Insulin-like Growth Factor 2 Is Required for Progression to Advanced Medulloblastoma in patched1 Heterozygous Mice

Ryan B. Corcoran, Tal Bachar Raveh, Monique T. Barakat, Eunice Y. Lee, and Matthew P. Scott

Abstract

Medulloblastoma (MB) can arise in the cerebellum due to genetic activation of the Sonic Hedgehog (Shh) signaling pathway. During normal cerebellum development, Shh spurs the proliferation of granule neuron precursors (GNP), the precursor cells of MB. Mutations in the Shh receptor gene patched1 (ptc1+/-) lead to increased MB incidence in humans and mice. MB tumorigenesis in mice heterozygous for ptc1+/- shows distinct steps of progression. Most ptc1+/- mice form clusters of preneoplastic cells on the surface of the mature cerebellum that actively transcribe Shh target genes. In ~15% of mice, these preneoplastic cells will become fast-growing, lethal tumors. It was previously shown that the loss of function of insulin-like growth factor 2 (igf2) suppresses MB formation in ptc1+/- mice. We found that igf2 is not expressed in preneoplastic lesions but is induced as these lesions progress to more advanced MB tumors. Igf2 is not required for formation of preneoplastic lesions but is necessary for progression to advanced tumors. Exogenous igf2 protein promoted proliferation of MB precursor cells (GNP) and a MB cell line, PZp53MED. Blocking igf2 signaling inhibited growth of PZp53MED cells, implicating igf2 as a potential clinical target.

Introduction

Mutations in the Sonic hedgehog (Shh) signaling pathway that cause constitutive Shh target gene transcription are associated with the formation of many human tumors. Medulloblastoma (MB), the most common malignant brain tumor in children, can arise when such mutations occur in developing cerebellar granule neuron precursors (GNP; ref.1). Loss of function of the PATCHED1 (PTCH, ptc1 in mice) tumor suppressor gene, which occurs in sporadic and hereditary MB, is the most common of these mutations (2, 3). PTCH encodes a 12-transmembrane receptor (Ptc1) for Shh. In the absence of Shh signal, Ptc1 inhibits the downstream transducers of the Shh pathway, preventing activation of target gene transcription by Gli transcription factors. Shh activates target gene transcription by binding to and inhibiting Ptc1. Shh and Ptc1 are therefore antagonists. Ptc1 and gli1 are Shh target genes. Induction of ptc1 and gli1 by Shh creates a negative feedback loop (through Ptc1) that restrains ongoing Shh signaling and a positive feedback loop (through Gli1) that acts as an amplifier of the initial Shh signal. The balance between negatively acting Ptc1 and positively acting Gli proteins determines the functional state of Shh pathway activity (4–7).

In the developing cerebellum, Shh signal produced by Purkinje neurons stimulates proliferation of GNPs in the external germinal layer (EGL) on the surface of the cerebellum (8–10). Shh target genes, such as N-myc and cyclin-D1, become active and promote cell cycle entry (11–13). In mice, expansion of GNPs ceases within 3 weeks after birth, as GNPs migrate to the internal granule cell layer and differentiate to form mature granule neurons (Fig. 1A). By the end of this period, the EGL has disappeared.

Mice heterozygous for ptc1 spontaneously form MB and rhabdomyosarcoma (RMS), a tumor derived from muscle (14, 15). These tumors exhibit constitutive Shh target gene expression due to reduced Ptc1 activity. Small molecule Shh pathway inhibitors can reduce tumor formation in ptc1+/− mouse models of MB (16, 17), demonstrating the dependence of MB cells on Shh target gene expression. Therefore, knowledge of critical Shh target genes or genes that cooperate with the Shh pathway to promote tumorigenesis has potential clinical importance as a step toward identifying drug targets.

