INTRODUCTION

Medulloblastoma (MB), a malignant embryonal brain tumor with a peak incidence in childhood, exhibits considerable molecular heterogeneity with the existence of at least four distinct subgroups—Wingless (WNT), Sonic Hedgehog (SHH), Group3 (G3) and Group4 (G4).1,2 These subgroups have distinct characteristics with respect to age, gender, prognosis and response to therapy.3 The genetic and phenotypic differences of MB subgroups are in part attributable to differences in their cellular origin.4 Identification of the cellular origin of a tumor often relies on the molecular characterization of normal and tumor tissues; however, tumor-specific somatic alterations likely confound such analysis. Thus, the generation of genetically engineered animals bearing tumors arisen under physiological conditions provides a context for investigating oncogenic events underlying the transformation of a normal cell into a tumor cell in vivo.

Spatial and temporal specification of cell fate within the cerebellum is well characterized5 and has been integral to the identification of molecular signatures associated with cerebellar neuronal subtypes along the spectrum of neurogenesis.6 These neuronal lineages provide a means to assess the effect of genetic modifications (e.g. activation of oncogenes and deletion of tumor suppressor genes) on a specific neuronal sub-population, and thus to develop genetically engineered MB mouse models.7 A number of studies have uncovered distinct cells of origin for MB subgroups: genetic ablation of Ptc1 enables descendants of cerebellar stem cells (e.g. granule neuron precursors (GNPs) and Nestin-positive (+) cells) to form SHH MBs,8–10 whereas expression of a constitutively active mutant form of Ctnnb1 in dorsal hindbrain progenitors, generates WNT MBs.11 In contrast, forced activation of WNT signaling in GNPs impairs their proliferation and induces differentiation,12–14 which highlights the differences in cancer susceptibility among neural progenitors under the influence of the same oncogenic insult. Enforced expression of MYCN under the glutamate transporter 1 (Glt1) promoter or in neural cells positive for the glial fibrillary acidic protein (GFAP)15 induces MBs with high expression of Kcnal1, a known marker of G4 MBs (MBG4),15 whereas MYCN overexpression in Trp53-null GNPs or embryonic cerebellar stem cells triggers MBG4.15–17 These studies demonstrate that genetic insults to specific cerebellar cell types can influence the subgroup-specific characteristics of MBs.

MYC-driven MBs are mainly classified as G3 (MBG3) and represent one of the most aggressive subgroup.2,3 They are characterized by frequent metastasis at diagnosis and are often associated with a dismal outcome.18 So far, several orthotopic models of MYC-driven MBG3 were developed in mice by forced activation of MYCN in GNPs or cerebellar stem cells.19–21 We and another group independently...
reported the first orthotopic mouse model of MBG3 by over-expressing Myc in combination with functional loss of Trp53 in postnatal cerebellar cells selected by fluorescence-activated cell sorting for the basic helix-loop-helix transcription factor Atoh1-positive (Atoh1+) and Prominin/CD133-positive (Prom1+) cells. Nevertheless, while human MBG3 have been found in relatively young children, the role of Myc expression in transforming embryonic stem/progenitors into malignant cells under physiological conditions has not yet been tested. We here report the first MBG3 model from embryonic cerebellar cells by Myc activation and loss of Trp53 function using in utero electroporation (EP)-based in vivo gene transfer combined to a
Cre/LoxP-mediated technology. The present study also provides the opportunity to trace tumor growth with bioluminescence and fluorescent proteins, which will help in the future to not only understand cellular and molecular mechanisms of tumorigenesis but also to undertake further preclinical trials.

