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Failure of human rhombic lip differentiation underlies medulloblastoma formation

A full list of authors and affiliations appears at the end of the article.

Abstract

Medulloblastoma (MB) comprises a group of heterogeneous paediatric embryonal neoplasms of the hindbrain with strong links to early development of the hindbrain 1-4. Mutations that activate Sonic hedgehog signalling lead to Sonic hedgehog MB in the upper rhombic lip (RL) granule cell lineage 5-8. By contrast, mutations that activate WNT signalling lead to WNT MB in the lower RL 9,10. However, little is known about the more commonly occurring group 4 (G4) MB, which is thought to arise in the unipolar brush cell lineage 3,4. Here we demonstrate that somatic mutations that cause G4 MB converge on the core binding factor alpha (CBFA) complex and mutually exclusive alterations that affect *CBFA2T2*, *CBFA2T3*, *PRDM6*, *UTX* and *OTX2*. *CBFA2T2* is expressed early in the progenitor cells of the cerebellar RL subventricular zone in *Homo sapiens*, and G4 MB transcriptionally resembles these progenitors but are stalled in developmental

Corresponding Author: Correspondence and request for materials should be addressed to T.E.W.-O., K.J.M., or M.D.T. Human material provided by the Joint MRC/Wellcome (MR/R006237/1) Human Developmental Biology Resource (www.hdbr.org), and the Birth Defects Research Laboratory (NIH-R24-HD000836 to IAG) was covered by a material transfer agreement between SCRI and HDBR/BDRL but samples may be requested directly from the HDBR/BDRL. Michael D. Taylor, The Hospital for Sick Children, 555 University Avenue, M5G1X8, Ontario Canada, Ph: 416-813-6427, mdt.cns@gmail.com.

L.D.H., P. Haldipur and O. Saulnier designed, performed and analysed the majority of the experiments in this study. J.M., A.H.S. and J.G. contributed to in situ hybridization and cell count analyses. A.W.E. contributed to bioinformatics analysis and SCENIC workflow. W.O. and D.S.S. contributed to cell line preparation for single-nucleus sequencing. W.O. and M.C.V. contributed to primary tissue isolation, single-cell preparation and sequencing. V.G. and L.C.-M. contributed to TurboID experiments and OTX2-KD experiments. A.L.M. contributed to GCP isolation and design and overexpression of the DN-CBFA2T3 construct. M.S. and T.P.-B. contributed to fusion CBFA2T2-TurboID construct design. M.L., R.A.S. and D. S. Scott contributed to CBFA2T2 mutation validation. S.B. contributed to western blot validations. H.F. contributed to SNP6 copy number analyses. T.N. and S.N. contributed to mutation analysis. A.B. contributed to pathway analysis. J.J.D.-M. contributed to CNV analysis of the scRNA-seq data. K.B. contributed to analysis and model design. P.S. and E.Y.W. contributed to RNA-seq fusion calling and analysis. O. Sirbu, S.A.K., P.B., A.V., J.J.Y.L., R.A., Xin Chen, Xiaodi Chen and K.L.M. contributed to experimental, statistical, bioinformatics analysis and/or interpretation. B. Luu, P. Bérubé, Y.C.W., A.S.M. and X.W. contributed to RNA-seq library preparation. S.M.P., N.J., S.-K.K., F.D., O.D., F.B., J.M.-P., W.A.G., J.L., P.D., M.F.-M., A.J., P.J.F., J.M.K., K.Z., S.D.B., C.G.E., A.A.N.R., C.G., J.M.O., M.G., P. Hauser, J.J.P., Y.S.R., C.d.T., J. Mora, K.K.W.L., H.-K.N., W.S.P., I.F.P., E.L.-A., G.Y.G., T.E.V.M., T.S., R.V., R.C.T., M.K.C., J.B.R., T.K., S.J., B. Lach, A.I., V.F., P.d.A., M.Z., G.C., S.R., D. S. Stearn, E.G.V.M., P.P., G.F., M.M., C.G.C., C.C.F., M.F.R., F. Boop, J.A.C., R.E.M., E.M.T., M.A., M.L.G., F.C., P.E. and M.P. provided patient tumour material and helped design the study. K.A.A., A.S.M., F.M.G.C., J.N.R., S.J.M.J., R.A.M., M.A.M., X.H., J.R., P.H.S., R.J.W.-R., W.A.W., T.J.P., L.G., C.L.K., L.D.S., N.J., D.M. and O.A. helped analyse data and provided expert advice. F.R. and E.S. provided late-gestation human samples. X.W. and C.D. provided reagents, equipment and expert advice. D.W.E. and J.A.G. provided expert neuropathology analysis and images. T.E.W.-O. and B.D. contributed to and provided expert advice on TurboID and OTX2-KD experiments. V.R. provided MRI images and provided expert advice on their interpretation. H.S., T.E.W.-O., K.J.M. and M.D.T. jointly supervised the project. L.D.H., P.H. and M.D.T. prepared the figures and wrote the manuscript.

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ADDITIONAL INFORMATION

Extended data is available for this paper.

Supplementary Information is available for this paper.

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These authors contributed equally to this work

^{*}These authors jointly supervised this work

AUTHOR CONTRIBUTIONS

time. Knockdown of *OTX2* in model systems relieves this differentiation blockade, which allows MB cells to spontaneously proceed along normal developmental differentiation trajectories. The specific nature of the split human RL, which is destined to generate most of the neurons in the human brain, and its high level of susceptible EOMES⁺ KI67⁺ unipolar brush cell progenitor cells probably predisposes our species to the development of G4 MB.

Keywords

Medulloblastoma; group 3; group 4; cerebellum; rhombic lip; CBFA2T2; CBFA2T3; OTX2

INTRODUCTION

MB comprises a group of malignant paediatric cerebellar embryonal neoplasms with extensive intertumoural and intratumoural heterogeneity^{1,2}. Mutations in genes in the Sonic hedgehog (SHH) signalling pathway lead to SHH MB in the granule cell lineage^{5–8}. Meanwhile, mutational activation of WNT signalling leads to WNT MB in the lower RL^{9,10}. Although group 4 MB (G4 MB, 40% of patients) and group 3 MB (G3 MB, 20% of patients) constitute the majority of MB cases, and the most deaths, far less is known about their specific cellular origin or somatic driver events. G4 MB is proposed to arise in the unipolar brush cell (UBC) lineage, whereas G3 MB appears to arise from an earlier population of stem cells^{3,4}. Activation of MYC is seen in many G3 tumours; however, the convergence and/or significance of various low-incidence driver mutations in G4 MB remains unclear¹¹.

The developing *H. sapiens* RL displays specific features compared to other mammals such as mice and macaques¹². At around 11 post-conception weeks (PCW), the human RL splits into two molecularly and structurally distinct zones—the ventricular RL (RL^{VZ}) and the subventricular RL (RL^{SVZ})—that are separated by a vascular plexus¹². The RL^{VZ} and RL^{SVZ} are transcriptionally distinct¹³, with the RL^{VZ} primarily composed of stem cells and the RL^{SVZ} primarily composed of proliferative progenitor cells. Expression of the classical UBC marker gene *EOMES* (also known as TBR2) in the RL^{SVZ} implies human-specific aspects of UBC development¹⁴. The human-specific features of RL development necessitate analyses of the developing H. *sapiens* cerebellum to deter- mine the true cellular origins of G4 MB.

We report new G4 MB mutations in genes that encode CBFA family proteins, including *RUNX1T1* (also known as *CBFA2T1*), *CBFA2T2* and *CBFA2T3*. These adaptor proteins form a large complex that recruits epigenetic modifiers and transcription factors (TFs) to chromatin^{15–19}. Apical G4 MB tumour cells resemble a specific, human-expanded, EOMES+KI67+ UBC progenitor cell population of the RL^{SVZ}, where members of the CBFA family are specifically expressed during human cerebellar development. We propose that the CBFA complex potentiates normal differentiation of EOMES+KI67+ RL^{SVZ} progenitors, the failure of which results in G4 MB.

RESULTS

The mutational landscape of G3 and G4 MB

We transcriptionally profiled G3 and G4 MB samples (n = 545) using bulk RNA sequencing (RNA-seq)^{2,11,20} (Fig. 1a). Recurrently mutated genes converged on epigenetic modifiers, the cell cycle and four gene families: ELP^{21} , $FANC^{11}$, CHD^{22} and members of the CBFA polyprotein complex. Genes from these four gene families are proximally clustered on human chromosomes, particularly within regions affected by copy number aberrations (CNAs) in G4 MB (Extended Data Fig. 1a–e). The combined deletion of multiple physically proximate drivers suggested a reason for specific chromosome arm deletions in G4 MB (Extended Data Fig. 2a–c). We identified new G4 MB alterations that targeted the CBFA complex, including the histone methyltransferase $PRDMo^{16}$, the histone demethylases KDM6A and KDM2B and the transcriptional corepressors RUNX1T1, CBFA2T2 and CBFA2T3. This is consistent with a model in which CBFA driver mutations result in the failed differentiation of G4-MB-initiating progenitor cells, which accumulate to form G4 MB, an embryological remnant.

CBFA2T2 and CBFA2T3 alterations are enriched in G4 MB

G4-MB-specific loss-of-function mutations that target *CBFA2T2* (Fig. 1b) were enriched in the NHR1 domain (Fig. 1c), which interacts with the SET and PR domains of PRDM proteins, including *PRDM6*¹⁶. *CBFA2T2* mutations tended to occur independently of high *PRDM6* expression, which indicated that there is a complementation group (Extended Data Fig. 3a). To uncover other members of this complementation group, we compared CNAs between G4 MB samples with *CBFA2T2* or *PRDM6* alterations, or neither (Fig. 1d). Focal chromosome 16q24 (*CBFA2T3*) deletions occurred in a mutually exclusive pattern with *CBFA2T2* and *PRDM6* mutations (Fig. 1d and Extended Data Fig. 3b). They also drove significantly reduced *CBFA2T3* expression (Extended Data Fig. 3c) and universally spanned *CBFA2T3* (Extended Data Fig. 3d–g). Although 16q deletions are detected in both G3 and G4 MB, they are rarely seen in SHH MB^{20,23}. Indeed, both *CBFA2T2* and *CBFA2T3* showed significantly higher expression in SHH MB, which indicates that the role of the CBFA complex may differ between the MB cells of origin (Extended Data Fig. 3h).

Chromosome 16q contains 3 additional G4 MB recurrently mutated genes (*FANCA*, *ZFHX3* and *CHD9*), and mutations in these genes were mutually exclusive with 16q deletions (Fig. 2a). Tight genomic clustering of these G4 MB driver genes along human chromosome 16 explains the bias towards large deletions of *CBFA2T3* and the relative absence of somatic single-nucleotide variants (SNVs) in *CBFA2T3* (Fig. 2b). *CBFA2T2* and *CBFA2T3* interact with *GFI1B* and other epigenetic modifiers ^{19,24}, and G4 MB with *GFI1B* enhancer hijacking events were mutually exclusive (Fig. 2c). Taken together, we propose that *CBFA2T2*, *PRDM6*, *CBFA2T3*, *GFI1* and *GFI1B* represent a new G4 MB complementation group.

CBFA complex disruptions underly G4 MB

As *CBFA2T2*, *PRDM6* and *CBFA2T3* form a complementation group, and are known to physically interact^{16,19}, we propose that the polyprotein CBFA complex contains additional G4 MB drivers. We performed in vitro TurboID²⁵ in the G3 MB cell line HDMB03

(Extended Data Fig. 4a,b), in which *MYC* is amplified. *CBFA2T2* interacted with *KDM6A*, a known G4 MB driver gene (Extended Data Fig. 4c,d). Other notable interactors included *RUNX1T1*, *KDM2B* and *SMARCA1*. *CBFA2T2* also interacted with *GLI2*, a recurrently amplified SHH MB oncogene. We combined our new *CBFA2T2* protein interactions with known interactions between G3 and G4 MB driver genes (Extended Data Fig. 4e). Most G4 MB driver genes were within two steps of *CBFA2T2*, which suggests that they might be part of the CBFA complex. Alterations that disrupted the CBFA complex were found in at least 57% of G4 MB samples and 39% of G3 MB samples (Fig. 2d and Extended Data Fig. 5a).

A subgroup-specific analysis of microglial expression differences demonstrated stark variances in subgroup-specific microenvironments (Fig. 2e). G4 MB microglia, but not other subgroups, expressed the ERBB4 ligands *HBEGF* and *EREG* (Fig. 2f). Tumours that expressed high levels of ERBB4 ligands were less likely to have mutations in the CBFA complex (Extended Data Fig. 5b). These data provide a possible explanation for activation of ERBB4 in G4 MB, as previously demonstrated through unbiased proteomics²⁶. As ERBB4 activation has been suggested to inhibit the activity of CBFA2T3²⁷, it appears that a subset of G4 MB tumours receive important and sustaining signals from their microenvironment that might be targetable for intervention. Collectively, we propose a model in which driver genes of G3 and G4 MB converge to inhibit the physiological CBFA complex through various mechanisms (Extended Data Fig. 5c).

CBFA2T2 demarcates the human RLSVZ

The role of the CBFA complex in cerebellar development is unknown. In humans, the RL is long-lived and seen throughout gestation and it undergoes a series of morphological and structural changes (Fig. 3a and Extended Data Figs. 5d and 6a). Notably, the human RL develops a substructure and expands into two distinct zones at 11 PCW: the RL^{VZ} and the RL^{SVZ} (Fig. 3a and Extended Data Fig. 5d). These zones are separated by a vascular plexus, which expresses *HBEGF* (Extended Data Fig. 6b,c). The RL^{VZ} was enriched for SOX2, whereas the RL^{SVZ} demonstrated strong cell division typical of progenitor cells (Fig. 3b). The developing RL became decreasingly proliferative with time (Fig. 3b,c and Extended Data Fig. 6d) until it disappeared around birth (Extended Data Fig. 6e).

CBFA2T2 was specifically expressed in the RL^{SVZ} starting at 11 PCW (Fig. 3d). At 14 and 17 PCW, *CBFA2T2* and *CBFA2T3* remain expressed in the RL^{SVZ} but not the RL^{VZ} (Fig. 3d,e and Extended Data Fig. 6f,g). At each timepoint, *CBFA2T2* and *CBFA2T3* were also expressed in the external granule layer (EGL), where *Cbfa2t2* and *Cbfa2t3* expression is observed in mice (Extended Data Fig. 6h,i). The EGL is composed of granule cell precursors (GCPs), the cell of origin for SHH MB. By contrast, the expression of both genes decreased along the ventral exit from the RL, where differentiating *LMX1A*+ UBCs migrated away from the proliferative RL (Extended Data Fig. 6j). *CBFA2T2* and *CBFA2T3* expression was retained by GCPs (and SHH MB), but not UBCs, which suggests that these genes have a role in this cell fate decision during normal development.

G3 and G4 MB mirror embryonic human RL

CBFA2T2 expression in RL^{SVZ} progenitor cells indicates that G4 MB might arise from the RL^{SVZ} because of dysfunctional *CBFA2T2*. An analysis of single-nucleus RNA-seq (snRNA-seq) data from the developing human cerebellum¹³ demonstrated the expected cell types and lineages descending from the apical RL^{VZ} (Fig. 4a and Extended Data Fig. 7a,b). *CBFA2T2* and *CBFA2T3* were highly expressed in the RL^{SVZ} and in GCP clusters (Extended Data Fig. 7a,c).

