Wnt activation as a therapeutic strategy in medulloblastoma

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Medulloblastoma (MB) is defined by four molecular subgroups (Wnt, Shh, Group 3, Group 4) with Wnt MB having the most favorable prognosis. Since prior reports have illustrated the antitumorigenic role of Wnt activation in Shh MB, we aimed to assess the effects of activated canonical Wnt signaling in Group 3 and 4 MBs. By using primary patient-derived MB brain tumor-initiating cell (BTIC) lines, we characterize differences in the tumor-initiating capacity of Wnt, Group 3, and Group 4 MB. With single cell RNA-seq technology, we demonstrate the presence of rare Wnt-active cells in non-Wnt MBs, which functionally retain the impaired tumorigenic potential of Wnt MB. In treating MB xenografts with a Wnt agonist, we provide a rational therapeutic option in which the protective effects of Wnt-driven MBs may be augmented in Group 3 and 4 MB and thereby support emerging data for a context-dependent tumor suppressive role for Wnt/β-catenin signaling.
MDM2 by IHC (Supplementary Fig. 6a). Using a similar approach, a conserved downstream Wnt target gene was noted in SU_MB002 cells cultured in Wnt3A-conditioned medium. A significant increase in Axin2 expression, a conserved downstream Wnt target gene, was noted in SU_MB002 cells cultured in Wnt3A-conditioned medium (Supplementary Fig. 5a). Wnt3A-mediated pathway activation also reduced secondary tumor sphere size (Supplementary Fig. 5b–c), frequency (Supplementary Fig. 5d), and cell proliferation in non-Wnt MBs (Supplementary Fig. 5e). These findings support a transient alteration in the oncogenic phenotype of aggressive MBs following Wnt activation.

Since developmental processes may be carried forward from ontogeny into oncology, we investigated their clinical utility in a cohort of 377 Group 3 and 4 MBs. Using a Bmi1 gene signature that predicts metastasis, tumor progression, and death from cancer, we found a reduced overall survival advantage compared with Group 3 MBs for xenografts generated from both 1.0 × 10^5 (Supplementary Fig. 4g) and 5.0 × 10^5 cells (Supplementary Fig. 4h). The observed in vivo growth differences are most likely reflective of the in vitro differences in proliferation and self-renewal between Wnt and Group 3 MBs. These findings support the clinical observations of prognostic variations among MB subgroups.

**Methods**

**MB BTICs retain subgroup affiliation.** To assess the biological validity of our MB BTIC model, we asked if gene expression differences between subgroups in bulk MB manifested themselves in our model. We performed differential expression analysis from bulk MB data to identify upregulated genes specific to each subgroup, and scored both the bulk MB data and our MB BTIC lines for these upregulated gene expression signatures using single-sample gene set enrichment analysis (ssGSEA). As expected, upregulated genes associated with Wnt (n = 6326), Group 3 (n = 7491), and Group 4 (n = 6510) showed strongest positive enrichment in bulk Wnt MB, Group 3, and Group 4 MB samples, respectively, validating the specificity of the signatures (Supplementary Fig. 1a). Interestingly, the Wnt, Group 3, and Group 4 MB signatures showed strongest relative enrichment in our Wnt MB line (BT853), Group 3 (SU_MB002), and Group 4 (ICB1299) MB BTIC lines, respectively (Supplementary Fig. 1b), showing preservation of subgroup affiliation among our MB BTIC lines.

Using principal component analyses (PCA), we examined the inter-subgroup heterogeneity between MB BTIC lines from Wnt, Group 3, and Group 4 MBs (Supplementary Fig. 2a). RNA-seq data from these lines yielded distinct gene expression profiles between our Wnt, Group 3 and, Group 4 MB lines (Supplementary Fig. 2b–e, Supplementary Data 1). Specifically, several genes implicated in self-renewal of malignant stem cell populations and previously described by our group to be enriched in poor-outcome MBs were more highly expressed in Group 3 and 4 MBs when compared with Wnt MB (Supplementary Fig. 2f). Bmi1 and Sox2 are hallmark master regulatory stem cell genes essential to BTIC self-renewal, and thus enriched in experimental BTIC models. Gene set enrichment analysis (GSEA) of non-Wnt MB (Group 3 MB line (Supplementary Fig. 3a) and Group 4 MB line (Supplementary Fig. 3b)) showed enrichment of genes in the Bmi1 signature, which function as key epigenetic regulators of fate determination and self-renewal in normal and malignant cerebellar stem cells. By contrast, cell cycle checkpoint and apoptosis gene signatures were more active in Wnt MB lines (Supplementary Fig. 3b) when compared with Group 4 MBs. No differences were identified in cell cycle checkpoint and apoptosis gene signatures between Wnt and Group 4 MB lines. Pathway network analysis further identified an increase in DNA replication, transcriptional regulation, ribosomal processing, and translational regulation in Group 3 and 4 MB lines (Supplementary Fig. 3c, Supplementary Data 2), suggestive of a hyperproliferative state.

