**Abstract**

Medulloblastoma is the most common malignant pediatric brain tumor. Despite its prevalence and importance in pediatric neuro-oncology, the genes and pathways responsible for its initiation, maintenance, and progression remain poorly understood. Genetically engineered mouse models are an essential tool for uncovering the molecular and cellular basis of human diseases, including cancer, and serve a valuable role as preclinical models for testing targeted therapies. In this review, we summarize how such models have been successfully applied to the study of medulloblastoma over the past decade and what we might expect in the coming years.

**Key words** Genetically engineered mouse, medulloblastoma, mouse model

As the most common malignant childhood brain tumor, medulloblastoma (MB) is a development-associated embryonal tumor of the cerebellum, a component of the central nervous system (CNS) essential for motor coordination in vertebrates. Within two years of diagnosis, approximately 10%–15% of patients die of the disease [1]. Although ~60% of MB patients can be cured with current therapeutic regimens, the majority of survivors suffer from severe long-term neurologic, endocrinologic, and cognitive sequelae [1]. Biological heterogeneity is a major property of MB, however, in the clinic the disease is segregated into standard risk and high risk groups. Completely resected tumors from patients older than 3 years with no leptomeningeal macroscopic dissemination at diagnosis are considered standard risk, whereas all others are designated high risk [2]. In 2007, the disease was histologically classified by the WHO into classic MB (approximately 50% of the disease) and four other subgroups including desmoplastic MB, anaplastic MB, large-cell MB, and MB with extensive nodularity (MBEN) [3].

Enormous efforts have been devoted to classify and subgroup MB in order to improve prognosis and to reduce treatment-associated side effects. Recently, employing cytogenetic and molecular biomarkers, MB have been classified into distinct subgroups [1,2]. Multiple deregulated signaling pathways and cytogenetic aberrations have been identified in MBs; however, many questions about the pathogenesis of the disease remain to be answered.

Over the past two decades, mouse models have proven to be a very effective tool for answering questions about human cancers. Here, after a brief review of modelling human cancers in the mouse, we will discuss many of the prominent MB mouse models (categorized according to signaling pathway) published since 1997, when the first Ptc" mouse model was generated. Upon conclusion, we will describe future perspectives on modelling human MB in the mouse.

**Modelling Human Cancers in the Mouse**

For the past two decades, conventional and advanced conditional genetically engineered mouse models (GEMMs) as well as xenografts passed in the mouse have been an indispensable tool in basic cancer biology and translational cancer research [4]. As mammals, mice are closely related to humans at a physiological, biochemical, and genetic level. In addition, mice are less expensive to maintain than other mammals and are particularly amenable to genetic manipulation. During the last decade, enhanced knowledge of human and mouse genomes has facilitated genetic manipulation, permitting a better understanding of gene function and allowing more accurate modelling of human diseases. Ideally, a faithful recapitulation of human disease in genotype and phenotype is an essential element of mouse modelling. Four basic mouse models are briefly described below.
Transplantation/xenograft models

After in vivo or in vitro manipulation, human or mouse tissues or cells can be transplanted into either immuno-compromised or wild-type mice subcutaneously, or orthotopically for studying gene function and screening therapeutic molecules. These models allow speedy, flexible, and relatively inexpensive manipulation of tissues or cells with multiple genetic and molecular approaches, such as gene knockout, ectopic expression of mutated genes, and RNA interference.

Conventional GEMMs

Classically, GEMMs are germline models, in which mice carry genetic modification in their germline and maintain the modification through breeding. In gain-of-function studies, pronuclear injection into a fertilized zygote is the most popular method to over-express or mis-express a gene in the mouse germline. In these models, depending on the regulatory elements chosen for expression, the transgene can be expressed either ubiquitously or in a tissue and/or time specific manner. In contrast, gene targeting in mouse embryonic stem (ES) cells is employed to generate null or “knockout” mutations in order to study the loss-of-function.