RESULTS

EP of embryonic cerebellar neuroepithelium with Myc and Trp53DN induces MBG3

Human MBG3 is typically restricted to infants and younger children,20 prompting us to assess the predisposition of embryonic cerebellar progenitors to initiate MBG3. Previous orthotopic MBG3 models from postnatal progenitors required loss of Trp53 function to overcome cell death caused by excess Myc expression.12 Consistent with these results, the oncogenic transcriptional repessor growth factor independent 1, known to antagonize Trp53,21 was subsequently shown to cooperate with Myc to induce MBG3 in an orthotopic transplant model.22 Indeed, MYC-TP53 dysfunction has been reported in relapsed human MBG3.23 We reasoned that we could use Myc expression and a dominant-negative form of Trp53 to model MBG3 using embryonic cells. In vivo EP of plasmids into mouse embryos is an effective method to transfer genes into cerebellar progenitors under physiological conditions.11,24 To avoid dilution of plasmid-driven gene expression as cells divide, we used the Tol2 transposon-mediated genomic integration system.25 We constructed two LoxP (LRL)-Myc-IRES-Luciferase (Luc) plasmids: Tol2-CAG-LoxP-DsRed2-LoxP (LRL)-Myc-IREs-Luciferase (Luc)-Tol2 (pT2K LRL-Myc-IREs-Luc) that strongly express both Myc and Luc after excision of the LRL cassette in the presence of the Cre recombinase protein (Figures 1a and b) and Tol2-CAG-EP1-EP2-Flp,encoding plasmids (pCAggs T2P) that lack Tol2 cis elements to prevent multiple gene hopping (Figure 1c). At E13.5, Cre expression was restricted to the cerebellar ventricular zone (VZ) and the external granule layer (EGL), the two germinal zones of the cerebellum (Supplementary Figures 1a and b). Lineage tracing analysis in Blbp-Cre-IREs-nlacZ; R26-LSL-EYFP mice identified broadly distributed enhanced yellow fluorescence protein-positive (EYFP)1 cells at P7 (Supplementary Figures 1c and d). While Calb1+ Purkinje cells lack EYFP (Supplementary Figure 1e), the great majority of cerebellar cells, including inhibitory cerebellar interneuron progenitors (Pax2+), granule cells (Pax6+) as well as Bergmann glia and multipotent progenitors (Sox2+), expressed EYFP (Supplementary Figures 1f–h). We electroporated the LRL-EGFP gene27 into the cerebellar neuroepithelium of [Blbp-Cre-IREs-nlacZ] embryos by EP at E13.5 to confirm that Cre-mediated recombination of transfected plasmids occurred in vivo. Two days after EP (E15.5), most of the labeled cells expressed enhanced green fluorescent protein (EGFP) rather than DsRed2 (Supplementary Figures 1i and j), which is consistent with the distribution of Cre+ cells revealed by X-gal staining (Supplementary Figure 1k). We also observed a small population of EGFP+ cells in the choroid plexus (CP) (Supplementary Figure 1j). Immunohistochemistry (IHC) revealed successful recombination of the transfected genes within Pax6+ cells in the EGL, Sox2+ cells in the VZ and Pax2+ cells migrating inwards deep into the cerebellum (Supplementary Figures 1l–n).

We next performed in utero EP of LRL-Myc-IREs-Luc and T2TP with or without Trp53DN-copGFP in [Blbp-Cre-IREs-nlacZ] embryos at E13.5 and subsequently monitored Luc activation in neonatal mice by in vivo bioluminescence. Strong Luc signal was detected in the head of Cre-carrying, but not wild-type (Cre-negative) mice (Figure 1d). In the absence of Trp53DN-copGFP, the bioluminescence signal disappeared by 5 weeks after birth and no MBs developed within 5 months after birth (n=6, Supplementary Figure 2a). These data indicate that overexpression of Myc alone is insufficient to induce MBs in vivo. Intriguingly, in one out of six cases, a CP carcinoma (CPC) was induced by Myc overexpression in the absence of Trp53 loss (Supplementary Figure 2b), which is consistent with the activation of EGFP expression downstream of LRL in electroporated CP cells (Supplementary Figure 1j). In contrast, triple EP of LRL-Myc-IREs-Luc, Trp53DN-copGFP and T2TP resulted in cerebellar tumors with 100% penetrance. Luc signals increased steadily (Figure 1e) and all mice were killed within 8 weeks of birth due to progressive tumor burden (Figure 1f). Six out of 14 mice developed only CPCs in the fourth ventricle (Supplementary Figures 2d–f and Table 1). All the other tumors were diagnosed as MBs with large-cell anaplastic characteristics (hereafter called Blbp-MYC MB) (Figures 1e–h). Tumors were highly proliferative (Ki67+) with MBG3 characteristics, namely Npr3 expression (Figures 1i and j). Importantly, tumor cells strongly expressed copGFP (Figure 1k), indicating that Trp53 inactivation is a prerequisite for MBG3 formation. EP-induced Blbp-MYC MBs also showed strong expression of known stem cell marker genes (Figure 1l). These tumors displayed gene expression profiles for the most part different from Myc-driven CPCs; however, some genes were shared with MBG3 (e.g. Npr3) (Supplementary Figures 2e and f). Hematoxylin and eosin (H&E)-stained sections confirmed leptomeningeal metastasis of the tumors (Supplementary Figures 3a and b). The luciferase signal expanded from the hindbrain to the cerebral cortex as tumors developed, invading the olfactory bulb (Supplementary Figures 3c–h), implying a high tendency of Blbp-MYC MBs for metastatic dissemination as reported for human MBG3.28 No tumors developed without Myc overexpression (Figure 1f), excluding the possibility that Blbp-MYC MBs were accidentally caused by transposition of Tol2-flanked transgenes.
rather than due to Myc oncogenic properties. Thus, in vivo EP generated MBs from Blbp<sup>+</sup> neural stem/progenitor cells under physiological conditions.