Insulin-like growth factor 2 (igf2) is expressed at high levels compared with normal tissue in both MB and RMS in ptc1+/− mice (15) and in humans (18–20, 22). Hahn and colleagues (21) showed that igf2 is critical for the formation of both MB and RMS in ptc1+/− mice. When ptc1+/− mice are crossed into an igf2-deficient background, they no longer form MB or RMS such as their igf2-wild-type littermates. Igf2 has also been shown to synergize with the Shh pathway to promote MB, as retroviral transfer of Shh and Igf2 expression constructs into postnatal cerebella causes a higher incidence of MB formation than delivery of a Shh expression construct alone (23). Igf2 protein can stimulate proliferation of cultured MB cell lines and their precursor cells, GNPs, suggesting a mechanistic basis for the role of igf2 in MB tumorigenesis (24).

Important questions remain about the involvement of igf2 in MB. The mechanism of increased igf2 expression in MB and RMS from ptc1+/− mice is not clear. Two independent analyses of igf2 expression from maternal and paternal alleles showed that increased igf2 levels do not result from loss of imprinting in MB or RMS or from gene amplification (21, 24). Rather, increased igf2 RNA levels seemed to be induced at a purely transcriptional level. Hahn and colleagues (21) proposed that igf2 might be a Shh target gene, overexpressed in ptc1+/− tumors due to constitutive Shh pathway target gene transcription. In support of this hypothesis, the authors showed that compared with ptc1+/+ embryonic day 8.5 embryos, igf2 transcript levels were higher in ptc1+/− embryos and highest in ptc1−/+ embryos. C3H/10T1/2 cells transfected with the Shh pathway–activating transcription factor Gli1 have increased igf2 RNA levels after 72 hours (25). However, cultured GNPs treated with Shh do not have increased igf2 RNA levels at the 6-hours time point tested (24). Therefore, it remains unclear...
whether constitutive Shh pathway transcriptional activity causes the increased levels of igf2 transcript found in MB. It is also not clear what role igf2 plays in MB tumorigenesis. igf2 could contribute to tumor initiation, either as a critical Shh target gene or an independently regulated gene. Alternatively, igf2 could cooperate with the Shh pathway to cause tumor progression at a specific stage of tumorigenesis.

Here, we evaluate igf2 expression at multiple stages of MB tumorigenesis. We also test the requirement for igf2 during early and late stages of tumorigenesis. We show that igf2 is not required for tumor initiation but is required for progression to advanced MB lesions. Modulation of igf2 signaling can regulate proliferation of MB cells in vitro, suggesting that igf2 is important not just for tumor progression but also for tumor maintenance.

Materials and Methods

Mouse strains. Igf2 knockout mice were kindly provided by Dr. Argiris Efstatiadis (Columbia University, New York, NY). Because igf2 is an imprinted gene for which only the paternal gene is expressed, male mice heterozygous for igf2 were mated with ptc1+/−/− mice (108B2; ref. 14) with or without the Math1-gfp transgene. Math1-gfp mice were kindly provided by Dr. Jane Johnson (University of Texas Southwestern Medical Center, Dallas, TX). Offspring heterozygous for the igf2 knockout allele by PCR, using primers specific for wild-type and mutant igf2 alleles, were effectively igf2 null. Offspring homozygous for the wild-type igf2 allele were considered wild-type.

Math1-gfp ptc1+/−/− mice and igf2−/− littermates were sacrificed at 3 wk to evaluate for early MB formation. Cerebella were visualized under a fluorescent dissecting microscope for development of green fluorescent protein (GFP)-positive lesions. Adult ptc1+/−/− and ptc1+/−/−/− littermates were assessed for MB formation at 20 to 24 wk by visual inspection of cerebella and X-gal staining, as previously described (14).