Additional differences between Wnt, Group 3, and Group 4 MBs were determined using in vitro and in vivo tumorigenic assays. TCF reporter assays showed a significant increase in endogenous Wnt activity in Wnt compared with Group 3 and 4 MB lines (Supplementary Fig. 3d). Similar to the Wnt-mediated inhibition of cerebellar stem cell self-renewal, proliferation (Supplementary Fig. 4a–b) and self-renewal (Supplementary Fig. 4c–d) were impaired in our Wnt MB line (BT853) when compared with Shh (Daoy), Group 3 (SU_MB002, HD-MB03, D425, D458), and Group 4 (ICB1299) MB lines. It should be noted that the Daoy cell line contains a homozygous TP53 deletion, which is not seen in Shh MB, and results must therefore be reviewed in context for data with the Daoy cell line. Xenografts generated with 5.0 × 10^5 Wnt MB cells led to tumor formation in 4/5 mice, whereas no tumors formed with 1.0 × 10^5 cells (0/5 mice). By contrast, all Group 3 MB xenografts formed tumors at both numbers (10/10 mice) (Supplementary Fig. 4e). Overall tumor volume was higher in Group 3 than Wnt MB xenografts (Supplementary Fig. 4f). Wnt MBs also contained a marked survival advantage compared with Group 3 MBs for xenografts generated from both 1.0 × 10^5 (Supplementary Fig. 4g) and 5.0 × 10^5 cells (Supplementary Fig. 4h). The observed in vivo growth differences are most likely reflective of the in vitro differences in proliferation and self-renewal between Wnt and Group 3 MBs. These findings support the clinical observations of prognostic variations among MB subgroups.
signature\(^{21}\) identified Sox2\(^{\text{high}}\) patients to have a significant reduction in overall survival (Supplementary Fig. 6b). To further understand factors associated with patient outcome, we performed univariate and multivariate survival analyses, which included the index, age, metastatic status, and subgroup. Bmi1 signature genes (Fig. 1a, Supplementary Table 1) showed a significant association with patient outcome in a univariate analysis (\(p = 0.0113\)), which remained significant (\(p = 0.0287\)) when adjusted for age, metastatic status, and subgroup. Metastatic status alone was not significantly associated with patient outcome in both univariate and multivariate analyses, but it was significantly associated with the index (\(p = 0.03339\)). The Sox2 signature index (Fig. 1a, Supplementary Table 2) showed a significant association with patient survival in univariate analysis (\(p = 0.00441\)) but became a trend when adjusted for age, metastatic status, and subgroup for the multivariate analysis (\(p = 0.0924\)). An association was noted between metastatic status and subgroup affiliation (\(p = 0.03888, p = 0.02943\) respectively). The expression of both self-renewal genes Bmi1 and Sox2 were reduced in Wnt MB BTICs (BT853) when compared with Shh (Daoy), Group 3 (SU_MB002, HD-MB03, RCMB40, D425, Med8A, D458) and Group 4 (ICB1299) MB BTICs (Fig. 1b and Supplementary Figs. 7 and 8), indicating differences in self-renewal activity across MB subgroups.

Considering that CTNNB1, which encodes β-catenin, is mutated in 86% of Wnt MBs, we ectopically expressed a stabilized β-catenin mutant in Group 3 and 4 MB lines. Axin2 expression was used to determine sufficient overexpression (Fig. 1c, d). Bmi1 and Sox2 were reduced following β-catenin overexpression (Fig. 1c, d). The self-renewal capacity of β-catenin-overexpressing lines was markedly reduced when compared with controls (Fig. 1e); it should be noted that no direct comparison was made to native Wnt models in these experiments. Orthotopic injections of Group 3 MB cells revealed a significant reduction in overall tumor burden (Fig. 1f, Supplementary Fig. 9) and increase in survival (Fig. 1g) following the ectopic expression of β-catenin.

Non-Wnt MBs contain Wnt-active cells that are less tumorigenic. While genetic heterogeneity exists within individual MB subgroups\(^{10}\), general subgroup affiliation does not change at recurrence\(^{22}\), metastasis\(^{23}\), or within different regions of a tumor\(^{24}\). However, much of this work has been done on bulk tumor samples without considering subclonal variations observed at a single-cell level. Given that emerging data in other malignancies have implicated subclonal genetic drivers in disease progression and relapse\(^{25,26}\), we performed single-cell RNA-seq on Wnt, Group 3, and Group 4 MB lines (Fig. 2a, Supplementary Figs. 10–14) to understand the functional relevance of subclonal Wnt activity in MB cells. As expected, cells from the Wnt MB line had the highest enrichment for the Wnt subgroup gene signature (Fig. 2b–d). Intriguingly, we found that a small population of cells in Group 3 and 4 MB lines exhibited high Wnt subgroup signature scores, similar to cells of Wnt subgroup origin (Fig. 2b). These results support the presence of a rare population of cells in Group 3 and 4 MBs that resemble Wnt subgroup cells. We validated our findings in four independent datasets consisting of scRNA-seq\(^{27,28}\) (Supplementary Figs. 15 and 16) and bulk tumor RNA-seq data\(^{1,10}\) (Supplementary Figs. 17 and 18). We then wondered if these Wnt-active cells maintained the reduced tumorigenic potential observed in Wnt MB. Using a lentiviral Wnt reporter (7XTCF-GFP)\(^{29}\), we identified rare Wnt-active cells (TGP+) from Group 3 MB cells (Fig. 3). Both Wnt-active (TGP+) and Wnt-inactive (TGP−) cells were isolated using flow cytometric cell sorting (Supplementary Fig. 1a–b). When TGP+ and TGP− cells were cultured in vitro, we observed TGP+ convert to TGP− cells and vice versa, but the majority of them repopulated to TGP− cells (Supplementary Fig. 19c). Wnt activity in TGP+ cells was validated using the TCF Wnt reporter assay (Fig. 3a) and Axin2 transcript levels (Fig. 3b). TGP+ cells contained reduced levels of Bmi1 and Sox2 transcript (Fig. 3b) and protein expression (data not shown) when compared with TGP− cells. Decreased proliferative (Fig. 3c) and self-renewal (Fig. 3d) indices were also noted in TGP+ compared with TGP− cells. Separate xenografts generated from orthotopic injections of TGP+ or TGP− cells revealed a reduced tumor burden (Fig. 3e, Supplementary Fig. 20a–b) and enhanced survival (Fig. 3f) in TGP+ xenografts compared with TGP−, with some TGP+ cells reverting back to TGP− cells as observed by immunostaining for GFP positivity in these xenografts (Fig. 3g). These data highlight the intrinsic tumor suppressive role of activated Wnt signaling in MB. To examine the clinical utility of our model systems, we used the Wnt hallmark gene signature to probe a clinically annotated dataset of 113 Group 3 human MBs\(^{10}\) for differences in overall survival. Group 3 patients with high expression of the Wnt hallmark signature maintained a longer overall survival compared with those with low expression (Fig. 3h). As a result, endogenous Wnt activity in Group 3 MBs appears predict improved survival in MBs that are otherwise metastatic and refractory to current treatment. To further understand factors associated with patient outcomes, we performed univariate and multivariate survival analyses, which included the index, age and metastatic status. The index based on the Wnt signature (see Supplementary Table 3) was the only factor that showed a significant association with patient survival in univariate analysis (\(p = 0.0158\)) and when adjusted for other factors, such as age and metastatic status (\(p = 0.0103\)).

