, McGill University, McMaster University, Memorial Sloan Kettering Cancer Centre, Miami Children's Hospital, Portugese Cancer Institute, Queensland Children's Tumor Bank, Seattle Children's Hospital Fred Hunchinson Cancer Research Centre, Seoul National University Children's Hospital, Stanford University School of Medicine, the Chinese University of Hong Kong, Tohoku University, University of California San Francisco, University Health Network, Universitats Kinderklinik, Universite de Lyon, University of Arkansas, University of Calgary, University of Debrecen Medical and Health Science Centre, University of Pittsburgh, University of Ulsan Asan Medical Centre, University of Warsaw Children's Memorial Health Institute, Vanderbilt Medical Centre and Wolfson Children's Hospital. Statistical methods were not used to predetermine the sample size. Age, sex, subgroup and subtype information for used tumors are available in Supplementary Table 1. Primary tumor tissues were snap-frozen in liquid nitrogen and stored at -80°C until use.

Mouse housing and husbandry. All mouse breeding and procedures were performed as approved by the Toronto Centre for Phenogenomics.

Method details

G3 medulloblastoma cell lines and xenograft line. D425 and D283 cell lines were derived at Duke University (Supplementary Table 2) and verified with short tandem repeats before being used for experiments. MB051 patient-derived xenograft line was generated at the Hospital for Sick Children and passaged only by serial intracranial injection in NSG mice without expansion in vitro.

Source of NOD-SCID-IL2R γ null mice. NOD-SCID-IL2R γ null (NSG) mice were obtained from the Toronto Centre for Phenogenomics in-house breeding colony.

Intracranial injection of G3 medulloblastoma tumor cells. Intracranial injection was performed on NSG mice (age range of 6–10 weeks, ~50% males and females for all conditions) using D425 and MB051 xenograft lines as previously described⁴² using slightly modified stereotactic coordinates—2 mm posterior to λ , 1 mm lateral and 2 mm deep. In total, 2,000 Green fluorescent protein luciferase-tagged D425 cells transduced with BFP empty vector or ZIC1/ZIC4 vector were

injected per mouse. In total, 4,000 GFP luciferase-tagged MB051 cells transduced with BFP empty vector or ZIC1/ZIC4 vector were injected per mouse. Humane endpoint was called independently by staff at the Toronto Centre for Phenogenomics based on physiological conditions exhibited by the injected mice. These staff were blinded from construct information. Mice that did not exhibit any BLI signal above the background ($2.5 \times 10^4 \text{ p s}^{-1} \text{ cm}^{-2} \text{ sr}^{-1}$) by the third week after injection were excluded from the cohort.

Bioluminescence measurement. Bioluminescence was measured in NSG mice injected with GFP Luciferase-tagged tumor cells as previously described⁴². For D425, measurements were taken on week 1 (6–7 days after injection), week 2 (13–14 days after injection) and week 3 (20 days after injection). For MB051, measurements were taken on week 1 (7 days after injection) and week 2 (14 days after injection).

RNA-scope on developing human cerebellum slides. Manufacturer-recommended protocols were used for RNA-scope *in situ* hybridization (ISH) assays as previously described³⁷ using RNA-scope 2.5 High Definition-RED Assay (ACDBio, 322350). Briefly, RNA-scope was performed on mid-sagittal sections of the developing vermis, fixed in 10% formalin for 4 weeks. Manufacturer-recommended protocols (ACDBio/Bio-Techne) were used to assay the following probes: Hs-ZIC4 (525661) and Hs-ZIC1 (542991). All sections were counterstained with hematoxylin or methyl green. Stained slides were imaged using the Nanozoomer Digital Pathology slide scanner (Hamamatsu).

ZIC1 mutant construct generation. WT ZIC1 was cloned into pCDH-mCherry or pCDH-GFP empty lentiviral vector using the In-Fusion Snap Assembly Starter Bundle (Takara). Mutagenesis, or N-terminal FLAG tagging of ZIC1, was also done using the In-Fusion kit.

Isolation of cerebellar granule cells or GNPs. Cerebellar cells were isolated from the cerebellum as described previously⁴³. Briefly, cerebellum from postnatal day 5 (P5) mice was digested with high glucose Dulbecco's Phosphate Buffered Saline (DPBS) (Thermo Fisher Scientific) containing 10 U ml^{-1} papain (Worthington), $200 \mu\text{g ml}^{-1}$ L-cysteine and 250 U ml^{-1} DNase (Sigma) for 30 min. Tissue was triturated to obtain a single-cell suspension and then centrifuged through a 35% and 65% Percoll gradient (Sigma). Cells in the layer between 35% and 65% Percoll were washed once with DPBS containing 0.02% BSA and resuspended in GNP culture medium (neurobasal supplemented with B27 (50 \times), sodium pyruvate (100 \times), penicillin–streptomycin (100 \times) and glutamax (100 \times)). Granule cells or GNPs were enriched by depleting the adherent cells through two incubations in poly-D-lysine (PDL)-coated plates for 20 min each time. Enriched granule cells and GNPs were cultured with GNP culture medium supplemented with $3 \mu\text{g ml}^{-1}$ SHH (Peprotech) in PDL-coated plates. For the isolation of pure GNPs, cerebellar cells were isolated from Atoh1-GFP mice at P5 as described above. After washing once with DPBS containing 0.02% BSA, cells were suspended with DPBS containing 5% FBS (Thermo Fisher Scientific). GNPs with strong GFP expression (~40%) were sorted and cultured with the GNP culture medium as described above.

5-ethynyl-2'-deoxyuridine (EdU) assay in GNPs. GNPs isolated from P5 Atoh1-GFP mice, as described above, were infected with control (pCDH-mCherry) or ZIC1 viruses (pCDH-mCherry_ZIC1 WT/mutants) in triplicates. Cells were cultured in a GNP culture medium with SHH in PDL-coated 48-well plates. At each time point, cells were treated with $10 \mu\text{M}$ 5-ethynyl-2'-deoxyuridine (EdU) for 6 h and then dissociated for EdU staining (Click-iT Plus EdU Pacific Blue Flow Cytometry Assay Kit) and flow cytometry analysis. For data analysis, cells were first gated for mCherry $^+$ cells. The percentage of proliferating cells (EdU $^+$) was then calculated for each sample.

Quantification and statistical analysis

ChIP-seq data processing. Raw ChIP-seq reads were aligned to hg19 genome assembly using bowtie2 (v2.2.1)⁴⁴. PCR duplicates were removed using Picard MarkDuplicates. Reads with mapping quality lower than 20 were removed. Reads from nonchromosomal contigs, mitochondria or ENCODE blacklist regions were also filtered out before peak calling. H3K27ac peaks were identified using MACS2 (v2.1.1.20160309) with the following code: MACS2 callpeak -t IP_bam_file -f BAMPE -g hs --nomodel -B -q 1e-2 (ref. 45). H3K27me3 peaks were identified using the following parameters: MACS2 callpeak -t 27me3_IP_bam_file -c input_bam_file -f BAMPE -g hs --nomodel --broad -B -q 1e-5 --broad-cutoff 1e-4. Peaks that could not be identified in at least two primary medulloblastomas were excluded from any further analysis. Library sizes for samples in H3K27ac and H3K27me3 samples were calculated using SAMtools⁴⁶ and average fragment sizes of three different batches of H3K27ac and H3K27me3 were evaluated by deep-tools⁴⁷ (v3.1.3). H3K27ac and H3K27me3 peaks in each sample were annotated according to their closest genes and then categorized into different classes based on their distributions over different types of features, for example, promoter, exon, intron and distal intergenic. The distance between peaks and their assigned genes was calculated by using the center of the peak and the transcription start site as coordinates.

For ChIP-seq data from D283 cells transduced with FLAG-tagged *ZIC1* constructs, peaks were called using *Q* value threshold of 1×10^{-5} . For ChIP-seq data from GNP cells transduced with FLAG-tagged *ZIC1* constructs, peaks were called using a *Q* value threshold of 0.05.

SNP inference from ChIP-seq libraries. For samples harboring both H3K27ac and H3K27me3 peaks on the *ZIC1/ZIC4* locus, ‘H3K27ac–H3K27me3 hemizygous region’ was defined for each sample with bedtools (v2.27) intersect on the called peaks⁴⁸. From the bivalent region containing the *ZIC1/ZIC4* locus, allelic frequencies were calculated for each dbSNP151 annotated heterozygous SNP positions from H3K27ac and H3K27me3 library reads using bedtools multicov. Heterozygous SNPs were identified by first calculating allelic frequency r = absolute value of (reference (REF) alternate (ALT) allelic frequency). Afterward, SNPs with $r \geq 0.6$ in both H3K27Ac and H3K27me3, but biased for different alleles in each, were used to infer heterozygous SNPs (ex, H3K27ac enriched for REF allele and H3K27me3 enriched for ALT allele). Alternatively, SNPs with $r < 0.6$ in either H3K27ac or H3K27me3 libraries were also used to identify SNPs. Only SNPs that are supported by at least ten reads from each library were used.

SEs analysis and subgroup consensus peak sets. SEs were defined using the Rank Ordering of Super Enhancers (v0.1) algorithm using H3K27ac peaks as input⁴⁹. For all samples, the stitching distance was fixed at 12.5 kb to facilitate comparisons between samples. All other parameters used the default setting. Once SEs were generated for each sample, SEs were merged from samples within the same subgroup using GenomicRanges Bioconductor package⁵⁰. Only SEs that were present at least two times per subgroup were considered for merging.

RNA-seq data processing. Custom hs37d5 genome assembly generated in previous study²⁷ was used to align raw RNA-seq reads using STAR aligner (2.7.4) with the following parameters: --outFilterMultimapNmax 20 --alignSJoverhangMin 8 --alignMatesGapMax 200000 --alignIntronMax 200000 --alignSJDBoverhangMin 10 --alignSJstitchMismatchNmax 5 -1 5 5 --outSAMmultNmax 20 --twopassMode Basic⁵¹. Gene expression level was quantified using HTSeq (0.6.0) based on Gencode v19 annotations with the argument ‘-stranded reverse -m union’⁵². Differential gene expression analysis between subgroups was performed using the R Bioconductor package DESeq2 (v1.26.0)⁵³. An adjusted *P* value of 0.05 was used for differentially expressed gene identifications.

H3K27ac HiChIP data process and loop call. Raw HiChIP reads were aligned using bowtie2 (2.3.4) and HiC-pro (2.9.0) using the default parameters in HiC-pro⁵⁴. Output directory was used as input for hicchipper (v0.7.3) to call significant loops using the following parameters: min-dist 5000, max-dist 20000000, read-length 150, ‘macs2-string -q 0.01 --extsize 315 --nomodel’⁵⁵. Intrachromosomal loops with *Q* value less than 0.01 and read counts greater than 5 were used for downstream enhancer gene interactome analysis.

WGS data processing and germline variants calling. WGS data were aligned to the ‘hs37d5’ reference genome from 1000 Genomes Project Phase II as previously described²⁸, using Burrows–Wheeler aligner–MEM (v0.7.8) with the ‘-T 0’ parameter⁵⁶. For germline variant call, variants identified in both normal and tumor DNA from Platypus (v0.8.1) run with default parameters were used (<https://github.com/andyrimmer/Platypus>). To have the final heterozygous SNP list for each sample in WGS data, we only selected those passed Platypus quality control (minBaseQual and minMapQual: 20; alleleBias and strandBias: 0.001 and badReadsWindow: 11). Second, we retained SNPs with allele depth in tumor samples ≥ 10 , allele depth in paired blood samples ≥ 7 , allele ratio in blood between (0.3, 0.7) and allele ratio in tumor between (0.2, 0.8). Third, only bi-allelic sites and InDels shorter than three nucleotides were used. The final heterozygous SNP candidates were retained in the following allele imbalance analysis. We used EAGLE2 for haplotype phase estimation on bcftools (v1.9)⁵⁷ normalized variants, using a phased reference panel in 1000 Genomes Project⁵⁸.

Affymetrix SNP6 array data processing. SNP6 Affymetrix array data were mapped to hg19 and processed using Affymetrix Power Tools (v1.18.2) as previously described²⁷.