We performed single-cell RNA-seq (scRNA-seq) and integrative clustering on G3 MB (five patient samples), G4 MB (11 patient samples) and SHH MB (three patient samples) (Extended Data Fig. 7d). Non-tumour cells were identified through the expression of known marker genes and a paucity of CNA detection and were excluded from further analysis (Extended Data Fig. 7e-g). Using two distinct methods, a transcriptional comparison between the developing human cerebellum and MB cells consistently revealed that apical G3 and G4 MB cells were most similar to the RLSVZ (Fig. 4b and Extended Data Fig. 8a.b), whereas the more deadly G3 gamma subtype $(G3\gamma)^2$ displayed enrichment for the earlier RLVZ (Fig. 4c and Extended Data Fig. 8c). SHH, G3 and G4 MB all displayed a differentiation block, with few cells mapping to late granule neurons (GNs) and late UBCs. G3 MB displayed the lowest similarity to normal cerebellar cells, overall (Extended Data Fig. 8d). Mutations in the CBFA complex were highly enriched in the OTX2-depleted subtypes of G4 MB (G4 α and G4 β), but not commonly found in G4 γ , in which OTX2 levels were high and KBTBD4 mutations were found (Fig. 4d and Extended Data Fig. 8e). Whether the transcriptional differences between $G4\alpha$, $G4\beta$ and $G4\gamma$ are due to them arising from discrete developmental cell states, compared to the effects of specific somatic mutations, will require additional consideration. BARHL1 and DDX31, for which enhancers are known to drive aberrant GFI1B expression through enhancer hijacking²⁸, were expressed only in the RLSVZ, whereas *GFI1B* itself was not expressed (Extended Data Fig. 8f-k). These data are consistent with a model in which G4 MB and some G3 MB tumours arise in the RL^{SVZ} owing to the specific human RL split.

Human-evolved predisposition to G3 or G4 MB

The RL^{SVZ} is composed of two distinct populations: nascent GCPs, which express PAX6, but not EOMES; and nascent UBCs, which express both PAX6 and EOMES (Fig. 4e and Extended Data Fig. 9a). At around 11 PCW, PAX6⁺EOMES⁻ cells form a stream of cells between pockets of EOMES⁺ cells, which connect the tail-like RL with the EGL. However, this conspicuous subcompartmentalization was short-lived and no longer visible following RL internalization at 14 PCW. The frequency of PAX6⁺EOMES⁻ cells decreased after 11 PCW, which suggests that the early tail-like RL produces more GCPs relative to later stages (Fig. 4f). We propose that GCPs produced by the tail-like RL build the nascent EGL of all three cardinal lobes, and following internalization at 14 PCW, the RL seeds only the posterior lobe^{12,29,30}. Consistently, the RL^{SVZ} shifts towards UBC production, as shown by an increase in EOMES⁺ cell numbers after 14 PCW (Fig. 4g).

EOMES is currently thought of as a marker of post-mitotic UBCs^{31,32}, and previous comparisons of G4 MB to the developing mouse cerebellum suggested that G4 MB arises

in post-mitotic EOMES+ UBCs^{3,4}. However, in humans, EOMES⁺ cells in the RL^{SVZ} are predominantly proliferative, as shown by labelling for both EOMES and KI67 (Fig. 4h and Extended Data Fig. 9b,c). Furthermore, we identified the following additional features in the RL specific to humans compared to mice: (1) further compartmentalization of the RL^{SVZ} into an inner SVZ that contained EOMES⁺KI67⁺ cells and an outer SVZ that contained differentiated nascent or early UBCs (Fig. 4h, top right); (2) temporal expansion of the EOMES⁺KI67⁺ population late into gestation (Fig. 4h, bottom left, and Extended Data Fig. 9d); and (3) significant human-specific expansion of EOMES⁺KI67⁺ progenitors compared to mice (Fig. 4i and Extended Data Fig. 9e–h). Thus, there is a spatiotemporally expanded pool of MB-susceptible EOMES⁺KI67⁺ UBC progenitors throughout human gestation in comparison to mice, thereby providing a statistically larger risk for G3 and G4 MB to arise (Extended Data Fig. 9i).

OTX2 inhibits CBFA2T2 to retain RL state

We computationally inferred the activity of TFs in both the developing human cerebellum and our scRNA-seq MB samples (Fig. 5a). TFs highly active in G3 and G4 MB were also highly active in the RLSVZ. OTX2 activity was particularly high in G3 and G4 MB and the RL. OTX2 is frequently amplified in both G3 and G4 MB^{33,34} (Fig. 1a). OTX2 was highly expressed in the RLVZ and the RLSVZ, and preceded CBFA2T2 and CBFA2T3 expression (Fig. 5b and Extended Data Fig. 10a,b). OTX2 was also specifically expressed in the posterior cerebellar lobes (Fig. 1c). Consistent with our proposed cell of origin, G3 and G4 MB—but not SHH or WNT MB— always presented on the inferior surface of the cerebellum (Fig. 1d and Extended Data Fig. 10c-f). At 14 PCW, the RL was internalized into the nascent nodulus of the cerebellar flocculonodular lobe (Extended Data Fig. 6a). The location of the embryological remnant of the internalized RL explains the near universal localization of G3 and G4 MB in the inferior midline of the cerebellum (Fig. 1d). OTX2 was highly expressed in G3 MB and G4γ, whereas CBFA2T2, CBFA2T3, KDM6A, PRDM6 and other alterations to the CBFA complex tended to occur in OTX2^{low} G4a and G4B MB (Fig. 4d). Indeed, OTX2 expression was significantly reduced in G3 and G4 MB with alterations in the CBFA complex (Fig. 1e). This result suggests that either OTX2 overexpression or CBFA complex dysfunction can impair RLSVZ differentiation.

OTX2 bound the *CBFA2T2* but not the *CBFA2T3* promoter, suggesting transcriptional control of the CBFA complex (Extended Data Fig. 11a). We performed siRNA-mediated *OTX2* knockdown (OTX2-KD) on tumourspheres from two G3 MB lines harbouring *MYC* amplifications, which were profiled by bulk and snRNA-seq (Extended Data Fig. 11b–d and Supplementary Fig. 1). Expression of both *CBFA2T2* and *CBFA2T3* were significantly upregulated following OTX2-KD (Fig. 1f). Analysis of single OTX2-KD cells revealed a large increase in similarity to more differentiated cell types, which was dominated by GCPs and GNs (Fig. 1g). Pseudotime mapping of OTX2-KD cells revealed a sequential differentiation trajectory that mirrored normal GN development (Fig. 1h, Extended Data Fig. 11e–j and Supplementary Fig. 1). *CBFA2T2* and *CBFA2T3* were among the first genes to show increased expression following OTX2-KD. They were also highly expressed in GCP-like cells and preceded GN-like cells. This result provides further evidence for a role of *OTX2* in RL^{SVZ} fate decisions, GN differentiation or both (Fig. 1h and Extended

Data Fig. 11k). Previous studies have demonstrated that injection of significantly more OTX2-KD cells was required to generate G3 MB tumours *in vivo*, which may reflect the depletion of tumour-initiating cells through induced GN differentiation³³. Overexpression of CBFA2T2 significantly reduced the number of live cells in tumoursphere culture, without affecting viability. This result provides evidence for a role for CBFA2T2 in the promotion of differentiation in G3 and G4 MB (Extended Data Fig. 11l–p and Supplementary Fig. 1).

Collectively, these data support a model in which *OTX2* maintains RL identity by inhibiting the CBFA complex until cells exit the RL and differentiate. Disruptions to the CBFA complex, or overexpression of *OTX2*, according to this model, result in failed RL^{SVZ} differentiation. The resulting ball of RL^{SVZ} progenitor cells are retained with the RL in the nascent nodulus of the developing cerebellum, where ongoing mitotic activity eventually results in a mass lesion diagnosed as G3 or G4 MB (Extended Data Fig. 11q).

DISCUSSION

The RL gives rise to most of the neurons in the human brain, which provides a proximate explanation for why MB (cancer of the RL) is the most common embryonal neoplasm in humans and the most common malignant paediatric brain tumour. The physical splitting, interposed vascular plexus and dramatic expansion of the human RL, compared to that of mice and macaques, suggests that *H. sapiens* may be specifically and highly predisposed to the development of G4 MB. The teleological reasons and evolutionary benefits of the expansion of the human cerebellum are not immediately evident, although a predisposition to RL cancer appears to be a possible cost. Notably, the presence of atypical cell rests in the postnatal nodulus have been recognized since the 1940s and were already speculated to be a precursory lesion to MB on the basis of histological similarity^{35,36}. However, without knowledge of the MB subgroups, the inconsistent anatomical presentation of MB in both the cerebellar hemispheres and the midline probably led to these hypotheses being forgotten. Further work in the 1960s, principally by Lucy Rorke, found an association between the presence of cerebellar heterotopias and various chromosomal trisomies, including trisomy of commonly gained chromosomes in G3 and G4 MB such as trisomy 17^{37,38}. We propose that these heterotopias in the nodulus are more aptly named persistent RL (PeRL), as they are probably neither permanent nor ectopic. Our findings add support to the 80-year-old hypothesis that PeRLs may represent a premalignant lesion for G3 and G4 MB, although presumably most PeRLs spontaneously regress in the absence of further genetic insult. Cancers arising from different anatomical regions of the RL, and different points in developmental time, account for much of the heterogeneity seen between MB subgroups (Fig. 6). Future studies using modern techniques will be necessary to further examine the probable link between PeRLs and MB.

The clustering of G4 MB driver genes along human chromosomes suggests a reason for the predilection of CNAs over SNVs in G4 MB. It also explains the specific patterns of genomic gain and loss and emphasizes that G4 MB is a disease of the human genome. G4 MB cells appear to be stalled in developmental time, with release of the developmental blockade resulting in the resumption of pseudo-normal developmental differentiation cascades. MB is a cancer of the human RL, with G4 MB probably arising from the embryological remnant

of the RL^{SVZ} (that is, PeRLs) secondary to *OTX2* overactivity or CBFA complex failure. We propose that the detection of a PeRL in the postnatal period, through either imaging or biochemical detection of oncofetal antigens in the serum, could enable closer monitoring and potential early intervention to prevent the emergence of full-blown G3 or G4 MB.

METHODS

DATA REPORTING

No statistical methods were used to predetermine sample size. The experiments were not randomized. Investigators were not blinded to allocation during experiments and outcome assessment, except in the case of performing IHC-based cell counts where the investigator performing the count was blinded to experimental conditions such as sample age and antibodies used.

ACQUISITION OF PATIENT SAMPLES AND HUMAN TISSUE

Samples used to generate the bulk RNAseq cohort were obtained from the Medulloblastoma Advanced Genomics International Consortium (MAGIC). Samples used to generate the single-cell RNA sequencing cohort were obtained in compliance with the ethical regulations of the Hospital for Sick Children and McGill University Health Centre. All patient material was collected after receiving informed consent, under approval and oversight by their respective internal review boards. Relevant clinical metadata and overlapping data types for the 819 primary medulloblastoma samples used in this study are presented in Supplementary Data Table 2.

All human cerebellar samples used in this study were obtained using protocols approved by the Seattle Children's Research Institute IRB. Samples used for histological analysis were collected by the Human Developmental Biology Resource (HDBR) located at University College London, and Newcastle University, United Kingdom and Birth Defects Research Laboratory (BDRL) at the University of Washington. Third trimester samples were part of an archival collection at the Hôpital Necker-Enfants Malades in Paris, France. All samples were collected with previous patient consent and in strict accordance with institutional and legal ethical guidelines. Formalin-fixed-paraffin-embedded (FFPE) sections of the cerebellum were collected at 4 microns thickness along the sagittal plane and mounted on Superfrost plus slides (Thermofisher Scientific). Slides were refrigerated to preserve antigenicity and to prevent RNA degradation.

ANIMALS

Embryonic mouse tissue was collected in accordance with the guidelines laid down by the Institutional Animal Care and Use Committee (IACUC) of Seattle Children's Research Institute, Seattle, WA, USA, and to the National Institutes of Health guidelines on the care and use of laboratory animals and are in accordance with the applicable portions of the Animal Welfare Act. Mice were maintained on a corn cob bedding with a 10/14 dark/light cycle, ambient temperature and humidity were monitored and maintained within the recommended ranges by the Guide (Guide for the Care and Use of Laboratory Animals, 8th edition). CD1 mice were crossed, and the day of plug was taken as embryonic day (E)

0.5. Embryos were dissected out between E14.5 and E16.5, fixed in 4% paraformaldehyde (PFA) for 1-2 hours, washed in PBS and transferred to a solution of 30% sucrose overnight. Samples were then embedded in optimum cutting temperature (OCT) matrix and midsagittal cryosections of 11 μ m thickness were collected.

CELL LINES

Two group 3 medulloblastoma cell lines were used in this study, HDMB03 and MB3W1. HDMB03 was kindly provided by Dr. Till Milde (described in ref. ⁴⁶) MB3W1 was kindly provided by Dr. Matthias Wölfl (described in ref. ⁴⁷).

HISTOLOGY

FFPE sections were de-paraffinized in xylene and rehydrated in a gradient of ethanol prior to processing. Hematoxylin and Eosin (H&E) staining were carried out as previously described³⁰.

IMMUNOHISTOCHEMISTRY (IHC)

IHC was performed as described previously¹². Primary antibodies used in the study were as follows: KI67 (Agilent, M7240, mouse, 1:50; Thermofisher, MA5–14520, rabbit, 1:100), SOX2 (Thermofisher, PA1–094, Rabbit, 1:200), PAX6 (Biolegend, 901301, rabbit, 1:300), TBR2 (*EOMES*) (Thermofisher, 14–4875-82, Rat, 1:200), GFAP (Agilent, Z0334, rabbit, 1:1000). Fluorescent dye-labelled secondary antibodies from Thermofisher were used at a dilution of 1:1000 (anti-rabbit: Alexa Fluor 488, A-11034, Goat; Alexa Fluor 568, A-11011, Goat; anti-mouse: Alexa Fluor 488, A-11001, Goat; Alexa Fluor 568, A-11004, Goat). Following secondary antibody incubation, sections were counterstained with DAPI (4',6-diamidino-2-phenylindole) using Vectashield mounting medium (Vector laboratories, H-1200).

IN SITU HYBRIDIZATION (ISH)

ISH assays were run using commercially available probes from Advanced Cell Diagnostics. Manufacturer-recommended protocols were used without modification. The following probes were used in the study LMX1A (#540661), OTX2 (#484581), CBFA2T2 (#410331), CBFA2T3 (#406001), MKI67 (#591771), HBEGF (#431651), Cbfa2t2 (#491601), Cbfa2t3 (#434601), Hbegf (#437601). All sections were counterstained with hematoxylin or methyl green.

MICROSCOPY

All slides from fluorescent immunohistochemical assays were imaged using the Zeiss LSM-Meta confocal microscope and ZEN 2009 software (Zeiss). Nanozoomer Digital Pathology slide scanner (Hamamatsu; Bridgewater, New Jersey) was used for brightfield microscopy. Barring minor adjustments of contrast and brightness no additional image alteration was performed. Figures were prepared on Adobe Illustrator.

CELL COUNTS

Cell counts were performed manually using Image J. For all counts a minimum of 3 sections over 3 samples were used. For KI67, PAX6 and TBR2 (*EOMES*) counts, the total number of KI67 or TBR2 (*EOMES*) or PAX6+ cells relative to the total DAPI count in the RL was determined.

CBFA2T2 TURBO-ID

3xHA-TurboID-NLS_pCDNA3 was a gift from Dr. Alice Ting (Addgene plasmid # 107171). pCMV6-AC-CBFA2T2-GFP was purchased from Origene (RG202013). PCR-amplified TurboID and CBFA2T2 coding sequences were introduced into the KpnI/ClaI-linearized pCAG-H2B-mAG-P2A vector by using NEBuilder HiFi assembly (New England Biolabs). Non-vector sequences in resultant plasmids pCAG-H2B-mAG-P2A-3xHA-TurboID-CBFA2T2 and pCAG-H2B-mAG-P2A-3xHA-TurboID were verified by Sanger sequencing. For TurboID experiments, 3×10^6 HDMB03 cells were seeded equally across a 6-well ultra-low attachment plate and transfected with 12 μ g of plasmid using Lipofectamine 3000 (Thermo Fisher) according to manufacturer's guidelines. Cells were grown in suspension for 72 hours in StemPro NSC Serum-Free Medium (Life Technologies). After 72 hours of incubation, biotin-D (Sigma, B4639) was added to all wells to a final concentration of 500 μ M and the cells incubated for 15 min. Tumorspheres were then collected, washed in 1xPBS, snap-frozen and stored at -80° C until all runs were completed.