Location of in utero EP-engineered murine MB<sub>G3</sub> was similar to their human counterparts

In a subset of patients enrolled in the SJCRH protocol, SJMB03, we measured the location of MB<sub>G3</sub> in magnetic resonance imaging (MRI)-acquired images and compared this localization to in utero EP-engineered murine tumors (Figure 2). All human MB<sub>G3</sub> tumors were located near the midline in the fourth ventricle, and the median anterior–posterior location was distinct from MB<sub>WNT</sub>MBs, but not MB<sub>G3</sub> (Figures 2a–d). MB<sub>WNT</sub> tumors clustered proximal to MB<sub>G3</sub>, reflecting an origin from hindbrain progenitors in the brain stem. MB<sub>Shh</sub> tumors were dispersed distal to MB<sub>G3</sub> in the cerebellar hemispheres, reflecting their origin from GNPs in the EGL. Immunoreactive for Atoh1 (Figures 3o–q). The intermediate distribution of human MB<sub>G3</sub> tumors is consistent with an origin in the VZ or upper rhombic lip, with subsequent growth into the ventricle as observed in the Blbp-MYC mice (Supplementary Figures 4a and b). Nevertheless, it might not allow proper evaluation of the possibility that Myc overexpression and loss of Trp53 function could transform committed inhibitory interneuron progenitors due to rapid growth of MB<sub>G3</sub> from GNPs. We therefore used pancreas specific transcription factor 1a ([Ptf1a]-Cre) and glutamic acid decarboxylase 2 ([Gad2]-IRESCre) mice that specifically target committed inhibitory neuron progenitors. In parallel, [Atoh1-Cre] mice were used to control for neuronal subtype since Cre is expressed in committed excitatory neuron progenitors, including GNPs. Targeted gene expression/disruption within specific cerebellar cell populations are well characterized in [Atoh1-Cre] and [Ptf1a-Cre] mice, whereas those in [Gad2-IRESCre] animals are less well characterized. EP of the LRL-EGFP plasmid into E13.5 [Gad2-IRESCre] embryos confirmed that GFP<sup>+</sup> cells were Pax2<sup>+</sup> (Supplementary Figures 5a–f), but negative for PCNA and Pax6 (Supplementary Figures 5g–j) at E15.5. Furthermore, lineage-tracing analysis using the [R26-LSL-tdTomaTo] strain confirmed strong expression of tdTomato in Purkinje cells (Calb1<sup>+</sup> / Parv<sup>+</sup>) and cerebellar inhibitory interneurons (Parv<sup>+</sup>) (Supplementary Figures 5k–l), but not Bergmann glia (Sox2<sup>+</sup>) and granule cells (Pax6<sup>+</sup>) (Supplementary Figures 5m–n) in the adult mice. Thus, the [Gad2-IRESCre] and [Ptf1a-Cre] mice can be used to target inhibitory neuronal progenitors.

| Cre transgenic mouse | Targeted cerebellar cells | Tumor latency | Tumor incidence |
|----------------------|---------------------------|--------------|-----------------|
| Atoh1-Cre<sup>a</sup> | Granule neural progenitors (Pax6<sup>+</sup>) | 4–14 weeks | n = 12/14 |
| Prom1-Cre<sup>a</sup> | Multipotent cerebellar stem cells (Sox2<sup>+</sup>) | 6–11 weeks | n = 8/10 |
| Blbp-Cre<sup>b</sup> | Granule neural progenitors (Pax6<sup>+</sup>/Atoh1<sup>+</sup>) Inhibitory interneurons (Pax2<sup>+</sup>) Bergmann glia/multipotent cerebellar stem cells (Sox2<sup>+</sup>) Choroid plexus progenitors | 5–8 weeks for MBs n = 14/14 (n = 8/14 for MBs, n = 6/14 for CPCs) | |
| Atoh1-Cre<sup>b</sup> | Granule neural progenitors (Pax6<sup>+</sup>) Choroid plexus progenitors | 7 weeks for MBs 4 weeks for CPCs n = 3/3 (n = 1/3 for MBs, n = 2/3 for CPCs) | |
| Gad2-IRESCre<sup>b</sup> | Inhibitory interneurons (Pax2<sup>+</sup>) | 9–13 weeks | n = 4/11 |
| Ptf1a-Cre<sup>b</sup> | Inhibitory interneurons (Pax2<sup>+</sup>) | 9–11 weeks | n = 3/7 |