Identification of focal recurrent CNAs from SNP6 array. To identify recurrent focal copy gains and losses for each subgroup, SNP6 array-derived segmentation files were used as input for GISTIC2 (v2.0.23) from gene pattern with the following options: refgene file = Human_Hg19.mat, maxspace = 10,000, gene gistic = yes, confidence = 0.90, *Q* value threshold = 0.25, run broad analysis = no, max sample segs = 10,000, arm peel = yes, gene collapse method = extreme, amplification threshold = 0.5, deletion threshold = -0.5, focal length cutoff = 0.5, armlevelpeel = on, confidence level = 0.95, *Q* value = 0.25, run broad analysis = no, max sample segs = 10,000 (ref. 31). Other parameters were left as default.

Single-cell RNA-seq (scRNA-seq) data analysis. Publicly available scRNA-seq data were analyzed as previously described with minor modifications^{3,59}. Specifically, RL-SVZ cells from the glutamatergic lineage cells were further divided into three smaller cell clusters using the following criteria: RL-SVZ (KI67 high, EOMES+) – RL-SVZ residing UBC progenitor cells; RL-SVZ (KI67 high, ATOH1+) – RL-SVZ cells more committed to GCP lineage; RL-SVZ (KI67 low, EOMES+) – RL-SVZ residing UBC progenitor cells likely mixed with some early UBC.

Pathway enrichment analysis. Enriched pathways for differentially expressed genes were identified by using g-profiler at default parameters, using *Q* value threshold of 0.05 (ref. 60). Gene Ontology-biological term outputs were used for the final list of pathways. Top ten enriched/depleted pathways were identified for *ZIC1* mutant construct experiments using G3 medulloblastoma cell lines or GNP cells *in vitro* and G3 medulloblastoma xenograft experiments *in vivo*.

Calling CNA events from WGS data. Copy number information was derived from WGS data using Control-FREEC (v10.3)³² as previously described with the following parameters: breakPointType = 4, ploidy = ‘2,3,4’, step = 10,000, window = 50,000 (ref. 28).

Before focal CNA call from WGS data for known medulloblastoma driver genes, ploidy for all WGS samples was predicted with Control-FREEC. For samples with inferred ploidy greater than 3.5, pileup ratio was used from ploidy = 4 output. All other samples used pileup ratio from ploidy = 2 output. Median ratio values for each segmented genomic locus were used to generate a segmented (.seg) format for each sample. Merged seg file for each subgroup was used as input for GISTIC2 (v2.0.23) from gene pattern with the following options: refgene file = human_Hg19.mat, maxspace = 10,000, gene gistic = yes, confidence = 0.90, Qvalue threshold = 0.25, run broad analysis = no, max sample segs = 10,000, arm peel = yes, gene collapse method = extreme, amplification threshold = 0.25, deletion threshold = -0.25, focal length cutoff = 0.5, armlevelpeel = on, confidence level = 0.95, Q value = 0.25, run broad analysis = no, max sample segs = 10,000 (ref. 31). Other parameters were left as default. Output from focal_data_by_genes was used for genes previously identified to undergo recurrent CNA gain in G3/G4—*MYC*, *MYCN*, *OTX2* and *CDK6*, which have been previously reported^{14,26}.

For CNA identification from WGS data for the *ZIC1*/*ZIC4* locus, both broad chromosomal events and focal CNA were identified using the seg files generated above. An amplification threshold of 0.25 and a copy loss threshold of -0.25 were used to estimate the proportion of samples with copy number changes in SHH or G3/G4 samples, respectively.

Oncoplot generation. Highly expressed genes were identified by performing *k*-means clustering on size factor normalized RNA-seq counts with *k* = 2 for the following genes: *GFI1*, *GFIIB* and *PRDM6*. Group with higher expression of genes were categorized as highly expressing. Somatic SNVs, InDels, CNA amplifications and high expression samples for each gene were annotated for all samples using complexheatmap (v2.2.0) R package⁶¹.

Statistics and reproducibility

No statistical method was used to predetermine the sample size. Randomizing and blinding were not used for the experiments. For experiments involving the injection of mice with medulloblastoma cell lines or patient-derived xenograft lines, independent staff at the Toronto Centre for Phenogenomics were blinded from the experimental arm conditions before calling the endpoints. For mouse BLI experiments, mice that failed to reach the minimal detectable signal of 2.5×10^4 $\text{p s}^{-1} \text{cm}^{-2} \text{sr}^{-1}$ by the third week postinjection were removed from the cohort (failure to engraft).

Reporting summary

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

Data availability

The FLAG ChIP-seq, RNA-seq data generated from *ZIC1* mutant construct transduced G3 medulloblastoma cell lines and granule cells have been deposited in the Gene Expression Omnibus (GEO) database under the accessions [GSE217639](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE217639), [GSE217571](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE217571) and [GSE217638](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE217638). Bulk H3K27ac, H3K27me3 ChIP-seq, RNA-seq, WGS and H3K27ac hicup data generated from primary medulloblastoma tumor samples in this study have been deposited in the European Genome–Phenome Archive (EGA) database under the accession code [EGAS00001006741](https://www.ebi.ac.uk/ega/study/EGAS00001006741). The published medulloblastoma bulk RNA-seq data referenced in this study are available in the EGA database under the accessions [EGAS00001001953](https://www.ebi.ac.uk/ega/study/EGAS00001001953), [EGAD00001004347](https://www.ebi.ac.uk/ega/study/EGAD00001004347), [EGAS00001004435](https://www.ebi.ac.uk/ega/study/EGAS00001004435), [EGAS00001005826](https://www.ebi.ac.uk/ega/study/EGAS00001005826), [EGAD00001001899](https://www.ebi.ac.uk/ega/study/EGAD00001001899), [EGAD00001004958](https://www.ebi.ac.uk/ega/study/EGAD00001004958) and [EGAD00001008458](https://www.ebi.ac.uk/ega/study/EGAD00001008458). The published medulloblastoma WGS data referenced in this study are available in the EGA database under the accessions [EGAS00001001953](https://www.ebi.ac.uk/ega/study/EGAS00001001953), [EGAD00001003125](https://www.ebi.ac.uk/ega/study/EGAD00001003125) and [EGAD00001004347](https://www.ebi.ac.uk/ega/study/EGAD00001004347). The published medulloblastoma H3K27ac ChIP-seq data referenced in this study are available in the EGA database under the accessions [EGAS00001001953](https://www.ebi.ac.uk/ega/study/EGAS00001001953).

The Affymetrix SNP 6.0 data referenced during the study are available in the GEO database under the accession [GSE37385](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE37385). The expression array used for transcript abundance comparison between medulloblastoma subtypes is available in the GEO database under the accession [GSE132269](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE132269). Multiple databases were used for annotation of SNPs and promoters, which were referenced in this study. These include the GRCh37 dbSNP151 (https://ftp.ncbi.nlm.nih.gov/snp/organisms/human_9606_b151_GRCh37p13/VCF/), Gencode (v.19; https://www.gencodegenes.org/human/release_19.html), the hg19 reference genome (<https://hgdownload.soe.ucsc.edu/goldenPath/hg19/bigZips/>), the hs37d5 reference genome (https://ftp-trace.ncbi.nih.gov/1000genomes/ftp/technical/reference/phase2_reference_assembly_sequence/), ERCC spike-in sequence (<https://www.encodeproject.org/files/ENCFF908UQN/>) and Caltech profile 3 spike-in sequence (<https://www.encodeproject.org/references/ENCSR193ZXE/>). snRNA-seq data from the developing human cerebellum were obtained through correspondence from ref. 59 and are available through the Human Cell Atlas (<https://explore.data.humancellatlas.org/projects/85a9263b-0887-48ed-ab1a-ddfa773727b6>), the UCSC Cell Browser (<https://cbl-dev.cells.ucsc.edu>) or from Database of Genotypes and Phenotypes (dbGaP; accession [phs001908.v2.p1](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC8800000/)). Bulk RNA-seq data from the developing human cerebellum were obtained through correspondence from ref. 37 and are available through the dbGaP (accession [phs001908.v2.p1](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC8800000/)). Source data are provided with this paper. Human material provided by the Joint MRC/Wellcome (MR/R006237/1) Human Developmental Biology Resource (HDBR; www.hdbi.org) and the Birth Defects Research Laboratory (BDRL; NIH-R24-HD000836 to I.A.G.) was covered by a material transfer agreement between SCRI and HDBR/BDRL, but samples may be requested directly from the HDBR and BDRL. Please see the Supplementary Information for full lists of the reagents, resources and bioinformatics tools used for the study (Supplementary Tables 1–16). Requests for additional information or resources and reagents should be directed to and will be fulfilled by M.D.T.

Code availability

Original codes used for the study are available at <https://doi.org/10.5281/zenodo.1394024> (ref. 62). Full details of methods used for the study can be found in Supplementary Note.

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Acknowledgements

M.D.T. is a Cancer Prevention and Research Institute of Texas (CPRIT) Scholar in Cancer Research (CPRIT—RR220051); is the Cyvia and Melvyn Wolff Chair of Pediatric Neuro-Oncology at Texas Children's Cancer and Hematology Center; is supported by the National Institutes of Health (NIH) (R01NS106155, R01CA159859 and R01CA255369), the Pediatric Brain Tumor Foundation, the Terry Fox Research Institute, the Canadian Institutes of Health Research, the Cure Search Foundation, the Matthew Larson Foundation (IronMatt), b.r.a.i.n.child, Meagan's Walk, SWIFTY Foundation, the Brain Tumour Charity, Genome Canada, Genome BC, Genome Quebec, the Ontario Research Fund, Worldwide Cancer Research, V-Foundation for Cancer Research and the Ontario Institute for Cancer Research through funding provided by the Government of Ontario and is also supported by a Canadian Cancer Society Research Institute Impact grant, a Cancer Research UK Brain Tumor Award and by a Stand Up To Cancer (SU2C) St. Baldrick's Pediatric Dream Team Translational Research Grant (SU2C-AACR-DT1113) and SU2C Canada Cancer Stem Cell Dream Team Research Funding (SU2C-AACR-DT-19-15) provided by the Government of Canada through Genome Canada and the Canadian Institutes of Health Research, with supplementary support from the Ontario Institute for Cancer Research through funding provided by the Government of Ontario. Stand Up to Cancer is a program of the Entertainment Industry Foundation administered by the American Association for Cancer Research. P.A.N. is supported by the American Lebanese Syrian Associated Charities (St. Jude), the Brain Tumor Charity (Quest for Cures) and the National Cancer Institute (P01CA096832-16A1 and 1R01CA270785-01A1). L.F.H. is supported by a DoD PRCRP award (CA191188) and start-up funds from the Mayo Clinic Foundation. J.J.Y.L. is supported by the University of Toronto Fellowship and Ontario Graduate Scholarship. The tumor bank at the University of Debrecen was supported by the 2017-1.2.1-NKP-2017-00002 National Brain Research Program NAP 2.0 grant. The authors

would also like to thank A. Bondoc (Brain Tumour Biobank at SickKids) and recognize the Labatt Brain Tumour Research Centre and the Michael and Amira Dan Brain Tumour Bank Network. Human cerebellar tissue used in this study was provided by the Joint MRC/Wellcome (MR/R006237/1) HDBR (www.hdbr.org) and covered by a material transfer agreement between SCRI and HDBR. The authors also thank K. Lowe (Flow Cytometry Core Laboratory of the Developmental Neurobiology Department at St. Jude Children's Research Hospital) for flow cytometry analysis. Additionally, the authors would like to thank the staff at the Sickkids-University Health Network Flow Cytometry Facility for aid with the flow sorting of cells and Ben Pakuts for support with graphic design and illustration.