Replicate samples were thawed on ice and lysed with 1 ml of 1 x RIPA [20 mM Tris-HCl, pH 7.5; 150 mM NaCl; 1 mM EDTA-Na2; 1 mM EGTA; 1% NP-40; 1% sodium deoxycholate; 2.5 mM sodium pyrophosphate; 1 mM β-glycerophosphate; 1 mM Na3VO4, 1X HaltTM Protease/Phosphatase inhibitor (Thermo ScientificTM)] for 10 minutes. Samples were then sonicated over 2 × 10s pulses at 30% load to shear DNA (Fisher Scientific Sonicator FB50 with microprobe). Lysed samples were clarified by centrifugation at 16 000 x g for 10 minutes at 4°C. Each supernatant was quantified using the Bradford protein assay. Following quantification, 50 µl aliquots of streptavidin-sepharose bead slurry for each sample was washed once in 1 ml of 1X RIPA. Washed bead aliquots were then resuspended in 1 ml of 1X RIPA containing 2 mg of quantified protein supernatant and rotated overnight at 4°C. Samples were then centrifuged at 1000 x g for 5 minutes, and bead pellets were washed twice in 1 ml 2% SDS solution (in ddH2O) followed by three times in 1 ml Wash Buffer (50 mM Tris-HCl pH 7.4, 8 M Urea), rotating for 8 minutes at room temperature each time. Samples were then resuspended in Storage Buffer (285 µl of ammonium bicarbonate (50 mM) and 15 µl of 1 mM biotin) to saturate streptavidin binding and to prevent peptide recapture during on-bead digestion. Samples were stored on ice for analysis.

MASS SPECTROMETRY

Mass spectrometric (MS) data were obtained with an Orbitrap Exploris 480 instrument (Thermo Fisher Scientific, Bremen, Germany). All MS raw files were processed with Proteome Discoverer (v2.20.388) and searched for tryptic peptides against the human Uniprot protein database (Dec, 2020) using SEQUEST with standard Orbitrap settings: up to 2 missed cleavages were permitted, with a parent and fragment mass tolerance of 0.02

Da and 15 ppm, respectively. A fixed modification of cysteine carbamidomethylation was applied, and variable modifications including N-terminal acetylation, deamidation (at N and Q), phosphorylation (at S, T and Y), oxidation (at M and W), ubiquitylation (at K), double oxidation (at M and W) and biotinylation (at K) were permitted. The results were filtered by 1% False discovery rates (FDRs) at both the protein and peptide levels. SAINTexpress (v1.0.0) was then used to calculate the probability of each potential proximal protein interaction from background contaminants using default parameters ^{48,49}. Three replicates were used for control and experimental samples. A full list of CBFA2T2 interacting proteins can be found in Supplementary Data Table 3.

OTX2 KNOCKDOWN

OTX2 knock down in HDMB03 and MB3W1 G3 MB cells (2×10^5 cells/well) was performed as previously described 33,34 . Briefly, OTX2 was silenced using 30 nM Silencer Select siRNAs 9931 or 9932 (Life Technologies). A non-silencing (scramble) served as the negative control. For bulk RNA sequencing, OTX2 was knocked down in three independent biological replicates for each cell line and silencing was confirmed by western blot (OTX2, Abcam, ab21990, rabbit, at 1:500, and β -actin, Sigma-Aldrich, A2228, mouse, at 1/1000 was used as a loading control) 72 hours following transfection. Total RNA was extracted from all samples using the Norgen RNA extraction kit (Norgen Biotek), and bulk RNA sequencing (RNAseq) was performed by StemCore laboratories at the Ottawa Hospital Research Institute (Ottawa, ON, Canada). For single-nucleus RNA sequencing (snRNAseq), the above was repeated but using 4.5×10^5 cells/well for MB3W1 and only one replicate was performed. Granule neuron differentiation was validated by western blot for RBFOX3 (NeuN) (Cell Signaling Technology, D4G4O, at 1:1000).

CBFA2T2 OVEREXPRESSION

HDMB03 Group 3 MB cells were plated at a density of 2×10^5 cells/well (in a 6-well format) 24 hours prior to infection. The following day, Precision LentiORF CBFA2T2 viral particles (OHS5898–219582524) or RFP control lentiviral particles (Horizon Discovery Biosciences Limited, Cambridge, UK) were added to the cells in OptiMEM (Life Technologies) at an MOI=1. Twenty-four hours after infection, OptiMEM was aspirated and replaced with EMEM/10% FBS. Positively transduced cells were stably selected in blasticidin starting at day 5. Increased CBFA2T2 expression was confirmed by qPCR and immunoblotting (CBFA2T2, A303–593A-M, Bethyl Laboratories at 1:500, with β-actin, Sigma-Aldrich, A2228, mouse, at 1/1000 used as a loading control).

Following stable selection and cell expansion, assessment of total cell number and viability was performed as previously described 34 . Briefly, HDMB03 RFP control and CBFA2T2 OE cells were dissociated and cultured as tumorspheres in 6-well ultra-low-attachment plates at a density of 2×10^5 cells/well (in triplicate) in StemPro NSC Serum-Free Medium (Life Technologies). The total number of cells was counted at day 5 and viability was assessed by Trypan Blue exclusion.

MAGNETIC RESONANCE IMAGING (MRI)

All MRI scans were performed on medulloblastoma patients from the Hospital of Sick Children, Toronto, Ontario, Canada. Tumors were subsequently subgrouped by methylation array.

SANGER SEQUENCING

CBFA2T2 mutations were validated using Sanger sequencing (data not shown). Briefly, nested PCR was performed using the following reagents; Master mix (2.5 μ L 10X PCR Buffer (Invitrogen, #52724), 1 μ L 50 mM MgCl2 (Invitrogen, #52723), 0.5 μ L 10mM dNTP (Bio Basic, DD0056), 0.5 μ L F primer, 0.5 μ L R primer, 2 μ L loading dye, 0.2 μ L Taq polymerase (Invitrogen, #100021278), 15.8 μ L ddH2O (Wisent, #809–115-CL) and 2 μ L DNA. Secondary PCR products were run on a 2% agarose gel + TAE buffer to confirm expected product size. For each mutation, the band was excised, cleaned (with Geneaid PCR Clean-up Kit, Geneaid, #DFC100), and Sanger sequenced. Additionally, we TA cloned each mutation using the TA Cloning kit (ThermoFisher, #450641), followed by Mini prep (Geneaid Presto Mini Plasmid kit, Geneaid, #PD100) and Sanger sequencing.

BULK RNA SEQUENCING

Sample processing, mRNA library construction, and sequencing—Samples were processed and sequenced as previously described²³.

Alignment—The hs37d5 reference genome FASTA (1000 Genomes Project Phase II) was appended to the C1_2 ERCC spike-in sequences used for C1 Fluidigm, as well as Caltech profile 3 spike-ins sequences by ENCODE. A STAR assembly was then built with this reference and GENCODE (v19) gene annotations using parameter `--sjdbOverhang 124`. RNA-seq library reads were then mapped with the built assembly using STAR (v2.5.1b) and parameters '--outFilterMultimapNmax 20 --alignSJoverhangMin 8 --alignMatesGapMax 200000 --alignIntronMax 200000 --alignSJDBoverhangMin 10 --alignSJstitchMismatchNmax 5 -1 5 5 --outSAMmultNmax 20 --twopassMode Basic'. For alignment of OTX2-KD group 3 cell lines, raw sequence data were processed with fastp v0.20.1⁵⁰ to perform automatic adapter trimming and read quality filtering, retaining reads with at least 60% of bases having Q 15, and no more than 5 'N' bases. Transcripts from scramble and OTX2-KD samples were quantified with salmon v.1.4.0⁵¹ against an index built from the GENCODE v35 reference assembly with inclusion of genomic decoy sequences.

G3 and G4 MB subtype identification—Subtypes were determined using Similarity Network Fusion (SNF) as previously described² using RNA sequencing expression data in place of microarray expression data²³. Briefly, the full expression and methylation matrices were input into the SNF function of the SNFtool R package (v2.3.0) with parameters 'K = 20, alpha = 0.5, T = 100'. Spectral clustering was then performed, and the clusters obtained at k = 6 corresponded to the 6 group 3 and 4 subtypes.

G3 and **G4** MB subtype differentially expressed genes—Differential expression analysis was performed using the DESeq2 R package (v1.28.1)⁴⁵, comparing each subtype

to the other five, and controlling for subgroup as a covariate. Only genes with an FDR adjusted p-value < 0.05 were considered in downstream analysis. Full lists of subtype specific differentially expressed genes can be found in Supplementary Data Table 4.

RNA-seq mutation analysis—RNA-seq mutation calls were performed as previously described^{23,52}. Briefly, GATK (v3.8.0)⁵³ was used to detect variants which were then annotated with ANNOVAR (Feb 2016)⁵⁴. SNPs present in the gnomAD database were discarded, and SNPs with frequencies greater than 0.01 in 1000 Genomes, dnSNP138, RADAR, Exome Aggregation Consortium database, NHLBI-ESP project, Kaviar Genomic Variant Database, Haplotype Reference Consortium database, Greater Middle East Variome, Brazilian Genomic Variants database, and from an inhouse SNP database (356 sequenced whole genomes) were discarded. Significantly mutated genes (q < 0.05) were identified using MutSigCV (v1.41)⁵⁵ and OncodriveFML (v2.3.0)⁵⁶. A full list of mutations detected in G3 and G4 MB can be found in Supplementary Data Table 5.

Fusion calling and filtering—Gene fusions were called and filtered as previously described²³. Briefly, three fusion calling algorithms were run to maximize sensitivity, namely; Star-fusion (v0.8.0)⁵⁷, InFusion (v0.7.3)⁵⁸, and Trans-Abyss (v2.0.0)⁵⁹. Only putative fusions called by at least 2 algorithms, not detected in normal, and occurring in known mutated genes were retained.

SNP 6.0 ARRAYS

GISTIC analysis and determination of copy number responsive genes—

Samples were pre-processed as previously described²³. Processed SNP6 segment files were then input into GISTIC 2.0 (v2.0.23)⁴⁰ and run with the following changes to default parameters: '-conf 0.90 -broad 1 -ta 0.25 -td 0.3 -js 10 -rx 0 -brlen 0.7 -armpeel 1 -gcm extreme -genegistic'. Significantly amplified and deleted regions were then extracted, manually inspected in IGV, and categorized into either broad or focal events depending on if the segment spanned more than 12 Mb or equal to and less than 12 Mb, respectively. To determine if genes falling in significantly amplified or deleted regions showed concomitant expression changes, gene expression was categorized (where applicable) by amplification, neutral, or loss and a Kruskal-Wallis test was performed for greater than 2 conditions, and a Mann-Whitney U test in the case of only two conditions. Significance was adjusted using post-hoc Dunn's test with Benjamini-Hochberg correction.

SINGLE-CELL RNA SEQUENCING ON MB TUMORS

Preparation of single-cell suspensions—Fresh patient tumors were collected at the time of surgical resection. Tumor tissue was mechanically dissociated followed by collagenase-based enzymatic dissociation as previously described³.

Single-cell RNA library preparation and sequencing—Single cell suspensions were assessed with a trypan blue count. We aimed to load 10,000–15,000 cells per sample using the Chromium Controller in combination with the Chromium Single Cell 3' V3 and V3.1 Gel Bead and Chip kits (10X Genomics). Individual cells were partitioned into gel beads-in-emulsion (GEMS), followed by reverse transcription of barcoded RNA and cDNA

amplification. Individual single cell libraries with indices and Illumina P5/P7 adapters were generated with the Chromium Single Cell 3' Library kit and Chromium Multiplex kit. The libraries were sequenced on an Illumina Novaseq6000 sequencer.

Alignment of raw reads—Gene level counts were obtained using the 10X CellRanger pipeline (v3.1.0)⁶⁰. Briefly, raw base call (BCL) files were demultiplexed into fastqs using the *mkfastq* function. Fastq files were then aligned to the reference human genome hg19 v3.0.0 (from 10X Genomics) using the *count* function to generate raw gene-barcode count matrices. Alignment quality control metrics for all single-cell and single-nucleus RNAseq samples can be found in Supplementary Data Table 6.

Quality control and normalization—Quality control was performed at an individual sample level prior to sample integration and normalization. Briefly, low-quality cells were determined and excluded from further analysis based on outlier mitochondrial content (indicative of cellular stress or damage) or gene counts, using the R package Seurat $(v4.0.2)^{61}$. Genes expressed in less than 10 cells were also removed. High quality cells from each tumor sample were merged and normalized together using SCTransform $(v0.3.2)^{62}$, with parameters; *variable.features.n* = 3,000 and regressing unwanted variance associated with mitochondrial content.

Clustering analysis and visualization—Single-cell clustering was performed by first using principal component analysis (PCA) to determine statistically significant principal components (PCs) which were then used to construct a uniform manifold approximation and projection (UMAP) embedding in Seurat (v4.0.2)⁶¹. Cell clusters were identified using Seurat's shared-nearest neighbor algorithm following modularity optimization using the Louvain algorithm with multilevel refinement.

Identification of cell types—Non-tumor cell types present in the tumor single cell samples were first identified by examining expression of known cell-type specific markers (Extended Data Fig. 7g). As expected, these clusters displayed the most overlap between different samples. To confirm the remaining sample-specific cell clusters were comprised of tumor cells, inferCNV (v1.4.0)⁶³ was employed using the non-tumor cells as a reference and using the parameters *HMM_type* = i6, *noise_filter* = 0.1, *cutoff* = 0.1, and *sd_amplifier* = 0.15 (Extended Data Fig. 7e). All tumor cell clusters were confirmed to have CNVs characteristic of their subgroup¹¹, with only one sample (SHH MB3862) having no detected CNVs (Extended Data Fig. 7f). Given the high proportion of tumor cells in other SHH MB samples compared to non-tumor microenvironment cells, and the distinct clustering from microenvironment cells, we labelled these as tumor cells.

Cell to cell ligand receptor analysis—To identify putative ligand-receptor interactions occurring between tumor cells and microenvironmental non-tumor cells in the scRNAseq data, we employed CCInx (v0.4)⁴³, as previously described⁶⁴. Briefly, scaled expression level of both subgroup-specific ligands in microglia and receptors in G4 MB tumor cells was calculated using the function *BuildGeneStatList*, and interactions were determined using the built-in curated database of ligand-receptor interactions (https://baderlab.org/

CellCellInteractions). Edges represent putative interactions and they are ordered and coloured by the magnitude of the average expression of the ligand and receptor.

SINGLE-NUCLEUS RNA SEQUENCING (HUMAN SAMPLES)

Sample collection and processing—Human cerebellar tissue from 9-21 PCW were previously processed and the data was obtained¹³. Samples were aligned to reference human genome hg19 as for the tumor samples.

Clustering analysis and visualization—Clustering analysis and visualization for the entire human cerebellum dataset were performed as previously stated¹³. For more discrete cell-type similarity mapping a glutamatergic subset of the object was created by extracting cell clusters 02-RL, 03-GCP, 05-eCN/UBCs, and 04-GN and reclustering by the same methods as previously stated for tumor single-cell RNAseq samples.

Pseudo-time trajectory analysis—To recapitulate the expected developmental trajectories¹³ in the glutamatergic cell subset we employed the trajectory inference method Slingshot (v1.6.1)⁶⁵, as previously described⁶⁴. Briefly, a three-dimensional diffusion map embedding was constructed using the *DiffusionMap* function from the R package destiny (v3.2.0)⁶⁶. Slingshot analysis was then performed on the diffusion map to determine per-cell pseudotime estimates and mapped back to the UMAP embedding.

