Analysis of hedgehog signaling in cerebellar granule cell precursors in a conditional Nsdhl allele demonstrates an essential role for cholesterol in postnatal CNS development

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Abstract

NSDHL is a 3β-hydroxysterol dehydrogenase that is involved in the removal of two C-4 methyl groups in one of the later steps of cholesterol biosynthesis. Mutations in the gene encoding the enzyme are responsible for the X-linked, male lethal mouse mutations bare patches and striated, as well as most cases of human CHILD syndrome. Rare, hypomorphic NSDHL mutations are also associated with X-linked intellectual disability in males with CK syndrome. Since hemizygous male mice with Nsdhl mutations die by midgestation, we generated a conditional targeted Nsdhl mutation (Nsdhltm1.1Hrm) to investigate the essential role of cholesterol in the early postnatal CNS. Ablation of Nsdhl in radial glia using GFAP-cre resulted in live-born, normal appearing affected male pups. However, the pups develop overt ataxia by postnatal day 8–10 and die shortly thereafter. Histological abnormalities include progressive loss of cortical and hippocampal neurons, as well as deficits in the proliferation and migration of cerebellar granule precursors and subsequent massive apoptosis of the cerebellar cortex. We replicated the granule cell precursor proliferation defect in vitro and demonstrate that it results from defective signaling by SHH. Furthermore, this defect is almost completely rescued by supplementation of the culture media with exogenous cholesterol, while methylsterol accumulation above the enzymatic block appears to be associated with increased cell death. These data support the absolute requirement for cholesterol synthesis in situ once the blood-brain-barrier forms and cholesterol transport to the fetus is abolished. They further emphasize the complex ramifications of cholesterogenic enzyme deficiency on cellular metabolism.
Introduction

Cholesterol is an essential component of all mammalian cell membranes and is a major determinant of plasma membrane fluidity. It is enriched in the mammalian central nervous system (CNS) where myelin contains ~80% of the cholesterol of the adult brain (1). The cholesterol biosynthetic pathway is also intimately tied to a variety of important cellular functions, including signal- ing in lipid rafts and the formation of steroid hormones, bile acids, vitamin D, meiosis-activating sterols and oxysterols. Iso- prenoid intermediates in the first half of the pathway serve as precursors for the synthesis of modified tRNAs, dolichol, ubiquinone and farnesyl and geranylgeranyl moieties (reviewed in 2). Finally, active hedgehog proteins are modified by the covalent addition of cholesterol during their intracellular processing and are involved in numerous developmental pathways (3). Perturbations of cholesterol metabolism have been implicated in a variety of human CNS disorders ranging from autism (4) to Alzheimer disease (5,6).

Cholesterol is synthesized in a series of ~30 enzymatic reactions (7,8). The condensation of the 30 carbon isoprenoid squa- lene forms the first sterol intermediate, lanosterol; subsequent enzymatic reactions define the ‘post-squalene’ half of the path- way (Fig. 1A). Human disorders and/or mouse models have now been described for each step in post-squalene cholesterol biosyn- thesis (9) and serve as a unique resource to help to understand the role of cholesterol in the developing CNS. All of the disorders are associated with major malformations and intellectual dis- ability (ID), providing further evidence for the essential role of cholesterol in the developing fetus. However, the pathogenic mechanisms responsible for the defects in these disorders remain unclear. In particular, it has been suggested by us and others that cholesterol deficiency during critical periods of embryonic