Microarray analysis. Early and intermediate MB cells were isolated from cerebella of math1-gfp ptc1+/−/− mice at 3 to 6 wk or 10 to 20 wk, respectively. Tumors were identified by inspection of cerebella for GFP fluorescence. Tumors were dissected and GFP-expressing cells were isolated by fluorescence-activated cell sorting (FACS). Advanced MBs were isolated from symptomatic ptc1+/−/− mice and tumor cells were purified as previously described (10). RNA was isolated using Trizol reagent (Life Technologies). RNA was amplified using Riboamp kit (Arcturus). Probe was previously described (10). RNA was isolated using Trizol reagent (Life Technologies). RNA was amplified using Riboamp kit (Arcturus). Probe was previously described (10). RNA was isolated using Trizol reagent (Life Technologies). RNA was amplified using Riboamp kit (Arcturus). Probe was previously described (10). RNA was isolated using Trizol reagent (Life Technologies). RNA was amplified using Riboamp kit (Arcturus). Probe was previously described (10). RNA was isolated using Trizol reagent (Life Technologies). RNA was amplified using Riboamp kit (Arcturus).

Quantitative real-time reverse transcription-PCR. Igf2 and gli1 transcript levels were measured by real-time PCR using RNA isolated with Trizol reagent (Life Technologies). Gene expression assays for mouse igf2, gli1, and pgk1 and gapdh were purchased from Applied Biosystems. Igf2 and gli1 transcript levels were normalized to gapdh transcript levels in tumor cell, fibroblast, and PZp53MED cell experiments and to pgk1 transcript levels in GNP cell experiments.

Results

Igf2 transcription increases at a late stage of MB tumorigenesis. To observe developing MBs, we crossed ptc1+/−/− mice with mice expressing GFP from a Math1 enhancer that is specifically expressed in immature GPNs (26). Math1 is a critical regulator of GNP development that is required for the formation of GPNs (27). Because Math1 expression is preserved throughout MB tumorigenesis (28, 29), tumor cells from multiple stages of MB progression could be readily detected and isolated by virtue of their GFP expression. RNA from these cells was analyzed using cDNA microarrays to identify changes in gene expression that occur at each stage of MB tumorigenesis. We collected three tumor cell samples and analyzed them using cDNA microarrays. We found that igf2 transcription increases at a late stage of MB tumorigenesis. To observe developing MBs, we crossed ptc1+/−/− mice with mice expressing GFP from a Math1 enhancer that is specifically expressed in immature GPNs (26). Math1 is a critical regulator of GNP development that is required for the formation of GPNs (27). Because Math1 expression is preserved throughout MB tumorigenesis (28, 29), tumor cells from multiple stages of MB progression could be readily detected and isolated by virtue of their GFP expression. RNA from these cells was analyzed using cDNA microarrays to identify changes in gene expression that occur at each stage of MB tumorigenesis. We collected three tumor cell samples and analyzed them using cDNA microarrays.
populations for this analysis (Fig. 1A): (a) preneoplastic “early” MB lesions that are present in >85% of 3-week-old mice, the majority of which regress by 6 weeks; (b) intermediate, asymptomatic MB lesions that are present in ~15% of 10-week-old mice, the majority of which seem to progress to advanced MBs; and (c) advanced, symptomatic MB that occur in ~15% of mice, with peak incidence at ages ~20 weeks. For comparison, we used RNA from a purified population of GNPs, the precursor cells of MB, isolated from the EGL of normal postnatal day 7 (P7) cerebella.

We found that \( \text{igf2} \) transcript levels are not significantly increased in early MB lesions relative to normal GNPs. \( \text{igf2} \) transcript levels are significantly increased only in intermediate MBs and are even higher in advanced MBs (Fig. 1B). Only 20% of individually tested early MBs have significantly increased levels relative to normal GNPs, compared with 36% of intermediate and 100% of advanced MBs (Fig. 1C). To confirm these microarray results, \( \text{igf2} \) transcript levels from individual early, intermediate, and advanced MBs were measured by quantitative real-time PCR. Advanced MB samples showed a statistically significant increase (~20- to 50-fold) in \( \text{igf2} \) transcript levels (Fig. 1D). \( \text{igf2} \) transcript levels were not statistically increased in any of the early or intermediate MBs tested. Increased \( \text{igf2} \) transcription seems to be a late event in MB tumorigenesis and is correlated with the critical transition from early and intermediate to advanced MBs.