Conditional inducible GEMMs

To overcome limitations associated with conventional GEMMs, such as embryonic or early postnatal lethality, conditional GEMMs have been created, in which genetic events can be tightly controlled spatially and temporally. In these systems, the site-specific bacterial Cre recombinase enzyme and its modified version Cre-ER are used to control the expression, or the “knockout” of a gene at a specific location, or at a specific time in development. Additional elements of temporal control can be gained through the use of tetracycline (Tet) inducible systems: Tet-off or Tet-on systems to conditionally over-express an oncogene or knock out a tumor suppressor.

shRNAi and transposon-based GEMMs

Recently, short hairpin RNA interference (shRNAi) technology has been applied to mouse models in order to knockdown genes in vivo by introducing promoter-shRNA constructs into the mouse through conventional or conditional approaches. Meanwhile, DNA transposon systems such as Sleeping Beauty (SB) have been employed to induce tumors through genetic engineering in the mouse and have been proven insightful in functional cancer genomics studies.

Modelling Human MB in the Mouse

As the most common malignant pediatric brain tumor, MB has been extensively studied in pediatric neuro-oncology research. The different mouse modelling technologies described above have been applied to studies of MB biology, genetics, and preclinical research. Table 1 summarizes many of the genetically engineered MB mouse models currently published in the literature.
A signature feature of childhood MB is its occurrence in the developing cerebellum. Development of the cerebellum begins during embryogenesis and continues after birth in both mice and humans. Two distinct germinal zones, the primary and secondary, containing stem and/or progenitor cell populations, give rise to the various cell types of the mature cerebellum. The primary germinal zone, the ventricular zone (VZ), is located in the roof of the fourth ventricle, giving rise to GABAergic neurons, Purkinje cells (a specialized neuronal population of the cerebellum), and Golgi neurons. Progenitor cells of the secondary germinal zone originate in the rhombic lip and give rise to cerebellar granule neuron precursors (CGNPs) that migrate rostrally across the cerebellum to form the external granule/germinial layer (EGL). The EGL persists until postnatal day 21 (P21) in mice and into the second year of life in humans. As the cerebellum develops, CGNPs forming the EGL undergo a period of rapid and massive clonal expansion with a peak at P5–7 in the mouse before migrating inward, across the Purkinje cell layer, to eventually form the post-mitotic neurons of the internal granule layer (IGL). Extensive studies have shown that multiple signal transduction pathways are involved in normal cerebellar development. These pathways play a critical role in the expansion of neural precursor populations such as those found in the EGL and VZ and deregulation of these pathways are believed to contribute to MB pathogenesis. Among various developmental signaling pathways, the Wingless (WNT) and the Sonic hedgehog (Shh) pathways have been the primary area of focus for studies related to MB biology. Not surprisingly, the majority of published MB mouse models have thus been associated with these developmental signaling pathways.

**Shh signaling-based MB mouse models**

The morphogenetic factor Shh was initially discovered in *Drosophila* in 1980 and since then it has been intensively studied. Shh signaling plays pivotal roles in the development of multiple organs and tissues in vertebrate counterparts. As shown in Figure 1, major components in the pathway have been identified, including the negative regulators patched 1 (PTC1) and suppressor of fused (SUFU), the activating mediator smoothened (SMO), and the downstream effectors, the GLI transcription factors. There are many unanswered questions about the regulation and targets of Shh signaling. Early studies linking Shh signaling to disease identified heritable or somatic PTC mutations in patients with the basal cell nevus syndrome (BCNS, also known as Gorlin syndrome) and sporadic basal cell carcinoma (BCC) as well as in sporadic MBs and other cancers.