![Figure 1](https://academic.oup.com/hmg/article-abstract/24/10/2808/623190) Generation of a conditional Nsdhl allele. (A) Schematic diagram of the cholesterol biosynthesis pathway from lanosterol to cholesterol, with sterols listed in shaded boxes and the enzymes that catalyze each step shown next to the arrows. Nsdhl, along with SC4MOL and HSD17B7, is required for the removal of two C-4 methyl groups from 4,4-dimethylcholesta-8,24-dien-3-β-ol to generate zymosterol. Reduction of the C-24 double bond by DHCR24 can occur at multiple points along the pathway, but is shown only as the last step for simplicity. Ketoconazole inhibits CYP51A1 in the demethylation of lanosterol at C-14. Abbreviations: CYP51A1, cytochrome P450 lanosterol 14-demethylase; DHCR14, 3β-hydroxysterol-Δ7-reductase, LBR, lamin B receptor; SC4MOL, sterol C-4 methyloxidase-like; Nsdhl, NADH steroid dehydrogenase-like; HSD17B7, 7-dehydrocholesterol reductase; DHCR24, 3β-hydroxysteroid-Δ7-desaturase; DHCR7, 7-dehydrocholesterol reductase; T-MAS, 4,4-dimethylcholesta-8,24-dien-3-β-ol. (B) Experimental strategy for generating the NsdhlΔ knockout. The top line presents the mouse wild type Nsdhl gene from exon 4 to exon 8, indicating the position of diagnostic restrictions sites for Asp718 and BglI. The exons and introns are not drawn to scale. The position of probe sequences A and B used on Southern blots to detect homologous integration of the targeting construct are indicated. Below is a simplified map of the targeting construct generated in the vector pL451, showing loxP sites (black arrowheads) flanking Nsdhl exon 5, the neomycin-resistance gene (Neo) flanked by FRT sites (white arrowheads) for positive selection, and the thymidine kinase gene (TK) for negative selection. Homologous integration of the NsdhlΔ construct into the Nsdhl locus following electroporation into ES cells results in altered sizes of the Asp718 and BglI restriction fragments, as shown in the third diagram. Finally, the Neo cassette was excised by FLPo-mediated recombination in ES cell clones to generate the NsdhlΔ allele (bottom diagram). (C) Southern blots of genomic DNA from a WT and representative NsdhlΔ/Y neo-resistant ES clone digested with Asp718 and BglI, and hybridized with probes A and B, respectively, that lie outside of the sequence in the Δas construct. The targeted clone showed the expected changes in size of the diagnostic restriction fragments, demonstrating homologous integration into the Nsdhl locus. (D) A Western blot of total protein prepared from E9.5 male embryos from a NsdhlΔ/Y x Sox2-Cre cross probed with antibodies against Nsdhl (top) and β-tubulin (bottom) as a loading control. The NsdhlΔ control sample was from pooled cre-negative male embryos, and showed the expected 38 kDa wild type band for Nsdhl. No Nsdhl signal was detectable in the NsdhlΔ sample from pooled cre-positive male embryos. The higher level of β-tubulin signal in the NsdhlΔ sample is due to more total protein loaded than in the NsdhlΔ control.
or early postnatal development and/or accumulation of toxic sterol intermediates above an enzymatic block may be responsible, although convincing evidence remains lacking (9–11).

The CNS may be particularly susceptible to perturbations of cholesterol synthesis because of its high cholesterol content and complete dependence on de novo synthesis postnatally. Since cholesterol does not cross the blood-brain-barrier, a functional cholesterol biosynthetic pathway is necessary within the brain itself (starting at ~E10–11 in the mouse). At birth, ~90% of the cholesterol in the mouse brain is made within the CNS (12,13). During pre- and early postnatal neurogenesis, cholesterol synthesis occurs in both glia and neurons. Subsequently, it is generally accepted that most cholesterol destined for neurons is synthesized in astrocytes, with efflux from these cells mediated via apoE and ABCA1 and/or ABCG1 transporters. Neurons internalize the cholesterol as lipoprotein particles via LDL and related receptors (14).

NSDHL (NADH steroid dehydrogenase-like) is a 3β-hydroxysterol dehydrogenase involved in the removal of two C4 methyl groups from 4,4-dimethylcholesta-8,24-dien-3β-ol, a sterol intermediate derived from lanosterol (see Fig. 1A). Mutations in the murine Nsdhl gene are responsible for the X-linked dominant, male lethal mouse mutations bare patches (terol dehydrogenase involved in the removal of two C4 methyl fl)

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Nsdhllox-/+Y males from the cross of (Nsdhllox-/+Nsdhllox-) females X (hGFAP-cre+/) males were present in expected numbers (284/1114 or 25.5% of live-born pups) and appeared normal at birth. However, by P8 to P10, they exhibited overt ataxia, difficulty with ambulation, and subsequent weight loss (Supplementary Material, Fig. S1D). They died between P12–P18, probably secondary to defects in the cerebral cortex. On routine histology, at P0, we noted no NSDHL expression by IHC in most cortical and hippocampal neurons (Supplementary Material, Fig. S2E-G). We conclude that the phenotype in mice lacking exon 5 replicates the original Bpa1 null allele.

Ablation of Nsdhl in radial glia using GFAP-cre produces a dramatic cerebellar phenotype in affected males

Mice with the Nsdhllox-/+ allele were mated to transgenic animals that express cre-recombinase under the control of the human glial fibrillary acidic protein (GFAP) promoter, (FVB-Tg(GFAP-cre)25Mes, called hGFAP-cre) (21). In these mice, beginning at ~E13.5, cre-recombinase is expressed in radial glia in the ventricular zone of the developing CNS. These cells give rise to most differentiated neurons and glia of the cerebral cortex and cerebellum, although GFAP and, hence, cre-recombinase are not expressed in cerebellar Purkinje cells or epithelial cells of the choroid plexus. After early postnatal day (P) 7, cre expression occurs mostly in astrocytes. We established matings of homozygous Nsdhllox-/+ females and heterozygous hGFAP-cre males and monitored the pups for the presence of the cre transgene by PCR genotyping. The experiments described below were performed on progeny from homozygous Nsdhllox-/+ females derived by brother-sister mating of mice from 2 of the founder lines, derived from different ES cell clones, at the N2 or N3 backcross generation onto C57BL/6J. No phenotypic differences have been observed between the founder lines or using mice from different backcross generations. We performed our phenotypic analyses using deleted male mice (Nsdhllox-/+Y), to avoid variability related to random X-inactivation in heterozygous females. Floxed (Nsdhllox-/+Y), but not deleted, male littermates served as controls, avoiding potential sex differences that could be related to steroid hormones.