![Figure 1](image-url)

**Figure 1.** \( \text{igf2} \) transcription increases at a late stage of MB tumorigenesis. A, multiple stages of MB tumorigenesis in the \( \text{ptc1}^{+/−} \) mouse. GNPs (green) in the normal developing postnatal cerebellum proliferate in response to Shh produced by underlying Purkinje cell neurons (red). Throughout the first 3 wk of postnatal life, GNPs stop responding to Shh, differentiate, and migrate into the internal granule cell layer to become mature granule neurons (blue). GNPs express \( \text{Math1} \), whereas mature granule neurons do not. From age 3 wk to adulthood, no GNPs remain in the normal cerebellum, and \( \text{Math1} \) expression is lost. However, in the majority of \( \text{ptc1}^{+/−} \) mice (>85%) at age 3 wk, small clusters of \( \text{Math1} \)-expressing GNP-like cells persist as early, preneoplastic MB lesions. The majority of these lesions regress spontaneously by age 6 wk. A subset of these lesions (~15% of total) progress to form more aggressive-appearing intermediate, asymptomatic MBs by 10 wk and eventually advanced, symptomatic MBs by age ~20 wk. B, pooled averages of \( \text{igf2} \) transcript levels by microarray analysis of preneoplastic early MBs (early), intermediate MBs (inter), and advanced MBs (MB). Values are relative to normal P7 GNPs. Note that \( \text{igf2} \) transcript levels are not increased until late in tumorigenesis. *, \( P < 0.01 \). C, fold-enrichment of \( \text{igf2} \) transcript in individual MB lesions relative to normal GNPs. Each measurement is the average of five different \( \text{igf2} \) probes. *, \( P < 0.05 \). D, quantification of \( \text{igf2} \) transcript levels by quantitative real-time PCR from individual early MBs, intermediate MBs, and advanced MBs. Values are relative to normal P7 GNPs. *, \( P < 0.01 \). Columns, mean; bars, SD (B-D). \( P \) values were calculated using Student’s \( t \) test relative to normal P7 GNP controls.
In situ hybridization of early and advanced MB with a probe specific for *igf2* showed absent or low levels of *igf2* transcript in early MB but very high levels of transcript in advanced MBs (Fig. 2A and B). These data are consistent with our microarray data, confirming that increased *igf2* transcription is a late event in MB tumorigenesis. High levels of *igf2* transcript were also detected in the meninges of normal cerebella and cerebella containing tumors (Fig. 2A and B, arrows; Fig. 2C).

**Igf2** transcription can be regulated by the Shh pathway. The temporal pattern of *igf2* expression during MB tumorigenesis raises the question of how the induction of *igf2* expression is regulated. The expression pattern of *igf2* is different from what is expected and observed for most Shh target genes. Shh target genes such as ptc1 and gli1 are typically expressed at high levels in GNPs of the outer EGL, which proliferate in response to Shh produced by Purkinje neurons. In situ hybridization studies of developing P7 cerebella show high levels of *igf2* transcript in the meninges and low levels in the EGL (Fig. 2C). Conversely, gli1, a Shh target gene in all tissues, is absent from the meninges but abundantly transcribed in the outer EGL. We conclude that *igf2* transcript is minimally produced in areas of Shh target gene transcription in the developing cerebellum.

To further investigate the relationship between Shh pathway activity and *igf2* transcription, we examined the effect on *igf2* transcript levels of activating or inhibiting the Shh pathway in cultured GNPs and MB cells. Shh-treatment of primary GNP cultures does not increase *igf2* transcript levels after 6 hours (24). However, treatment of GNPs with Shh for 24 hours increased *igf2* transcription relative to untreated controls (Fig. 3A). Similarly, treatment of PZp53*MEF* cells, a MB cell line derived from a ptc1+/−; p53−/− mouse MB, with the Shh pathway inhibitor, cycloamine reduced *igf2* transcript levels relative to untreated controls (Fig. 3B). These results show that activation or inhibition of Shh target gene transcription can influence *igf2* mRNA levels in cultured MB cells and their precursors.