Abbreviations: CPC, choroid plexus carcinoma; MB, medulloblastoma; n, number of animals. *Tamoxifen administration at P0 and P1. †In utero electroporation at E13.5.
recombinase in CP cells as well as GNPs (Supplementary Figure 5o). Unlike Blbp-MYC and Atoh1-MYC MBs, the bioluminescence signal intensity from transfected [Gad2-IRES-Cre] and [Ptf1a-Cre] mice gradually decayed during the first several weeks after birth, but subsequently showed a marked increase in intensity with mice succumbing to tumor burden between 9 and 14 weeks after birth (Figures 4a, c and d; hereafter called Gad2-MYC and Ptf1a-MYC MBs). Intriguingly, peak of Luc signals were detected in several brain regions in mice from both models (Figures 4c and d), implying multifocal tumor development. Histopathological analysis revealed that Ptf1a-MYC and Gad2-MYC MBs, as well as one Atoh1-MYC MB have a large-cell anaplastic MB phenotype and high expression of Npr3 (Figures 4e–j). Interestingly, hyperproliferative lesions were not found in the cerebellum of [Gad2-IRES-Cre] and [Ptf1a-Cre] mice, but only in [Atoh1-Cre] neonates (Figures 4k and l). These data suggest that absolute Luc intensity was much lower in [Gad2-Cre] and [Ptf1a-Cre] mice compared with [Blbp-Cre] and [Atoh1-Cre] animals (Figures 1e and 4b–d), possibly because hyperplasia had not been initiated at neonatal stages in these mice. The delay of hyperplasia formation could explain the longer latency of Gad2-MYC and Ptf1a-MYC MBs. High RNA expression of Npr3 and stem cells markers were observed in most of Gad2-MYC and Ptf1a-MYC MBs (Figure 4m), suggesting that overexpression of Myc and Trp53DN induced MBG3s from Ptf1a+/Gad2+ inhibitory neuron progenitors as well as from GNPs targeted by Blbp-Cre animals.

Figure 2. Localization of the Blbp-MYC mouse model and human MBG3. (a–d) Tumor-center location of human G3 (n = 16, yellow) tumors in the standard atlas space, plotted along with SHH (n = 22, red), WNT (n = 13, blue) and G4 (n = 19, green) tumors. Data are shown on atlas planes at the approximate center of the MBG3 tumor distribution: (a) coronal at y = −56, (b) sagittal at x = 5 and (c) transverse at z = −35. Coordinate units are mm. The anterior–posterior location (vertical axis in c) varied with subtype (d, P < 10−15) and G3 location was distinct from SHH (P = 0.0005) and WNT (P = 0.008), but not G4 (P = 0.063). (e–g) Heat map of Blbp-MYC MB tumors (n = 6). Tumors were outlined on representative H&E-stained sections (e and f) and overlayed on a parasagittal plane at x = 0.12 mm (g) from the Franklin and Paxino brain atlas. Coordinates units are mm relative to Bregma, and color indicates cumulative number of tumors. The rostral–caudal orientation of the atlas slice is reversed for comparison with the human atlas. (h–m) Representative MRI imaging of 4-year-old human MBG3 patient (h–j) and 57-day-old mouse bearing a tumor (k–m). Colored dots highlight points for comparison of the anatomic similarity of tumor localization that was apparent in 2/6 Blbp-MYC MB studied with MRI.

EP-induced Myc-driven MBs molecularly recapitulate human MBG3

Using different Cre drivers, we generated EP-based MBG3 models from different cell types (Table 1). We compared gene expression profiles of these tumors with orthotopic Myc-driven MBG3 models engineered from postnatal cerebellar progenitors. We here developed [Atoh1-CreER; Trp53flox−/−] and [Prom1-CreER; Trp53flox−/−] mice that lacked one allele of Trp53 in the germline. Mouse pups were administered tamoxifen at P0 and P1 to genetically delete
the LoxP-flanked Trp53 in CreER-expressing cells, then killed at P7. Lineage tracing analysis under the same condition confirmed that Atoh1" GNPs and Sox2" are targeted by Atoh1-CreER and Prom1-CreER drivers, respectively (Supplementary Figure 6).