Pharmacological Wnt activation in MB xenografts improves survival. To develop a therapeutic strategy employing Wnt activation, we initially used the small molecule GSK-3 inhibitor, CHIR99021, which acts as a competitive inhibitor of ATP binding\(^{30}\). GSK-3 functions to inhibit Wnt/β-catenin signaling by phosphorylating the downstream Wnt effector, β-catenin, rendering it nonfunctional. TCF reporter measured Wnt activation following exposure to CHIR99021 in Group 3 and 4 MB lines (Supplementary Fig. 21a–b). Axin2 transcript levels (Supplementary Fig. 21c–d) and increase in nuclear and cytoplasmic β-catenin levels (Supplementary Fig. 22) provided further validation of Wnt activation. Bmi1 and Sox2 expression were decreased following CHIR99021 treatment (Supplementary Fig. 21c–d). Although CHIR99021 also reduced the proliferative and self-renewal abilities (Supplementary Fig. 21e–f) of Group 3 and 4 MB lines, we were unable to proceed to preclinical studies due to limited blood–brain barrier permeability.

In an attempt to find a suitable preclinical molecule, we identified L807mts, a GSK-3 inhibitor that functions through a substrate-to-inhibitor conversion mechanism within the GSK-3 catalytic site\(^{31}\). TCF reporter activity confirmed Wnt activation in L807mts-treated cells (Supplementary Fig. 23a–b), which was further corroborated with Axin2 expression (Fig. 4e, f) and β-catenin expression (Supplementary Fig. 22). PCA was used to examine changes in the transcriptional machinery following L807mts treatment (Fig. 4a). RNA-seq data from L807mts-treated cells showed diverging gene expression profiles (Fig. 4b, Supplementary Fig. 23c–e, Supplementary Data 2). In order to assess the activation of Wnt by L807mts treatment, we examined enrichment of the top 50 and top 100 signatures in the samples treated with L807mts relative to the matching samples treated with PBS. Enrichment analysis was performed
Fig. 1 Ectopic expression of β-catenin in treatment-refractory MBs reduces self-renewal gene expression and tumor burden while increasing overall survival. a Multivariate survival analysis including index, age, metastatic status, and subgroup shows significantly associated with patient survival using the Bmi1 (p = 0.0004774) and Sox2 (p = 0.00193) signatures. b Bmi1 and Sox2 protein levels are significantly reduced in Group 3 (HD-MB03, D425) MB lines when compared with Wnt MB (BT853) (n = 3, independent experiments per MB line, 1 line per subgroup). Differential Axin2 (Group 3: p = 0.0014, Group 4: p = 0.001), Bmi1 (Group 3: p = 0.0018, Group 4: p = 0.000087), and Sox2 (Group 3: p = 0.0011, Group 4: p = 0.0012) transcript levels following β-catenin overexpression in (c) Group 3 and (d) Group 4 MB lines (n = 3, independent experiments per MB line, 1 line per subgroup, all samples normalized to GAPDH). e Tumor sphere formation is impaired following β-catenin overexpression in Group 3 (p = 0.0005) and Group 4 MBs (p = 0.0058) (n = 3, independent experiments per MB line, 1 line per subgroup). f Representative histology images showing significant reduction in tumor formation in xenografts generated from Group 3 β-catenin-overexpressing lines (n = 5) compared with control (n = 5). g β-catenin overexpression xenografts (n = 5, median survival 22.0 days) display a significant increase in overall survival when compared with control xenografts (n = 5, median survival 22.0 days) (p = 0.0043). Histology image scale bar = 5000 μm. Panels (c–e) contain error bars expressed as mean ± standard error (mean) using two-tailed, unpaired Student’s t test. Panel (g) analyzed using log-rank (Mantel-Cox) test. ****p < 0.0001.
with GSEA. Top genes upregulated in Wnt MBs were found to be enriched in corresponding Group 3 and 4 samples treated with L807mts. This provides supportive evidence for the observed activation of Wnt signaling in Group 3 and 4 MBs treated with L807mts (Supplementary Fig. 23f–g). The Bmi1 gene signature was also reduced in L807mts-treated samples as per GSEA (Fig. 4c). Additional GSEA identified an increase in the expression of apoptotic and G2M checkpoint-related gene signatures in L807mts-treated samples (Fig. 4c). Our single-cell RNA-seq data also revealed that several Wnt activation genes that were highly enriched in endogenous single cells from Group 3 and 4 MBs, including NFATC4 and GALNT14, were also upregulated in L807mts-treated Group 3 and 4 MB lines (NFATC4 fc = 4.33, GALNT14 fc = 2.43, Supplementary Data 2). These findings support the activation of Wnt-driven tumor suppressive pathways following L807mts treatment. Given the enhanced expression of apoptotic and cell cycle inhibition pathways in L807mts-treated cells and prior reports that have indicated TP53-mutated Wnt MBs to be more radiosensitive than TP53-mutated non-Wnt MBs, we investigated the potential to radiosensitize treatment-refractory MBs with L807mts. An increase in radiosensitization was observed with L807mts treatment in vitro (Fig. 4d) and in vivo (Supplementary Fig. 24). L807mts treatment also decreased Bmi1 and Sox2 (Fig. 4e, f) levels and impaired the proliferation and self-renewal (Fig. 4g, h) of Group 3 and 4 MB lines. Preclinical studies were performed on three separate patient-derived MB lines (Shh: BT992; Group 3: SU_MB002, RCMB040). A reduction in overall tumor burden (Fig. 5a–c, Supplementary Fig. 25a–b) and increase in overall survival (Fig. 5d, e, Supplementary Fig. 25c) were observed in xenografts treated with L807mts. These preclinical studies provide evidence in support of a highly specific small molecule Wnt agonist with potential clinical utility for treating MB.