![Figure 1. Major components of Sonic hedgehog (Shh) signaling pathway.](image)

In the absence of Shh, the negative regulator patched 1 (PTC1), localized at the plasma membrane, inhibits signaling through the activating mediator smoothened (SMO). In the presence of Shh ligand, Shh binds to PTC and derepresses its negative regulation on SMO, activating signaling by inducing the expression of transcription factors including MycN and GLI in the nucleus. The Shh pathway can be negatively regulated by suppressor of fused (SUFU) in the cytosol.
The first Ptc\(^{+/+}\) mouse model  To study functions of PTC in development and tumorigenesis, the first Ptc\(^{+/+}\) mouse model was generated by homologous recombination in embryonic stem cells, in which part of Ptc exon 1 (including the putative start codon) and all of exon 2 were replaced with lacZ and a neomycin resistance gene [15]. Homozygous deletion of Ptc in the mouse was embryonic lethal. In addition to an alteration in neural cell fates, 14% of Ptc\(^{+/+}\) mice (B6D2F1 background) developed MB as early as 5 weeks after birth, with a peak between 12 and 25 weeks, consistent with tumors seen in BCNS patients [15]. Figure 2 shows the histology of a typical MB from a Ptc\(^{+/+}\) mouse. By employing the Ptc\(^{+/+}\) mouse model in 1999, Wechsler-Reya et al. [16] were the first to show that Shh is secreted by Purkinje cells, and that Shh is responsible for regulating CGNP proliferation during normal cerebellar development. After further characterizing the first generation of the Ptc\(^{+/+}\) mouse model [17], by crossing Ptc\(^{+/+}\) mice with Tp53 deficient mice (Tp53\(^{−/−}\)), Weetmore et al. [18] dramatically increased MB incidence to over 95% and reduced the latency to 12 weeks after birth, demonstrating that genomic instability may contribute to the pathogenesis of MB. The high incidence of MB in this model may make it especially useful for preclinical studies.

Ptc\(^{+/+}\)-Ink4c mouse model  Disruption of the RB (retinoblastoma) pathway, as a result of loss of cyclin-dependent kinase (CDK) inhibitor INK4 proteins (INK4A and INK4C) or RB itself, or over-expression of D-cyclins and their associated kinases (CDKs), is a hallmark feature of different types of human cancer [19]. To assess whether Ink4c loss could collaborate with Ptc\(^{+/+}\) inactivation to accelerate MB formation, Ptc\(^{+/+}\) mice were crossed with Ink4c null mice [16]. Either homozygous or hemizygous deletion of Ink4c on a Ptc\(^{+/+}\) background greatly increased MB incidence (from 7% to 30%) with shortened latency although deletion of Ink4c itself was not able to induce MB formation. This result demonstrated that deregulation of the Rb pathway contributes to the progression of MB.

Ptc\(^{+/+}\)-Kip1 mouse model  Recently, Kip1, another cyclin-dependent kinase inhibitor important for cerebellar development, was tested in a mouse model on a Ptc\(^{+/+}\) background [19]. Analogous to Ink4c deletion, either homozygous or hemizygous deletion of Kip1 in Ptc\(^{+/+}\) mice substantially increased MB incidence (from 40% to 60%−67%) with a trend towards shortened latency.

Ptc\(^{+/+}\)-Hic1\(^{+/+}\) mouse model  LOH of human chromosome 17p13.3, a locus distal to Tp53, is among the most frequent genetic events in sporadic human MBs with an incidence as high as 30%−50% [20]. Searching for tumor suppressor genes in this region has revealed hypermethylated in cancer 1 (HIC1), a frequent target of promoter hypermethylation and epigenetic gene silencing in MB and several other cancers [21]. To address the role of HIC1 in MB, Ptc\(^{+/+}\) mice were crossed with Hic1\(^{+/+}\) mice on a C57Bl/6 background [21]. Although the latency of MB genesis in Ptc\(^{+/+}\)-Hic1\(^{+/+}\) double heterozygous mice was similar to those in Ptc\(^{+/+}\) mice, homozygous deletion of Hic1 in Ptc\(^{+/+}\) mice had a 4-fold increase in MB incidence.