Author contributions

J.J.Y.L. and M.D.T. conceptualized and led the study. J.J.Y.L. and R.T. designed, performed and analyzed the majority of the experiments in the study. J.J.Y.L. and Z.Y. did most of the bioinformatic analysis in the study. P.H. contributed to performing RNA-scope on developing human cerebellum slides. Z.Y. and L.F.H. designed the method for mono-allelic analysis by integrating multi-omics profiles with Bayesian inference. H.F. contributed to processing Affymetrix SNP6 array data for copy number alteration analysis. L.D.H. contributed to processing published scRNA-seq data from the developing human cerebellum. N.A., C.M.R. and J.P. contributed to intracranial injection of cells into NSG mice. E.Y.W. contributed to using Cytoscape to generate a pathway analysis schematic. X.W. contributed to experimental designs. A.W.E., L.D.H., A.R., S.B., M.L., F.M.G.C. and S.M. contributed to bioinformatic analysis. L.G., K.A.M. and A.V. contributed to experimental design. V.F. and B.G.L. contributed to the validation of overexpression constructs. R.T. and N.D.G. contributed to the generation of *ZIC1* mutant constructs. R.T., J.H. and M.B. contributed to processing mouse cerebellum for GNP assays. R.S., B.L., J.L. and C.D. contributed to managing the tumor bank, tissues and resources at Sickkids. A.B., P.J.F., J.M.K., A.K., M.K., F.C.P.d.L., M.P.-D., B.L., S.K.S., S.E.S.L., B.-K.C., S.-K.K., K.-C.W., J.-Y.L., T.T., W.A.W., J.J.P., G.Z., A.G.S., B.L., A.K., I.F.P., R.L.H., Y.-s.R., W.A.G., M.P.-P., R.C.T., A.M.K. and M.K.C. provided patient tumor material and helped design the study. C.W.M. contributed to interpreting H&E staining results. S.D. contributed to generating the AlphaFold predicted *ZIC1* structure schematic. S.C.M., N.J., M.L., M.C., M.L.S., H.S. and K.M. provided expert advice. H.S. contributed to calling germline and somatic mutations from WGS data. L.F.H., P.A.N. and M.D.T. jointly supervised the project and provided funding support. J.J.Y.L., R.T. and Z.Y. prepared the figures. J.J.Y.L. and M.D.T. wrote the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

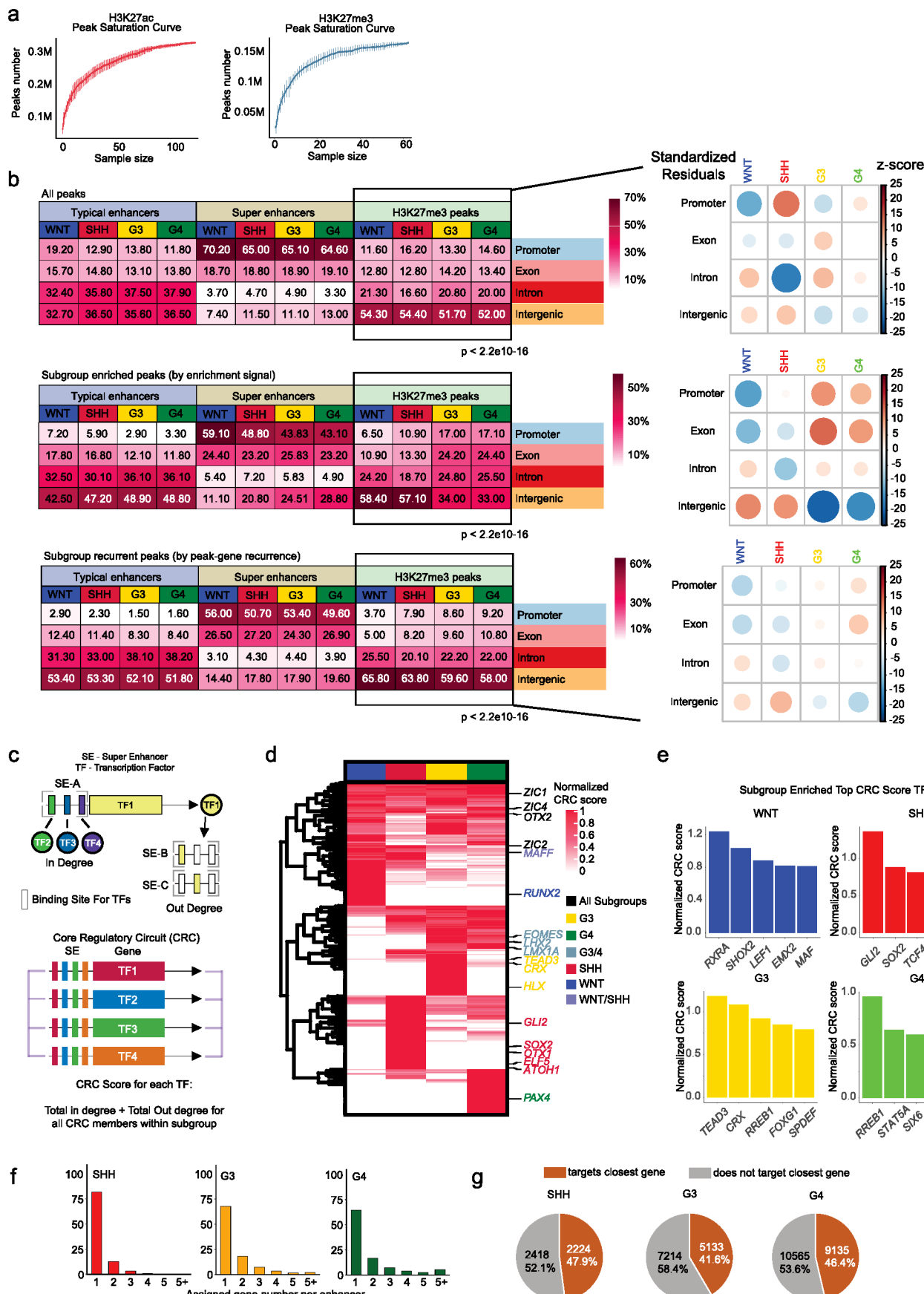
Extended data is available for this paper at <https://doi.org/10.1038/s41588-024-02014-z>.

Supplementary information The online version contains supplementary material available at <https://doi.org/10.1038/s41588-024-02014-z>.

Correspondence and requests for materials should be addressed to L. Frank Huang, Paul A. Northcott or Michael D. Taylor.

Peer review information *Nature Genetics* thanks Baoli Hu and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

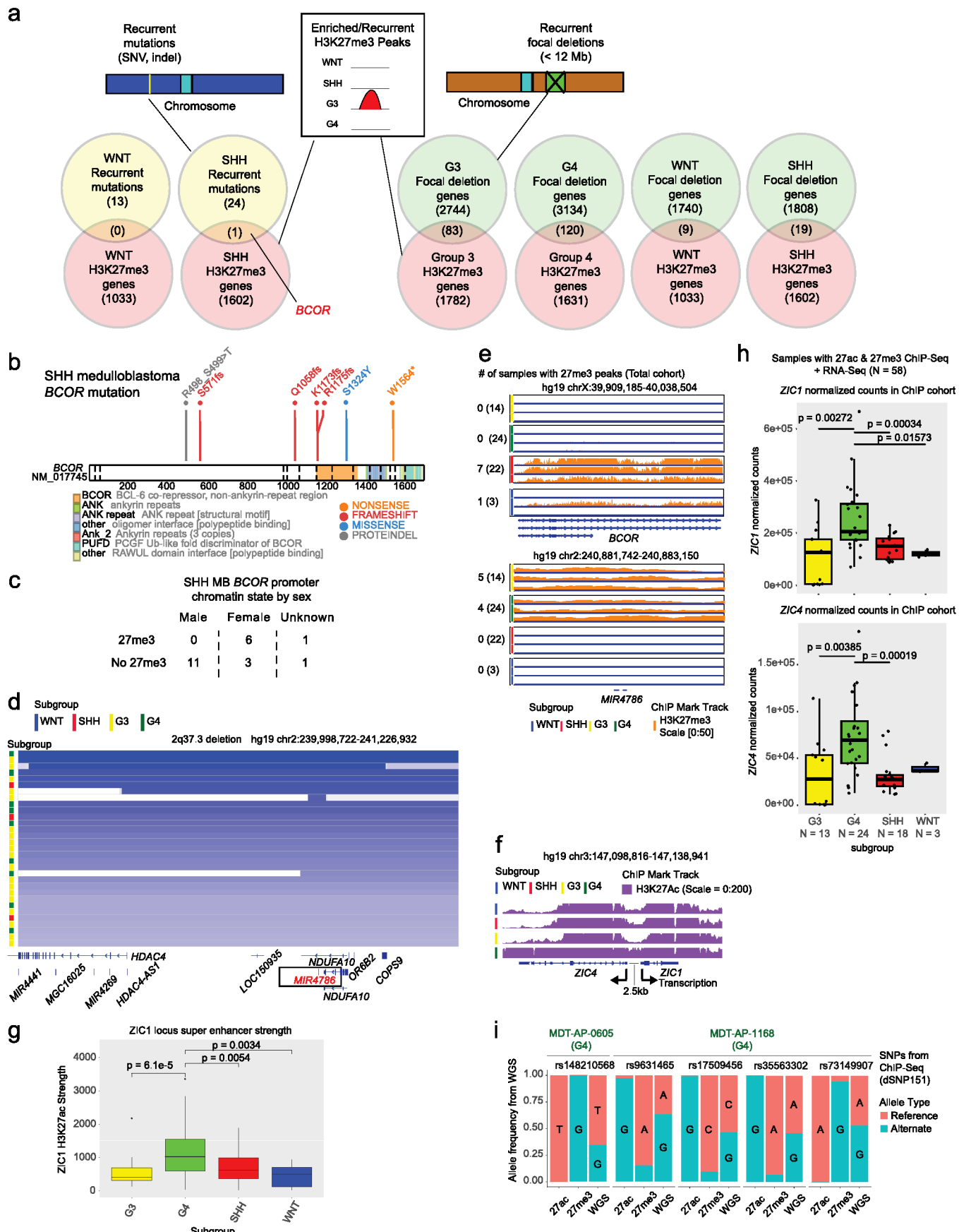
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Extended Data Fig. 1 | See next page for caption.

Extended Data Fig. 1 | Medulloblastoma exhibits subgroup-specific master transcription factors (TFs) and chromatin landscape. **a**, Saturation analysis for H3K27ac and H3K27me3 peak identification. For each number of samples shown on the x axis, a subset of total cohort of ChIP-seq samples corresponding to this number was randomly selected. Number of non-overlapping peaks identified from this subset were recorded for each iteration of random sampling. Average and standard deviation for 10 iterations were plotted for each number up to total cohort size. Number of peaks identified starts to plateau toward the end of the curve, suggesting that addition of new samples will likely lead to diminishing returns. **b**, Annotation for typical enhancers, super-enhancers (SE) and H3K27me3 peaks that are classified as (1) all peaks found in the subgroup, (2) subgroup-enriched peaks (defined in Fig. 1c) and (3) subgroup-recurrent peaks (defined in Fig. 1c). P values were calculated by performing two-tailed chi-square