*Igf2* also behaves as a Shh-responsive gene in fibroblasts derived from ptc1+/+ or ptc1+/− embryos. When ptc1+/+ fibroblasts are transfected with Shh target gene–activating transcription factors gli1 or gli2, *igf2* transcription is induced >1,500-fold (Fig. 3C). Ptc1+/− embryos have reduced Ptc1 function and exhibit increased transcription of Shh target genes compared with ptc1+/+ embryos (14). Fibroblasts derived from ptc1+/− embryos have a 4-fold higher basal level of *igf2* transcription than ptc1+/+ embryonic fibroblasts (Fig. 3D). Cycloamine treatment does not change *igf2* transcript levels in ptc1+/+ fibroblasts but decreases *igf2* transcript levels in ptc1+/− fibroblasts to levels similar to those seen in ptc1+/− fibroblasts.

In summary, regulation of *igf2* by the Shh pathway is complex. Although *igf2* is a Shh-responsive gene in some cell types, it seems that *igf2* is refractory to induction by Shh in the developing cerebellum and in early MBs that express other Shh target genes.

**Igf2** is required for MB tumor progression but not tumor initiation. Hahn and colleagues (21) showed that *igf2* is required for formation of advanced MBs in ptc1+/− mice, but which stage of tumorigenesis requires *igf2* is unknown. Because *igf2* transcription is induced during the transition between early and advanced MBs, we tested whether *igf2* is required for early MB formation. The low level of *igf2* expression in early MBs suggests that early MB formation would not require *igf2* to be active within the pretumor cells. However, it is still possible that *igf2* might play a role in early MB tumor initiation because paracrine *igf2* signal produced in the neighboring meninges could stimulate GNPs or MB cells during early tumorigenesis. In this way, early lesions might be dependent on *igf2*, despite their lack of *igf2* expression. Later in tumorigenesis, when lesions become too large to subsist on meningeal *igf2* alone, *igf2* produced by the tumor cells themselves may supplement the meningeal source through autocrine signaling.

To distinguish between these possibilities, we crossed Math1-gfp ptc1+/− females with *igf2*+/− males and analyzed Math1-gfp
Igf2 could act at a specific point in the tumor progression decision or as an ongoing promoter of growth. We tested whether igf2 plays a role in maintaining the growth of fully developed MB cells. Proliferative and survival-promoting signals are important for maintaining tumor growth. It was shown previously that Igf2 protein can induce proliferation in GNP s (24). Similarly, we find that Igf2 increases the fraction of proliferating GNPs 3-fold (Fig. 4). We also show that Igf2 promotes cell survival of cultured GNPs. If GNPs are cultured in minimal medium, the majority undergoes apoptosis. GNPs cultured in minimal medium with Igf2 alone show enhanced survival similar to GNP s grown in fully supplemented medium (Fig. 4F).

We tested the effects of augmenting or inhibiting Igf2 signaling on cell proliferation of the MB cell line, PZp53MED. Addition of exogenous Igf2 increased proliferation of MB cells in a dose-dependent manner (Fig. 5A).

Igf2 ligand acts through the Igf1 receptor (Igf1R), which also can serve as a receptor for Igf1 and Insulin. We used an Igf1R-blocking antibody to inhibit the autocrine activity of endogenously produced Igf2. PZp53MED cells derived from a MB have low basal levels of Igf1R phosphorylation and phosphorylation of Igf pathway target Akt (Fig. 5B). Phosphorylation of Igf1R and Akt increased dramatically when treated with exogenous Igf2 protein. In contrast, only a minimal increase in phosphorylation was observed in Igf2-treated HE La controls when cells were treated with anti-Igf1R antibody and Igf2, levels of Igf1R and Akt phosphorylation are reduced to levels equal to or below baseline. Treatment of PZp53MED cells with anti-Igf1R antibody can therefore block the effects of even superphysiologic levels of Igf2.

When PZp53MED cells or HE La controls were cultured in the presence of anti-Igf1R antibody, PZp53MED cells showed a 40% decrease in cell number relative to untreated PZp53MED cells after 48 hours (Fig. 5C). There was no significant difference in cell number between treated and untreated HE La cells.