Cerebellar cells were purified by percoll density gradient centrifugation, infected in vitro with retroviruses carrying Myc and red fluorescent protein and transplanted into the cortices of naive recipient nude mice as reported previously. Tumors from Atoh1-CreER" and Prom1-CreER" cells (hereafter referred to as

![Figure 4](image_url). MBG3 from cerebellar inhibitory interneuron progenitors. (a) Kaplan–Meier survival curve for mice developing Atoh1-MYC (n = 3/3, red curve), Gad2-MYC (n = 4/11, blue curve) and Ptf1a-MYC (n = 3/7, green curve) MBs. Atoh1-MYC tumors developed significantly faster than Gad2-MYC MBs (P < 0.0001, a two-tailed P-value). (b–d) Serial bioluminescence images of representative (b) glutamatergic (Atoh1-Cre) and (c and d) GABAergic (Gad2-IRE-Cre (c) and Ptf1a-Cre (d)) mice subjected to EP with LRL-Myc-Luc, Trp53DN-copGFP and T2TP. (e–j) Tumors stained with H&E (e, g and l) and an anti-Npr3 antibody (f, h and j). Scale bar = 25 μm. (k and l) Distribution of GFP" cells in electroporated cerebella of P2 Atoh1-Cre (k) and P3 Gad2-IRE-Cre (l) animals. (sagittal sections, scale bar = 100 μm). (m) Heat maps of subgroup-specific genes differentially expressed across EP-based MBG3 models.

![Figure 3](image_url). Hyperplasia in early development of Blbp-MYC MBs. (a–h) Posterior region of P1 Blbp-Cre-IRE-nlacZ cerebellum electroporated with LRL-Myc-IRE-Luc, Trp53DN-copGFP and T2TP genes (a–g and j–n) and with LRL-IRE-Luc, Trp53DN-copGFP and T2TP genes (h). (a) copGFP expression. (b and c) H&E staining on the ipsilateral (electroporated, b) and contralateral (non-electroporated, c) sides. Note: b is the neighboring section of a. Arrow indicates ectopic accumulation of cells on the VZ corresponding to where copGFP-positive cells form a cluster in a. Scale bar in a is 100 μm for a–c. (d–h and j–n) Confocal microscopy images of the hyperplasia stained with antibodies against Luc (d–e), Ki67 (f–h), Pax2 (j), Pax6 (k–l) and Sox2 (m–n). GFP" cells are devoid of Sox2 expression in the margin of the VZ (arrowheads in n). (e’, g’, I’ and n’) High magnification view of the area outlined by white squares in (e, g, I and n), respectively. Scale bar in j = 100 μm for (e, f, h, j, k and m) and scale bar in n’ = 25 μm for (e’, g’, I’ and n’). (i) Quantification of the percentage of Ki67" and Pax6" cells in copGFP" cells distributed in the IX and X lobules of neonatal cerebella (n = 3). The bars represent normalized mean value ± s.d. (o–t") Confocal microscopy images the most posterior region of P4 Blbp-Cre-IRE-nlacZ cerebellum electroporated with LRL-Myc-IRE-Luc, Trp53DN-copGFP and T2TP genes. (q–q") and (t’–t") are high-power view of the area q and t, respectively. (o–s) GFP expression in sagittal sections of the cerebellum counterstained with 4',6-diamidino-2-phenylindole (DAPI). IHC with Atoh1 (p) and Tbr2 (s) antibodies. Scale bar = 100 μm in (p and s). Scale bar = 25 μm in (q" and t"). Cerebellar surface is outlined by dashed lines.
Atoh1ER-MYC and Prom1ER-MYC MBs, respectively) developed within 3 months after birth (Supplementary Figure 7). These tumors as well as our previous model 16 exhibited large-cell anaplastic characteristics and high expression of *Npr3*, a faithful marker of MBG3 (Supplementary Figure 7).

Unsupervised hierarchical clustering using the top 1000 signature genes from the 430 v.2 Affymetrix microarray chip (Affymetrix, Santa Clara, CA, USA) revealed that EP-based MBG3 and orthotopic MBG3 tumors clustered separately from each other and from published murine MBWNT and MBSHH models (Figure 5a). Consistent with these results, principal component analysis using all genes measured on the arrays revealed that the MBG3 models reported here group together, were similar to our previous orthotopic models16 and were clearly separated from MBWNT and MBSHH models (Figure 5b). Of note, EP-MYC tumors did not form subclustering in a cellular origin-dependent manner (Supplementary Figure 8a). Rather, they were intermingled with each other, suggesting that dysfunction of MYC and TP53 is a strong determinant driving MBG3, overriding their origins. Consistent with this idea, cross-species analysis using orthologs...
between human and mouse further revealed significant similarity of EP-mediated tumors to human MBG3 (Figure 5c and Supplementary Figures 8b–d). Irrespective of the cell types examined in this study, EP-based tumors generated by aberrant activation of Myc combined with the loss of Trp53 function mimicked human MBG3.