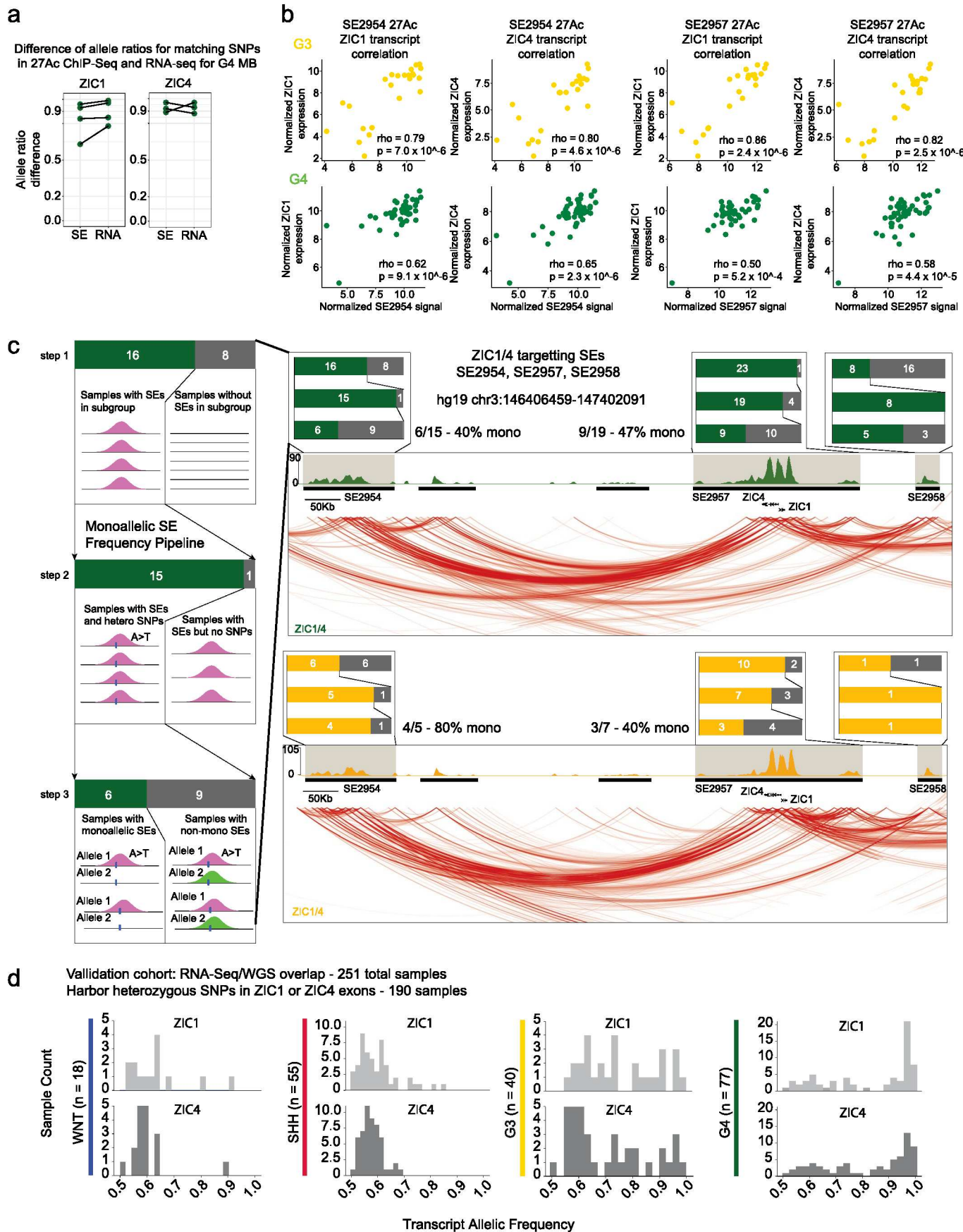
test on H3K27me3 peaks. Standardized residuals for chi-square tests performed on H3K27me3 peak distributions were also calculated. **c**, Strategy used to define core regulatory circuit (CRC) score for each transcription factor for each subgroup. In degree (number of TFs that target the TF of interest) and out degree (number of TF promoters targeted by the TF of interest) were calculated for each TF to identify subgroup-specific and pan-subgroup core TFs. **d**, Heatmap summarizing pan subgroup and subgroup-specific core TFs crucial for shaping core circuitry landscape for each subgroup. **e**, Top 5 subgroup-specific master transcription factors identified for each subgroup according to CRC score. **f**, Number of genes assigned for each enhancer across enhancer–promoter interactions identified using HiChIP and 27ac ChIP-seq data. **g**, Proportion of enhancers that target the closest genes for SHH, G3 and G4 subgroups.



Extended Data Fig. 2 | See next page for caption.

Extended Data Fig. 2 | Overlap between recurrent copy number deletions and subgroup enriched/recurrent H3K27me3 peaks for group 3 (G3)/group 4 (G4) medulloblastoma. **a**, Venn diagram depicting overlap between subgroup-enriched H3K27me3 peaks with recurrently mutated genes in WNT, SHH as well as genes recurrently affected by focal deletion (<12 Mb) in all 4 subgroups (Supplementary Table 13). **b**, *BCOR* mutation pattern identified in SHH medulloblastoma. **c**, Breakdown of *BCOR* H3K27me3 pattern in SHH medulloblastoma. Highly female-enriched pattern is observed, suggesting that X inactivation may have a role in the observed chromatin phenomenon. **d**, Showcase of recurrent deletion of 2q37.3 locus identified in G3 and G4. *MIR4786* locus exhibits a G3/G4-enriched copy loss pattern (Supplementary Table 13). **e**, Representative H3K27me3 ChIP-seq signal patterns for all subgroups on *BCOR* and *MIR4786* locus, which exhibit SHH-enriched and G3/G4-enriched H3K27me3 signal, respectively (Supplementary Tables 12 and 13).

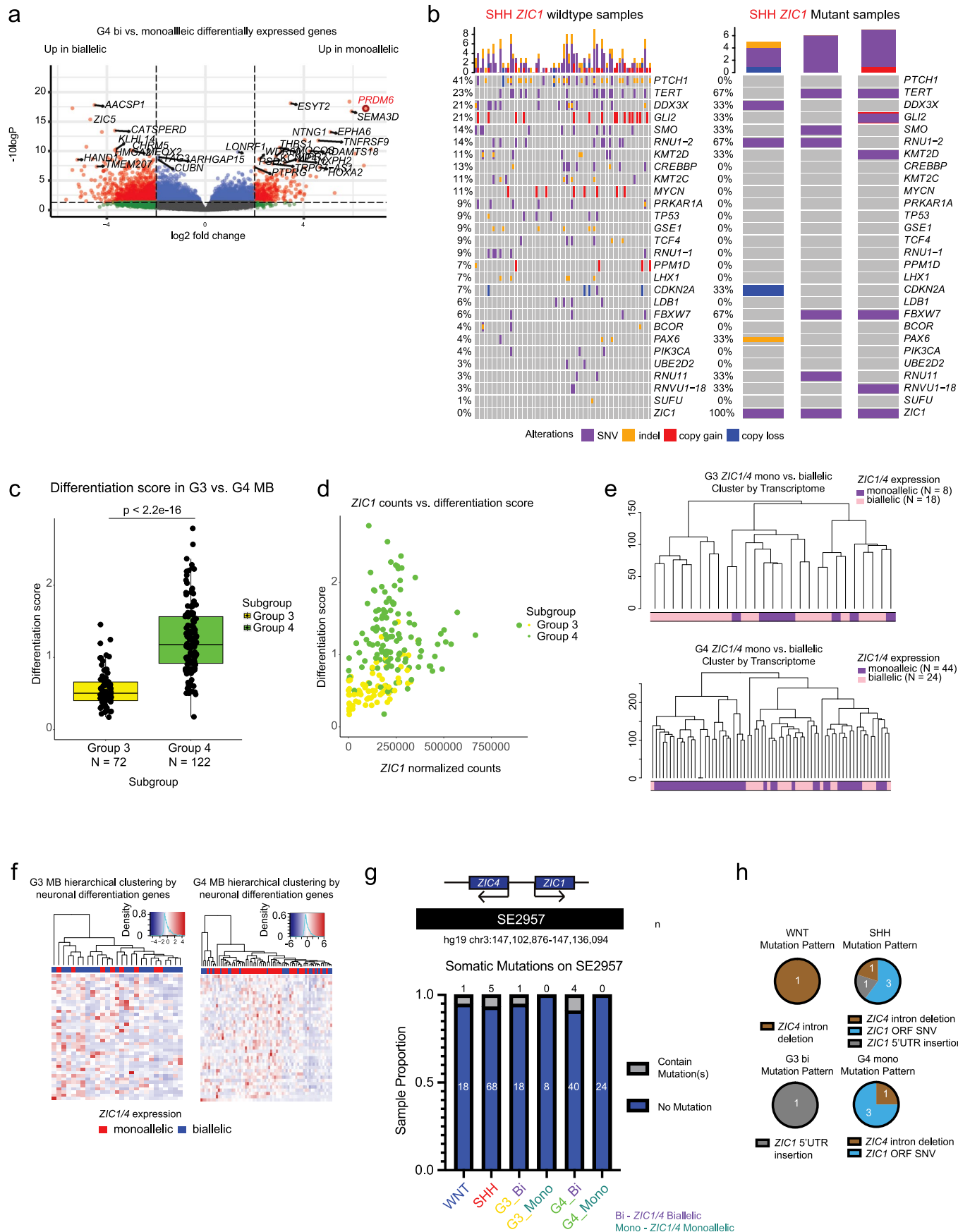
f, Read depth normalized 27ac bigwig tracks for a representative sample from each subgroup. Bidirectional promoters regulating *ZIC1* and *ZIC4* transcription are regulated by a common super-enhancer identified across all subgroups. **g**, H3K27ac signal strength of SE overlapping *ZIC1/4* promoter across MB subgroups. Biological sample size: G3/G4/SHH/WNT = 27/47/39/10. Center of box—median. Bounds of box—25% and 75% percentile. Whiskers show minimum and maximum values within the 1.5× interquartile range. P values from two-tailed Mann–Whitney *U* test. **h**, *ZIC1*- and *ZIC4*-normalized transcript count levels in ChIP cohort samples with matching H3K27ac, H3K27me3 and RNA-seq data (N = 58). Biological sample size: G3/G4/SHH/WNT = 13/24/18/3. Box plot parameters same as **g**. P values from two-tailed Wilcoxon rank-sum test. **i**, Allelic frequencies for the inferred heterozygous single-nucleotide polymorphisms (from Fig. 2h,i) in 2 G4 samples with matching WGS data.



Extended Data Fig. 3 | See next page for caption.

Extended Data Fig. 3 | *ZIC1/4* locus is regulated by multiple super-enhancers (SE) that are recurrently epigenetically repressed on single alleles. **a.** Allelic frequencies for heterozygous SNPs present in both H3K27ac ChIP-seq reads on *ZIC1/4* SE as well as RNA-seq reads on *ZIC1/4* exons. Identical schematic to dot plots from Fig. 3a, but only the exact match heterozygous SNPs identified in both H3K27ac ChIP-seq and RNA-seq data were used. Matching samples are connected by lines between SE and RNA columns. Y axis shows difference in pooled allelic frequency between SNPs from the two different alleles. *ZIC1/4* RNA and SE exhibit bias for the same alleles from the heterozygous single-nucleotide polymorphisms (SNPs), suggesting that the monoallelic SE drives monoallelic expression. **b.** Correlation between H3K27ac reads on two SEs that target *ZIC1/4* locus (from Extended Data Fig. 2g), SE2954 and SE2957, and *ZIC1/ZIC4* transcript levels in group 3 (G3) and group 4 (G4) medulloblastoma. P values generated

from two-tailed Spearman correlation analysis. **c.** *ZIC1/4* targeting SEs, their interaction maps with *ZIC1/4* locus and frequency of their monoallelic status in G3 and G4 medulloblastoma. SE directly on top of *ZIC1/4* genes (SE2957) was monoallelic in 9 out of 19 samples in G4 and 3 out of 7 samples in G3. SEs upstream (SE2954) and downstream (SE2958) of *ZIC1/4* locus are also recurrently monoallelic and were identified as high-confidence enhancer–promoter interactions with HiChIP, H3K27ac ChIP-seq and RNA-seq data. While most samples harbored SE2957, a smaller proportion of G3 and G4 samples harbored SE2954 and SE2958. **d.** Allelic frequency distribution of heterozygous germline SNPs for *ZIC1* and *ZIC4* transcripts in RNA-seq within the validation cohort (total of 251 samples with both WGS and RNA-seq data). A total of 190 samples contain heterozygous SNPs within *ZIC1/4* exons in both normal control and tumor DNA.



Extended Data Fig. 4 | See next page for caption.