Blocking Igf1R can potentially also block the effects of Igf1 and Insulin. To more specifically block the effects of MB cell-produced Igf2, we used a soluble Igf2-Receptor (slg2R) construct that has been used previously to block the effects of Igf2 on cells or in vivo tissues (30, 31). When slg2R is expressed, it is secreted from the cell where it binds and sequesters the Igf2 ligand, blocking its effect. Because slg2R specifically binds Igf2, it does not interfere with the actions of insulin, Igf1, or Igf1R. When PZp53MED cells were transfected with slg2R, they showed a 60% reduction in proliferation, measured by BrdUrd incorporation, relative to CFP-transfected cells (Fig. 5D). These results indicate that Igf2 is specifically required for MB cell proliferation. In summary, we observe that activation or inhibition of Igf2 signaling can promote or prevent MB cell growth, respectively.

**Discussion**

The mechanism of igf2 induction in MB. Here, we present evidence that igf2 is a Shh-responsive gene in cultured GNP s, MB cells, and fibroblasts. It might seem that igf2 is expressed at high levels in MB s because these tumors have increased Shh target gene transcription. However, increased igf2 transcription is not seen until late stages of MB tumorigenesis, although constitutive Shh target gene transcription begins at early stages. Furthermore, increased igf2 transcription is not seen in developing GNPs of the outer EGL in vivo (Fig. 2B), which respond to Shh signal and express Shh target genes (10). Previous studies have indicated that neither genomic imprinting nor gene amplification are the causes of increased igf2 expression (21, 24). Rather, increased igf2 transcription seems to be due purely to transcriptional regulation.

One explanation for why igf2 transcription might be induced by Shh in cultured GNPs but not in GNPs or early tumor cells in vivo is that the level of Shh pathway activation achieved in culture may be higher than in vivo. The concentration of Shh protein used in our cultured assays causes maximal activation of Shh target gene transcription (10). If igf2 transcription requires very high levels of Shh pathway activity, then perhaps this threshold activity is not achieved until late in tumorigenesis. Alternatively, the presence of some inhibitory factor in the culture medium, or the absence of some activating factor that is present in the intact developing cerebellum may account for the differences in the response of GNPs to Shh in vitro versus in vivo. For example, restrictive chromatin
structure or the presence of repressive transcription factors in cells in vivo may not be reproduced when these cells are cultured in vitro.

Another explanation for the pattern of igf2 regulation could be that Shh target gene transcription alone cannot account for increased igf2 transcription in MB. Shh target gene transcription likely plays a role in increased igf2 transcription because igf2 transcript levels are decreased in cultured MB cells treated with the Shh pathway inhibitor cyclopamine, albeit only 2-fold. Another regulator may cooperate with the Shh pathway to induce igf2 transcription.

What might be responsible for the dramatic increase in igf2 transcription that occurs late in MB tumorigenesis? Increased igf2 expression due to altered transcriptional regulation in human tumors is a common finding, and perhaps such a mechanism cooperates with the Shh pathway to cause increased igf2 transcript levels in MB. Igf2 expression is induced by overexpression of the early growth response gene 1 protein in prostate cancer (32). Igf2 expression may be induced due to mutation of its flanking regulatory sequences. Changes in the 3’ untranslated region of igf2 are associated with 100- to 1,000-fold increases in igf2 transcript levels in colorectal cancer (33). Altered methylation of the igf2 promoter causes increased igf2 transcription in hepatoblastoma (34).

PTEN activity can modulate igf2 expression in hepatoma (35). Increased PTEN activity leads to decreased igf2 transcription, and decreased PTEN activity causes increased igf2 transcription. Because igf2 signaling antagonizes PTEN activity byactivating PI3-K (36), increases in Igf2 production may induce progressively increased autotranscription in feed-forward fashion. This hypothesis could explain why Hartmann and colleagues (24) did not see induction of igf2 transcription in GNPstreated with Shh alone for 6 hours but did see low-level induction of igf2 transcription in GNPstreated with Shh and Igf2. Shh target gene transcription and Igf2 production, first from the meninges and later from tumor cells themselves, may cooperate to induce igf2 transcription during MB tumorigenesis. A better understanding of the mechanism of increased igf2 expression in MB could yield opportunities to reverse this induction and inhibit the oncogenic effects of Igf2.