Second generation Ptc mouse model  Recently the Cre-loxP system was employed to generate second-generation Ptc mouse models. In an attempt to identify the cell of origin of Shh signaling-associated MB, Yang et al. [22] generated conditional Ptc-knockout mice, Math1-Cre/Ptc\(^{+/+}\) mice and Gfap-Cre/Ptc\(^{+/+}\) mice, in which homozygous deletion of Ptc was selectively carried out in granule neuron precursors (GNPs) and neuronal stem cells (NSCs), respectively. All of the conditional Ptc

Figure 2. Histology of medulloblastoma (MB) from Ptc\(^{+/+}\) mouse. Compared to non-tumor cells, Ptc\(^{+/+}\) MB tumor cells are stained by hematoxylin (blue) with higher intensity, showing a high ratio of nucleus to cytoplasm (stained by eosin, pink) in the poorly differentiated tumor cells. A, Hematoxylin and Eosin (H&E) stained MB at the surface of cerebellum (>10); B, H&E stained MB tumor cells (>40).
mice, either Math1-Cre mice or Gfap-Cre mice, gave rise to MB incidence at the surface of cerebellum. However, the MB latency was different between the two genotypes of mice, with a latency of 3–4 weeks for Gfap-Cre/Ptc1 mice and 8–12 weeks for Math1-Cre/Ptc1 mice. Moreover, the authors also generated Math1-Cre-ER/Ptc1 mice to specifically delete Ptc during post-natal development by controlling Cre expression through the administration of tamoxifen.

**Sufu−/−/Tp53−/− mouse model** In addition to Ptc, the negative regulator Sufu and the positive regulator Smo have also been employed for the establishment of Shh-dependent mouse models of MB. A subset of MB patients has been shown to carry germline and somatic SUFU mutations, implicating SUFU as a tumor suppressor [24]. To confirm the role of SUFU during development and its deregulation in MB, Lee et al. [25] employed a gene-trap approach to generate genetically modified mice carrying a Sufu mutation. Similar to Ptc null mice, Sufu null mice died around E10. Sufu−/− mice did not develop any tumors, whereas Sufu+/−/Tp53−/− double-mutant mice developed MBs and other types of cancer. Compared with Ptc+/−/Tp53−/− mice, in which the incidence of MBs was ~93% with a latency of 2 months, ~58% of Sufu+/−/Tp53−/− double-mutant mice developed MB by 4 months.

**Mouse models with constitutively active Smo** In order to obtain a MB mouse model with a tumor incidence higher than the first generation Ptc mouse model, Hallahan et al. [26] generated a ND2-SmoA hemizygous mouse model, in which a constitutively active Smo was specifically expressed in GNPs under regulation of the NeuroD2 promoter. MBs were observed in 48% of the ND2-SmoA mice at median age of 25.7 weeks, indicating selectively activating Smo does increase MB incidence in the mouse. More recently, a ND2-SmoA homozygous mouse model (Smo/Smo) was generated by the same group [27]. The homozygous model greatly increased MB incidence to ~94% with a much earlier onset (4–8 weeks). More interestingly, the authors also observed leptomeningeal metastases in this mouse model that are very similar to the pattern of metastases seen in human children with MB [27]. The increased incidence, early onset, and leptomeningeal metastasis should render the Smo/Smo model useful for preclinical research on Shh signaling associated MBs.

**Tp53 mutation-based MB mouse models**

Although studies have demonstrated aberrant activation of the Shh pathway plays an essential role in a subset of MBs, mutations in this pathway, including PTC, SUFU, SMO as well as amplification of GLI1 and GLI2, are observed in only 25%–30% of human MB cases [28], indicating other genetic and/or epigenetic events are involved in MB pathogenesis. Mutations in WNT, TP53, and RB pathways as well as genes controlling histone lysine methylation have been identified in human MB [29,29]. To fully model the spectrum of human MB, there has been great interest in the development of Shh-independent MB mouse models. Attempts to generate a mouse model independent of Shh pathway activation have been focused on using a Tp53 null background as described below.