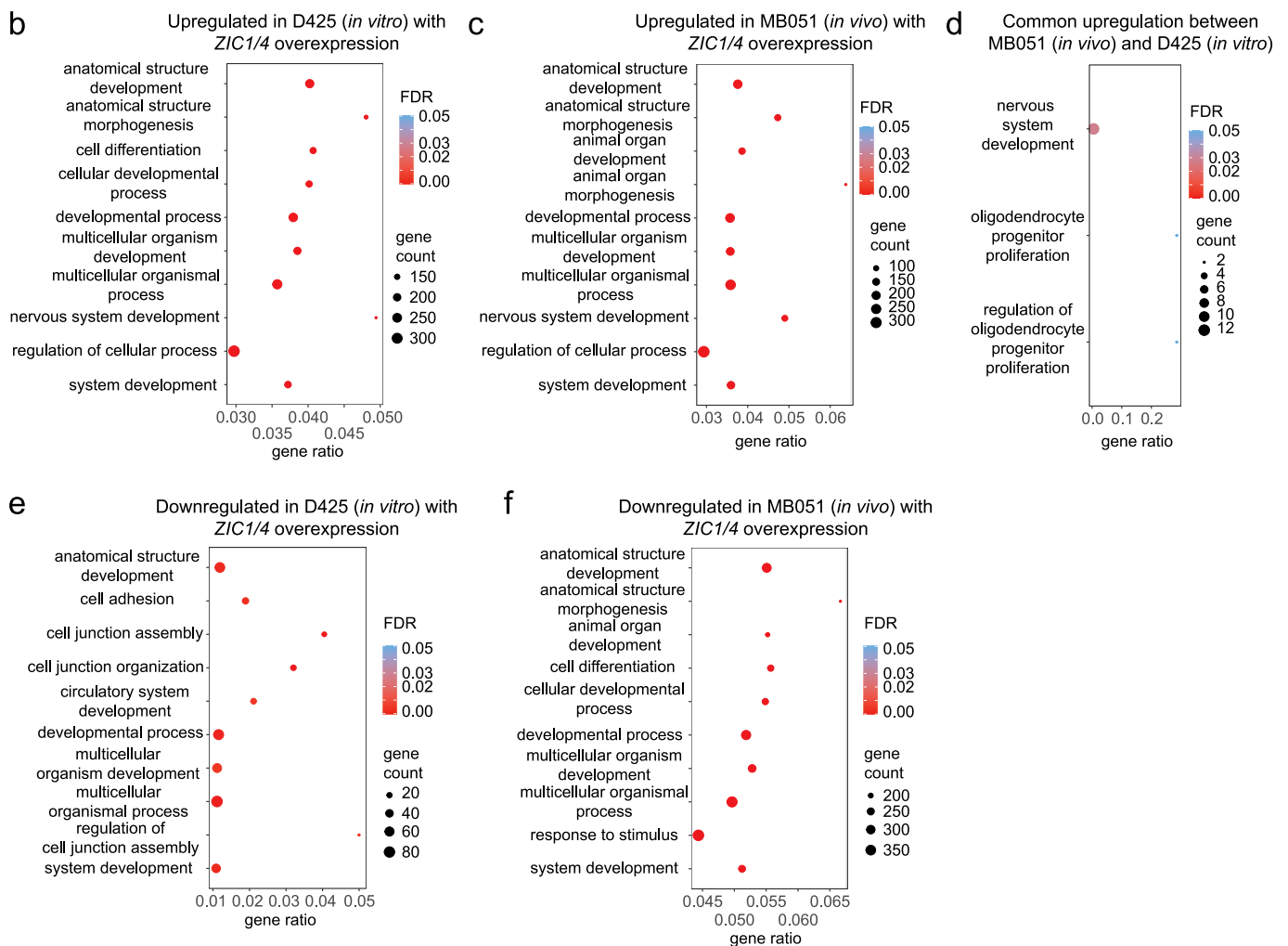
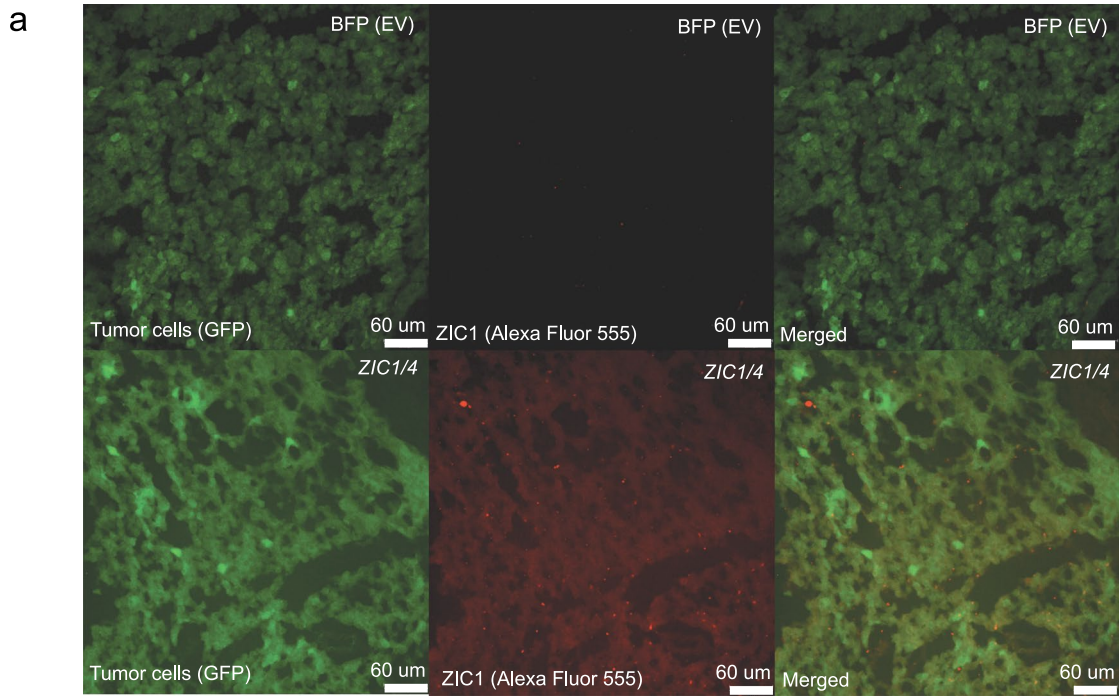
Extended Data Fig. 4 | Genetic and transcriptional patterns associated with biallelic and monoallelic status of *ZIC1/4* across medulloblastoma.

a, Volcano plot summarizing differentially expressed genes between *ZIC1/4* monoallelic and biallelic group 4 (G4) samples. Q value threshold of 0.01 and \log_2 (fold change) threshold of 2 were used. **b**, Oncoplot summarizing the mutational landscape of SHH tumors with or without *ZIC1* mutations.

U1 snRNA mutations were always mutated together (RNU1-2, RNVU1-18) with *ZIC1*. **c**, Whisker box plot summarizing neuronal differentiation score for group 3 (G3) and G4 medulloblastoma tumors. Previously published 39 G3/G4 neuronal differentiation signature genes (Supplementary Table 14) were used to calculate the overall differentiation score for each tumor. Biological sample size: G3/G4 = 72/122. P value was calculated by two-tailed Mann–Whitney U test.

Center of box—median. Bounds of box—25% and 75% percentile. Whiskers show minimum and maximum values within the $1.5 \times$ interquartile range. **d**, Scatter plot showing expression level of *ZIC1* across G3 and G4 medulloblastoma tumors vs. differentiation score in the same tumors. **e**, Hierarchical clustering of G3/G4 samples by top 10,000 variable genes from transcriptome. *ZIC1/4* monoallelic G3/G4 samples do not form distinct clusters from the biallelic samples.

f, Hierarchical clustering of G3/G4 samples by expression level of the neuronal differentiation signature genes from **c**. *ZIC1/4* monoallelic G3/G4 samples do not form distinct clusters from the biallelic samples. **g**, Frequency of somatic mutations on super-enhancer (SE) on top of *ZIC1/4* locus (SE2957) across WNT, SHH, G3 *ZIC1/4* biallelic, monoallelic, G4 *ZIC1/4* biallelic and monoallelic samples. **h**, Breakdown of somatic mutation patterns on SE2957 for all subgroups.



Extended Data Fig. 5 | See next page for caption.

Extended Data Fig. 5 | MB051 exhibits similar transcriptional changes as D425 upon *ZIC1/4* overexpression in vivo. **a**, Immunofluorescence showing tumor cells (GFP+) and *ZIC1* protein level (Alexa Fluor 555), both separately and merged, for BFP (empty vector) or *ZIC1/4*-transduced MB051 patient-derived group 3 (G3) medulloblastoma xenograft intracranially injected into NOD SCID γ (NSG) mice. One biological replicate for BFP-transduced MB051, and two biological replicates for *ZIC1/4*-transduced MB051. Two fields of views captured for BFP, and four fields of views captured for *ZIC1/4*-transduced MB051 (3 for one biological replicate and 1 for another). All views exhibited identical observations.

b, Top 10 pathways upregulated in D425 in vitro upon overexpression of *ZIC1/4* compared to BFP empty vector. **c**, Top 10 pathways upregulated in MB051 in vivo upon overexpression of *ZIC1/4* compared to BFP empty vector. **d**, Pathway analysis depicting commonly upregulated pathways between D425 in vitro and MB051 in vivo. While there was a small overlap, neuronal differentiation pathway emerged as a commonly upregulated pathway between two different models. **e**, Top 10 pathways downregulated in D425 in vitro upon overexpression of *ZIC1/4* compared to BFP empty vector. **f**, Top 10 pathways downregulated in MB051 in vivo upon overexpression of *ZIC1/4* compared to BFP empty vector.

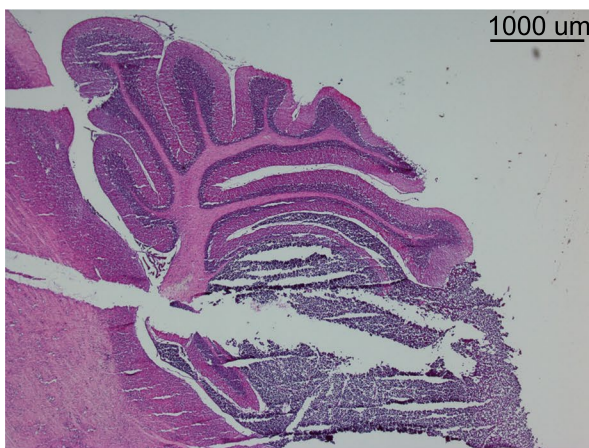
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MB051 xenograft H&E results from *in vivo*

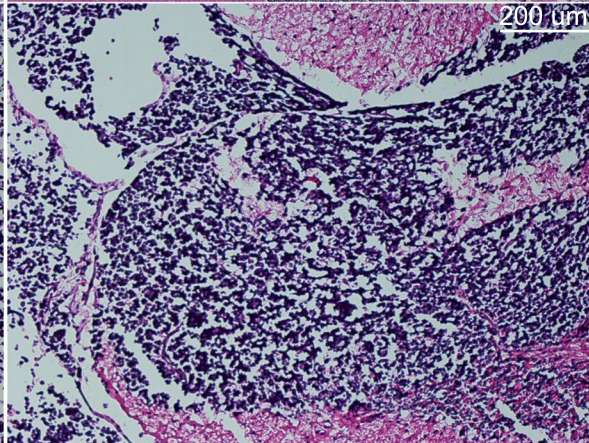
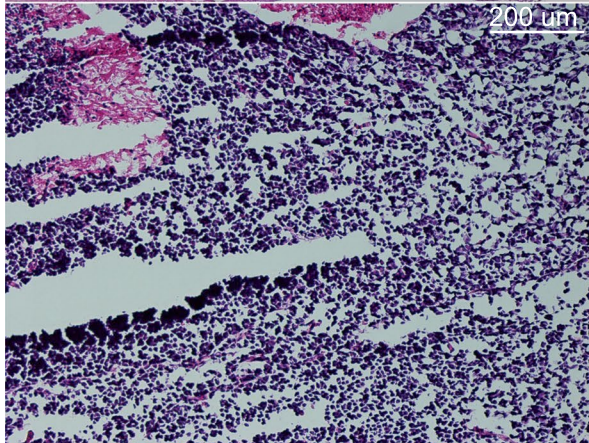
BFP (EV)