**DISCUSSION**

Identifying the cellular origin of solid tumors is of utmost importance to better understand the transformation of a normal cell into a cancer cell, especially regarding the lineage dependency driving tumor subtype diversity. The in utero EP-based gene transfer system enabled us to examine the susceptibility of distinct cerebellar progenitors to transformation by Myc overexpression and Trp53 loss into malignant MBG3. Hyperplasia formation in Blbp-Cre and Atoh1-Cre mice at neonatal stages clearly indicated that cerebellar cellular transformation began during embryogenesis. The expression of Atoh1 but not Sox2 within the hyperplastic cells suggests a commitment of neural stem cells to GNPs before transformation ensues. In addition, the lack of hyperplasia in electroporated Ptf1a-Cre and Gad2-Cre neonatal cerebella, as well as incomplete penetration of MBG3 from these animals, implies resistance of inhibitory interneuron progenitors to Myc-induced transformation. Thus, our study suggests that there are critical periods during which distinct neuronal progenitors are susceptible to hyperproliferation and/or transformation in response to the same oncogenic insults. Tumor localization within the caudal cerebellum of Blbp-Cre animals was consistent with our measurements of human MBG3 tumors around the caudal midline of the fourth ventricle in tight clusters as well as with subgroup-specific categorical descriptions of human MBs based on diagnostic MRI studies. We previously showed that quantitative tumor localization within the posterior fossa reflects the molecular and cellular context of tumorigenesis for MBs and MBG3 in both mice and human. The distribution of human MBG3 tumors suggests that tumors arise from the caudal vermis, which develops from the medial cerebellar primordium in human and mouse, and postnatal lateral-to-medial migration of postmitotic EGL granule cells that populates the expanding caudal vermis, at least in the mouse. Convergence toward the midline of granule cells from an extended segment of the upper rhombic lip might afford opportunities for transformation of MYC-induced cells and help to explain the low variability of MBG3 tumors location in humans.

Our results show that Myc overexpression with functional inactivation of Trp53 developed tumors from multiple cellular origins. However, Myc and Trp53DN expression also induced CPCs from Blbp-Cre and Atoh1-Cre mice that target CP cells as well as neural cells. Although CP cells partly share the same origin (e.g. Gdf7+ cells) as GNPs, and postnatal lateral-to-medial migration of postmitotic EGL granule cells that populate the expanding caudal vermis, at least in the mouse. Convergence toward the midline of granule cells from an extended segment of the upper rhombic lip might afford opportunities for transformation of MYC-induced cells and help to explain the low variability of MBG3 tumors location in humans.

Plasmids and virus production

For Tol2 transposon-mediated stable expression, we used pCAGGS T2TP and pT2K LRL-EGFP. EGFP of pT2K LRL-EGFP was replaced with either IRES-luc or Myc-IRES-luc, yielding pT2K LRL-IRES-luc and pT2K LRL-Myc-IRES-luc, respectively. The gene encoding Trp53DN was fused to the T2A peptide with copGFP (Trp53DN-copGFP) and was inserted into the pT2K-CAGGS plasmid, yielding pT2K Trp53DN-copGFP. Retroviruses were produced as described previously.

**MATERIALS AND METHODS**

Animal husbandry

Genetically engineered mice used here were [Blbp-Cre-IRES-nLucZ] (01XMA, NC), [Atoh1-CreER] (007684, Jax), [Prom1-CreER] (017743, Jax), [Trp53Fl/−] (009642, Jax), [R26-LSL-EYFP] (006148, Jax), [R26-LSL-tdTomato] (007908, Jax), [Atoh1-Cre] (011104, Jax), [Gad2-IRES-Cre] (010802, Jax), [Ptf1a-Cre] (RES184, BCBC), Trp53-null (008652, Jax) and [Trp53−/−; Cdkn2c−/−]. Cre-carrying male were bred with CD-1 female mice (Charles River, Wilmington, MA, USA) for in utero EP. Day E0.5 was defined by the observation of a vaginal plug and the day of birth as P0. Conditional deletion of the Trp53 gene was performed by administration of tamoxifen (Sigma, St Louis, MO, USA) at P0 and P1 (4 mg/40 g body weight). All experiments were conducted in strict accordance with the National Institute of Health guidelines for the Care and Use of Laboratory Animals and according to the guidelines established by the SJCRH Institutional Animal Care and Use Committee. All procedures in the protocol were approved by the Animal Care and Use Committee (ACUC) of SJCRH (Animal Assurance Number: A3077-01) and were approved by the responsible authorities in Germany (G176/13 and G48/14).