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hippocampal neurons present, particularly in the dentate gyrus (Supplementary Material, Fig. S2A and B). At P7, the Nsdh\(^{Δ5/Y}\) brains showed obvious defects in the cerebellum, hippocampus and cerebral cortex (Supplementary Material, Fig. S3). The cerebellum and hippocampus were smaller than normal, with reduced numbers and organization of granule neurons. More pycnotic nuclei were evident in the outer layer of the Nsdh\(^{Δ5/Y}\) cerebral cortex, suggesting elevated cell death.

In the mouse, most cerebral cortical and hippocampal neuronal proliferation, differentiation and migration are complete prior to birth. Cerebellar GCPs migrate from the rhombic lip adjacent to the fourth ventricle beginning at \(\sim\)E13.5 and cover the entire cerebellar surface prior to birth (24,25). However, unlike cortical neurons, they undergo extensive proliferation postnatally, as well as subsequent differentiation and migration. We focused our phenotypic and mechanistic analyses on the cerebellum because much of its well-studied growth and development occurs synchronously within the first 3 weeks after birth and involves relatively few cell types that are arranged in morphologically distinct layers. Moreover, relatively large, pure populations of GCPs can be isolated for in vitro analysis of both their proliferation and differentiation into neurons. These properties enabled us to perform studies examining the relative roles of cholesterol depletion and methylsterol accumulation in disease pathogenesis.

In the cerebellar cortex of Nsdh\(^{Δ5/Y}\) males, early postnatal development (P0–P4) appeared normal by routine histology, although Nsdh expression remained only in Purkinje cells, which do not express GFAP (21,26) (Supplementary Material, Fig. S3I and J). However, by P5, the external granule cell layer (EGL) in Nsdh\(^{Δ5/Y}\) males appeared thinner with fewer granule cell precursors (GCPs) present (Fig. 2A and B). The magnitude of this difference progressed such that by P8, there was a single layer of GCPs in the EGL in the most severely affected mutant pups. Although mutant cerebellar Purkinje cells expressed NSDHL, they did not form extensive dendritic trees compared with their wild-type littermates, and they eventually degenerated (Fig. 2C, Supplementary Material, Fig. S4A). Bergmann glia, which are specialized...
astrocytes of the cerebellar cortex, were fewer in number, shorter and appeared disorganized in cerebella from mutant males (Fig. 2D). These results are consistent with previous studies demonstrating that contact with (normal) granule cells and Bergmann glia is required for the proper alignment and orientation of the Purkinje cells, as well as for their dendritic arborization (26,27).

To determine whether the developmental timing of the EGL phenotype corresponds to a concurrent decrease in cholesterol levels, gas chromatograph-flame ionization detection (GC-FID) sterol analysis was performed on freshly isolated GCPs from Nsdh<sup>fl/fl</sup>/Y and Nsdh<sup>Δ/Δ</sup>/Y pups at P3, P5 and P7 (Fig. 2E). In P3 samples, cholesterol levels were not significantly different between mutants and controls. However, the Nsdh<sup>Δ/Δ</sup>/Y samples showed a 25% reduction in cholesterol concentration relative to controls at P5 and P7. At all three stages, the mutant GCPs showed greatly elevated concentrations of sterol intermediates, identified using full scan MS data as 4α-methylcholest-7(8)-en-3β-ol and 4α-methylcholesta-8(9),24-dien-3β-ol (see Supplementary Material, Fig. S5A for sample GC-FID chromatograms and Fig. S5D for sample mass spectrometry scan data), consistent with loss of NSDHL function. Both compounds were nearly undetectable in control samples. The relative content of desmosterol, which contains a residual C24 double bond and is present in high concentrations in the developing CNS (28), was similar in control and mutant samples at all three stages (Fig. 2D). This discrepancy may reflect differences in the turnover of pre-existing, maternally derived cholesterol versus desmosterol in vivo. Indeed, Jansen et al. demonstrated that desmosterol accumulation in the mouse brain peaks in the first postnatal week; they suggest that post-transcriptional repression of 3β-hydroxysterol 24-reductase may be responsible for this accumulation (28).