ZIC1/4 construct

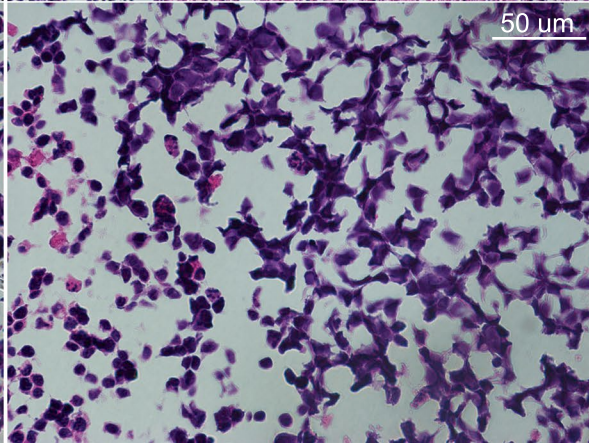
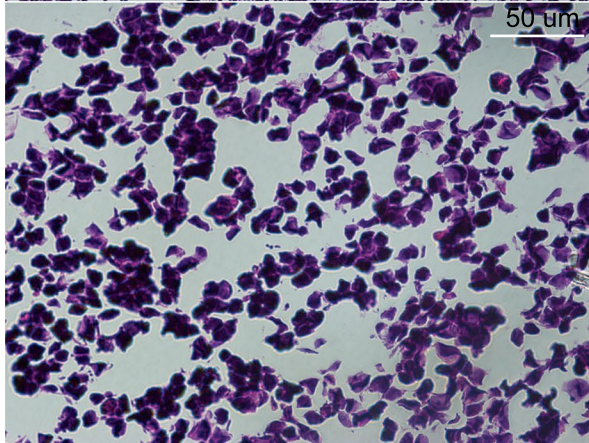
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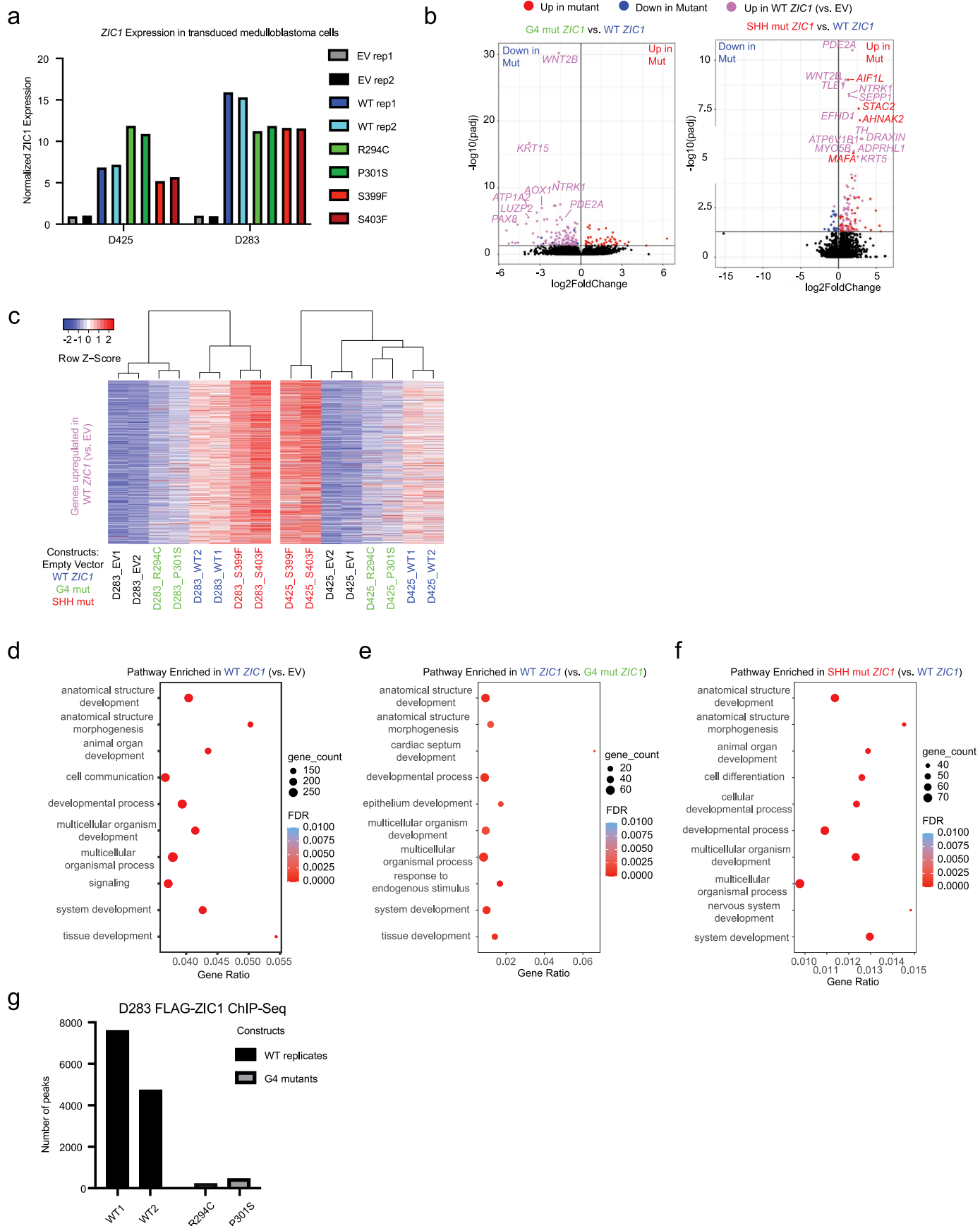


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Extended Data Fig. 6 | *ZIC1/4* overexpression does not result in morphological differences for MB051 at the H&E level. Representative H&E results at various magnifications generated from injecting MB051 into NOD SCID γ (NSG) mice. Magnifications are shown on the left side of the panels. MB051 was transduced with BFP (empty vector) or *ZIC1/4* overexpression construct prior to injection. Minimal morphological differences were observable between the different

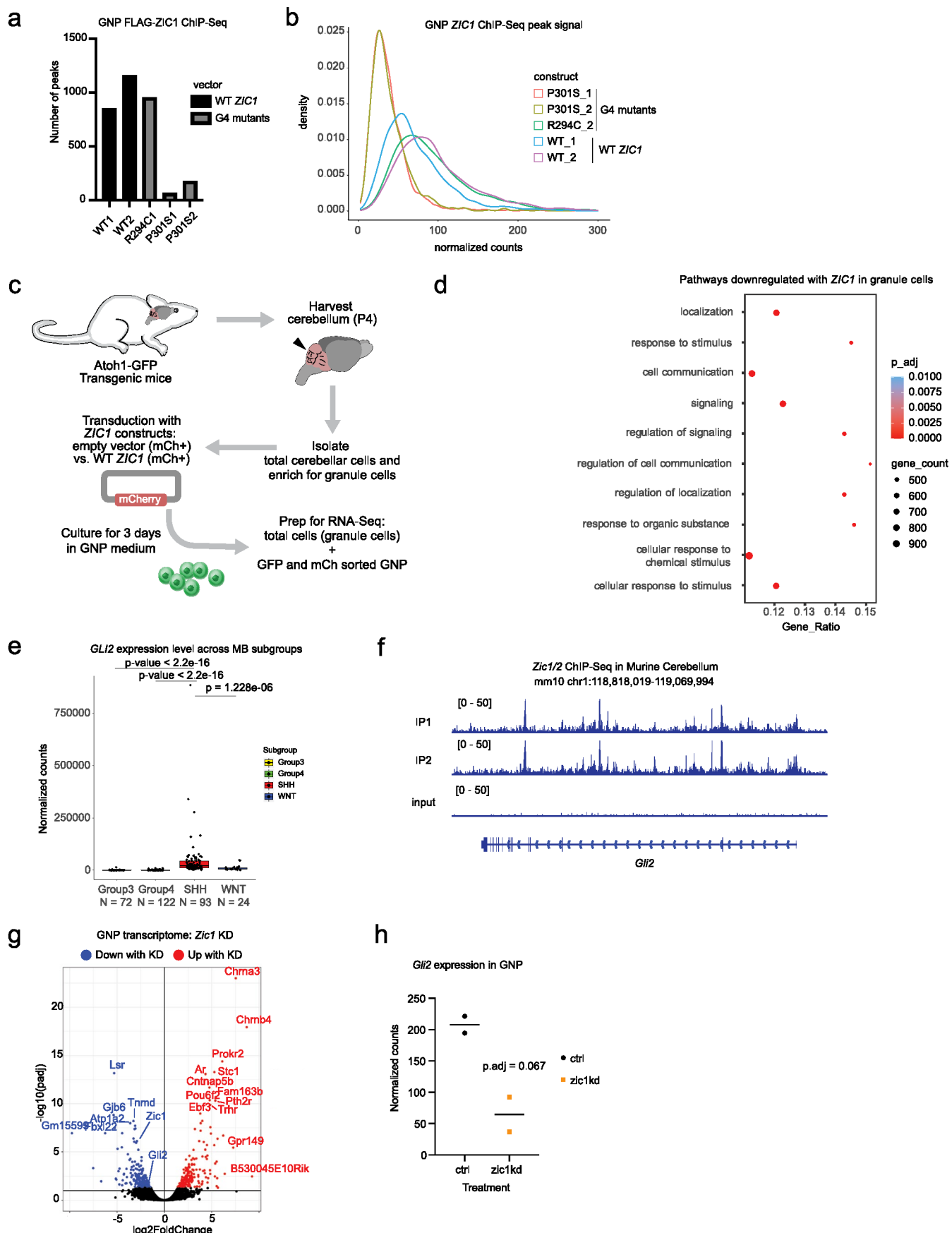
constructs. One biological replicate for BFP-transduced MB051, and two biological replicates for *ZIC1/4*-transduced MB051. Three fields of views captured for BFP and each biological replicate of *ZIC1/4*-transduced MB051. Twenty-one fields of views for BFP-transduced MB051, 20 fields of views for one replicate of *ZIC1/4*-transduced MB051 and 27 fields of views for the other replicates. Images were captured at varying magnifications ranging from $\times 2$, $\times 10$, and $\times 40$.



Extended Data Fig. 7 | See next page for caption.

Extended Data Fig. 7 | Group 4 (G4) and SHH medulloblastoma *ZIC1* mutant overexpression result in distinct transcriptional changes in group 3 (G3) cells. **a**, *ZIC1* transcript levels (qRT-PCR) across the biological and technical replicates of G3 cell lines transduced with *ZIC1* constructs. Primers used are in Supplementary Table 1. **b**, Volcano plot summarizing genes differentially expressed in G4 medulloblastoma mutant vs. wild-type (WT) *ZIC1* and SHH medulloblastoma mutant vs. WT *ZIC1*-transduced G3 medulloblastoma cells (D425 and D283). Genes that are upregulated with WT *ZIC1* compared to empty vectors are highlighted in purple. *P*adjusted threshold of 0.05 was used. **c**, Heatmap showcasing expression pattern of all WT *ZIC1*-induced genes across all *ZIC1* mutation construct overexpressing cells. G4 medulloblastoma *ZIC1*