Identification of cell types—Re-clustering analysis of the glutamatergic cell clusters from the developing human cerebellum snRNAseq dataset revealed additional heterogeneity within each original cluster. The RL cells clustered into two distinct transcriptional clusters, consistent with previous histological findings^{12,13}. We performed differential gene expression analysis using the FindAllMarkers methods with the following non-default parameters; *only.pos* = *T, test.use* = "*MAST*"⁴². We annotated these two clusters as RL^{VZ} and RL^{SVZ} based on expression of characteristic markers of these distinct cell types^{12,13} (Extended Data Fig. 7a). The eCN/UBC cluster also split into two distinct transcriptional clusters, which we annotated as "Early" and "Late" UBCs based on pseudotime results. The GN cluster split into several transcriptionally distinct clusters which we labelled as "Early" and "Late" GN based on pseudotime results. A list of the top 50 differentially expressed gene markers of each cluster can be found in Supplementary Data Table 7.

Mapping transcriptional similarity between medulloblastoma and the human cerebellum—To compare the transcriptional similarity between MB cells and the developing human cerebellum, we first used the SingleR package $(v1.2.4)^{67}$. A human cerebellum development reference was created using the function aggregateReference, with the parameter power = 0. We confirmed that the new reference was able to successfully self-classify the human cerebellar cells. This reference was then used to classify tumor cells from each MB sample by transcriptional similarity. This method was repeated with an additional step of removing genes associated with cell cycle^{61,68} from the expression matrices to confirm there was no spurious cell type correlation based on medulloblastoma cell cycle state. As a control, we found that SHH MB cells were most similar to GCPs, consistent with their presumed cell of origin^{3,5,6,69}.

Deconvolution analysis—As a parallel method to determine transcriptional similarity and in a much larger cohort of bulk RNAseq MB samples, we used CIBERSORT⁷⁰ to deconvolute the relative proportions of each glutamatergic cell type in the developing human cerebellum. Briefly, the CIBERSORT algorithm was called in R using the amritr package (v0.1.0), with parameters perm = 10, QN = F. As a control, we again found that the dominant cell type proportion in SHH MB was GCPs, as expected.

Transcription factor activity analysis—The activity of specific TFs in each cell type was inferred in both the human development cells and the MB cells using the package pySCENIC (v0.10.0)^{44,71}, implemented in python (v3.7.6), with the parameter *-min_genes* 10. For each TF the cellular activity was determined and binarized into an "on" or "off" state using the function *binarize* with default parameters.

SINGLE-NUCLEUS RNA SEQUENCING (G3 MB CELL LINES)

Preparation of single-nuclei suspensions—Nuclei were isolated from fresh, snap frozen tumor tissues as previously described ⁷². Frozen pellets were dounced in 1 mL of chilled lysis buffer (lysis buffer; 10 mM Tris-HCl (pH 7.4), 10 mM NaCl, 3 mM MgCl₂, 0.05% NP-40 detergent) 5 times with a loose pestle, 10 times with a tight pestle and lysed for 10 minutes on ice. Chilled wash buffer (5 ml, wash buffer; 5% BSA, 0.04 U/ μ l Rnase inhibitor, 0.25% glycerol) was added to the sample, passed through a 40 μ m cell strainer and centrifuged at 500 x g for 5 minutes at 4°C. After pelleting, the nuclei were resuspended in 5–10 ml of wash buffer. After two washes, single-nuclei suspensions were passed through a 20 μ m cell strainer, pelleted, and resuspended in PBS with 0.05% BSA. Library preparation and sequencing where then conducted as stated above for single-cell RNA sequencing of medulloblastoma samples.

Alignment of raw reads—Alignment was performed as previously stated for human medulloblastoma tumors, with the following differences. When invoking CellRanger's (v6.0.1) *count* method, the additional parameter *--include-introns* was specified as recommended for single-nucleus sequencing data.

Quality control, normalization, and clustering—Quality control, normalization, and clustering were performed as previously stated for human MB tumors, with the following difference. Following QC, there remained a number of cells with notably low number of features detected per RNA count, indicative of ambient RNA background or low complexity cells, these cells were filtered out on a per sample basis.

Mapping transcriptional similarity between cell lines and the human cerebellum—We observed differentiation in our cultures of OTX2-KD cells from both HDMB03 and MB3W1. Using the previously stated method above for comparing primary MB tumor cells to the human developing cerebellum, we used SingleR with the same reference dataset to classify individual cells from each scramble and OTX2-KD sample.

Identification of cellular differentiation state—In the OTX-KD samples for both HDMB03 and MB3W1 we noted a gradient of *OTX2* expression, rather than a consistent

decrease in expression. We suspected this could be due to inefficient OTX2-KD in certain cells. Using the differentially expressed genes (LFC > 2, p < 0.05) obtained from bulk RNAseq of OTX2-KD and scramble samples, we generated two gene signatures called "Scramble signature" and "OTX2-KD signature", which represent gene expression programs of the dominant cellular populations in each condition. Using the *AddModuleScore* function in Seurat, we scored the single cells from each condition for these signatures (Extended Data Fig. 11g), and found cluster specific enrichment for the scramble signature, likely indicating incomplete knockdown in a small subset of cells. If these cells indeed had inefficient OTX2-KD, we would expect those cells to retain features of wild-type cells, namely a lack of differentiation. To test this, we applied the CytoTRACE (v0.3.3) algorithm which estimates the differentiation state of single cells using the number of genes expressed per cell⁷³. As expected, these putatively inefficient knockdown cells were inferred to be the least differentiated cells (Extended Data Fig. 11h).

Pseudo-time trajectory analysis—We then conducted pseudotime trajectory analysis using Slingshot, as described for tumor samples above, however specifying the start of the lineage (or, the 'root') as the cluster with the highest CytoTRACE scores. To identify genes whose expression as significantly correlated with the determined lineages, we first fit each gene with a negative binomial general additive model as implemented in the R package TradeSeq $(v1.2.01)^{74}$. We then tested whether these gene level models were correlated with inferred pseudotime using the *associationTest* function with the parameter *lineages* = T, and filtered out genes with FDR corrected p-values > 0.05. Finally, we overlapped these significantly lineage associated genes with significant cell type marker genes from the human developing cerebellum dataset and used the *predictSmooth* function to bin the expression of these genes along OTX2-KD and normal human cerebellar development pseudotime.

SURVIVAL ANALYSIS

Overall survival for each of the subtypes of G3 and G4 MB was analyzed by the Kaplan-Meier method and p-values were determined using the log-rank test. Survival analysis was conducted using the R package survival (v3.2–11)⁷⁵.

CBFA2T2 PROTEIN STRUCTURAL MODEL

The structure of the NHR1 domain of CBFA2T2 was accessed from SWISS-MODEL (accession number O43439). This structure was previously inferred using homology to the known structure of the RUNX1T1 NHR1 domain⁷⁶. The full protein structure was predicted using i-TASSER³⁹. The optimal CBFA2T2 prediction was selected for visualization based on which model's NHR2 domain exhibited the highest degree of similarity to the known SWISS-MODEL structure. This model was then visualized and coloured in PyMOL (v2.4.2). However, the high degree of disorder in CBFA2T2 makes the predicted model imprecise and is provided only for context of the NHR1 domain.

MUTUAL EXCLUSIVITY ANALYSIS

Mutual exclusivity of various alterations was statistically assessed using the DISCOVER R package $(v0.9.3)^{41}$. All *p*-values derived from multiple testing were corrected using built-in false discovery rate (FDR).

PROTEIN-PROTEIN INTERACTION NETWORK

To determine the extent of protein-protein interactions (PPIs) that are known to occur between G3 and G4 MB driver genes (Fig. 1a), and with novel CBFA2T2 interacting proteins (TurboID), we employed STRING analysis (v11.5)⁷⁷. Briefly, gene lists of driver genes and genes encoding CBFA2T2 interacting proteins were queried for known interactions. The resulting network was further filtered for only interactions with experimental evidence and a confidence score of at least 0.250. The network was then imported into Cytoscape^{78,79} for visualization. Each node corresponds to a protein and was coloured by the relative frequency that the corresponding gene is altered (i.e. mutated, deleted, amplified, enhancer hijacked, etc.) across G3 and G4 MB. Edges correspond to known PPIs and their weight is proportional to the STRING confidence score. The nodes of CBFA2T2 prey proteins as determined by TurboID were assigned a diamond shape, but not all edges between diamond-shaped nodes and CBFA2T2 were drawn to reduce complexity. Nodes were then manually arranged to group proteins in the same gene family, and thus edge length is arbitrary, however the degree of connectivity between groups guided organization of the nodes.

ILLUSTRATIONS

Oncoprint landscape figures were generated in R (v4.0.2) using the ComplexHeatmap (v2.4.3) library 80 . Gene mutation summary lollipop figures were generated using ProteinPaint 81 .

DATA AVAILABILITY

The bulk and single-cell RNAseq data generated from MB tumor samples in this study has been deposited in the European Genome-Phenome Archive (EGA) database under the accession code EGAS00001005826. The bulk and single-cell RNAseq data generated from the G3 MB cell lines HDMB03 and MB3W1 in this study has been deposited in the Gene Expression Omnibus (GEO) database under the access codes GSE189238 and GSE200791, respectively. The published MB bulk and single-cell RNA-seq data referenced in this study is available in the EGA database under the accessions EGAD00001006305, EGAD00001004435, EGAD00001004958, EGAS00001003170, and EGAS00001003368. The referenced GTEx normal cerebellum RNAseq controls were acquired from the NCBI public repository phs000424.v6.p1. The Affymetrix SNP 6.0 data referenced during the study are available in the Gene Expression Omnibus (GEO) under the accession GSE37385. The whole-genome sequencing data referenced during the study are available in EGA under the accessions EGAD00001003125 and EGAD00001004347. The Illumina 450k methylation data referenced during the study are available in GEO under the accession GSE85218. The G3 tumorsphere ChIP-seq data referenced during the study is available in GEO under the accession GSE132269. There were multiple databases used for annotation

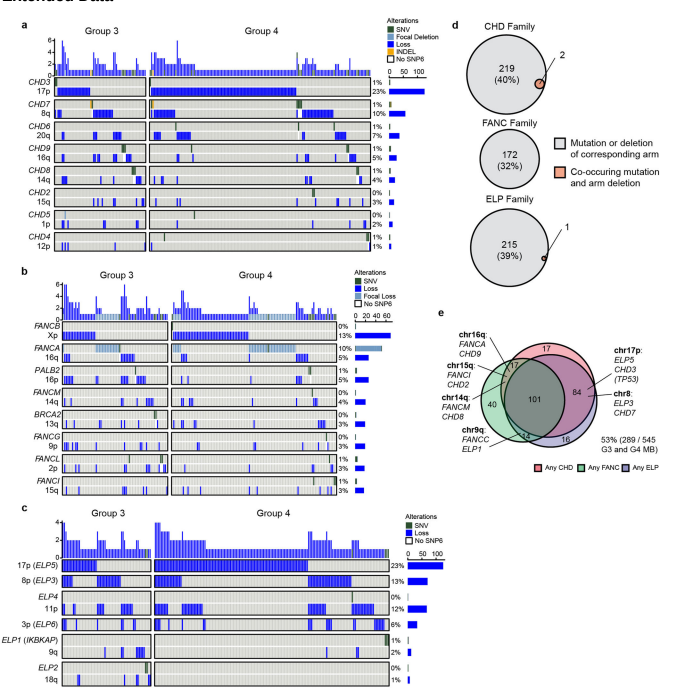
and filtering referenced in this study. These include the Exome Aggregation Consortium / gnomAD [https://gnomad.broadinstitute.org/downloads], the NHLBI-ESP project [https://esp.gs.washington.edu/drupal/], the Kaviar Genomic Variant Database [http://db.systemsbiology.net/kaviar/#:~:text=Kaviar%20Genomic%20Variant%20Database%20%7C%20SNP,and%20frequency%20of%20observed%20variants.], the Haplotype Reference Consortium [http://www.haplotype-reference-consortium.org/], the Greater Middle East Variome [http://igm.ucsd.edu/gme/], the Brazilian Genomic Variants Database [http://abraom.ib.usp.br/], RADAR [http://rnaedit.com/], GENCODE (v19) [https://www.gencodegenes.org/human/release_19.html], the hs37d5 reference genome [https://ftp-trace.ncbi.nih.gov/1000genomes/ftp/technical/reference/phase2_reference_assembly_sequence/], ERCC spike-

release_19.html], 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/]. Single-nucleus RNA sequencing data from the developing human cerebellum was obtained through correspondence from Aldinger et al., 13 and is also available through the Human Cell Atlas [https://www.covid19cellatlas.org/aldinger20], the UCSC Cell Browser [https://cbl-dev.cells.ucsc.edu], or from Database of Genotypes and Phenotypes (dbGaP) (accession number: phs001908.v2.p1). Bulk RNA sequencing data from the developing human cerebellum was obtained through correspondence from Haldipur et al. 12 and is available through the dbGaP (accession number: phs001908.v2.p1).

CODE AVAILABILITY

No custom code was used in this study. Only open-source algorithms were used, and their applications are detailed in the methods. Further details on how these algorithms were used are available from the corresponding author upon request.

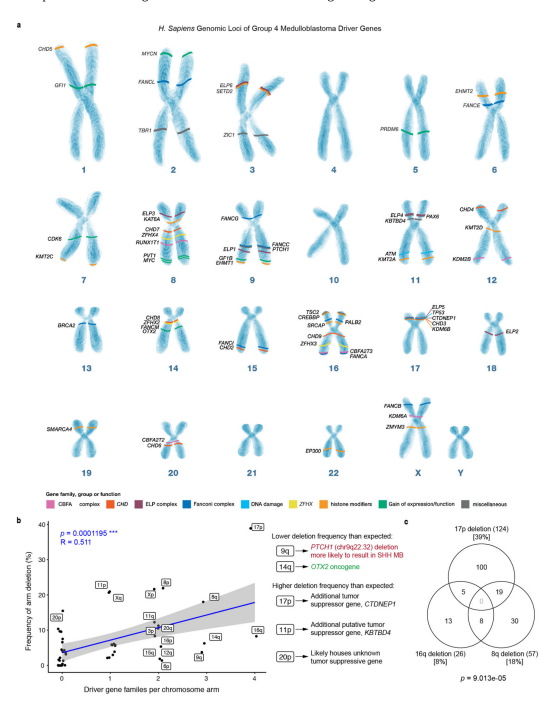
Extended Data



Extended Data Fig.1 \mid . Mutations and copy number losses targeting the same genes are mutually exclusive in G3 and G4 MB.

a, b, c, Oncoprint summarizing gene mutations and copy number losses in their corresponding genomic loci (chromosome arm) for CHD (**a**), FANC (**b**) and ELP (**c**) driver families. Mutations are less frequent than copy number losses but tend to occur independently suggesting they are targets of the deletions. **d,** Mutual exclusivity of mutation and copy number loss events targeting genes in the CHD, FANC, and ELP families. **e,**

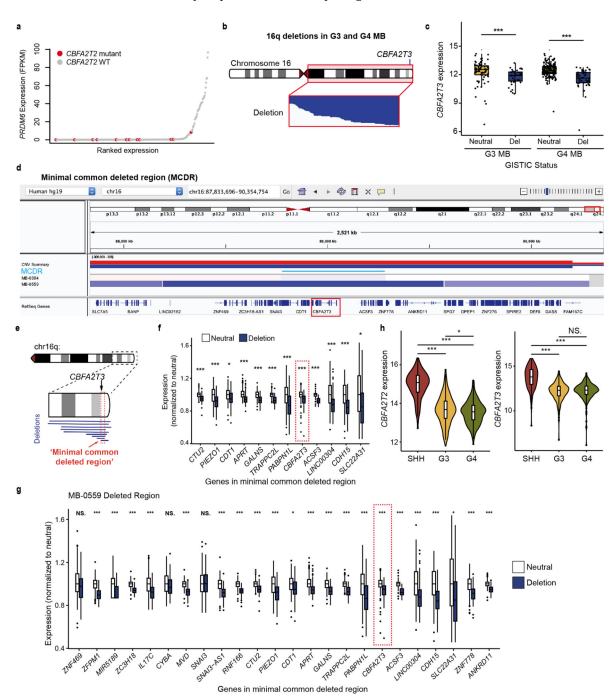
Overlap between events targeting CHD, FANC, or ELP genes. Most G4 MB tumors are haploinsufficient for genes in at least two families through a single deletion event.