Discussion
Current MB clinical trials have based risk-adapted therapy on molecular subgroups. Trials focused on Wnt-driven MB are deescalating chemo/radiotherapy (NCT01878617, NCT02724579, NCT02212574). By contrast, trials for Group 3 and 4 MBs are escalating therapy with the hope of improving survivorship with recurrent MB patients treated with palliation alone. Given the few targeted treatment options for Group 3 and 4 MBs, our work highlights a rational therapeutic option in which the protective effects of Wnt-driven MBs may be augmented in Group 3 and 4 MBs through targeted Wnt activation. While our more limited findings in Shh MB (Supplementary Figs. 4b, 4d, 7, 23) align with prior reports describing an antitumorigenic role for Wnt activation in murine models of Shh MB, the presence of endogenous...
Fig. 3 Rare endogenous Wnt-active MB cells from treatment-refractory MBs have reduced tumorigenic properties. a SU_MB002 cells transduced with a Wnt reporter show enhanced TCF reporter activity in TGP+ cells compared with TGP– cells (p = 0.0035) (n = 3, independent experiments). b Differential Axin2 (p = 0.0007), Bmi1 (p = 0.0005), and Sox2 (p = 0.0028) transcript levels in TGP+ and TGP– cells (n = 3, independent experiments, all samples normalized to GAPDH). c Proliferative (p = 0.0012) and d self-renewal capacity (p = 0.0031) of TGP+ cells are significantly reduced when compared with TGP– cells (n = 3, independent experiments). e Representative histology images illustrating reduced tumor burden in TGP+ (n = 10) compared with TGP– xenografts (n = 10). f TGP+ xenografts (n = 10, median survival 39.0 days) display a significant increase in overall survival compared with TGP– xenografts (n = 10, median survival 29.0 days) (p = 0.000012). g TGP+ tumors revert to TGP– as deduced by IHC for GFP expression. h Treatment-refractory MB patients (n = 113) with high expression of the Wnt hallmark signature have an improved overall survivorship (median survival undetermined) when compared with those patients with low Wnt hallmark signature expression (median survival 54.0 months) (p = 0.0126). Xenograft H/E histology image scale bar = 5000 μm. IHC histology image scale bar = 50 μm. Panels (a–d) contain error bars expressed as mean ± standard error (mean) using two-tailed, unpaired Student’s t test. Panels (f, h) analyzed using log-rank (Mantel-Cox) test. ****p < 0.0001.
Wnt activity in Shh MBs may confound the therapeutic effects of Wnt activation and thus requires further investigation. This work illustrates an emerging paradigm in Wnt biology in which functions of the pathway may be conserved in a context-dependent manner, such as a hindbrain-specific function in which self-renewal is impaired from ontogeny to oncology. However, before considering Wnt activation as a treatment paradigm in childhood MB, it is important to identify the mechanisms by which Wnt/β-catenin signaling function to target self-renewal genes that could impede growth of these aggressive MBs. As molecular oncology trials continue to develop, approaches to overcome the dependence of tumors on malignant pathways are warranted and as such we provide a therapeutic rationale with clinical appeal that may alter our approach to cancer by reactivating anti-oncogenic programs that have been developmentally silenced in a tissue-specific manner.
**Fig. 4** Pharmacological activation of Wnt signaling in MB impairs stem cell properties. 

- **a** Visualization of the first three principal components from PCA of Group 3 and 4 lines treated with small molecule Wnt activator (L807mts) or control (PBS) (n = 3, independent samples per MB line, 1 line per subgroup).
- **b** Heatmaps of differentially expressed genes between L807mts- and PBS-treated Group 3 (SU_MB002, left panel) and Group 4 (ICB1299, right panel) MB lines (n = 3, independent samples per MB line, 1 line per subgroup).
- **c** GSEA enrichment plots showing that Bmi1 associated genes are significantly reduced while apoptosis and cell cycle inhibitors are enriched in L807mts-treated cells compared with control.
- **d** L807mts-treated cells are much more radiosensitive than control cells (1 Gy p = 0.01145; 2 Gy p = 0.0013; 3 Gy p = 0.0004; 4 Gy p = 0.00008; 5 Gy p = 0.00004) (n = 3, independent experiments).
- **e** Transcript levels in L807mts-treated (SU_MB002: p = 0.0006, Group 3: p = 0.0008) and (f) Group 4 MB lines (RCMB040: p = 0.0003, Group 4: p = 0.0008) transcript levels in L807mts-treated (SU_MB002: p = 0.0008, Group 3: p = 0.0008) and (f) Group 4 MB lines (n = 3, independent experiments per MB line, 1 line per subgroup).

**Fig. 5** Pharmacological activation of Wnt signaling in MB reduces tumor formation and improves overall survival. 

- **a** Representative histology images of xenografts generated from patient-derived treatment-refractory MBs following L807mts treatment (n = 6 per MB line) contain a reduction in overall tumor burden when compared with xenografts treated with PBS control (n = 6 per MB line).
- **b** SU_MB002 (n = 12, p = 0.0001) and **c** RCMB40 (n = 12, p = 0.00001) treated with L807mts (n = 6, individual mice per MB line) contain a significant reduction in overall tumor volume when compared with PBS control-treated xenografts (n = 6, individual mice per MB line).
- **d** SU_MB002 (n = 12; median survival L807mts 49.5 days, PBS 32.0 days; p = 0.0008, n = 12), and **e** RCMB40 (n = 12; median survival L807mts 52.9 days, PBS 38.0 days; p = 0.0005) treated with L807mts display a significant survival advantage when compared with control PBS-treated mice. Histology image scale bar = 5000 μm.
Methods