**Proliferation and migration defects in mutant cerebellar granule cells**

GCPs in the outer EGL proliferate extensively between P0 and P14. One granule cell generates ~250 progeny, and they account for ~50% of all CNS neurons in the adult (reviewed in 25). This proliferation peaks at P6–8 and requires sonic hedgehog (SHH) secreted by Purkinje cells (29,30). Post-mitotic granule cells begin to migrate inward from the surface and establish an inner EGL. Migration requires additional secreted factors, including BDNF (31). The mechanisms responsible for the switch of cerebellar GCPs from proliferation to differentiation and migration are not completely understood. They do require and are guided by the Bergmann glia, which also differentiate in response to SHH signaling (29,32).

The presence of a thinner EGL after P4 in Nsdh<sup>Δ/Δ</sup>/Y males suggested that there could be reduced proliferation and/or early differentiation and migration of mutant cerebellar GCPs, as found in several other mouse mutants with cerebellar phenotypes (26,33,34). As measured by phosphohistone H3 (PHH3) IHC (Fig. 3A and B) and BrdU incorporation (see Supplementary Material, S4C and D), we found progressively reduced proliferation of GCPs in mutant cerebella starting at P5. When apoptosis was measured using the TUNEL assay, a significant increase in TUNEL(+) cells was first observed at P7 in the Nsdh<sup>Δ/Δ</sup>/Y males relative to controls (Fig. 3C). Thus, the diminished EGL in the mutant cerebella was the result of reduced proliferation beginning at ~P5 followed by increased cell death at ~P7. By immunofluorescence with antibodies to marker proteins of differentiating neurons, such as TAG-1 (contactin 2) (35) (Supplementary Material, Fig. S4B), and DCX (doublecortin) (36) (not shown), some mutant GCPs did become post-mitotic, began to differentiate, and migrating to reside within the inner layer of the EGL and inner granule cell layer. However, Nsdh<sup>Δ/Δ</sup>/Y males also had a defect in GCP migration as shown by BrdU pulse-labeling experiments (Fig. 4). Mice were injected with BrdU at P6, and brains were collected either 2 or 48 h later and stained for BrdU incorporation. Labeling in the 2-h samples was mainly in the outer EGL (oEGL) of both Nsdh<sup>fl/fl</sup>/Y and Nsdh<sup>Δ/Δ</sup>/Y cerebella, as expected (Fig. 4A). In the samples collected after 48 h, a majority of the labeled cells had migrated to the inner EGL (iEGL) or IGL in the Nsdh<sup>fl/fl</sup>/Y samples, while most of the BrdU(+) cells remained in the oEGL in the Nsdh<sup>Δ/Δ</sup>/Y cerebella. Plotting the ratio of labeled cells in the IGL versus the oEGL in mutant and wild type cerebella at different labeling intervals showed that granule cell migration was not significantly different for cells born on P4, but migration of Nsdh<sup>Δ/Δ</sup>/Y cells was reduced at P5 and P6 (Fig. 4B).

A decreased response to SHH underlies the proliferation defect in mutant granule cell precursors

A clear advantage to studies involving the early postnatal cerebellum is the ability to replicate most aspects of neuronal proliferation, differentiation, and migration in vitro (37,38), including the proliferation of primary GCPs in response to exogenous SHH (39–41). GCPs isolated from mutant Nsdh<sup>Δ/Δ</sup>/Y mice at P4 and cultured for 48 h without added cholesterol demonstrated reduced proliferation in the presence of exogenous SHH compared with control Nsdh<sup>fl/fl</sup>/Y littermates as measured by BrdU incorporation (Fig. 5A and B). Furthermore, there were many fewer and thinner neurite extensions in the mutant cells compared with the controls after 24 h (not shown) or 48 h in culture (Fig. 5A). The addition of cholesterol to the primary GCP cultures led to almost complete restoration of the proliferative response to SHH (Fig. 5B) and much improved cell morphology (Fig. 5A).

Hedgehog signaling in recipient cells via the canonical pathway involves binding of ligand to the transmembrane receptor PTCH1, relieving its inhibition of smoothened (SMO). Release of SMO inhibition is associated with movement of the protein into the primary cilium and subsequent activation of GLI transcription factors (42,43). The major effector of cerebellar granule cell proliferation via canonical hedgehog signaling is GLI2 (32,44), although induction of GLI1 gene expression is more robust in vitro assays (45). SHH signaling also induces Nmyc expression which in turn activates D-type cyclins and progression through the G<sub>1</sub> phase of the cell cycle (39,40,46). Nmyc induction is independent of GLI expression, but downstream of smoothened (SMO) (40). Phosphorylation of multiple sites on NMYC via GSK-3 in vivo promotes its degradation, enabling cell cycle exit in the presence of continuous SHH (47,48).