Extended Data Fig.2 \mid . Clustering of G4 MB driver genes in the human genome predisposes humans to develop MB.

a, Cartoon of the *Homo sapiens* genome with the locations of known and newly identified G3 and G4 MB candidate driver genes demonstrating clustering of genes at locations known to be deleted in G3 and G4 MB. **b,** Frequency of whole chromosome arm loss is significantly correlated with the number of driver gene families—as detailed in (**a**)—

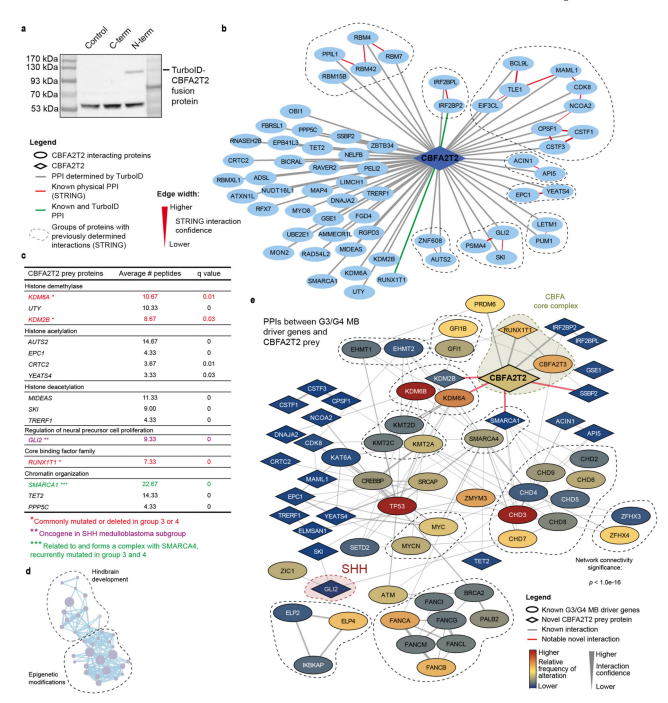
contained on the arm. Significance was assessed by a two-sided linear regression model; grey shaded area denotes the 95% confidence interval. **c**, Mutual exclusivity of copy number losses of chromosome arms 17p, 16q, and 8q. Significance was assessed using the impurity test for mutual exclusivity, implemented in the R package DISCOVER⁴¹.



Extended Data Fig.3 |. CBFA2T3 is a G4 MB tumor suppressor gene.

a, *PRDM6* expression in *CBFA2T2* mutant (red) and *CBFA2T2* WT (grey) G3 and G4 MB samples demonstrates that enhancer hijacking mediated *PRDM6* expression is largely

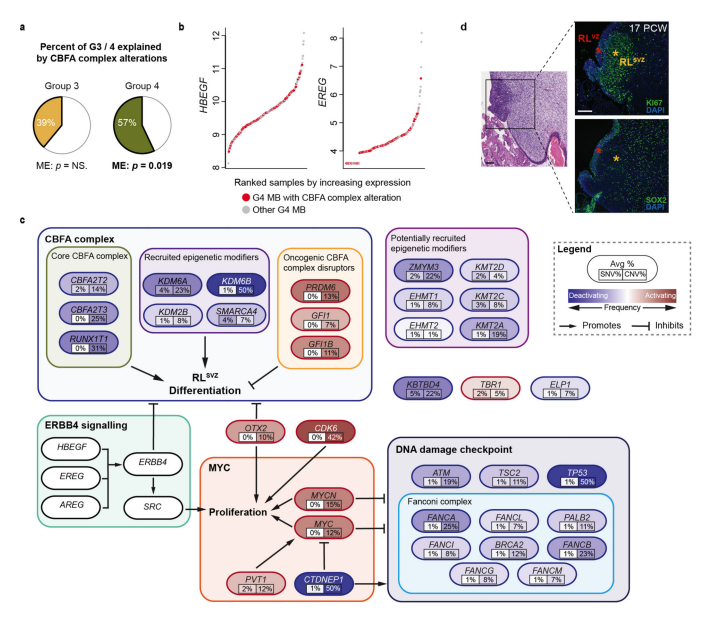
limited to CBFA2T2WT cases. b, Density of regions of chromosomal gain and loss along human chromosome 16q in G3 and G4 MB cases, demonstrating that deletions are biased towards the telomeric end of 16q, the location of known drivers, particularly CBFA2T3. c, CBFA2T3 expression differences between samples with and without CBFA2T3 deletions, split by subgroup. Statistical significance was assessed using Kruskal-Wallis rank-sum test (FDR < 0.05), *** p < 0.0005, G3, p = 2.88 $e^{-0.05}$; G4, p = 2.60 $e^{-0.09}$. G3, n = 112; G4, n = 206. CBFA2T3 is a copy-number responsive tumor suppressor gene in G4 MB. d, IGV analysis showing focal deleted region in two G4 MB samples MB-0364 and MB-0559. MB-0364, which is the minimal common deleted region (MCDR) on 16q in G3 and G4 MB, though does not quite achieve statistical significance in the GISTIC analysis. MB-0559 is the MCDR achieving statistical significance in GISTIC analysis. CBFA2T3 is identified with a red box. e, Cartoon illustrating the MCDR concept. f, Expression differences between copy neutral or hemizygously deleted G3 and G4 MB samples for genes within the MB-0364 MCDR on chr16q24.3. Statistical significance was assessed using two-sided Mann-Whitney U tests with FDR adjustment, * p < 0.05, *** p < 0.0005. Deletion, n = 86; Neutral, n =232. e, Expression differences between copy neutral or hemizygously deleted G3 and G4 MB samples for genes within the MB-0559 MCDR on chr16q24.3. Statistical significance was assessed using two-sided Mann-Whitney U tests with FDR adjustment, * p < 0.05, *** p < 0.0005. Deletion, n = 86; Neutral, n = 232. A full list of p values for genes presented in (f) and (g) can be found in Supplementary Data Table 1. h, CBFA2T2 (left) and CBFA2T3 (right) expression in SHH, G3, and G4 MB by bulk RNAseq. Statistical significance was assessed by Kruskal-Wallis rank-sum test (FDR < 0.05), * p < 0.05, *** p< 0.0005. For *CBFA2T2*: SHH-G3, $p = 2.29e^{-47}$; SHH-G4, $p = 4.42e^{-73}$; G3-G4, p = 0.035. For *CBFA2T3*: SHH-G3, $p = 7.10e^{-42}$; SHH-G4, $p = 1.13e^{-46}$; G3-G4, p = 0.61. G3, n = 0.61219; G4, n = 326; SHH, n = 250. While CBFA2T2 and CBFA2T3 are recurrently targeted and have low expression in G3 and G4 MB, high expression of both genes and an absence of alterations are observed in SHH MB. CBFA2T2 and CBFA2T3 likely have different roles in SHH MB compared to G3 and G4 MB. For c, f, g, and h box plots show the median and interquartile range, and whiskers show the data range. Points outside this range are outliers and are plotted individually.



Extended Data Fig.4 \mid . The CBFA polyprotein complex contains multiple known and novel G4 MB driver genes.

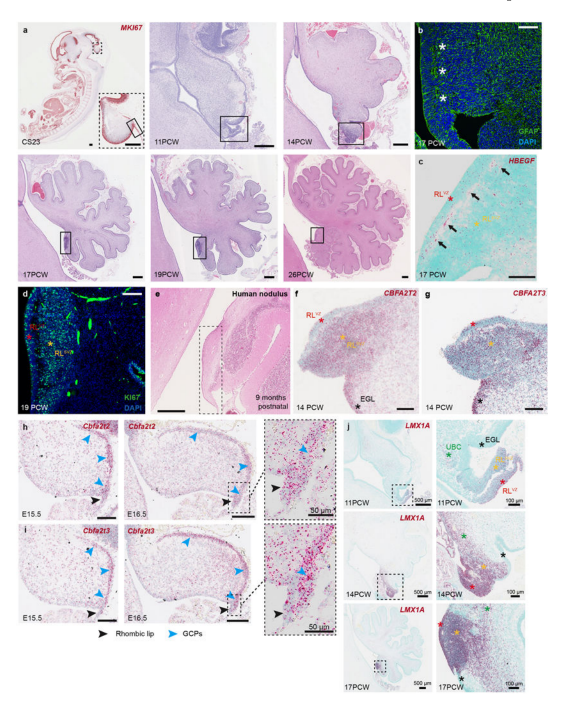
a, Western blot showing successful expression of the TurboID-CBFA2T2 fusion protein when the TurboID construct is fused to the N-terminal of CBFA2T2, but not to the C-terminal. **b,** Protein-protein interaction (PPI) network of novel CBFA2T2 interacting proteins. Each node represents a protein and edges between the proteins represent known or novel PPIs. Edges in red represent known interactions between CBFA2T2 interacting proteins, and edges in green represent known interactions with CBFA2T2 that were

recapitulated in our TurboID screen. Proteins are grouped with dashed lines if they contain known interactions between each other. **c**, Significant CBFA2T2 prey proteins enriched in each indicated biological process. GLI2 is a SHH oncogene and has been recently shown to maintain GCP proliferation and identity, implicating the CBFA complex⁸². **d**, Enrichment map of biological processes (GO:BP) enriched in CBFA2T2 prey proteins by TurboID. Each node represents a significantly enriched pathway and edges represent shared genes between nodes. Nodes are grouped and labelled with a common biological theme. Significance was assessed using G:Profiler⁸³ with FDR correction. **e**, Protein-protein interaction (PPI) network of CBFA2T2 TurboID proteins and G3/G4 MB driver genes (Fig. 1a) using STRING⁴¹. Edges between CBFA2T2 and diamond-shaped nodes are not drawn for simplicity. Connectivity significance was assessed by STRING, $p < 0.1e^{-16}$.



Extended Data Fig.5 |. Disruption to the CBFA complex explains most G4 MB tumors.

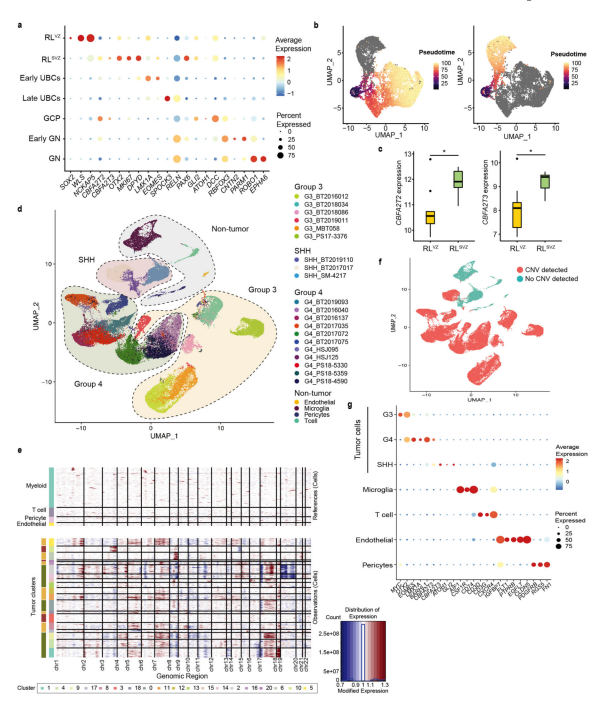
a, Percent of G3 and G4 MB in our cohort (*n* = 545) explained by alterations in genes connected to *CBFA2T2* with a known or novel PPI (one step in the network presented in Extended Data Fig. 4e). Significance assessed using the impurity test for mutual exclusivity implemented in the R package DISCOVER⁴¹. **b,** Ranked expression of *HBEGF* (left) and *EREG* (right) in G4 MB (*n* = 326). Points are coloured by the presence (red) or absence (grey) of known CBFA complex alterations. Samples with the highest expression of *HBEGF* and *EREG* typically do not have CBFA complex alterations, suggesting an alternate mechanism of CBFA complex inhibition. Data presented in **a** were not performed in replicates. **c,** Summary of disrupted pathways in G3 and G4 MB. Altered genes are grouped by pathway and labelled with alteration frequency. **d,** (Left) H&E–stained midsagittal section from 17 PCW human cerebellum. (Right) Fluorescence immunohistochemistry (IHC) showing *KI67* and *SOX2* expression in the human RL compartments. Data presented in **d** were not performed in replicates. RL^{VZ} and RL^{SVZ} are denoted by red and yellow asterisks, respectively. Scale bars: 100 μm.



Extended Data Fig.6 |. LMXIA expression distinguishes the two downstream lineages of the RLSVZ.

a, (Top left) *In situ* hybridization (ISH) showing *MKI67* expression. In-set highlights the developing cerebellum, and the RL is indicated by the black box. (Other images) Hematoxylin and eosin (H&E)–stained midsagittal sections of the developing human cerebellum. In each, the rhombic lip is indicated by the black box. Scale bars: 500 μ m. **b,** GFAP expression in the developing human RL at 17 PCW. Scale bar: 100 μ m. The RL^{VZ} and RL^{SVZ} are physically divided by a vascular plexus, as indicated with white

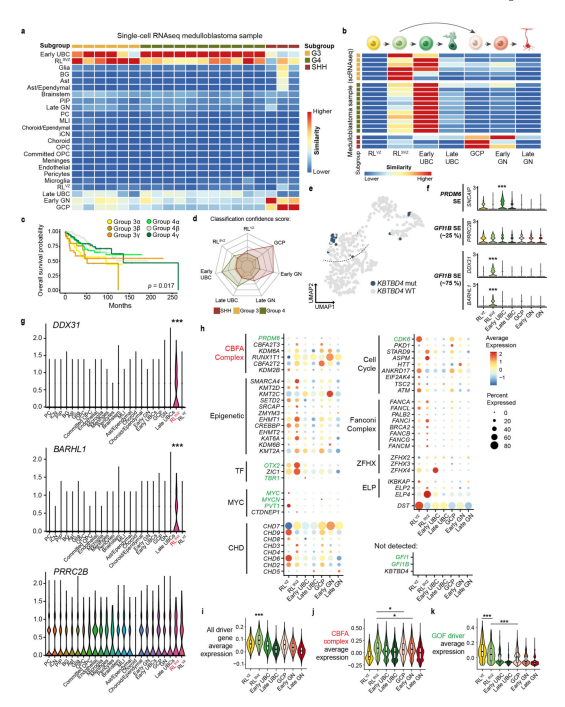
asterisks. c, ISH showing spatially resolved RNA expression of HBEGF in the developing human cerebellum at 17 PCW. Scale bar: 50 µm. HBEGF foci are enriched along the RL vascular plexus. d, KI67 expression in the developing human RL at 19 PCW. Scale bar: 100 μm. e, H&E-stained midsagittal sections of the 9-month postnatal human cerebellum. Scale bar: 500 µm. The RL is only present during gestation and disappears around birth. f, g, ISH showing spatially resolved RNA expression of CBFA2T2 (f) and CBFA2T3 (g) in the developing human cerebellum at 14 PCW. Scale bars: 100 µm. h, i, ISH showing spatially resolved RNA expression of Cbfa2t2 (h) and Cbfa2t3 (i) in the developing mouse cerebellum at E15.5 (Left) and E16.5 (Right). Scale bars: 100 µm. We do not observe a similar expression pattern of either gene in the mouse RL as we do in the human RL, and note an enrichment of expression in the EGL, similar to humans. j, ISH showing spatially resolved RNA expression of LMX1A in the developing human cerebellum at 11, 14, and 17 PCW. LMX1A is highly expressed in both the RLVZ and RLSVZ, but LMX1A expression is only retained in UBCs migrating away from the RL and is completely absent in GCPs that migrate to the EGL. Data presented in d is a representative image from three independent experiments with similar results, data in remaining panels were not performed in replicates.