Culturing primary MB samples. Primary human pediatric MBs, BT853, and BT992 were obtained from consenting patients and families as approved by the Hamilton Health Sciences/McMaster Health Sciences Research Ethics Board. BT853 is a primary Wnt subgroup MB, whereas BT992 is a primary Shh subgroup from a patient who later relapsed. Both samples were dissociated in PBS containing 0.2 Wünsch unit/ml of Liberase Blendzyme 3 (Roche), and incubated at 37 °C in a shaker water bath until dissociated tissue was aspirated through a 7.0 mm cell strainer and collected by centrifugation (450 g, 3 min). Tumor cells were resuspended in a serum-free BTIC enrichment medium, and replated on ultra-low attachment plates (Corning). Additional primary human pediatric MB cultures were obtained from collaborators as kind gifts. SU_MB002, a treatment-refractory Group 3 MB acquired at autopsy, was dissociated tissue was aspirated through a 7.0 mm cell strainer and collected by centrifugation (450 g, 3 min). Tumor cells were resuspended in a serum-free BTIC enrichment medium, and replated on ultra-low attachment plates (Corning).

Additional factors, such as age, metastatic status, and subgroup were used.

RNA-seq. Total RNA was extracted using the Norgen Total RNA isolation kit and quantified using a NanoDrop Spectrophotometer ND-1000. The RNA was sequenced using single-end 50 bp reads on the Illumina HiSeq platform (Illumina, San Diego CA, USA). Raw sequence data were exported to FASTQ format and were filtered to remove reads with mapping quality scores of Q20 and unique mapping reads for at least 90% of the bases in the sequence. The next reads were mapped to the UCSC mRNA transcript human database based on the GRCh38/hg38 version using HISAT2. The counts were obtained by using ht-seq count with the “intersection–strict” option26. Counts were transformed with TMM transformation and then normalized with VOOM (package limma) in R27. Distributions of samples were examined by performing PCA with “pg” package in R (https://cran.r-project.org/web/packages/pg/index.html) and by adding a dendrogram built with “hclust” function from “stats” package in R with Euclidean distance and average linkage.

Differential expression was obtained by using “limma” package. p values were adjusted with BH method28 and results values <0.05 were considered to be significant. Pathway analysis was performed by using the Reactome tool (https://reactome.org/PathwayBrowser/#/ TOOL = AT), and only terms yielding FDR <0.05 were considered to be significant and were used for further examination.

GSEA was performed by using Oncomine (C6) and Hallmark MsigDB collections of gene sets. Further GSEA analysis was performed using top 50 and top 100 genes belonging to the signatures obtained from the differential expression analysis. Survival analysis was performed by using the following signatures: (1) Sox2 signature (BenPorath_SOX2_TARGETS signature deposited in C2 MsigDB collection), (2) Wnt signature ( deposited in C2 MsigDB collection), and (3) BMP1 and Glinsky signature30. For each signature a signature score was calculated for each patient: \( s = \frac{x}{n} \), where \( x \) is the log2-transformed expression, \( R \) is the set of genes comprising the signature of interest, and \( n \) is the number of genes in that signature, similar to as described previously31,32, but using only the genes regulated in the same direction. For each signature, patients were divided into “High” and “Low” signature index groups using tertiles (1/3 of the patients were assigned to the “High” category and 2/3 were assigned to the “Low” category).

Additional factors, such as age, metastatic status, and subgroup were used. Univariate and multivariate survival analyses were performed using the “survival” package in R (https://cran.r-project.org/web/packages/survival/index.html).

Comparative analysis of transcriptomic data of bulk MB and MB BTIC lines. All analyses were performed in R 3.4.4. The RNA microarray data was downloaded from GEO (GSE68527)17. Before performing analyses with this data, the data were subsetted to include WNT, Group 3 and Group 4 samples (i.e. no SHH). In addition, ENSG identifiers were mapped to HGNC symbols using Ensembl version 77, with mappings downloaded using the BioMart package45,46. Differential expression was conducted using voom and limma for RNA-seq data (this study) and using limma for microarray data (Cavalli 2017 data). Differentially expressed genes were filtered at false discovery rate (FDR) 0.05 (Benjamini–Hochberg procedure). Differential expression was performed for the Cavalli data in three “one vs. all others” comparisons (one for each MB subtype), and differential expression was calculated in a similar manner for untreated MB stem cell lines from this study. For the Cavalli et al. microarray data17 and our MB BTIC samples were scored for gene expression signatures separately using ssGSEA as implemented in the Gsva package47, using default settings for Gsva-gsva (expr = expression.matrix, gset.idx).

Cell proliferation assay. Single cells were plated in 96-well plates, at a density of 1000 cells/200 μl per well in quads for each sample and incubated for four days. Twenty microlayers of Presto Blue (Life Technologies), a fluorescent cell viability indicator, was added to each well 4 h prior to fixation. Fluorescence was measured with a FLUOstar Omega Fluorescence 556 Microplate reader (BMG LABTECH) at an excitation and emission wavelength of 540 and 570 nm, respectively. Readings were analyzed using the Omega software.

Secondary sphere formation analysis. SU_MB002 MB tumorspheres were mechanically dissociated with a 1000 μl pipette tip. Cells were plated at 200 cells per well in 200 μl of BTIC enrichment medium in a 96-well plate. Tumorspheres were left undisturbed at 37 °C with 5% CO2. After 3 days, the number of secondary spheres per well was counted and used to estimate the mean number of spheres/2000 cells.

In vivo experiments. All in vivo studies were performed according to McMaster University Animal Research Ethics Board approved protocols. Intracranial injection of tumor cells was performed using the following human MB samples: BT853, BT992, SU_MB002, RCMB40. Briefly, the appropriate number of live cells, determined by trypan blue exclusion, was resuspended in 10 μl of PBS. NOD-SCID mice were anesthetized using isoflurane gas (5% induction, 2.5% maintenance) and cells were injected into the frontal lobe using a 10 μl Hamilton syringe.