Temporal requirement for igf2 in late tumorigenesis. Our results suggest that induction of igf2 transcription is a late event in MB tumorigenesis that occurs as the preneoplastic cells of early MBs acquire a malignant phenotype. The requirement for Igf2 activity arises in late tumorigenesis, as early MB lesion formation occurs in the absence of igf2. These data do not preclude that igf2 may contribute to early tumorigenesis, as high levels of igf2 transcription are observed in the overlying meninges. However, the indispensable role for igf2 in MB formation occurs only in the late malignant conversion of MB and not during tumor initiation. This hypothesis is consistent with the observation that activation of Igf2 signaling alone in the developing cerebellum by retroviral transfer of igf2 is not sufficient to initiate MB formation (23).

This temporal pattern suggests a cooperation of autocrine Igf2 and other Shh target genes in MB tumorigenesis. Recent data have supported a cooperative hypothesis for these pathways. If Shh and Igf2 are introduced into neural progenitors by retroviral delivery, the frequency of MB formation is 3-fold greater than with delivery of Shh alone (23). The Shh and Igf2 pathways converge to trigger the production of high levels of N-myc protein. The Shh pathway stimulates transcription of N-myc, whereas the Igf pathway leads to stabilization of N-myc protein through Akt-mediated effects on GSK-3β (12, 13, 37). Consistent with this hypothesis, retroviral delivery of stabilized N-myc along with Shh also causes a 3-fold increase in MB frequency compared with Shh alone (38).

Igf2 may also cooperate with the Shh pathway in MB tumorigenesis by activating cell survival pathways. We show that Igf2 can promote GNP survival under conditions that normally lead to apoptosis. Similarly, the Igf family members, insulin and Igf1, can promote survival in cultured GNPsthat undergo apoptosis. Thus, the Igf family of proteins, along with insulin and Igf1, may contribute to early tumorigenesis, as high levels of igf2 expression is not sufficient to initiate MB formation (23).

A model for MB tumorigenesis. Based on our findings, we propose the following model for MB tumorigenesis. Normal GNPsthat undergo apoptosis, and affect other component. Cells with active Shh targets would proliferate and would resist cues to arrest and differentiate.

### Table 1. Early MB formation persists in Math1-gfp ptc1+/- igf2−/− mice

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NOTE: One hundred percent of ptc1+/- igf2+ and ptc1+/- igf2−/− mice sacrificed at age 3 wk had early MB lesions. However, although ptc1+/- igf2+ mice developed advanced MBs, no MBs were observed in ptc1+/- igf2−/− mice (P < 0.05).

**Figure 4.** Igf2 promotes GNP proliferation and survival. A, the percentage of GNPsproducing Igf2 versus control (con) for 48 h. B, Igf2 increases GNP survival under conditions which normally lead to apoptosis. If P7 GNPspreserved in media without supplement (no sup) for 48 h, a majority undergo apoptosis compared with those cultured in supplemented media (sup). When Igf2 is added to unsupplemented medium, GNP survival is restored. Columns, mean; bars, SD.
leading to formation of early MBs. Most of these lesions resolve or remain as benign lesions that are noted in most asymptomatic adult ptc1+/− mice. In an ill-fated subset of these mice, the pre-MB cells would undergo additional changes, the most critical being those that cause increased transcription of igf2. Increased autocrine Igf2 production would allow persistence of these lesions and their conversion to a malignant phenotype. Such a further transformation would be consistent with the effects of Igf2 on cell survival and growth. These lesions then progress to form the advanced MB seen in ~15% of ptc1+/− mice.