SHH treatment of Nsdh<sup>fl/fl</sup>/Y cultured GCPs resulted in >500-fold and ~8-fold induction of GLI1 and GLI2 gene expression, respectively, while their induction was greatly reduced in Nsdh<sup>Δ/Δ</sup>/Y cultured cells (Fig. 6A). Growth of the SHH-treated mutant cells in the presence of exogenous cholesterol resulted in substantially increased GLI transcription, although the recovery of GLI1 expression was less dramatic than the proliferation response (compare Figs 5B and 6A). Transcription of GLI3 in GCPs is not responsive to SHH (49) and remained unchanged under all conditions in both wild type and mutant cells (not shown). Transcription of Ptc1 and Nmyc was induced ~2 and 1.3-fold, respectively, by SHH in control cells, but remained unchanged in mutant cultured cells. NMYC targets Ccnd1 and Ccnd2, encoding cyclins D1 and D2, respectively, were also induced upon SHH treatment with a significantly greater response in wild type versus mutant cells (Fig. 6A).
Addition of cholesterol to wild type or mutant cultured cells by itself did not affect the levels of transcription of any of the hedgehog-responsive genes tested (Fig. 6A). However, in the presence of added SHH, cholesterol resulted in variable, but uniformly, increased transcription of SHH and NMYC-induced genes in the mutant GCPs. The most dramatic response was found for N-Myc itself, where there was no longer a significant difference in levels of transcription between control and mutant cultured cells.

As with other cholesterol biosynthetic enzymes, NSDHL protein levels are high in neuronal cells in late gestation and the early postnatal period and then drop ∼6-fold by 10–12 weeks of age (50,51). There is also a surge in cholesterol synthesis in oligodendrocytes between 2 and 3 weeks of age, corresponding with axon myelination. As expected, the precise regulation of the cholesterol biosynthetic pathway resulted in upregulation of the two cholesterol biosynthetic genes examined, Hmgcr and Sc4mol, in mutant cells cultured without added cholesterol (Fig. 6A). Expression of these genes was significantly reduced in both WT and mutant cells with cholesterol supplementation (43% reduction in Sc4mol expression in WT and 26% in mutant cells; 26 and 17% reduction in Hmgcr expression in WT and mutant cells, respectively), indicating that the cells were able to incorporate and respond to free cholesterol. However, expression of Hmgcr and Sc4mol in the Nsdhl<sup>Δ5</sup>/Y GCPs remained well above that of WT cells, suggesting that their cellular cholesterol homeostasis had not been completely restored. Unexpectedly, SHH treatment alone resulted in lower expression of Hmgcr and Sc4mol in both WT and mutant cells compared with growth in serum-free media (SFM) alone. We attribute this finding to the differentiation of a majority of cultured GCPs into TuJ1-positive neurons that occurs in the absence of SHH, with more newly-formed membranes that require cholesterol as they form neurite extensions (not shown).

Although less dramatic, significant decreases in transcript levels of hedgehog response genes were also seen in experiments using whole cerebella isolated at P6 (Fig. 6B). The smaller in vivo effect sizes may be related to the presence of residual cholesterol available to the cells, as well as the presence of non-deleted cells within deep structures in mutant cerebella. Using mice that express lacZ under the control of the Gli1 promoter, we were also able to demonstrate variable, but reduced, induction of this SHH response gene in vivo (Supplementary Material, Fig. S6).

**Assessment of the effects of NSDHL deficiency on cholesterol homeostasis in cerebellar GCPs in vitro**

To gauge the relative contributions of cholesterol depletion and methylsterol accumulation to the cerebellar phenotypes associated with NSDHL deficiency, additional experiments were...
performed in which GCP cultures were treated with ketoconazole to block endogenous cholesterol biosynthesis. Ketoconazole inhibits lanosterol 14-alpha-demethylase (CYP51A1) in the conversion of lanosterol to 4,4-dimethylcholesta-8(9),14,24-trien-3β-ol, in a step upstream of NSDHL in the cholesterol biosynthetic pathway (see Fig. 1A). Thus, ketoconazole reduces both cholesterol synthesis and the accumulation of methylsterols associated with NSDHL inactivation. Human LDL was used as a soluble and easily measured source of exogenous cholesterol.

Initially, we performed dose response assessments to determine the minimal concentrations of LDL and ketoconazole necessary to observe an effect on the Gli1 response to an exogenous SHH signal. Gli1 transcript level was used as a sensitive measure of responsiveness to SHH because, of the genes that were tested, it showed the greatest range of response to exogenous SHH in cultured GCPs (Fig. 6A). For the experiments below, we used a concentration of ketoconazole of 2 μM, that results in >90% suppression of Gli1 expression and only minor alterations.