With regards to specific treatment groups and injected cell numbers, comparative tumor-initiating capacities of BT853 and SU_MB002 were performed at limiting dilutions consisting of 1.0 × 10^3 and 5.0 × 10^3 with five mice in each group (n = 10 BT853, n = 10 SU_MB002). Mice were assessed for both histological differences in tumor formation along with length of survival. Comparative histological and survival analyses of mice containing intracranial tumors generated from stable lines of SU_MB002 with β-catenin overexpression (n = 5) or negative-control construct expression (n = 5) were performed by using 1.0 × 10^3 cells from each line. Tumor-initiating capacity, histological differences in tumor size, and survival analysis of SU_MB002 TGP+ (endogenous Wnt-active cells, n = 10) compared with SU_MB002 TGP− (endogenous Wnt-inactive cells, n = 10) were performed using five mice per dilution at dilutions of 1.0 × 10^5, 5.0 × 10^5. Tumor-initiating capacity and histological differences of HD-MB03 TGP+ compared with HD-MB03 TGP− cells were performed using two dilutions of 5 × 10^5 and 1 × 10^6 cells with 4 and 12 mice, respectively. L807mts treatment and PBS controls were delivered via intranasal injection to mice following brief anesthetization with isoflurane gas (2.5% induction). Treatment was given at 50 μg/dose every other day, following a 2-week engraftment incubation. Treatment and control mice were injected with 1.0 × 10^6 cells for SU_MB002 (n = 12), 1.0 × 10^5 cells for BT992 (n = 12), and 1.0 × 10^5 cells for RCMB40 (n = 12). All mice were sacrificed at endpoint, brains were harvested, formalin-fixed, and paraffin-embedded for hematoxylin and eosin staining. Images were taken using the Aperio Slide Scanner and analyzed using ImageScope v11.2.7.60 software (Aperio).

For checking the radiosensitivity of L807mts in vivo, mice were intracranially xenografted with SU_MB002 cells at 60,000 cells per mouse (n = 16) and divided into four cohorts with n = 4/cohort: (i) Control, (ii) Radiotherapy; (iii) L807 and (iv) L807 + Radiotherapy. Ten days post injection, mice allocated into treatment cohorts (iii) and (iv) were administered L807 intrathecally at 50 μg/dose every other day for a total of six doses. Mice in groups (ii) and (iv) were given 2 Gy of cranial irradiation 14 days post injection. Following treatment completion, mice were monitored as per established animal utilization protocols and culled at endpoint.

Immunofluorescence on xenografts. Five micrometers patient-derived xenograft paraffin-embedded tissues were deparaffinized in xylene and processed through a graded series of alcohol concentrations. Antigen retrieval was performed at 95–100 °C for 10 min in citrate buffer pH 6.0. Samples were then incubated with blocking solution (1% BSA, 0.2% Triton 100X, and 5% goat animal serum in 1x TRIS-buffered saline) for 45 min at the room temperature followed by overnight incubation in primary antibody, rabbit anti-human polyclonal Sox2 antibody (1:100, Abcam #97999) at 4 °C. The next day samples were washed in 1×TBS and treated with secondary antibody, Alexa Fluor 488 goat anti-rabbit IgG (1:200, Life Technologies), for 1 h at room temperature. Slides were then washed extensively with 1×TBS and counterstained with prolong gold anti-fade mountant with DAPI (Life technologies).
Images were acquired using Olympus microscope and Volocity software. Nuclei from 20 high power fields were counted and positivity for Sox2 marker expression was quantified and expressed as % positive cells.

Immunohistochemistry (IHC) on xenografts. Immunohistochemical staining was performed on 5 μm sections of TCG-negative and TCG-positive harvested tumor sections. Briefly, tumor sections were deparaffinized by immersion in xylene followed by rehydration in increasing concentrations of alcohol (100, 95, 70%). Antigen retrieval was accomplished using a low pH solution (Vector laboratories) and a biotin-avidin-peroxidase incubation, endogenous peroxidases, biotin, biotin receptors, avidin binding sites, and nonspecific binding was blocked by treating slides with 3% hydrogen peroxide solution, Avidin/Biotin Blocking Kits (Vector laboratories) and Normal Donkey Serum respectively. Rabbit polyclonal GFP primary antibody (Abcam; ab6565; 1:1000) incubation was performed overnight at 4 °C in a humidified chamber followed by donkey anti-rabbit biotin secondary antibody (Jackson Immunoresearch Laboratories; 711-065-152; 1:1000) incubation at the room temperature for 2 h. Slides were washed in 1X TBS solution with or without 0.05% Tween-20 after each step. Following secondary antibody incubation, slides were stained with 3,3’-diaminobenzidine (Vector Laboratories), counterstained with hematoxylin, and differentiated with acid alcohol after which, slides were dehydrated in decreasing concentrations of alcohol (70, 95, 100%), cleared by immersion in xylene and mounted with PolyMount (Polysciences Inc.). All images were obtained using the Aperio Slide Scanner (Leica Biosystems).

Wnt3A-conditioned medium. L. Wnt3A cells (ATCC®-CRL-2647) transgenic mouse fibroblasts transfected with a Wnt3A expression vector, and control L cells (ATCC®-CRL-24) cultured according to ATCC’s recommendations. Briefly, L. Wnt3A cells were maintained in DMEM and 10% FBS with 0.4 mg/ml G418. Control L cells were maintained in DMEM and 10% FBS. Both, L. Wnt3A and control L cells were subcultured 1:10 in culture medium without G418 and grown until confluency. Culture media were then removed and filtered (0.2 μm). Wnt3A and control conditioned media were separately mixed with BTIC enrichment 1 Perfecta SybrGreen (Quanta Biosciences) and CFX96 instrument (Bio-Rad). CFX 30 min at 42 °C, 5 min at 85 °C, hold at 4 °C. qRT-PCR was performed using the were normalized to GAPDH expression with secondary validation normalized to ATGATAAAAGAT-3 until con.