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Generation of mouse MBG3 models

To develop EP-based MBG3 models, in utero EP was performed. Only the animals showing luciferase signal by P7 were selected for further analysis. Tumor growth was measured every 1–2 weeks by bioluminescence imaging of luciferase activity using a Xenogen IVIS system (PerkinElmer, Waltham, MA, USA). Orthotopic MBG3 mouse models were generated by cranial implants of purified GNPs from [Trp53−/−; Cdkn2c−/−; Atoh1-CreER], [Trp53−/−; Prom1-CreER] and [Trp53−/−; Ptf1a-Cre] mice, with infected with retroviruses carrying Myc and red fluorescent protein according to the previous study. No specific randomization or blinding was performed. Tumor cell purification and genomic DNA and total RNA extraction for genotyping and Affymetrix microarray analysis were described previously. For detection of non-recombined and recombined allelic of Trp53 in Atoh1-IER-MYC and Prom1-IER-MYC MBs, 1F/1R (370 bp) and 1F/10R (612 bp) primer sets were used, respectively. For internal control, endogenous Prom1 (563 bp) and Pch1 (220 bp) were detected. Comparison of survival curves was performed by calculation of two-tailed P-value using the GraphPad Prism software (GraphPad Software, Inc., La Jolla, CA, USA).
Gene expression analysis

Human Affymetrix U133 plus 2.0 expression array data for genes of interest were extracted from publicly available data sets (GSE10327, GSE12992, GSE37418 and GSE49243) from the R2 software tool for analysis and visualization of genomic data (http://r2.amc.nl) (Amsterdam, The Netherlands). Additional cases came from an unpublished data set generated at the DKFZ (Heidelberg, Germany) (M Kool et al., unpublished data). Total RNAs from murine tumors were extracted and analyzed with the Affymetrix 430 v.2 chip. The microarray data of murine SHH and WNT data). Total RNAs from murine tumors were extracted and analyzed with a heat map using Partek Genomics Suite 6.6.

RNA from four CPC and one MBG3 mouse tumors were extracted and analyzed by mogeno 2.0 ST arrays where the MBG3 RNA was a repeated measure of a pre-existing 430 v.2 array. Data were RMA normalized in Partek Genomics Suite 6.6 (St Louis, MO, USA) and imported in STATA/MP 14.2 (StataCorp LLC, College Station, TX, USA) in which unannotated and duplicated probe sets were removed such that only the highest measured probe sets was retained for any gene. The mouse 430 v.2 array data was also deduplicated in the same manner for MB443 and MBG3 data. These two data sets were then joined by gene symbol and a chip-based correction for each gene was calculated by two data sets were then joined by gene symbol and a chip-based correction for each gene was calculated by finding the simple difference in RMA signal in the 430 v.2 sample that matched the mogeno 2.0 ST Grp3 sample. Finally, the mogeno 2.0 samples data were corrected using this factor and the resulting matrix was principal component analysis visualized. Further selected matched genes were clustered and displayed in a heat map using Partek Genomics Suite 6.6.

Antibodies
All antibodies used in this study are shown in Supplementary Table 1.

Immunohistochemistry

IHC was performed with the antibodies shown in Supplementary Table 2 according to the previous study.16 For quantification of the stained sections (five sections per brain), the number of positive (+) markers (for example, Ki67 and GFP+ cells) in the most posterior lobules (IX and X) were counted and used to calculate the ratio of positive marker (for example, Ki67+) to green fluorescence positive (GFP+) cells. Cell counting was carried out in a double-blind manner whenever applicable.

Immunoblotting

Mycoplasma-free HEK293T cells from American Type Culture Collection (Manassas, VA, USA) were transfected with pT2K-empty (negative control), pT2K Myc-ires-luc (positive control), and pT2K LRL-Myc-ires-Luc with or without MSCV-Cre plasmids and harvested 2 days after transfection. Protein lysates were prepared from the transfected HEK293T cells using standard RIPA buffer (Sigma) and used 10 μg of the lysates for immunoblotting.

Neurosphere culture and differentiation assay

Cerebellar cells from postnatal (P) day P7 Prom1-CreER; R26-LSL-EYFP mice treated with tamoxifen at P0 and P1 were purified as published previously,16 followed by fluorescence-activated cell sorting sorting of EYFP expressing cells. Sorted cells were plated in serum-free Neurobasal media supplemented with B27 and N2 (Invitrogen, Carlsbad, CA, USA) (passage 0). Basic fibroblast growth factor (50 ng/ml); Peprotech, Rocky Hill, NJ, USA) and epidermal growth factor (EGF) (50 ng/ml; Peprotech) were added every 3 days and neurospheres were passaged every 7–10 days. After the second passage, cells were dissociated with trypsin and subsequently plated on Matrigel-coated coverslips. For differentiation assays, basic fibroblast growth factor and EGF were removed and platelet-derived growth factor-AA (10 ng/ml; Sigma) was added for 2 weeks. Cultured cells were fixed with 4% paraformaldehyde/phosphate-buffered saline, followed by immunofluorescence.