Given the apparent requirement of igf2 for progression to advanced MB, it is notable that when igf2 expression is analyzed in independent early MBs (Fig. 1C), a significant increase in igf2 transcript is observed in ~20% of lesions. This frequency is similar to the ~15% of mice that form advanced MB, suggesting that those early MBs that have induced igf2 transcription comprise the subset of lesions that will progress to advanced MB. Failure to induce igf2 transcription would cause these lesions to regress or remain benign. In summary, igf2 may be an important malignancy associated factor required for the progression to advanced MB, making it an attractive therapeutic target.

Igf2 as a therapeutic target in MB. Strategies for blocking Igf2 signaling for the treatment of human tumors, such as MB, might include inhibition of Igf1R with small molecule inhibitors or receptor-blocking antibodies. Blocking Igf1R offers the advantage of preventing signaling by both Igf1 and Igf2, which stimulate the same signaling cascades. Whether Igf1 contributes to ptc1+/− MB formation is unclear. Increased igf1 expression is noted sporadically in ptc1+/− MBs (data not shown) but at levels much lower than those of igf2. We do not observe substitution for Igf2 by Igf1 in igf2−/− ptc1+/− mice.

Our work shows the promise of specific targeting of the Igf2 ligand. We used a soluble Igf2R to inhibit MB cell proliferation, which presumably worked by sequestering Igf2. Recently, this same protein, produced from a transgene, was shown to inhibit tumor formation in a mouse model of intestinal adenoma associated with high Igf2 levels (31). Neutralizing antibodies specific to Igf2 could also achieve this end. Finally, manipulation of Igf-binding proteins (IgfBP) could block signaling by Igf ligands. For instance, IgfBP-3 can bind to and regulate the mitogenic activities of Igfs and inhibit their antiapoptotic effects (40).

Our data show a critical role for Igf2 in the progression and continued growth of MB in ptc1+/− mice. In human MB, dysregulation of the Shh pathway, often through loss of PTCH, is associated with a specific desmoplastic MB histology. Compared with classic MB, desmoplastic MBs have a distinct gene expression profile that includes increased expression of igf2 and Shh target genes (41). It is possible that Igf2 could hold similar importance in this class of human MB and that Igf2-directed therapies could provide clinical benefit.
Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Acknowledgments

Received 6/10/2008; revised 7/24/2008; accepted 8/21/2008.

Grant support: National Cancer Institute grant #RO1 CA 088060. R.B. Corcoran and M.T. Barakat were supported by the Stanford Medical Scientist Training Program NIH training grant. T. Bachar Raveh was supported by a Damon Runyan postdoctoral fellowship. M.T. Barakat was also supported by a National Science Foundation graduate research fellowship. E.Y. was supported by the NIH under Ruth L. Kirschstein National Research Service Award (F32CA117775-03) from the National Cancer Institute. M.P. Scott is an investigator of the Howard Hughes Medical Institute.

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We thank Drs. Argiris Efstratiadis and Jane Johnson for providing mouse strains, Dr. Susan McConnell for an in situ hybridization protocol, Dr. Bass Hassan for the sigfGR plasmid, Dr. Anna Penn for RNA from Gli1- and Gli2-transfected fibroblasts, Natalia Snarskaya for assistance with microarray and reverse transcription-PCR experiments, and Hermie Manuel for assistance with mouse husbandry.

References
2. Raffel C, Jenkins RB, Frederick L, et al. Sporadic postdoctoral fellowship. M.T. Barakat was also supported by a National Science Program NIH training grant. T. Bachar Raveh was supported by a Damon Runyan and M.T. Barakat were supported by the Stanford Medical Scientist Training Johnson RL, Rothman AL, Xie J, et al. Human homolog
3. 5.
4. Ingham PW, Taylor AM, Nakano Y. Role of the
5. 9.
6. Dahmane N, Ruizi Altaba A. Sonic hedgehog regulates
7. 12.
8. 10.
9. 11.
10. Kenney AM, Rowitch DH. Sonic hedgehog promotes
11. 17.
12. 16.
13. 15.

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Insulin-like Growth Factor 2 Is Required for Progression to Advanced Medulloblastoma in patched1 Heterozygous Mice

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