 ${\bf Extended\ Data\ Fig.7\ |.\ Characterization\ of\ single\ cells\ used\ in\ transcriptional\ mapping\ between\ MB\ and\ human\ cerebellum\ development.}$

a, Dot plot showing expression of characteristic marker genes across RL glutamatergic cell types in the developing human cerebellum¹³. **b,** UMAP embeddings coloured by pseudotime inferred from Slingshot⁶⁵, where the direction of pseudotime is from dark to light colours, for the granule cell lineage (Left) and the UBC lineage (Right). **c,** Expression of *CBFA2T2* (Left) and *CBFA2T3* (Right) in each zone of the developing human RL by bulk RNAseq¹². Statistical significance was assessed using a two-sided Mann-Whitney U test, * p < 0.05;

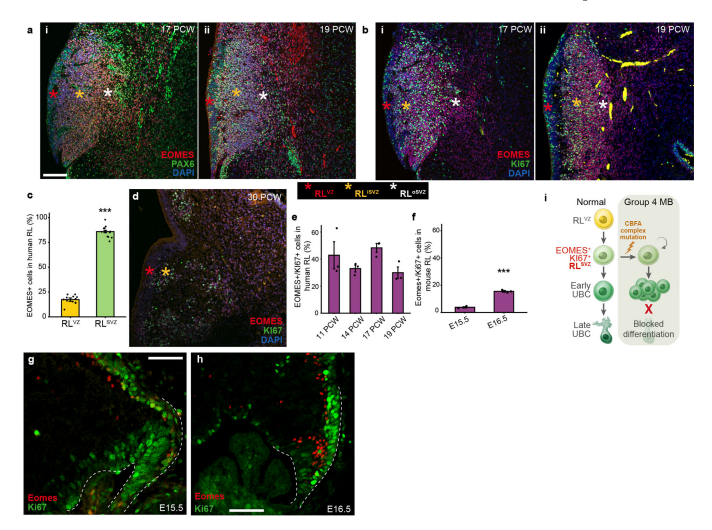
CBFA2T2, p = 0.0078; CBFA2T3, p = 0.0056. n = 9 biological samples, per zone, acquired between 9 and 19 PCW. Box plots show the median and interquartile range, and whiskers show the data range. Points outside this range are outliers and are plotted individually. **d**, UMAP embedding of 63,296 single cells derived from G3 (n = 6), G4 (n = 11), and SHH (n = 3) MB scRNAseq samples. Clusters of transcriptionally similar cells are colored and labeled by tumor sample or annotated cell type for non-tumor cells. **e**, Copy number variations detected in single cells inferred using inferCNV⁶³. (Top) Reference non-tumor cells are devoid of copy number variations. (Bottom) Tumor cell clusters were enriched for copy number variations characteristic of the sample subgroup. Cells containing CNVs were assigned as tumor cells for downstream analysis. **f**, UMAP embedding as in (**d**) coloured by the detection of copy number variations. **g**, Dot plot showing expression of characteristic marker genes of SHH, G3, G4 MB, and the non-tumor cell types identified. For **a**, **g**, colour indicates average expression and size of each dot indicates the percent of cells in that cluster that express the genes.



Extended Data Fig.8 \mid . G3 and G4 MB resemble specific components of the human RL, whose differentiation is stalled in time.

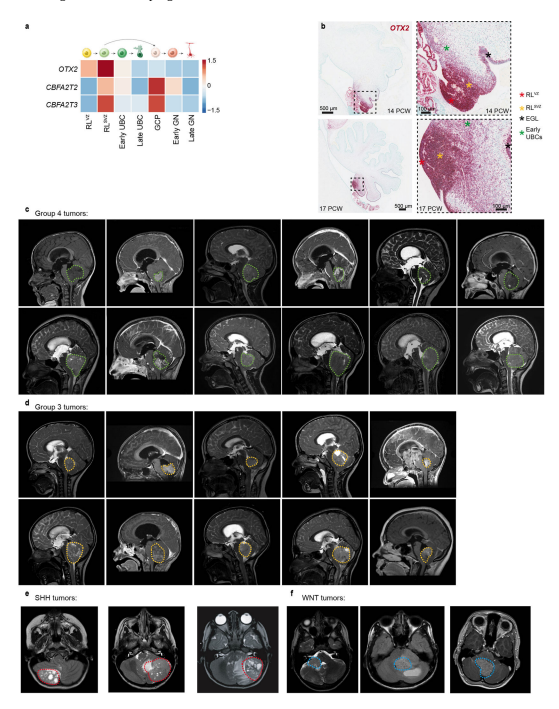
a, SingleR⁶⁷ classification of tumor cells from G3 (n = 6), G4 (n = 11), and SHH (n = 3) MB scRNAseq samples, by comparison to the entire developing human cerebellum¹³. As expected, MB cells are most similar to glutamatergic cells. **b,** SingleR classification of tumor cells from (**a**) by comparison to glutamatergic cell types. **c,** Kaplan-Meier plot showing overall survival of G3 and G4 MB subtypes in the current dataset. Significance assessed using a log-rank test. Censored cases, +. **d,** Relative confidence of per cell

classifications, calculated as the average similarity score per subgroup (b), minus median similarity scores from other subgroups per cell type. e, UMAP embedding of n = 545 G3 and G4 MB bulk RNAseq samples, coloured by presence of KBTBD4 mutations. G4y which are impoverished for CBFA complex mutations, and display high OTX2 expression, are also enriched for KBTBD4 mutations. f. Expression of super enhancer (SE) genes in the developing human cerebellum snRNAseq data. These gene promoters have been demonstrated to promote transcription of PRDM6 (SNCAIP) and GFI1B (DDX31/BARHL1 and *PRRC2B*) in G3 and G4 MB secondary to enhancer hijacking events ^{11,28}. Significance was assessed using a two-sided Wilcoxon Rank Sum test with FDR correction, *** p < 0.0005. SNCAIP, $p = 3.39e^{-261}$; DDX31, $p = 4.27e^{-71}$; BARHL1, $p = 4.91e^{-40}$. $n = 4.91e^{-40}$. 9,208 cells. g, Expression of DDX31, BARHL1, and PRRC2B across all cell types in the developing human cerebellum. DDX31 and BARHL1 exhibit correlated expression specific to the RLSVZ, while PRRC2B is non-specifically expressed. Significance was assessed using a two-sided Wilcoxon Rank Sum test with FDR correction, *** p < 0.0005. DDX31, p = $3.66e^{-113}$; BARHL1, $p = 6.26e^{-191}$. n = 59,608 cells. **h**, Expression of G3 and G4 MB driver genes (from Fig.1a) in the developing human cerebellum snRNAseq data. i, j, k, Average expression of all G3 and G4 MB driver genes (i), CBFA complex genes (j), and gain of function (GOF) driver genes (k) in the developing human cerebellum snRNAseq data. Significance was assessed using a two-sided Wilcoxon Rank Sum test, * p < 0.05, *** p < 0.0005. n = 9,208 cells. (i), $p = 1.5e^{-13}$; (j) GCP, p = 0.0085; Early GN, p = 0.047; (k) RL^{SVZ}, $p = 1.9e^{-0.5}$; GCP, $p = 3.2e^{-1.3}$. For i, j, and k, box plots show the median and interquartile range, and whiskers show the data range. Points outside this range are outliers and are plotted individually.



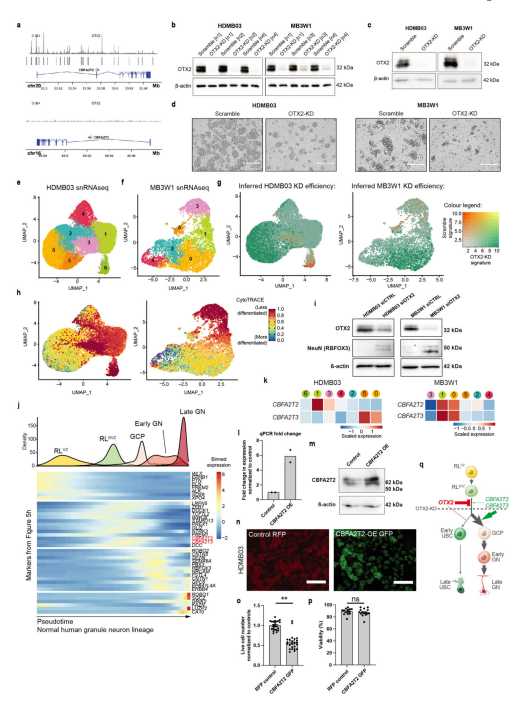
Extended Data Fig.9 |. Human EOMES^{+ve} RL cells are mitotically active UBC progenitors. a, EOMES and PAX6 expression in the developing human RL at 17 (i) and 19 (ii) PCW. Scale bars: 100 µm. b, EOMES and KI67 expression in the developing human RL at 17 (i) and 19 (ii) PCW. Scale bars as in a. Proliferating EOMES+ve UBC progenitors are common across all developmental timepoints assessed. c, EOMES+ cells in the human RL zones. The RLSVZ contains significantly more EOMES+ cells that the RLVZ. Significance was assessed using an unpaired two-tailed t-test, *** $p = 1.048e^{-18}$. n = 3 biological repeats, per N = 4 time points; error bars, SEM. d, EOMES and KI67 expression in the developing human RL at the late timepoint 30 PCW. Scale bar as in (a). Proliferating EOMES^{+ve} UBC progenitors can be found across fetal development, though at reduced frequency at later time points as KI67 expression is reduced (Fig.3c). e, Quantification of the number of EOMES+/KI67+ cells in the human RL across various developmental timepoints. All comparisons to 11 PCW were non-significant using two-tailed unpaired t-tests; 14PCW, p = 0.43; 17PCW, p = 0.65; 19PCW, p = 0.33. n = 3 biological repeats per timepoint; error bars, SEM. EOMES+/KI67+ UBC progenitors are a long-lived and dominant population of the RL, rather than a transient state preceding differentiation. f, Quantification of the number of Eomes+/Ki67+ cells in the mouse RL across various developmental timepoints. Significance was assessed using an unpaired two-tailed t-test, *** p = 0.00015. n = 3 biological repeats per timepoint; error

bars, SEM. Eomes+/Ki67+ UBC progenitor cells are a rare population in the mouse RL. **g**, **h**, Eomes and Ki67 expression in the mouse RL at E15.5 (**g**) and E16.5 (**h**). The RL boundaries are indicated with white dashed lines. Scale bars: 50µm. Eomes+ UBCs are rarely Ki67+. Data presented in **a**, **b**, **g**, **h** are representative images from three independent experiments with similar results, data in **d** were not performed in replicates. **i**, Oncogenic divergence of RL^{SVZ} progenitors from normal initiate G4 MB.



Extended Data Fig.10 \mid . The location of G3 and G4 MB tumors coincides with OTX2 expression and supports an RL cell of origin.

a, Scaled *OTX2*, *CBFA2T2*, and *CBFA2T3* expression by scRNAseq. **b**, Expression of *OTX2* at 14 and 17 PCW by ISH in the developing human cerebellum. **c**, T1 enhanced or T2 mid-sagittal MRI images of G4 MB (n = 12) tumors at initial diagnosis. **d**, T1 enhanced or T2 mid-sagittal MRI images of G3 MB (n = 10) tumors at initial diagnosis. Both G3 and G4 MB tumors present exclusively in the *OTX2*+ inferior cerebellum. **e**, Axial T1 enhanced, T2 or FLAIR images of SHH MB (n = 3) at initial diagnosis. SHH tumors occur in the cerebellar hemispheres, consistent with an EGL cell of origin. **f**, Axial T1 enhanced, T2 or FLAIR images of WNT MB (n = 3) at initial diagnosis. Data presented in **b** were not performed in replicates.



Extended Data Fig.11 \mid . OTX2 knockdown promotes G3 MB differentiation through intermediate upregulation of CBFA2T2 and CBFA2T3.

a, *OTX2* ChIP-seq³⁴ peaks are enriched at *CBFA2T2* gene locus, but not *CBFA2T3*. **b,** OTX2 protein expression is reduced following OTX2-KD. Samples were used for bulk RNA sequencing. Beta actin used as a loading control. **c,** OTX2 protein expression is reduced following OTX2-KD. Samples were used for single-nucleus RNA sequencing. Beta actin used as a loading control. **d,** Representative images of primary tumorspheres in OTX2-KD and scramble conditions for both HDMB03 and MB3W1 cultures. Scale bar: 300 μm. **e,**

f, Unbiased clustering of single nuclei following OTX2-KD in HDMB03 and MB3W1 G3 MB cells lines (c). g, Average expression of gene signatures derived from bulk RNAseq on OTX2-KD from HDMB03 and MB3W1 (b). Cells that are more orange than green indicate cells with higher expression of genes characteristic of the unchanged G3 MB cell lines, and vice-versa. Orange cells are likely cells where OTX2-KD was inefficient. h, Differentiation score as determined by CytoTRACE⁷³. Less differentiated cells are indicated in red and more differentiated cells are indicated in blue. The results support a model where cluster 6 in HDMB03 and cluster 3 in MB3W1 represent inefficient OTX2-KD cells that retain the most similarity to WT tumor cells. i, RBFOX3 (NeuN) protein expression is increased following OTX2-KD in both HDMB03 and MB3W1, validating GN differentiation following OTX2-KD. j, Expression of genes significantly correlated with granule neuron differentiation along pseudotime in normal human RL development. (Top) Density of cell along pseudotime in the granule neuron lineage (Extended Data Fig.6b, left). (Bottom) Binned gene expression of markers derived from the developing human cerebellum snRNAseq dataset (Fig.4a). The stepwise expression of granule neuron genes observed when OTX2 is knocked down in G3 MB (Fig.5i) strikingly mirrors that of normal granule neuron differentiation, suggesting that G3 and G4 MB arise from failed normal differentiation rather than alternate hypotheses, such as trans- or de-differentiation. k, CBFA2T2 and CBFA2T3 expression in HDMB03 (Left) and MB3W1 (Right). CBFA2T2 expression is strongly upregulated in cells where the CytoTRACE score drops below 0.8, and CBFA2T3 follows. The results suggest CBFA2T2 and then CBFA2T3 are upregulated early in response to efficient OTX2-KD. I, CBFA2T2 expression change in response to CBFA2T2 overexpression (CBFA2T2-OE) in HDMB03 by qPCR. m, CBFA2T2 protein expression is increased following CBFA2T2-OE. β-actin used as a loading control. n, Representative images of primary tumorspheres in CBFA2T2-OE GFP and Control RFP conditions. Scale bar: 600 µm. o, p, Live cell number (o) and viability (p) in response to CBFA2T2-OE. Live cell number is significantly reduced in response to CBFA2T2-OE, while viability is unchanged. Data are normalized to their respective controls and presented showing points from n = 3 technical replicates per N = 8 or N = 5 biological replicates, for live cell number and viability, respectively. Error bars indicate SEM. Significance assessed using a two-tailed paired t-test on biological replicates, ** p = 0.0047. **q,** OTX2 restrains differentiation of RL progenitors through CBFA complex inhibition. Data presented in **d**, **n** are representative images from 4 and 8 independent experiments, respectively, with similar results, data in **b**, **c**, **i**, **m** were not performed in replicates. For gel source data, see Supplementary Figure 1.