MRI imaging

In patients, preoperative imaging was acquired at the local hospital where each patient had tumor resection before referral for protocol-based therapy at SJCRH, and analyzed, as reported previously.11 The research protocol was approved by the SJCRH Institutional Review Board and all patients gave written informed consent to participate. The preoperative imaging examination varied among patients, but always included T1-weighted scans before and after administration of a gadolinium-based contrast agent. The slice thickness was 4 or 5 mm and the in-plane resolution was 0.8 to 1 mm. T1-weighted images from all patients were wrapped, or spatially normalized,46 to the standard brain space defined by the Montreal Neurological Institute (MINI) brain template.48,49 The MINI space approximates the proportional stereotaxic brain space of Talairach and Tournoux50 which was extended to the cerebellum in the atlas by Schmahman et al.51 Spatial normalization was performed with Statistical Parametric Mapping software (SPMS; Wellcome Institute of Neurology, London, UK) using default parameters, except that the brain bounding box was extended from 50 to 80 mm in the negative z-direction for sagittal images and all output images were resliced to 1 mm x 1 mm x 1 mm resolution. The T1-weighted images without contrast agent were used for spatial normalization because the tumor contrast was low, which would minimize adverse effects of the tumor on the accuracy of spatial normalization. To characterize the accuracy of spatial normalization we defined a set of landmarks from the MINI space that were identified in the normalized images from each patient. The landmarks and their MINI coordinates (x, y, z) included the center of the anterior commissure (0, 5, −4), the cerebral aqueduct (0, −30, −10), a point on the posterior aspect of the brain stem in the fourth ventricle (0, −34, −26), the apex of the cerebellum in the midline (0, −54, 4) and lateral extremes of the cerebellum on the right (59, −58, −39) and left (−59, −58, −39).52 The locations of landmarks in the normalized images of each patient were measured with the 3D Slicer software (version 3.4, http://www.slicer.org).52

Cross-species analysis

To classify mouse models molecularly, all analyses were performed in the R environment (v.3.1.2) with CRAN and Bioconductor (v.3.0) packages, including GEOquery (v.2.32), frma (v.1.18), sva (v.3.12), randomForest (v.4.6) and ggplot2 (v.1.0). Public expression data of human MBs were collated from previous studies.40,44,45,53,54 Raw expression data of human MBs on the Affymetrix Human Genome U133 Plus 2.0 array platform (Affymetrix) were retrieved from GEO accessions GSE10327, GSE12992, GSE19404, GSE35493 and GSE37418. This data set was integrated with an expression data set (on the Affymetrix Mouse 430 2.0 array platform) of previously published mouse models of MBG3, MB443 and MBG3 (accessions GSE33199 and GSE24628)11,16 in addition to expression data of mouse models generated in the current study. The raw data was normalized using FRMA,55 transformed into z-scores using Barcode,56 adjusted for center-specific biases and cross-species differences using CombBat, and converted into expression barcodes (non-expressed vs expressed) with a logarithm (base 10) of odds score threshold of 12. The human samples and the mouse samples of previously published models were used to train a random forest classifier.56 The expression barcodes of the data sets were visualized by principal component analysis. The integrated data set (expression barcodes) of human MBs and previously published mouse MB models was used to train a random forest classifier.57

For computational efficiency, genes determined to be non-expressed across all samples were removed before training. The trained classifier was subsequently used to predict the molecular subgroups (WNT, SHH, G3, G4) of each mouse tumor sample in the present study by its expression barcode. For each sample, the distribution of votes across the classes was interpreted as the posterior probability distribution of the sample belonging to each class. The posterior probability of a mouse model belonging to class, given the expression data of samples of the model, was calculated as

\[
p(c|x) = \frac{\prod_{i} p(c|x_i)}{\sum_{i} \prod_{i} p(d|x_i)}
\]

which follows from Bayes' rule and the assumption that samples of the model are conditionally independent.

**ACCESSION NUMBER**

All microarray data has been submitted to the US National Institute of Health GEO database under the accession numbers GSE65888 and GSE33199.
CONFLICT OF INTEREST
The authors declare no conflict of interest.

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**Supplementary Information** accompanies this paper on the Oncogene website (http://www.nature.com/onc)