Immunohistochemical staining was accomplished using a low pH solution (Vector laboratories) and a biotin-avidin-peroxidase incubation, endogenous peroxidases, biotin, biotin receptors, avidin binding sites, and nonspecific binding was blocked by treating slides with 3% hydrogen peroxide solution, Avidin/Biotin Blocking Kits (Vector laboratories) and Normal Donkey Serum respectively. Rabbit polyclonal GFP primary antibody (Abcam; ab6565; 1:1000) incubation was performed overnight at 4 °C in a humidified chamber followed by donkey anti-rabbit biotin secondary antibody (Jackson Immunoresearch Laboratories; 711-065-152; 1:1000) incubation at the room temperature for 2 h. Slides were washed in 1X TBS solution with or without 0.05% Tween-20 after each step. Following secondary antibody incubation, slides were stained with 3,3’-diaminobenzidine (Vector Laboratories), counterstained with hematoxylin, and differentiated with acid alcohol after which, slides were dehydrated in decreasing concentrations of alcohol (70, 95, 100%), cleared by immersion in xylene and mounted with PolyMount (Polysciences Inc.). All images were obtained using the Aperio Slide Scanner (Leica Biosystems).

Western immunoblotting. Ten micrograms of denatured total protein or 7.5 μg of nuclear and cytoplasmic extracts (NE PER reagents, Thermo Scientific) were mixed with NC gel beads, aiming for 1000 cap-

Flow cytometric analysis and cell sorting. Tumorspheres were dissociated and single cells resuspended in PBS. Samples were sorted using a MoFlo XDP Cell Sorter (Beckman Coulter). Dead cells were excluded using the viability dye 7AAD (1:10, Beckman Coulter) or a near IR Live/Dead fixable staining kit (Life Technologies). Compensation was performed using mouse IgG CompBeads (BD). GFP expression was defined as positive or negative based on the analysis of regions established by the isotype control. Cells were sorted into tubes containing 1 ml. BTIC enrichment medium and small aliquots from each sort tube were analyzed to determine the purity of the sorted populations. Cells were allowed to equilibrate at 37 °C for a few hours prior to experimentation. Intracellular levels of Bmi1 and Sox2 were determined following preparation using Fixation/Permeabilization Solution (BD Biosciences) along with antibodies anti-Bmi1 (1:11; Cat. 130-106-736, Miltenyi) and anti-Sox2 (1:20; Cat. 561610, BD Horizons), for which viable cells were stained using LIVE/DEAD® Fixable Near-IR Dead Cell Stain Kit (Cat. L10119, Thermo Scientific).

Small molecule Wnt activation. The competitive ATP inhibitor functions as a selective GSK-3β inhibitor CHIR99021, was used to activate the canonical Wnt pathway in SU_MB002 and ICB1299 cells. Similarly, the substrate-competitive peptide inhibitor of GSK-3, L807mts, was also used to activate the canonical Wnt pathway in SU_MB002 and ICB1299 cells. 2 × 105 cells were plated in a 24-well plate in triplicate at a volume of 500 μL/well at CHIR99021 concentrations of 1, 3, 5, 7, 10 μM. DMSO was used as a control. CHIR99021 and DMSO were replenished after 24 and 48 h and cultures were used for qRT-PCR, self-renewal, proliferation, or radiation assays after 3 days. 2 × 105 cells were plated in a 24-well plate in triplicate at a volume of 500 μL/well at L807mts concentrations of 1, 5, 10 μM. PBS was used as a control. L807mts and PBS were replenished after 24 and 48 h and cultures were used for qRT-PCR, self-renewal, and proliferation.

In vitro radiotherapy. Cells were plated at density of 1 × 106 cells/ml and treated with either one dose of 2 Gy radiation (Faxitron BX-650) and incubated for a week or a range of doses ranging from 0 to 5 Gy. Functional self-renewal and proliferation assays were performed following incubation.

Single-cell RNA-seq and library preparation. Single-cell suspensions of 1000 cells with a final viability of >80% was used. Single-cell library preparation was carried out as per the 10X Genomics Chromium single-cell protocol using the v2 chemistry reagent kit (10X Genomics, Piscataway, USA). Cell suspensions were loaded onto individual channels of a Chromium Single-Cell Chip along reverse transcription (RT) master mix and single-cell 3’ gel beads, aiming for 1000 capture per individual cell sample. Following isolation of gel beads by centrifugation, cDNA was underwent a two-stage purification process with Dynal MYOne Silane beads (Thermo Fisher Scientific), followed by SPRSelect beads (Beckman Coulter). RT and cDNA amplification were performed using a 96-well Veriti Thermocycler (Life Technologies). Amplified cDNA purified using SPRSelect beads and sequencing libraries with barcodes were generated using single uncleaved adapters. Dual sided SPRSelect cleanup was used to clean up samples and size select. Libraries were sequenced on an Illumina 2500 in Highoutput mode at the Princess Margaret Genomics Centre (Toronto, ON) using the 10X Genomics recommended sequencing parameters. Samples were quantified by Kapa Library Quantification kit (Roche) and normalized to achieve the desired median read depth per cell.

Single-cell RNA-seq data preprocessing. The Cell Ranger software pipeline (version 2.1.0) developed by 10X Genomics has been used to demultiplex cell barcodes, map reads to the transcriptome (GRCh38) using STAR aligner and downstream reads as required to generate normalized aggregate data across samples. The number of reads per cell barcode was calculated using the BamTab histogram function in the Drop-seq Alignment Cookbook. Subsequently, the number of cells per sample was calculated by summing the cumulative fraction of reads corresponding to each individual cell barcode in a library. Cell barcodes were sorted in decreasing order and the inflection point was identified using the R package DropRead (version 0.3.1) to distinguish between empty droplets with only ambient RNA and true droplets containing a cell. The raw matrix of gene counts versus cells from Cell Ranger output was filtered by cell barcodes identified from Dropbase. We processed the resultant unique molecular identifier (UMI) count matrix using the R package Seurat (version 2.3.4).

Single-cell RNA-seq quality control, normalization, and dimensionality reduction. For each cell, we quantified three quality control metrics: the number of genes with at least one UMI, the total number of UMIs detected and the percentage of UMIs belonging to expressed mitochondrial genes. We excluded all cells with >30% mitochondrial UMIs, potentially indicative of damaged cells with compromised cellular membranes.