Supplementary Material

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Authors

Liam D. Hendrikse^{1,2,3,*}, Parthiv Haldipur^{4,*}, Olivier Saulnier^{1,2,*}, Jake Millman⁴, Alexandria H. Sjoboen⁴, Anders W. Erickson^{1,2,5}, Winnie Ong^{1,2,5}, Victor Gordon⁶, Ludivine Coudière-Morrison⁶, Audrey L. Mercier⁷, Mohammad Shokouhian⁸, Raúl A. Suárez^{1,2}, Michelle Ly^{1,2,5}, Stephanie Borlase⁶, David S. Scott^{1,2}, Maria C. Vladoiu^{1,2,5}, Hamza Farooq^{1,2,5}, Olga Sirbu^{1,2,3}, Takuma Nakashima⁹, Shohei

Nambu⁹, Yusuke Funakoshi⁹, Alec Bahcheli^{10,11}, J. Javier Diaz-Mejia¹², Joseph Golser⁴, Kathleen Bach⁴, Tram Phuong-Bao⁸, Patryk Skowron^{1,2,5}, Evan Y. Wang^{1,2,3}, Sachin A. Kumar^{1,2,5}, Polina Balin^{1,2,5}, Abhirami Visvanathan^{1,2}, John J.Y. Lee^{1,2,5}, Ramy Ayoub³, Xin Chen^{1,2}, Xiaodi Chen^{1,2}, Karen L. Mungall¹³, Betty Luu^{1,2}, Pierre Bérubé¹⁴, Yu C. Wang¹⁴, Stefan M. Pfister^{15,16}, Seung-Ki Kim¹⁷, Olivier Delattre^{18,19}, Franck Bourdeaut^{18,19}, François Doz^{18,20}, Julien Masliah-Planchon²¹. Wieslawa A. Graikowska²². James Loukides¹. Peter Dirks^{1,2,10,23}. Michelle Fèvre-Montange^{24,25}, Anne Jouvet^{25,100}, Pim J. French²⁶, Johan M. Kros²⁷, Karel Zitterbart²⁸, Swneke D. Bailey^{29,30}, Charles G. Eberhart³¹, Amulya A.N. Rao³², Caterina Giannini³³, James M. Olson³⁴, Miklós Garami³⁵, Peter Hauser³⁵, Joanna J. Phillips^{36,37}, Young S. Ra³⁸, Carmen de Torres^{39,100}, Jaume Mora³⁹, Kay K.W. Li⁴⁰, Ho-Keung Ng⁴⁰, Wai S. Poon⁴¹, Ian F. Pollack⁴², Enrique López-Aguilar⁴³, G. Yancey Gillespie⁴⁴, Timothy E. Van Meter⁴⁵, Tomoko Shofuda⁴⁶, Rajeev Vibhakar⁴⁷, Reid C. Thompson⁴⁸, Michael K. Cooper⁴⁹, Joshua B. Rubin⁵⁰, Toshihiro Kumabe⁵¹, Shin Jung⁵², Boleslaw Lach^{53,54}, Achille Iolascon^{55,56}, Veronica Ferrucci^{55,56}, Pasqualino de Antonellis^{55,56}, Massimo Zollo^{55,56}, Giuseppe Cinalli⁵⁷, Shenandoah Robinson⁵⁸, Duncan S. Stearns⁵⁹, Erwin G. Van Meir⁶⁰, Paola Porrati⁶¹, Gaetano Finocchiaro⁶¹, Maura Massimino⁶¹, Carlos G. Carlotti⁶², Claudia C. Faria^{63,64}, Martine F. Roussel⁶⁵, Frederick Boop⁶⁵, Jennifer A. Chan⁶⁶, Kimberly A. Aldinger^{4,67}, Ferechte Razavi⁶⁸, Evelina Silvestri⁶⁹, Roger E. McLendon^{70,71}, Eric M. Thompson⁷¹, Marc Ansari^{72,73}, Maria L. Garre⁷⁴, Fernando Chico⁷⁵, Pilar Eguía⁷⁵, Mario Pérezpeña⁷⁶, A. Sorana Morrissy^{66,77,78}, Florence M.G. Cavalli^{79,80,81}, Xiaochong Wu^{1,2}, Craig Daniels^{1,2}, Jeremy N. Rich⁸², Steven J.M. Jones 13,83,84, Richard A. Moore 13, Marco A. Marra 13,83, Xi Huang 1,2,10. Jüri Reimand^{3,10,11}, Poul H. Sorensen^{85,86}, Robert J. Wechsler-Reya⁸⁷, William A. Weiss^{36,88,89}, Trevor J. Pugh^{3,11,12}, Livia Garzia⁹⁰, Claudia L. Kleinman^{91,92}, Lincoln D. Stein^{10,93}, Nada Jabado^{94,95}, David Malkin^{3,96}, Olivier Avrault⁷, Jeffrey A. Golden⁹⁷, David W. Ellison⁹⁸, Brad Doble⁸, Vijay Ramaswamy^{1,2,3,96}, Tamra E. Werbowetski-Ogilvie^{6,99,#}, Hiromichi Suzuki^{9,#}, Kathleen J. Millen^{4,#}, Michael D. Taylor^{1,2,3,5,23,#}

Affiliations

- ^{1.}The Arthur and Sonia Labatt Brain Tumor Research Centre, The Hospital for Sick Children, Toronto, Ontario, Canada
- ² Developmental & Stem Cell Biology Program, The Hospital for Sick Children, Toronto, Ontario, Canada
- 3. Department of Medical Biophysics, University of Toronto, Toronto, Ontario, Canada
- ⁴·Center for Integrative Brain Research, Seattle Children's Research Institute, Seattle, Washington, United States
- ^{5.}Department of Laboratory Medicine and Pathobiology, University of Toronto, Toronto, Ontario, Canada
- ⁶ Department of Biochemistry and Medical Genetics, University of Manitoba, Winnipeg, Manitoba, Canada

⁷·PSL Research University, Université Paris Sud, Université Paris-Saclay, CNRS UMR 3347, INSERM U1021, Institut Curie, Orsay, France

- ⁸ Department of Pediatrics and Child Health & Department of Biochemistry and Medical Genetics, Rady Faculty of Health Sciences, University of Manitoba, Winnipeg, Manitoba, Canada
- ^{9.}Division of Brain Tumor Translational Research, National Cancer Center Research Institute, Tokyo, Japan
- ^{10.}Department of Molecular Genetics, University of Toronto, Toronto, Ontario, Canada
- ¹¹·Computational Biology Program, Ontario Institute for Cancer Research, Toronto, Ontario, Canada
- ¹² Princess Margaret Cancer Centre, University Health Network, Toronto, Ontario, Canada
- ¹³·Canada's Michael Smith Genome Sciences Centre, BC Cancer Agency, Vancouver, British Columbia, Canada
- ¹⁴.McGill University Genome Centre, McGill University, Montreal, Quebec, Canada
- ¹⁵·Division of Pediatric Neurooncology, German Cancer Research Center (DKFZ), Heidelberg, Germany
- ¹⁶ Department of Pediatric Oncology, Hematology, Immunology and Pulmonology, University Hospital Heidelberg, Heidelberg, Germany
- ¹⁷ Department of Neurosurgery, Division of Pediatric Neurosurgery, Seoul National University Children's Hospital, Seoul, South Korea
- ^{18.}SIREDO Oncology Center (pediatric, adolescent and young adults oncology), Institut Curie, Paris, France
- ^{19.}INSERM U830, Institut Curie, Paris, France
- ^{20.}Université Paris Cité, Paris, France
- ²¹.Unit of Somatic Genetics, Institut Curie, Paris, France
- ^{22.}Department of Pathology, The Children's Memorial Health Institute, Warsaw, Poland
- ²³ Division of Neurosurgery, The Hospital for Sick Children, Toronto, Ontario, Canada
- ²⁴·INSERM U1028, CNRS UMR5292, Centre de Recherche en Neurosciences, Université de Lyon, Lyon, France
- ²⁵·Centre de Pathologie EST, Groupement Hospitalier EST, Université de Lyon, Bron, France
- ²⁶ Department of Neurology, Erasmus University Medical Center, Rotterdam, Netherlands

²⁷·Department of Pathology, Erasmus University Medical Center, Rotterdam, Netherlands

- ²⁸ Department of Pediatric Oncology, Masaryk University School of Medicine, Brno, Czech Republic
- ^{29.}Department of Surgery, Division of Thoracic and Upper Gastrointestinal Surgery, Faculty of Medicine, McGill University, Montreal, Quebec, Canada
- ^{30.}Cancer Research Program, Research Institute of the McGill University Health Centre, Montreal, Quebec, Canada
- ³¹ Departments of Pathology, Ophthalmology and Oncology, John Hopkins University School of Medicine, Baltimore, Maryland, United States
- ³² Division of Pediatric Hematology/Oncology, Mayo Clinic, Rochester, Minnesota, United States
- ³³ Department of Laboratory Medicine and Pathology, Mayo Clinic, Rochester, Minnesota, United States
- ³⁴·Clinical Research Division, Fred Hutchinson Cancer Research Center, Seattle, Washington, United States
- ³⁵·2nd Department of Pediatrics, Semmelweis University, Budapest, Hungary
- ³⁶ Department of Neurological Surgery, University of California San Francisco, San Francisco, California, United States
- ³⁷·Department of Pathology, University of California San Francisco, San Francisco, California, United States
- ^{38.}Department of Neurosurgery, University of Ulsan, Asan Medical Center, Seoul, South Korea
- ³⁹ Developmental Tumor Biology Laboratory, Hospital Sant Joan de Déu, Esplugues de Llobregat, Barcelona, Spain
- ⁴⁰ Department of Anatomical and Cellular Pathology, The Chinese University of Hong Kong, Shatin, New Territories, Hong Kong
- ⁴¹ Department of Surgery, The Chinese University of Hong Kong, Shatin, New Territories, Hong Kong
- ⁴² Department of Neurological Surgery, University of Pittsburgh School of Medicine, Pittsburgh, Pennsylvania, United States
- ⁴³·Division of Pediatric Hematology/Oncology, Hospital Pediatría Centro Médico Nacional century XXI, Mexico City, Mexico
- ⁴⁴·Department of Neurosurgery, University of Alabama at Birmingham, Birmingham, Alabama, United States
- ⁴⁵·Pediatrics, Virginia Commonwealthy University, School of Medicine, Richmond, Virginia, United States

⁴⁶·Division of Stem Cell Research, Institute for Clinical Research, Osaka National Hospital, Osaka, Japan

- ⁴⁷ Department of Pediatrics, University of Colorado Denver, Aurora, Colorado, United States
- ⁴⁸ Department of Neurological Surgery, Vanderbilt Medical Center, Nashville, Tennessee, United States
- ^{49.}Department of Neurology, Vanderbilt Medical Center, Nashville, Tennessee, United States
- ⁵⁰Departments of Neuroscience, Washington University School of Medicine in St. Louis, St. Louis, Missouri, United States
- ⁵¹·Department of Neurosurgery, Kitasato University School of Medicine, Sagamihara, Kanagawa, Japan
- ⁵²·Department of Neurosurgery, Chonnam National University Research Institute of Medical Sciences, Chonnam National University Hwasun Hospital and Medical School, Hwasun-gun, Jeollanam-do, South Korea
- ⁵³ Department of Pathology and Molecular Medicine, Division of Anatomical Pathology, McMaster University, Hamilton, Ontario, Canada
- ⁵⁴·Department of Pathology and Laboratory Medicine, Hamilton General Hospital, Hamilton, Ontario, Canada
- ⁵⁵·Dipartimento di Medicina Molecolare e Biotecnologie Mediche (DMMBM), University of Naples Federico II, Naples, 80131, Italy
- ⁵⁶.CEINGE Biotecnologie Avanzate, Naples, 80145, Italy
- ⁵⁷·Department of Pediatric Neurosurgery, Santobono-Pausilipon Children's Hospital, Naples, Italy
- ⁵⁸ Division of Pediatric Neurosurgery, Case Western Reserve, Cleveland, Ohio, United States
- ^{59.}Department of Pediatrics-Hematology and Oncology, Case Western Reserve, Cleveland, Ohio, United States
- ^{60.}Department of Hematology & Medical Oncology, Emory University School of Medicine and Winship Cancer Institute, Atlanta, Georgia, United States
- ⁶¹ Fondazione IRCCS Istituto Nazionale Tumori, Milan, Italy
- ⁶² Department of Surgery and Anatomy, Faculty of Medicine of Ribeirão Preto, University of São Paulo, São Paulo, São Paulo, Brazil
- ⁶³ Division of Neurosurgery, Centro Hospitalar Lisboa Norte (CHULN), Hospital de Santa Maria, Lisbon, Portugal
- ⁶⁴·Instituto de Medicina Molecular João Lobo Antunes, Faculdade de Medicina, Universidade de Lisboa, Lisbon, Portugal

- ⁶⁵ Department of Tumor Cell Biology, St. Jude Children's Research Hospital, Memphis, Tennessee, United States
- ⁶⁶.Charbonneau Cancer Institute, University of Calgary, Calgary, Alberta, Canada
- ⁶⁷ Brotman Baty Institute for Precision Medicine, Seattle, Washington, United States
- ⁶⁸·Assistance Publique Hôpitaux de Paris, Hôpital Necker-Enfants Malades, Paris, France
- ⁶⁹ Surgical Pathology Unit, San Camillo Forlanini Hospital, Rome, Italy
- ⁷⁰. Department of Pathology, Duke University, Durham, North Carolina, United States
- ⁷¹ Department of Neurosurgery, Duke University, Durham, North Carolina, United States
- ⁷²·Cansearch Research platform for pediatric oncology and hematology, Faculty of Medicine, Department of Pediatrics, Gynecology and Obstetrics, University of Geneva, Geneva, Switzerland
- ⁷³ Division of Pediatric Oncology and Hematology, Department of Women, Child and Adolescent, University Geneva Hospitals, Geneva, Switzerland;
- ⁷⁴·U.O. Neurochirurgia, Istituto Giannina Gaslini, Genova, Italy
- ⁷⁵ Department of Neurosurgery, Hospital Infantil de Mexico Federico Gomez, Mexico City, Mexico
- ⁷⁶ Instituto Nacional De Pediatría de México, Mexico City, Mexico
- ⁷⁷ Department of Biochemistry & Molecular Biology, Cumming School of Medicine, University of Calgary, Calgary, Alberta, Canada
- ⁷⁸ Alberta Children's Hospital Research Institute, Calgary, Alberta, Canada
- ⁷⁹ INSERM U900, Institut Curie, Paris, France
- ^{80.}PSL Research University, Institut Curie, Paris, France
- ⁸¹·CBIO-Centre for Computational Biology, PSL Research University, MINES ParisTech, Paris, France
- 82. UPMC Hillman Cancer Center, Pittsburgh, PA, United States
- ⁸³ Department of Medical Genetics, University of British Columbia, Vancouver, British Columbia, Canada
- ^{84.}Department of Molecular Biology & Biochemistry, Simon Fraser University, Burnaby, British Columbia, Canada
- ⁸⁵ Department of Pathology and Laboratory Medicine, University of British Columbia, Vancouver, British Columbia, Canada
- ⁸⁶ Department of Molecular Oncology, BC Cancer Agency, Vancouver, British Columbia, Canada

⁸⁷·Tumor Initiation and Maintenance Program, NCI-Designated Cancer Center, Sanford Burnham Prebys Medical Discovery Institute, La Jolla, California, United States

- ^{88.}Department of Neurology, University of California San Francisco, San Francisco, California, United States
- ^{89.}Department of Pediatrics, University of California San Francisco, San Francisco, California, United States
- ⁹⁰·Cancer Research Program, McGill University Health Centre Research Institute, Montreal, Quebec, Canada
- ^{91.}Department of Human Genetics, McGill University, Montreal, Quebec, Canada
- ⁹²·Lady Davis Research Institute, Jewish General Hospital, Montreal, Quebec, Canada
- ^{93.}Adaptive Oncology, Ontario Institute for Cancer Research, Toronto, Ontario, Canada
- ^{94.}Departments of Pediatrics and Human Genetics, McGill University, Montreal, Quebec, Canada
- ^{95.}The Research Institute of the McGill University Health Center, Montreal, Quebec, Canada
- ⁹⁶·Division of Haematology / Oncology, Department of Pediatrics, The Hospital for Sick Children, Toronto, Ontario, Canada
- ⁹⁷·Department of Pathology, Cedars-Sinai Medical Center, Los Angeles, California, United States
- ⁹⁸ Department of Pathology, St. Jude Children's Research Hospital, Memphis, Tennessee, United States
- ⁹⁹ CancerCare Manitoba Research Institute, Winnipeg, MB, Canada.
- ¹⁰⁰.Deceased.