In fact, other than Ptc, Tp53 is the most frequently targeted gene in MB mouse modelling, in spite of the fact that Tp53 mutation (both homozygous and hemizygous deletion) alone very rarely induces MB formation. Various compound mutations of Tp53 with other candidate genes, however, have been shown to be able to effectively induce MB in mice likely due to the genomic instability provoked by Tp53 mutation. Indeed, as mentioned earlier, Tp53 deficiency promotes MB formation in both Ptc−/− and Sufu−/− backgrounds. Other than Shh signaling-associated MB mouse models, Tp53 deficiency has been shown to cooperate with the following genes in published MB mouse models.

**Tp53 and Rb** With an initial intention to study the pathogenesis of gliomas, Gfap-Cre/Rblox/lox/TP53−/− and Gfap-Cre/Rblox/lox/TP53−/− mice were generated [30]. In this study, Gfap-driven expression of Cre was detectable in newborn mice not only in the developing forebrain and spinal ganglia, but also in precursor cells located in the EGL of the developing cerebellum. Instead of gliomas, conditional inactivation of both Tp53 and Rb led to MB formation in this study. Although the cohort was small (9 mice), over 80% of the double null mice developed MBs by 3 months.

**Tp53 and defective DNA damage sensor and repair genes** Lig4, a nuclear ligase, is a critical component of the non-homologous end joining (NHEJ) apparatus that is required for (D)J somatic recombination and repair of DNA double strand breaks (DSBs) [31]. *Lig4* homozygous mutations result in embryonic lethality in mice which however, can be rescued by Tp53 deficiency [31]. Interestingly, before the onset of pro-B lymphomas, all Lig4−/−/Tp53−/− mice developed MBs within 9 weeks and as early as 3 weeks of age [32]. The same group went on to generate Nestin-Cre/Xrc4−/− mice and obtained similar results as from *Lig4−/−/Tp53−/−* mice with a slightly lower incidence of medulloblastoma, but no incidence of lymphoma [33]. By selectively deleting *Xrc4* (another NHEJ pathway members) in neuronal precursor cells (NPCs) on a Tp53 null background, the authors found that, along with loss of Ptc, extensive chromosomal translocations occurred in the tumor. Characterization of the induced MBs showed the tumor was similar to Ptc−/−...
MB. More recently, to further characterize the tumor suppressor role of DNA DSB repair pathways in defined tissues, another group used the Cre-LoxP system to conditionally inactivate Lig4, Krc, and Brca2 in nestin-expressing cells [34]. In combination with a Tp53 homozygous deletion, rapid development of MBs occurred in all three mouse models, indicating that both defective NHEJ and homologous recombination (HR) DNA repair pathways predispose the developing cerebellum to MB initiation. Furthermore, the specific deletion of two major tumor suppressor genes Ptc and Pten were identified in mouse models, demonstrating their important role in the suppression of MB formation.

Poly-(ADP-ribose) polymerase (PARP1) is another DNA repair protein. To test the role of PARP1 in MBs, a mouse model in which Parp was homozygously deleted on a Tp53 null background was generated through crossing of Tp53 null mice with Parp null mice [35]. The Parp null mouse was generated by targeted Parp deletion in ES cells, and was healthy and fertile but impaired in response to environmental stress [36]. However, starting from 8 weeks, Parp1 homozygous deletion induced MBs in Tp53 null mice at an incidence of ~49%.

As described above, Parp1/Tp53-/- MB showed activated Shh signaling, such as an increased expression of Gli1, and severe chromosomal aberration.

Shh pathway-independent MB mouse models

As mentioned earlier, characterization of mouse MBs generated in the context of Tp53 deficiency showed aberrant Shh signaling (particularly loss of Ptc) in the tumors, indicating they are not independent of Shh pathways, likely because in all of the models, CGNPs were the cells targeted for tumor initiation. Most recently, two models have shed light on Shh-independent MB mouse models by targeting different cell populations during CNS development.