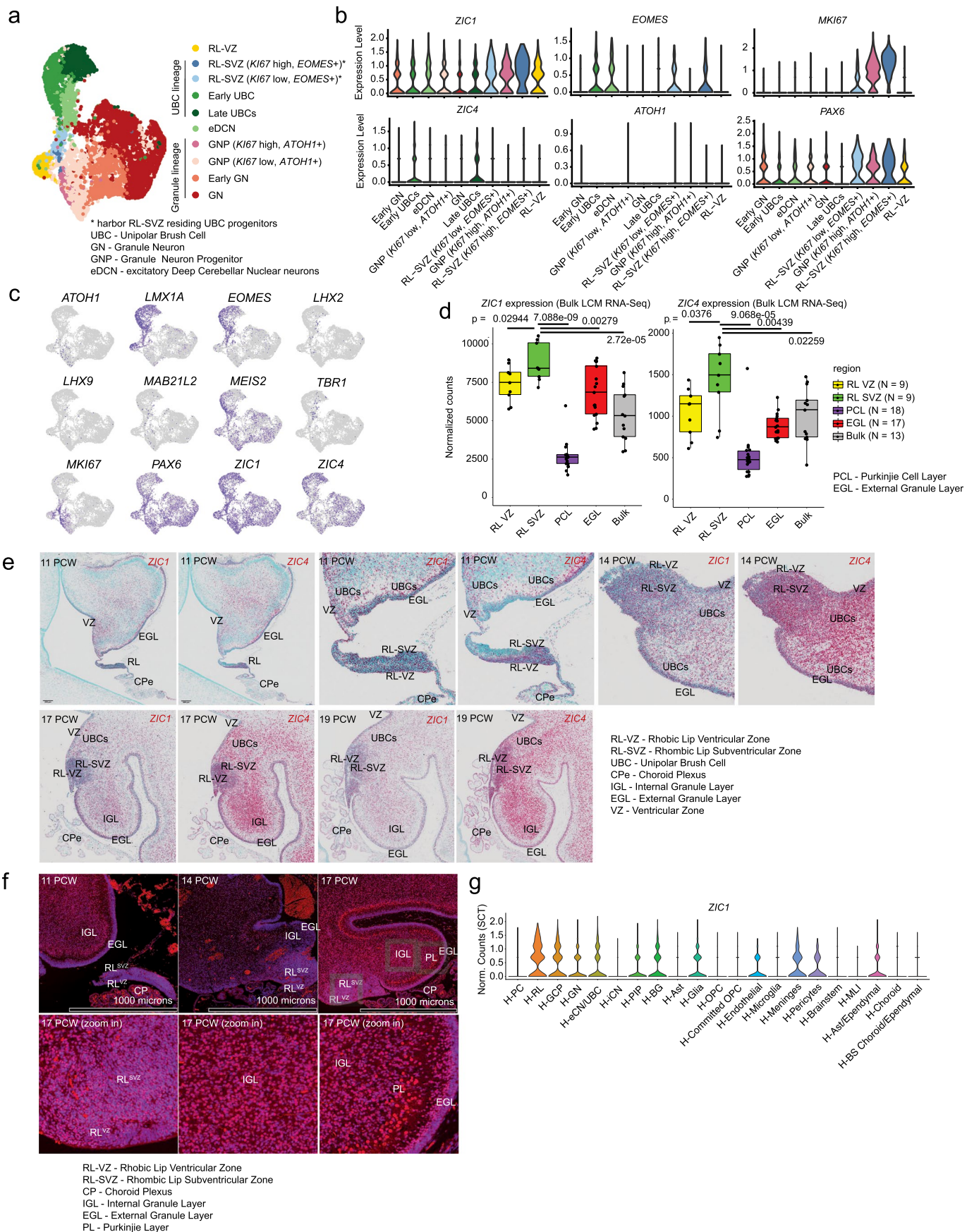
mutants exhibit reduced upregulation of the *ZIC1* target genes, whereas SHH medulloblastoma *ZIC1* mutants exhibit augmented upregulation of these genes. **d**, Pathway analysis of genes upregulated with WT *ZIC1* construct compared to empty vector. **e**, Pathway analysis of genes that are downregulated with G4 medulloblastoma *ZIC1* mutant compared to WT *ZIC1*. **f**, Pathway analysis of genes upregulated by SHH medulloblastoma *ZIC1* mutant compared to WT *ZIC1*. **g**, Number of ChIP-seq peaks identified from Flag-tagged *ZIC1* ChIP-seq in D283 cells transduced with WT *ZIC1* or G4 medulloblastoma *ZIC1* mutant. Two biological replicates were generated for each arm, using different constructs for the G4 medulloblastoma *ZIC1* mutants.



Extended Data Fig. 8 | See next page for caption.

Extended Data Fig. 8 | *ZIC1* regulates *Gli2* and cell cycle pathway genes in granule cells. **a**, Number of ChIP-seq peaks identified from Flag-tagged ZIC1 ChIP-seq in granule neuron progenitor (GNP) cells transduced with wild-type (WT) *ZIC1* or group 4 (G4) medulloblastoma *ZIC1* mutant. Two biological replicates were generated for WT *ZIC1* and three for G4 medulloblastoma *ZIC1* mutants. **b**, Distribution of normalized reads for WT vs. G4 medulloblastoma mutant Flag-tagged ZIC1-transduced GNP cells across peaks identified from FLAG ChIP-seq. **c**, Schematic summarizing the RNA-seq libraries generated from mouse granule lineage cells. **d**, Top 10 pathways downregulated by *ZIC1* overexpression compared to empty vector in bulk granule cells and GNPs. **e**, Expression level of *GLI2* across different medulloblastoma molecular subgroups. Plot was generated using the RNA-seq cohort used in the study

(N = 311). *GLI2* exhibits a highly SHH medulloblastoma-specific expression pattern. Center of box—median. Bounds of box—25% and 75% percentile. Whiskers show minimum and maximum values within the 1.5× interquartile range. P values calculated by two-tailed Mann–Whitney U test. **f**, *Zic1/2* ChIP-seq track demonstrating presence of peaks on the *Gli2* promoter in 2 immunoprecipitation replicates but not in input (data for **f–h** from [GSE60731](#)). **g**, Volcano plot summarizing genes differentially expressed by knocking down *Zic1* from mouse GNP. *P*adjusted threshold = 0.05. **h**, Normalized counts of *Gli2* transcript in control shRNA and *Zic1* shRNA treated GNP. Biological sample size = 2 for each arm. *P*adjusted value was obtained from DESeq2 differential expression analysis.



Extended Data Fig. 9 | See next page for caption.

Extended Data Fig. 9 | *ZIC1/4* are expressed throughout the rhombic lip, particularly in the rhombic lip ventricular zone (RL-VZ) and rhombic lip subventricular zone (RL-SVZ). **a**, Breakdown of glutamatergic neuronal cell lineage from developing human cerebellum (panel **a–c** from ref. 59 data). RL-SVZ cell populations were further subdivided according to expression pattern of *KI67*, *EOMES* and *ATOH1*. **b**, Violin plots summarizing expression level of *ZIC1*, *ZIC4*, *KI67* and other transcription factors critical for rhombic lip development throughout distinct glutamatergic lineage cell types. **c**, Feature plot summarizing expression levels for 12 developmental transcription factors across the developing human rhombic lip. **d**, Bulk RNA-seq quantification of *ZIC1* and *ZIC4* transcript levels across human rhombic lip regions isolated by laser capture microdissection (LCM; ref. 37 data). Center of box—median. Bounds of box—25%

and 75% percentile. Whiskers show minimum and maximum values within the 1.5× interquartile range. P values from two-tailed Mann–Whitney U test. **e**, RNA-scope visualization of *ZIC1* and *ZIC4* expression pattern across different regions of the rhombic lip in developing human cerebellum (11–19 postconception weeks). High expression level of both transcripts is observed across all regions, particularly in the RL-VZ and RL-SVZ. Biological sample size of 1 for 11, 14, 17 and 19 post-conception weeks (PCW). **f**, Immunofluorescence result showcasing *ZIC1* protein expression pattern across different regions of the rhombic lip in developing human cerebellum (11–17 postconception weeks). Biological sample size of 1 for 11, 14 and 17 PCW. Three different sections were used for each sample. Representative images are shown. **g**, Violin plots summarizing expression level of *ZIC1* transcript across different cells of the developing cerebellum (ref. 59 data).

Reporting Summary

Nature Portfolio wishes to improve the reproducibility of the work that we publish. This form provides structure for consistency and transparency in reporting. For further information on Nature Portfolio policies, see our [Editorial Policies](#) and the [Editorial Policy Checklist](#).

Statistics

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

n/a Confirmed

- 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
Only common tests should be described solely by name; describe more complex techniques in the Methods section.
- A description of all covariates tested
- A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
- A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
- 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 was used to collect data.
Data analysis	<p>Bowtie2 2.2.1 https://github.com/BenLangmead/bowtie2</p> <p>MACS2 2.1.1.20160309 https://pypi.org/project/MACS2/</p> <p>DeepTools 3.1.3 https://github.com/deeptools/deepTools</p> <p>Samtools 1.5 https://github.com/samtools/</p> <p>Bedtools 2.27 https://github.com/arq5x/bedtools2</p> <p>ROSE 0.1 https://bitbucket.org/young_computation/rose/src/master/</p> <p>GenomicRanges https://bioconductor.org/packages/release/bioc/html/GenomicRanges.html</p> <p>STAR 2.7.4 https://github.com/alexdobin/STAR</p> <p>HTSeq 0.6.0 https://github.com/simon-anders/htseq</p> <p>DESeq2 1.26.0</p>

<https://github.com/mikelove/DESeq2>
 FeatureCounts 1.6.2
<https://rdrr.io/bioc/Rsubread/man/featureCounts.html>
 CRC
<https://github.com/linlabcode/CRC>
 FIMO 5.0.5 51
<https://memesuite.org/meme/doc/fimo.html>
 Bowtie2 2.3.4
<https://github.com/BenLangmead/bowtie2>
 HiC-pro 2.9.0
<https://github.com/nservant/HiC-Pro>
 Hichipper 0.7.3
<https://github.com/aryeelab/hichipper>
 BWA-mem 0.7.8
<https://github.com/lh3/bwa>
 Platypus 0.8.1
<https://github.com/andyrimmer/Platypus>
 EAGLE2
<https://alkesgroup.broadinstitute.org/Eagle/#Xeagle2>
 Bcftools 1.9
<http://www.htslib.org/download/>
 Affymetrix Power Tools 1.18.2 ThermoFisherScientific
<https://www.thermofisher.com/ca/en/home/lifescience/microarrayanalysis/microarrayanalysis-partnersprograms/affymetrixdevelopersnetwork/affymetrixpower-tools.html>
 GISTIC 2.0.23
https://www.genepattern.org/modules/docs/GISTIC_2.0#gsc.tab=0
 G profiler
<https://biit.cs.ut.ee/gprofiler/Control-FREEC 10.3 32 http://boevalab.inf.ethz.ch/FREEC/>
 Genome Analysis Tool Kit 4.1.2.0
<https://github.com/broadinstitute/gatk/releases>
 Complexheatmap 2.2.0
<https://jokergoo.github.io/ComplexHeatmap-reference/book/>
 Homer 4.9
<http://homer.ucsd.edu/homer/>

Unique codes that were used in the study are available at the github page:
https://github.com/jjy-lee/ZIC_medulloblastoma

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 and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio [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 description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our [policy](#)

The FLAG-ChIP-Seq, RNA-Seq data generated from ZIC1 mutant construct transduced G3 MB cell lines and granule cells have been deposited in the Gene Expression Omnibus (GEO) database under the accession numbers GSE217639, GSE217571 and GSE217638.
 (Reviewer token: aladwoeqfhmphw).

Bulk H3K27ac, H3K27me3 ChIP-Seq, RNA-Seq, WGS and H3K27ac hiclip data generated from primary MB tumor samples in this study have been deposited in the European Genome-Phenome Archive (EGA) database under the accession code EGAS00001006741. The published MB bulk RNA-Seq data referenced in this study are available in the EGA database under the accessions EGAS00001001953, EGAD00001004347, EGAD00001004435, EGAS00001005826, EGAD00001001899 and EGAD00001004958. The published MB WGS data referenced in this study are available in the EGA database under the accessions EGAS00001001953, EGAD00001003125 and EGAD00001004347. The published MB 27ac ChIP-Seq data referenced in this study are available in the EGA database under the accessions EGAS00001001953. The Affymetrix SNP 6.0 data referenced during the study are available in the GEO database under the accession GSE37385. Expression array used for transcript abundance comparison between medulloblastoma subtypes are available in the GEO database under the accession GSE132269.

Multiple databases were used for annotation of SNPs and promoter, which were referenced in this study. These include the GRCh37 dbSNP151 (https://ftp.ncbi.nlm.nih.gov/snp/organisms/human_9606_b151_GRCh37p13/VCF/), GENCODE (v.19) (https://www.gencodegenes.org/human/release_19.html), the hg19 reference genome (<https://hgdownload.soe.ucsc.edu/goldenPath/hg19/bigZips/>), the hs37d5 reference genome (https://ftp943trace.ncbi.nih.gov/1000genomes/ftp/technical/reference/phase2_reference_assembly_sequence/), ERCC spike-in sequence (<https://www.encodeproject.org/files/ENCF908UQN/>) and Caltech profile 3 spike-in sequence (https://www.encodeproject.org/references/946_ENCSR193ZXE/). snRNA-seq data from the developing human cerebellum were obtained through correspondence from Aldinger et al. 2021 and are available through the Human Cell Atlas (<https://www.covid19cellatlas.org/aldinger20>), the UCSC Cell Browser (<https://cbi950dev.cells.ucsc.edu>) or from Database of Genotypes and Phenotypes (dbGaP)(accession number phs001908.v2.p1). Bulk RNA-seq data from the developing human cerebellum were obtained through correspondence from Halidpur et al. 2019 and are available through the dbGaP (accession number phs001908.v2.p1).

Original codes used for the study are available via GitHub(https://github.com/jjy-lee/ZIC_medulloblastoma/).

Research involving human participants, their data, or biological material

Policy information about studies with [human participants or human data](#). See also policy information about [sex, gender \(identity/presentation\), and sexual orientation](#) and [race, ethnicity and racism](#).