Figure 4. Migration of BrdU-labeled GCPs in Nsdhlflx5/Y and Nsdhldel5/Y cerebella. (A) Distribution of BrdU-labeled cells in the EGL of lobule 3 in Nsdhlflx5/Y and Nsdhldel5/Y cerebella at 2 and 48 h after BrdU injection on P6. The lighter staining of cells in the Nsdhlflx5/Y sample at P8 was likely due to dilution of incorporated BrdU by cell division that followed the initial labeling 48 h earlier. Note that a large majority of the labeled GCPs in the Nsdhlflx5/Y sample remained in the oEGL in the P8 sample, 48 h post-injection. Abbreviations: oEGL, outer external granule layer; iEGL, inner external granule layer; ML, molecular layer; IGL, inner granule cell layer. Scale bars: 100 μm. (B) The migration efficiency of Nsdhlflx5/Y and Nsdhldel5/Y GCPs at different ages. Migration was quantified by injecting pups with BrdU at P4, P5 or P6 and collecting cerebella for IHC 48 h later at P6, P7 or P8, respectively. The ratio of BrdU-labeled cells in the IGL to those remaining in the oEGL of lobule 3 was calculated for each sample (see Methods).
of morphology in isolated WT GCPs, and 5 µg/ml LDL, that fully restores Gli1 expression in mutant cells to WT levels (Fig. 7A) and produces much improved mutant cell morphology (Supplementary Material, Fig. S7). Conversely, treatment of WT GCPs with ketoconazole reduced Gli1 RNA by 10-fold, to a level similar to that of mutant cells. The presence of LDL completely abrogated the effect of ketoconazole on Gli1 levels in both WT and mutant cells. Ketoconazole alone at a concentration of 2 μM had no effect on the response of mutant cells to SHH, while higher doses of ketoconazole produced significant cellular toxicity even in WT cells (not shown).

We then determined the effect of ketoconazole and/or LDL on GCP viability by propidium iodide (PI) staining of live cells in situ after 48 h in culture. Mutant cells treated with SHH alone showed significantly lower survival (65%) than WT cells (90%) (Fig. 7B).

Figure 5. Response of cultured GCPs to SHH and exogenous cholesterol. (A) Immunofluorescent staining for BrdU (red) and neuronal marker TuJ1 (green), with DAPI (blue) counterstaining of Nsdhlflx5/Y and NsdhlΔ5/Y GCPs that were cultured for 48 h with either 1 μg/ml recombinant SHH alone or SHH plus 15 µg/ml cholesterol. BrdU was added to the samples 4 h before fixing. Note that there are fewer and thinner neurite extensions in the mutant NsdhlΔ5/Y cells than in the Nsdhlflx5/Y controls. Scale bars: 100 μm. (B) Four aliquots of isolated GCPs from individual Nsdhlflx5/Y and NsdhlΔ5/Y P4 cerebella were cultured for 48 h in either serum free medium (SFM) alone, SFM with 15 µg/ml cholesterol (cho), SFM with 1 µg/ml recombinant SHH, or SFM with 15 µg/ml cholesterol and 1 µg/ml SHH. BrdU was added to the cultures 4 h before fixing the cells. Cells were immunostained for BrdU and counterstained with DAPI. Values are the percentage of BrdU-labeled cells from the total (DAPI-stained) number of cells. Each bar represents the mean ± SEM of results from three independent experiments.

Figure 6. Expression of SHH target genes and genes for cholesterogenic enzymes in Nsdhlflx5/Y and NsdhlΔ5/Y cultured GCPs and cerebella. (A) qPCR analysis of gene expression in cultured GCPs isolated from Nsdhlflx5/Y and NsdhlΔ5/Y P4 cerebella that were cultured as described in Figure 5B. The relative quantity of mRNA from each gene was calculated from real-time PCR analysis of cDNA. Values were normalized to the untreated Nsdhlflx5 sample set equal to 1 for each gene. Each bar represents the mean ± SEM of results from three independent RNA samples that were each assayed in triplicate. (B) qPCR analysis of gene expression in Nsdhlflx5/Y and NsdhlΔ5/Y whole cerebella at P6, performed as in (A). Values represent the mean ± SEM of results for RNA samples from 3 cerebella that were each assayed in triplicate.

LDL supplementation increased the survival of mutant cells to a level comparable with that of WT GCPs. As was seen with pure cholesterol, LDL treatment also resulted in greatly improved
cell morphology in the mutant GCPs (Supplementary Material, Fig. S7). Ketoconazole treatment did not affect the viability of WT cells, but it significantly increased mutant cell viability (from 65 to 78%; Fig. 7B). Moreover, in the presence of ketoconazole, the mutant GCPs again showed a markedly improved morphology, with larger cell bodies and more extensive neurite outgrowth (Supplementary Material, Fig. S7). These data suggest that elevated methylsterol levels in mutant cells may directly affect cell viability.