Similar to the workflow in Lun et al.,52, likely multiple captures were removed if log library size exceeded 2 median absolute deviations above the median. We further discarded low-quality cells where below 500 genes were detected. We filtered out lowly expressed genes detected in less than 5 cells across the aggregated dataset, corresponding to 1% of cells in the smallest library. After applying these QC criteria, 3725 single cells and 18721 genes were remained and were included in downstream analysis. Expression quantification was performed in Seurat on the filtered matrix to obtain log-normalized counts scaled to library size. Relative gene expression was calculated by centering expression across all cells in the cohort.
Single-cell RNA-seq gene signature scoring and classification. Gene signature scores in individual cells were calculated with two methods. The first being the AddModuleScore() function in the R package Seurat (version 2.3.4). In brief, the average relative expression level for each gene signature was calculated on a single-cell level and subtracted by the aggregated relative expression of control gene sets. Control gene sets were defined by binning all 18,721 expressed genes into 25 bins of aggregate expression levels and then for each gene in the gene signature of interest, randomly selecting 100 genes from the comparable expression bin. As a result, control gene sets have comparable expression level distributions to the gene signature of interest. A second set of gene signature enrichment scores were calculated using GSVA with default parameters.

To estimate the significance of gene signature scores, a hundred sets of randomly selected genes sets with the same size as the gene signatures were generated and scored using both methods described above. Random sets were then used to define a 5% cutoff for the expected gene signature scores. Cells were classified as “enriched” for a given signature if they surpassed the 5% threshold.

Single-cell RNA-seq analysis of publicly available data. Normalized count matrices for human MB tumor scRNA-seq data for 25 samples (five Wnt, three Shh, eight Group 3, Group 4) accounting for 7745 malignant cells was downloaded from GSE119926. Normalized log2(TPM/10 + 1) expression was transformed into log(TPM/10 + 1) for input into Seurat to perform dimensionality reduction and clustering as described previously (6571 variable genes, 15 significant PCs). Cells were scored for gene signatures with GSVA and classified as previously described.

scRNA-seq data in the form of raw UMI count matrices for eight human MBs (two Shh, two Group 3, four Group 4) accounting for 28,268 cells were downloaded from Vladoiu et al. Raw counts were log normalized using Seurat (v2.3.4). Data scaling, dimensionality reduction, and clustering were performed as described previously (2814 variable genes, 12 significant PCs) using Seurat. Cells were scored for gene signatures with GSVA (v1.30.0) and classified as previously described. Microglia/monocyte (AIF1, CD14, CD68) and T cell (CD3G, CD8A, IL7R) clusters were annotated using known gene markers and cell labels as described in Vladoiu et al. Remaining cells labeled as malignant tumor cells (1155 microglia/monocytes; 188T cells; 26,925 tumor cells).

Gene set variance analysis of publicly available data. Microarray data for Cavalli et al. were obtained from GEO (GSE85217). ENSEMBL gene identifiers were converted to HGNC symbols with annotations downloaded from biomart. HGNC symbols mapping to an identical ENSG ID were handled by only taking 1 HGNC symbol per ENSG id, and ENSG IDs mapping to an identical HGNC symbol were handled only using 1 ENSG per HGNC symbol. GSVA 3 signatures were used for GSVA scoring: Group 3 (n = 7941 genes) and Group 4 (n = 6510 genes) defined in differential expression analyses (Group 3 vs. union of Wnt and Group 4, Group 4 vs. union of Wnt and Group 3, respectively), and the Broad Institute MSGBD HALLMARK_WNT_BETA_CATENIN_SIGNALING signature (n = 42 genes). Differential expression analyses were performed using limma version 3.38.3. GSVA was run using GSVA version 1.3.0. Microarray data from Northcott et al. were generously provided by the Northcott laboratory. ENSEMBL transcript ids (ENST) were mapped to HGNC symbols in a manner similar to that done for mapping ENSG ids to HGNC symbols for the Cavalli et al. data. GSVA scoring was done using the same methods as for the Cavalli et al. data.

Statistical analysis. At least three biological replicates were performed for each experiment. Data represent mean ± standard error (mean) with n values listed in figure legends. Student t test analyses were performed using GraphPad Prism™ with significance set to P < 0.05. Kaplan–Meier survival curves were visualized using GraphPad Prism™. All statistical analysis and data visualization for scRNA-seq was performed in R (version 3.5.0).

Data availability

The RNA-Seq data discussed in this publication have been deposited in NCBI’s Gene Expression Omnibus and are accessible through GEO Series accession number GSE131473. The scRNA-seq data have been deposited in CReSCENT (https://crescen.t.cloud/ CRESCENT) and Processed data is uploaded on BROAD institute site https://singlecell.broadinstitute.org/single_cell/study/SCP840. Source data are provided with this paper.

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**Author contributions**

B.M., C.V., B.W.D., and S.K.S. conceptualized and designed the experiments. B.M., C.V., D.B., A.A.A., L.R., M.M.K.S., O.W., A.D.G., M.S., N.S., N.T., D.K., B.B.A., and N.W. performed experiments and acquired data. B.M., C.V., L.R., O.W., A.D.G., N.W., R.H., G.D.B., T.J.P., B.W.D., S.K.S. analyzed and interpreted data. B.M. wrote the manuscript with revisions contributed by C.V., B.W.D., and S.K.S. B.Y., O.A. provided pediatric MB study samples. J.P.P. analyzed all clinical and preclinical histological samples. A.F. provided pediatric MB study samples. J.P.P. analyzed all clinical and preclinical histological samples. B.Y., O.A. provided pediatric MB study samples. J.P.P. analyzed all clinical and preclinical histological samples. B.M., C.V., B.W.D., and S.K.S. conceptualized and designed the experiments. B.M., C.V., B.W.D., and S.K.S. conceptualized and designed the experiments. B.M., C.V., B.W.D., and S.K.S. conceptualized and designed the experiments.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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