**GTML mouse model** Using a tetracycline repressor-based Tet-off system, in which the tetracycline controlled transactivator (TfTa) was specifically driven by the glutamate transporter 1 (Glt1) promoter, in combination with a bidirectional tetracycline response element (TRE) to express both MycN and luciferase, Swartling et al. [37] recently generated a novel MB mouse model, Glt1-TfTa and TRE-MycN/luciferase (Luc), which they dubbed as GTML. Although expression of N-myc in nestin-expressing cerebellar progenitors did not induce MB [38], by 200 days, over 40% and 75% of GTML hemizygous and homozygous mice developed MB, respectively. Tumors were characterized as Shh-independent classic and/or LCA MBs. Remarkably, some of the mice displayed leptomeningeal metastases. These animals likely represent the first model of non-Shh/non-Wnt MB, and should therefore prove invaluable in preclinical research. After “switching off” the MycN transgene using tetracycline, the authors were able to observe involution of the tumor, illustrating the power of tetracycline inducible models, and demonstrating the critical role of MycN in MB maintenance. We predict others will use this technology to test the role of various oncogenes not just in tumor initiation but more importantly in tumor maintenance.

**Blbp-Cre/Ctnnb1-flox/flox/Tp53-flox/- Wnt-subgroup MB mouse model** Most recently, in order to validate their observations from the human disease, Gibson and colleagues generated a mouse model to recapitulate human WNT-subgroup MBs, in which, Ctnnb1 was mutated in Blbp-expressing cells via Cre-LoxP system. In combination with Tp53-/-, 15% of the Ctnnb1 and Tp53 double mutants developed MBs (latency ~290 days) with a molecular profile similar to human WNT-subgroup MBs [39]. The authors used this model to great effect to identify a candidate cell of origin for Wnt MB that differs from the cell of origin for Shh MB.

**Non-GEM MB mouse models**

Similar to numerous other types of cancer, both ectopic flank and orthotopic intracranial xenografts into immuno-compromised mice, such as Nu/Foxn1Nu and NOD/SCID, have been widely used in MB research, especially for validation studies and preclinical research [40-43]. The extent to which human xenograft models of MB experience genetic or epigenetic “drift” overtime that causes them to lose similarity to the freshly resected tumor is not currently known.

To directly test whether the activation of the Shh pathway can lead to tumor formation, Weiner et al. [44] created a retroviral mouse model by injecting a Shh-expressing retrovirus into the embryonic mouse cerebellum using in utero ultrasound backscatter microscopy (UBM) image guidance. MB was observed in about 76% of the experimental mice at P14 and P21. Although this model can be technically demanding, further optimization might render it useful for preclinical drug screening. Similarly, the RCAS/fv/a system is a mouse model that integrates orthotopic cell transplantation into a transgenic mouse. Dan Fult’s group has been using RCAS/fv-a for simultaneously targeting multiple transgenes including MycN, MycC, Igf2. Akt. Gli1 into nestin-expressing cell populations to induce/modulate MB in neural progenitors of the cerebella of newborn mice [35]. In this system, RCAS, the replication-competent avian leukemia virus splice-acceptor subgroup-A (ALV-A) vectors that carry the transgenes, are transfected in vitro and the virus-producing cells are then injected into a transgenic
Summary and Future Perspectives

In summary, the majority of published MB GEM and related models are associated with the Shh signaling pathway using either a Ptc or a Tp53 mutant background. These models are based on observations from the human disease. The genetic, epigenetic and pathologic heterogeneity of MB strongly suggests other MB driver genes and pathways responsible for the initiation, maintenance and metastasis of MBs remain to be uncovered. Recent, SB transposon-based mouse models have been reported for functional cancer gene discovery\[15,40\].

Metastasis and recurrence of MB is the main cause of death in patients. Currently MB patients either with or without metastasis receive similar treatments due to scarce knowledge about the biology and molecular genetics of metastasis. Among the published models, metastasis was observed only in two of them (Smo/Smo and GTML), suggesting the necessity of developing additional mouse models that can mimic both the primary tumor and its metastases.

Current developments in genomics, particularly deep sequencing technologies, promise to deliver large amounts of genetic and epigenetic data from the various subsets of human MB in the very near future. Determining which events are driver mutations and among those, which are required for tumor maintenance and worth targeting in the clinic, will require sophisticated mouse models to be developed and tested in the near future.

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