In serum free media (SFM), in the presence of added SHH, measured cholesterol levels after 48 h in culture were 19.75 ± 2.9 SEM µg/mg protein in WT cell lysates and 15.05 ± 0.93 µg/mg protein in mutant GCPs lysates (74% of wild type, P = 0.13) (Fig. 7C). The lack of a significant difference in cholesterol levels may be attributable to the state of differentiation of WT versus mutant cells after SHH treatment and concomitant differences in cellular morphology. Specifically, since sterol levels were calculated relative to the total protein content of the samples and the dramatic reduction in neurites, cell surface membranes, and the size of the cell bodies of the mutant cells grown in SFM is accompanied by reduced levels of total proteins, the ratio of cholesterol levels to protein may underestimate the extent of cholesterol depletion (compare Supplementary Material, Fig. S7A and B, for example). Cholesterol levels for all of the cultures dropped significantly with ketoconazole treatment (Fig. 7C). As expected, ketoconazole-treated cultures showed increased levels of lanosterol as a result of CYP51A1 inhibition (Supplementary Material, Fig. S5C). The improved morphology in mutant cells (Supplementary Material, Fig. S7F) may reflect further reduced levels of methylsterols, since treatment with either cholesterol or ketoconazole resulted in significant decreases in the methylsterol levels (Fig. 7D). Finally, treatment with both LDL and ketoconazole produced the lowest levels of methylsterols in the mutant GCPs, similar to those in WT cells grown without either additive.

In contrast to the relatively small differences in desmosterol levels in freshly isolated WT versus mutant GCPs (Fig. 2E), we
noted much more dramatically reduced desmosterol levels in cultured mutant cells grown with or without LDL in comparison with cultured WT cells (Fig. 7E). WT desmosterol levels in the absence of exogenous LDL-cholesterol were similar to those seen in freshly isolated GCPs, suggesting a deficiency of desmosterol in the cultured mutant cells. We believe this deficiency reflects the high demand for cholesterol as the cultured mutant GCPs attempt to differentiate.

Taken together, these results demonstrate that cholesterol depletion is sufficient to account for virtually all of the reduced response of mutant GCPs to SHH in vitro. Although the reduction of methylsterol levels by ketoconazole treatment apparently improved the viability and cell morphology of the mutant GCPs, it had no effect on Gli1 expression (Fig. 7A).

Since sterols have been implicated in modulating the initial steps of hedgehog signaling in target cells by binding to PTCH1 and SMO (52,53), we asked whether the HH signaling defect in Nsdhl<sup>−/−</sup> Y GCPs could be bypassed by treating the cells with the smoothened agonist SAG, a synthetic small molecule that activates the HH signaling pathway by direct binding to SMO (54). Gli1 expression levels were measured in cultured GCPs that were isolated from individual cerebella at P4, split into 6 wells and treated with either SHH or SAG (Fig. 8). WT GCPs treated with SAG (100 nM) showed the same level of Gli1 expression as SHH-treated cells. Gli1 levels were reduced by 42% with ketoconazole treatment of SAG-stimulated WT GCPs, compared with a >10-fold reduction in SHH-stimulated cells. Thus, signaling mechanisms at or upstream of SMO appear to be more sensitive to depletion of cholesterol and/or oxysterols by ketoconazole than downstream steps. The response of Nsdhl<sup>−/−</sup> Y cells to SAG was more than 10-fold lower than that of wild type cells, similar to their diminished response to SHH. However, ketoconazole increased Gli1 levels by ~4-fold in SAG-treated mutant cells (P = 0.029), in marked contrast to the lack of an effect of ketoconazole on their response to SHH. Note that the Gli1 response of the mutant cells to stimulation with SAG was inversely proportional to the levels of C4 methylsterols that were measured in ketoconazole-treated cultures (Fig. 7D). This suggests that the deleterious effects of methylsterols, that include reduced viability and reduced neurite outgrowth, may also affect HH signaling at steps downstream of SMO. However, in the presence of ketoconazole, Gli1 levels were still more than 2-fold lower in mutant cells than WT cells. This may be due to the fact that in the presence of ketoconazole, while methylsterol levels were reduced in the mutant cells, they were still ~10-fold higher than in ketoconazole-treated WT cells (Fig. 7D). Finally, LDL treatment increased Gli1 expression in SAG-treated mutant cells to levels similar to those in WT cells, likely due to both restoring normal cholesterol levels and further reducing methyl sterols to the levels seen in untreated wild type cells (Fig. 7D). In summary, a comparison of results from SHH versus SAG stimulation of the HH signaling pathway in GCPs confirmed that steps at or above SMO are more sensitive to sterol depletion than downstream steps. Furthermore, by circumventing the sterol-sensitive steps with SAG treatment, we uncovered an effect of elevated methylsterol levels in Nsdhl<sup>−/−</sup> Y cells that also reduces the efficiency of HH signaling by unknown mechanisms downstream of SMO.