Reporting on sex and gender

N/A

Reporting on race, ethnicity, or other socially relevant groupings

Patients diagnosed with cerebellar pediatric brain tumors were recruited from McGill University Health Centre and the Hospital for Sick Children/The Arthur and Sonia Labatt Brain Tumour Research Centre Biobank.

Primary tumors used in the study were obtained from the Medulloblastoma Advanced Genomics International Consortium (MAGIC) and International Cancer Genome Consortium (ICGC). All materials were collected after receiving written informed consents, including consent to publish the generated data, as per guidelines from Research Ethics Board from the following institutes: Agostino Gemelli University Hospital, Children's Hospital of Minnesota, Cooperative Human Tissue Network, David Geffen School of Medicine at University of California Los Angeles, Duke University, Emory University, Erasmus University Medical Centre, German Cancer Research Centre (DKFZ), Hospital Cantonal De Geneve, Hospital Infantil de Mexico Federico Gomez, Hospital Sant Joan de Deu, Ludwig Maximilans University, Masaryk University, McGill University, McMaster University, Memorial Sloan Kettering Cancer 969 Centre, Miami Children's Hospital, Portugese Cancer Institute, Queensland Children's Tumor Bank, Seattle Children's Hospital Fred Hutchinson Cancer Research Centre, Seoul National University Children's Hospital, Stanford University School of Medicine, The Chinese University of Hong Kong, Tohoku University, University of California San Francisco, University Health Network, Universitats Kinderklinik, Universite de Lyon, University of Arkansas, University of Calgary, University of Debrecen Medical and Health Science Centre, University of Pittsburgh, University of Ulsan Asan Medical Centre, University of Warsaw Children's Memorial Health Institute, Vanderbilt Medical Centre and Wolfson Children's Hospital.

Human cerebellar samples were obtained under approval from the Seattle Children's Research Institute IRB. Samples were collected with consent and in accordance with institutional and legal ethics guidelines, from the Human Developmental Biology Resource (HDBR), University College London and Newcastle University, United Kingdom, the Birth Defects Research Laboratory (BDRL) at the University of Washington, USA, and the Hôpital Necker-Enfants Malades in Paris, France.

Population characteristics

N/A

Recruitment

N/A

Ethics oversight

REB MCH003-26 approved by McGill University Health Centre (Montreal). REB 0020020238 and 1000055059 approved by the Hospital for Sick Children (Toronto).

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

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 Behavioural & social sciences Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see nature.com/documents/nr-reporting-summary-flat.pdf

Life sciences study design

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

Sample size

Statistical methods were not used to predetermine the sample size. Sample sizes were chosen based on the availability of the primary tumors from the tumor bank.

Data exclusions

Tumors were excluded from the study if molecular classifier tumor identity turned out to be not medulloblastoma. Except these cases, no data were excluded from the study.

Replication

Experiments were performed in technical and biological replicates, and similar experiments performed across two different labs (Northcott lab, Taylor lab) lead to similar results.

Randomization

Randomization was not relevant to our study, as we were interested in interrogating molecular differences between known tumor identities.

Blinding

Blinding was not relevant to our study, as we were interested in interrogating molecular differences between known tumor identities.

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
<input type="checkbox"/>	<input checked="" type="checkbox"/> Antibodies
<input type="checkbox"/>	<input checked="" type="checkbox"/> Eukaryotic cell lines
<input checked="" type="checkbox"/>	<input type="checkbox"/> Palaeontology and archaeology
<input type="checkbox"/>	<input checked="" type="checkbox"/> Animals and other organisms
<input checked="" type="checkbox"/>	<input type="checkbox"/> Clinical data
<input checked="" type="checkbox"/>	<input type="checkbox"/> Dual use research of concern
<input checked="" type="checkbox"/>	<input type="checkbox"/> Plants

Methods

n/a	Involved in the study
<input type="checkbox"/>	<input checked="" type="checkbox"/> ChIP-seq
<input checked="" type="checkbox"/>	<input type="checkbox"/> Flow cytometry
<input checked="" type="checkbox"/>	<input type="checkbox"/> MRI-based neuroimaging

Antibodies

Antibodies used

H3K27ac Active Motif 39133
 H3K27me3 Diagenode C15410069
 H3K27me3 Cell Signalling Tech 9733
 FLAG Sigma-Aldrich F1804
 ZIC1 Sigma-Aldrich HPA004098
 ZIC4 ThermoFisher Scientific PA5-56392
 H3 Abcam 1791
 GAPDH Cell Signalling Tech 2118
 Actin Cell Signalling Tech 8457
 Rabbit IgG secondary antibody Thermofisher A27039

Validation

H3K27ac antibody has been validated by ChIP-Seq (PMID 29258295), ChIP-qPCR, immunofluorescence, western blot and dot blot analysis (Active Motif website).

H3K27me3 (Diagenode) antibody has been validated by ChIP-qPCR, ChIP-Seq (PMID: 24553142), dot blot and western blot (Diagenode website).

H3K27me3 (Cell Signaling) antibody has been validated by western blot, immunohistochemistry, immunofluorescence, flow cytometry, ChIP-Seq (PMID: 33259802) and cut&run (Cell Signaling Technology website).

FLAG antibody has been validated by immunoblotting, immunoprecipitation, immunohistochemistry, immunofluorescence, immunohistochemistry (Sigma-Aldrich website) and ChIP-Seq (PMID: 28215080).

Histone ChIP-Seq antibodies have also been validated with ChIP-qPCR within the laboratory, using primers against positive and negative control regions.

ZIC1 antibody has been validated by immunohistochemistry and immunofluorescence (Sigma-Aldrich website).

ZIC4 antibody has been validated by immunohistochemistry and immunofluorescence (Thermofisher website).

H3 antibody has been validated by western blot (Abcam website).

GAPDH antibody has been validated by western blot, immunohistochemistry, immunofluorescence and flow cytometry (Cell Signaling Technology website).

Actin antibody has been validated by western blot, immunofluorescence and flow cytometry (Cell Signaling Technology website).

Western blot antibodies have been validated with appropriate positive and negative control samples within the laboratory.

Eukaryotic cell lines

Policy information about [cell lines and Sex and Gender in Research](#)

Cell line source(s)

D283 (G3 MB cell line, male, PMID: 4056828), D425 (G3 MB cell line, male, PMID: 1904513), BT2019051 (patient derived G3 MB xenograft, male, derived from this study)D283 (G3 MB cell line, male, PMID: 4056828), D425 (G3 MB cell line, male, PMID: 1904513), BT2019051 (patient derived G3 MB xenograft, male, derived from this study)

Authentication

D283 and D425 were authenticated by STR profiling. BT2019051 was generated at the Hospital for Sick Children and passaged only in vivo.

Mycoplasma contamination

Cell lines were not tested for mycoplasma contamination.

Commonly misidentified lines
(See [ICLAC](#) register)

N/A

Animals and other research organisms

Policy information about [studies involving animals](#); [ARRIVE guidelines](#) recommended for reporting animal research, and [Sex and Gender in Research](#)

Laboratory animals	Mus Musculus, NOD scid gamma mouse, 6-10 weeks of age
Wild animals	This study did not involve wild animals.
Reporting on sex	Findings in the study do not apply to one sex. Sex was not considered in the study design. Sex based analysis was only performed for determining bias in H3K27me3 peak presence on BCOR promoter (Extended Figure 2). For other analysis, sex based analysis was not performed, as there was no apparent bias in the manifestation of the observed genetic/epigenetic phenomenon.
Field-collected samples	This study did not involve samples collected from the field.
Ethics oversight	All mouse breeding and procedures were performed as approved by The Centre for Phenogenomics.

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

Plants

Seed stocks	Report on the source of all seed stocks or other plant material used. If applicable, state the seed stock centre and catalogue number. If plant specimens were collected from the field, describe the collection location, date and sampling procedures.
Novel plant genotypes	Describe the methods by which all novel plant genotypes were produced. This includes those generated by transgenic approaches, gene editing, chemical/radiation-based mutagenesis and hybridization. For transgenic lines, describe the transformation method, the number of independent lines analyzed and the generation upon which experiments were performed. For gene-edited lines, describe the editor used, the endogenous sequence targeted for editing, the targeting guide RNA sequence (if applicable) and how the editor was applied.
Authentication	Describe any authentication procedures for each seed stock used or novel genotype generated. Describe any experiments used to assess the effect of a mutation and, where applicable, how potential secondary effects (e.g. second site T-DNA insertions, mosaicism, off-target gene editing) were examined.

ChIP-seq

Data deposition

Confirm that both raw and final processed data have been deposited in a public database such as [GEO](#).

Confirm that you have deposited or provided access to graph files (e.g. BED files) for the called peaks.

Data access links	https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE217639 https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE217571 https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE217638 (Reviewer token: aladwoeqfhmphwv)
	Bulk H3K27ac, H3K27me3 ChIP-Seq, RNA-Seq, WGS and H3K27ac hicchip raw data generated from primary MB tumor samples in this study have been deposited in the European Genome-Phenome Archive (EGA) database under the accession code EGAS00001006741.
Files in database submission	narrowPeak and broadPeak files generated from MACS2 as well as count matrices for G3 MB cells transduced with ZIC1 constructs
Genome browser session (e.g. UCSC)	https://genome.ucsc.edu/s/jjylee/hg19_ZIC1_locus_MB_ChIP

Methodology

Replicates	Primary tumors - single H3K27ac and/or H3K27me3 ChIP-Seq libraries were generated for different biological medulloblastoma tumors without technical replicates. 102 samples for H3K27ac, 63 samples for H3K27me3.
	Cell lines with FLAG tagged ZIC1 constructs - at least two biological replicates were generated for WT ZIC1 ChIP-Seq and Group 4 mutant ZIC1 ChIP-Seq.
Sequencing depth	Each ChIP-Seq library was sequenced with at least 30M reads, typically resulting in >25M uniquely mapped reads. Primary tumor samples were sequenced with 126 bp (27ac, 27me3) or 101 bp (27ac - active motif) paired end reads. For cell lines, samples were sequenced with 151 bp paired end reads.
Antibodies	H3K27ac Active Motif 39133 H3K27me3 Diagenode C15410069

H3K27me3 Cell Signalling Tech 9733
FLAG Sigma-Aldrich F1804

Peak calling parameters

```
H3K27ac samples with inputs
macs2 callpeak -t IP_bam -c input_bam -f BAMPE -g hs --nomodel -B -q 1e-2

H3K27ac samples without inputs
macs2 callpeak -t IP_bam -f BAMPE -g hs --nomodel -B -q 1e-2

H3K27me3 samples
macs2 callpeak -t -t IP_bam -c input_bam -f BAMPE -g hs --nomodel --broad -B -q 1e-5 --broad-cutoff 1e-4

FLAG tagged ZIC1 ChIP-Seq (D283)
macs2 callpeak -t IP_bam -c input_bam -f BAMPE -g hs --nomodel -B -q 1e-5

FLAG tagged ZIC1 ChIP-Seq (GNP)
macs2 callpeak -t IP_bam -c input_bam -f BAMPE -g hs --nomodel -B -q 5e-2
```

Data quality

For H3K27ac, on average, each sample exhibited 30k peaks with FDR < 1e-2 and fold enrichment > 5. For H3K27me3, on average, each sample exhibited 5k peaks with FDR < 1e-4 and fold enrichment > 5. Unsupervised hierarchical clustering lead to robust recapitulation of known molecular subgroups of medulloblastoma for both marks, suggesting that the generated data set are able to identify known biological identities in unbiased manner.

Software

```
Bowtie2 2.2.1 https://github.com/BenLangmead/bowtie2
MACS2 2.1.1.20160309 https://pypi.org/project/MACS2/
Deeptools 3.1.3 https://github.com/deeptools/deepTools
Samtools 1.5 https://github.com/samtools/
Bedtools 2.27 https://github.com/arq5x/bedtools2
ROSE 0.1 https://bitbucket.org/young\_computation/rose/src/master/
GenomicRanges https://bioconductor.org/packages/release/bioc/html/GenomicRanges.html
CRC https://github.com/lilnlabcode/CRC
FIMO 5.0.5 https://memesuite.org/meme/doc/fimo.html
Homer 4.9 http://homer.ucsd.edu/homer/
```

Original codes used for analysis of ChIP-Seq data are available at GitHub (https://github.com/jjy-lee/ZIC_medulloblastoma/).