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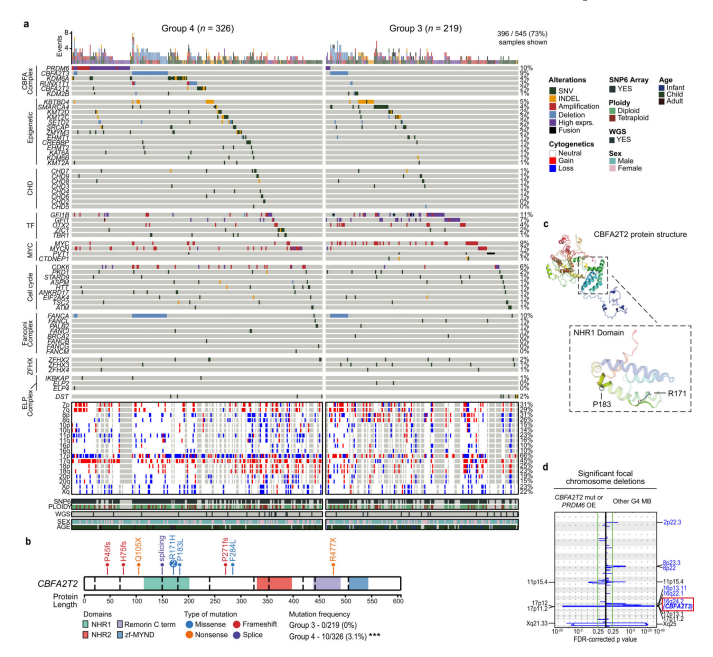


Fig.1 |. The landscape of oncogenic drivers in G3 and G4 MB.

a, Oncoprint summary of mutations, copy number variations, gene expression, and gene fusions in G3 and G4 MB (n = 545 tumors). 173 samples without recurrent alterations detected are not shown. **b,** Gene-level summary of *CBFA2T2* mutations in G4 MB. **c,** Structural model of CBFA2T2 protein NHR1 domain, highlighting positions affected by missense mutations. The structure of the NHR1 domain has been previously determined (Bottom), while the full protein structure was inferred using iTasser³⁹ (Top). **d,** Comparison of significant focal deletions in n = 206 G4 MB, either with *CBFA2T2* mutations, or *PRDM6* overexpression, versus tumors without *CBFA2T2* or *PRDM6* abnormalities. Significance assessed using GISTIC 2.0⁴⁰ (FDR < 0.25).

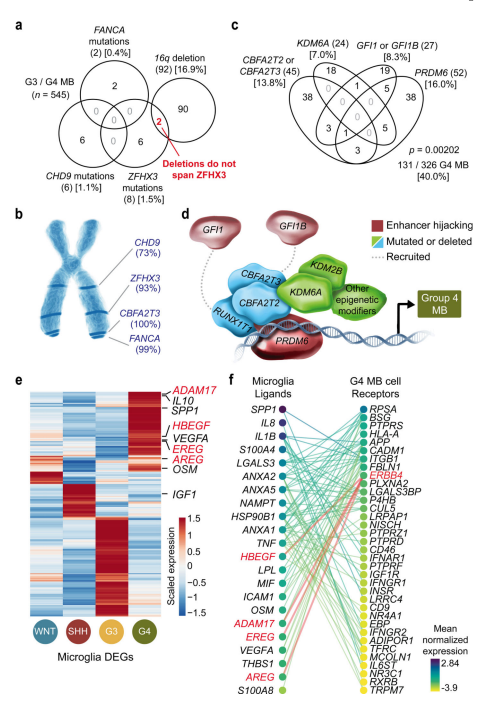


Fig.2 |. CBFA complex members are recurrently somatically mutated in G4 MB. **a,** Mutual exclusivity (ME) of somatic SNVs in 16q genes *FANCA*, *ZFHX3*, and *CHD9*.

Deletions of 16q do not typically co-occur with mutations in these genes. G3 and G4

MB prefer deletions to simultaneously disrupt several tumor suppressor genes (TSGs). **b,**Locations of G4 MB TSGs on chr16 and percentage of samples where TSGs are deleted to haploinsufficiency in G3 and G4 MB samples with 16q deletions. **c,** ME of *PRDM6* overexpression, *CBFA2T2* mutations, *CBFA2T3* deletions, *KDM6A* mutations, and *GFI1* or *GFI1B* enhancer hijacking in G4 MB. Significance assessed using the impurity test for

ME implemented by DISCOVER⁴¹. **d,** Cartoon of known or suspected CBFA complex members. **e,** Expression of significant differentially expressed genes (DEGs) between microglia from each subgroup using scRNAseq (MAST⁴², FDR < 0.05). **f,** Predicted receptor-ligand interactions between microglia and tumor cells in G4 MB (CCInx⁴³). Node colours represent mean normalized gene expression in each cell type, and edge colour represents the average of the node expression levels. Only significant DEGs from **g** were included, demonstrating the microenvironment specificity of G4 MB.

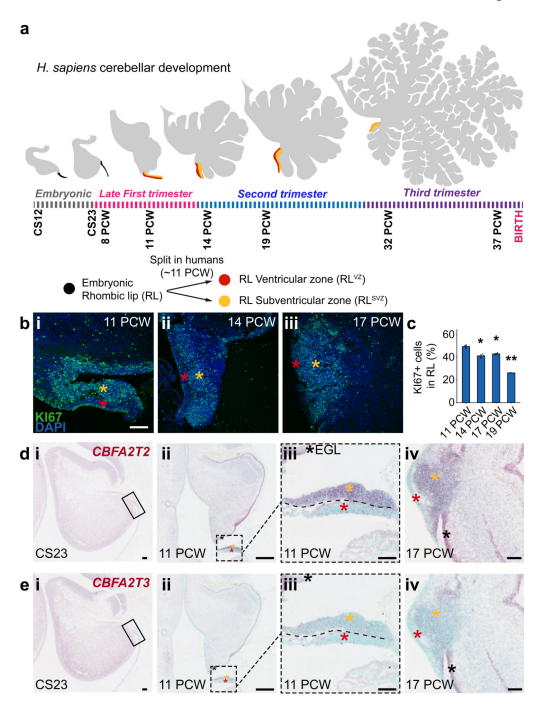


Fig.3 |. $\it CBFA2T2$ and $\it CBFA2T3$ define the RL VZ / SVZ boundary in the developing human cerebellum.

a, Schematic summarizing *Homo sapiens* cerebellar development. Prior to 11 PCW, the RL resembles that of *Mus musculus*. Around 11 PCW the human RL splits into a ventricular and subventricular zone leading to human-specific cerebellar expansion. **b,** KI67 expression in the developing human RL at 11, 14 and 17 PCW. **c,** Percent of human RL cells expressing KI67 across several developmental timepoints. Significance assessed using a paired two-tailed t-test compared to 11 PCW, ** p < 0.005, * p < 0.05. 14PCW, p = 0.0072; 17PCW,

p=0.012; 19PCW, p=0.0024. n=3 biological repeats per time point; error bars, SEM. **d**, **e**, *In situ* hybridization (ISH) showing spatially resolved RNA expression of *CBFA2T2* (**d**) and *CBFA2T3* (**e**) in the developing human cerebellum at CS23, 11, and 17 PCW. *CBFA2T2* and *CBFA2T3* expression is first observed at 11 PCW in the RL^{SVZ} but not the RL^{VZ}. Data presented in **b** are representative images from three independent experiments with similar results, data in **d** and **e** were not performed in replicates. RL^{VZ}, RL^{SVZ}, and EGL are denoted by red, yellow, and black asterisks, respectively. Scale bars: 100 µm.

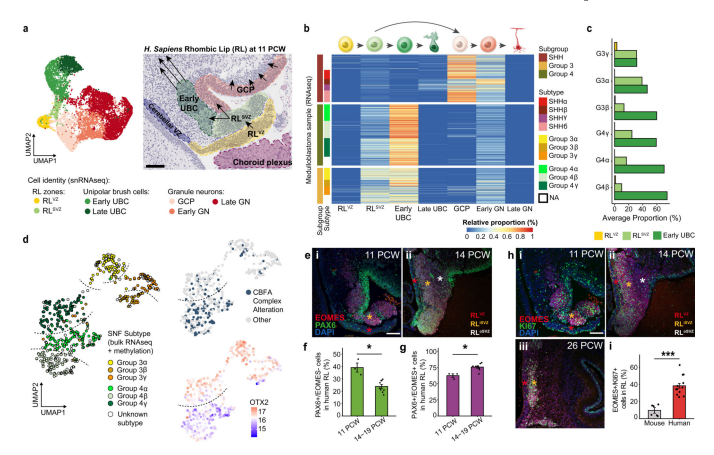


Fig.4 |. The expanded human RL is uniquely predisposed to G3 and G4 MB.

a, (Left) UMAP embedding of 9,208 single cells from the developing human cerebellum¹³. (Right) Midsagittal H&E section from 11PCW human cerebellum highlighting expected distribution of indicated cell types. b, Deconvolution of RNAseq expression profiles from G3 (n = 219), G4 (n = 326), and SHH (n = 250) MB by cell types in **a. c**, Deconvolution proportions along the RL to UBC lineage in G3 and G4 MB subtypes. d, UMAP embedding of G3 and G4 MB RNAseq samples (of n = 545), showing subtype, OTX2 expression, and CBFA complex alterations. e, EOMES and PAX6 expression in human RL at 11 and 14 PCW. **f**, **g**, Percent of RL cells expressing PAX6 only (**f**), *p = 0.036 or both PAX6 and EOMES (g), *p = 0.017. Significance assessed using unpaired two-tailed t-tests; n =3 biological repeats per time point; error bars, SEM. h, EOMES and KI67 expression at 11, 14, and 26 PCW. i, EOMES+/KI67+ population in humans and mice across timepoints. Significance assessed by two-sided Mann-Whitney U test, ***p = 0.000108. n = 3 biological repeats per time point; error bars, SEM. Data presented in e, h (11 and 14 PCW) are representative images from three independent experiments with similar results, data in a, **h** (26 PCW) were not performed in replicates. The RL^{VZ}, inner subventricular (RL^{iSVZ}), and outer subventricular (RLoSVZ) zones indicated with red, yellow, and white asterisks, respectively. Scale bars: 100 µm.

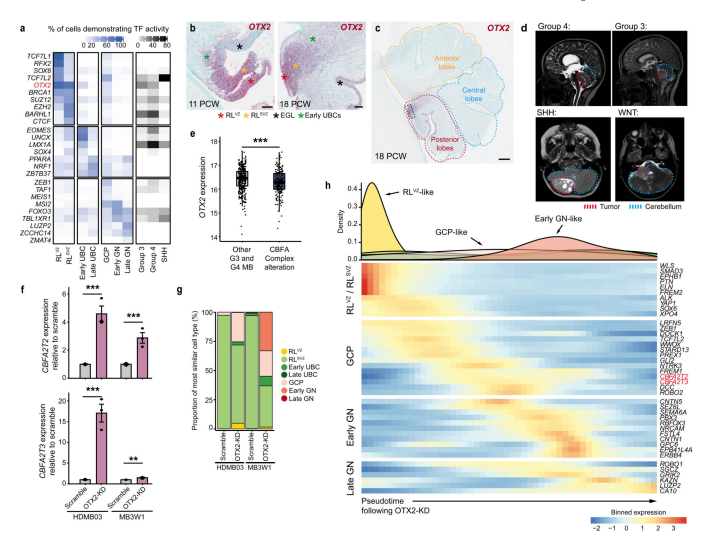


Fig.5 |. OTX2 restrains RL differentiation through inhibition of the CBFA complex.

a, Percentage of cells in normal cell types and MB cells which exhibit predicted TF activity (SCENIC⁴⁴). **b,** OTX2 expression in the developing human RL at 11 and 18 PCW. Scale bars: 100 µm. c, OTX2 expression at 18 PCW. High expression is observed in the RL (black box) and the posterior lobes. Scale bar: 500 µm. d, Representative T1 enhanced or T2 MRI scans showing typical locations of each MB subgroup at initial diagnosis. Mid-sagittal scans are shown for G3 and G4 MB, axial scans for SHH and WNT MB tumors. e, OTX2 expression in G3 and G4 MB samples by CBFA complex alterations (Fig. 4d). Significance assessed using two-sided Mann-Whitney U test, *** p = 0.0000162, n = 545 MBs. Box plots show the median and interquartile range, and whiskers show the data range. Points outside this range are outliers and are plotted individually. f, CBFA2T2 and CBFA2T3 expression following OTX2 knockdown in G3 MB tumorspheres. Data shown as mean fold change \pm SEM compared to scramble, with n=3 biological replicates. Significance assessed using DESeq2⁴⁵ (FDR < 0.05), *** p < 0.0005, ** p < 0.005. *CBFA2T2*: HDMB03, p = $1.20e^{-30}$; MB3W1, $p = 4.45e^{-15}$. CBFA2T3: HDMB03, $p = 2.90e^{-91}$; MB3W1, p = 0.0055. g, Single-cells show increased similarity to normal differentiated cell types upon OTX2-KD in G3 MB tumorspheres. h, Expression of granule neuron (GN) lineage marker genes along

pseudotime following OTX2-KD in MB3W1. (Top) Density of most similar developmental cell type (\mathbf{g}) along pseudotime. (Bottom) Binned expression of GN differentiation genes (Fig.4a). Data presented in \mathbf{b} , \mathbf{c} were not performed in replicates.

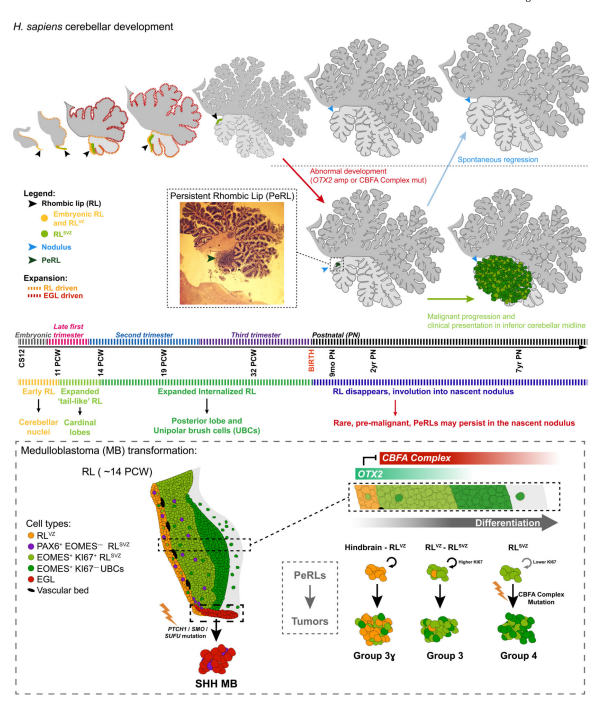


Fig.6 \mid G3 and G4 MB initiate in $\it utero$ from failed differentiation of components of the human RL.

Proposed model of the initiation of SHH, G3, and G4 MB. (Top) Normal human cerebellum development is marked by changing roles and shrinking of the RL until it dissipates within the nodulus around or shortly after birth. Rarely, this process can be disrupted, leaving residual disorganized RL termed PeRL, which appears as a non-neoplastic malformation (dyplasia) in the nodulus. (Bottom) Following the split of the human RL around 11 PCW, EOMES⁺ KI67⁺ RL^{SVZ} progenitors are born, which in the presence of chromosomal instability and/or driver alterations, fail to differentiate properly and give rise to PeRLs

which in the presence of further genetic insult give rise to G3 and G4 MB in the inferior cerebellum. PN, postnatal.