Finally, we asked whether commercially available testicular meiosis-activating sterol (T-MAS; 4,4-dimethyl-5α-cholesta-8(9),24-dien-3β-ol) would affect HH signaling in WT GCPs. T-MAS and the related follicular fluid meiosis-activating sterol (FF-MAS; 4,4-dimethyl-5α-cholesta-8(9),14,24-trien-3β-ol) are naturally occurring 4α,4-methyl sterols that accumulate upon inhibition of the C4-demethylase complex (see Fig. 1A). They were originally identified as compounds that produced resumption of meiosis in oocytes in vitro (55,56). Pure T-MAS was mixed with fatty acid-depleted BSA as a carrier to form a water-soluble complex that was added to the culture medium. T-MAS had no effect on the level of Gli1 expression in response to SHH (Fig. 9A). GC-FID sterol analysis of cell lysates indicated that T-MAS was incorporated by the cells in a dose-dependent manner (Fig. 9B) although its intracellular localization relative to endogenous sterol intermediates was not determined. The 2.5 μg/ml treatment resulted in a cellular T-MAS level similar to that of methylsterols in untreated Nsdhl<sup>−/−</sup> Y GCPs (3–4 μg/mg protein, Fig. 7D). The GCPs treated with 10 μg/ml T-MAS showed an abnormal morphology, with smaller cell bodies and thinner neurites, reminiscent of Nsdhl<sup>−/−</sup> Y cells (Supplementary Material, Fig. S8). Furthermore, T-MAS treatment at 10 μg/ml produced significantly reduced cell viability in WT GCPs (Fig. 9D). This effect was less dramatic than in mutant cells with elevated levels of endogenous methylsterols (Fig. 7B), perhaps because WT cells should be able to convert accessible T-MAS to less toxic intermediates, as well as cholesterol itself. In addition, the mutant cells that are cholesterol deficient may be more sensitive to methylsterol toxicity.

![Figure 8. Comparison of Gli1 relative expression levels in response to SHH or SAG between Nsdhl<sup>+/+</sup> Y and Nsdhl<sup>−/−</sup> Y GCPs cultured in vitro. GCPs were isolated from individual cerebella at P4, split into 6 culture wells with the indicated treatments and analyzed after 48 h in culture. Concentrations of ketoconazole and LDL were as in Figure 7. The results are from 4 independent samples of each genotype that were each assayed in triplicate by qPCR with Gapdh as the endogenous control and normalized to the untreated Nsdhl<sup>+/+</sup> sample set equal to 1.](https://academic.oup.com/hmg/article-abstract/24/10/2808/623190)

**Discussion**

We have developed the first conditional mouse model for a disorder of post-squalene cholesterol biosynthesis, although a conditional Flot1 allele encoding squalene synthase (SQS) has been reported (57). In our study, we tested the utility of the conditional Nsdhl allele by investigating the effects of defective cholesterol synthesis in vivo in the developing CNS. Ablation of Nsdhl in radial...
glia at E13.5 using the hGFAP-cre transgene resulted in postnatal lethality between P12 and P20. The expected loss of NSDHL in neuronal lineages and astrocytes of the cerebrum, hippocampus, and cerebellum was confirmed by IHC. Histologic differences among affected brain regions were evident at birth, with the mutant hippocampus already disorganized, while the cerebellum showed no obvious phenotype (Supplementary Material, Fig. S2). Subsequently, between P5 and P8, cerebellar GCPs underwent a rapid sequence of reduced proliferation, impaired migration and, finally, massive cell death. Abnormalities were also noted in the Bergmann glia (Fig. 2D) and in Purkinje cells that continue to express NSDHL. The latter displayed deficient arborization and dendrite outgrowth and eventual cell death (Fig. 2C and Supplementary material, Fig. S4A). We also noted some delays in cerebellar lobeule development by P7 (see Supplementary Material, Fig. S4C and D and Fig. 6), although we did not observe any obvious patterning defects in mutant cerebella. Folia number and patterning are SHH dependent; however, they are determined in late gestation (44,58), when sufficient cholesterol should still be available in the mutant CNS.

All of the known human disorders and mouse models of post-squalene cholesterol biosynthesis are associated with major somatic and/or CNS malformations, suggesting that perturbations of the pathway have potent teratogenic effects. Microcephaly and ID are also features in surviving infants with all of the human sterol synthesis pathway disorders, except for affected heterozygous females with X-linked CHILD syndrome and X-linked dominant chondrodysplasia punctata (9,18). We and others have proposed that decreased cholesterol or total sterols; accumulation of toxic sterol intermediates proximal to the enzymatic block; abnormal feedback regulation of earlier steps in the pathway; generation of abnormal bioactive oxysterols; and/or abnormal signaling by hedgehog proteins that normally contain bound cholesterol, may all play a role in disease pathogenesis, although exact mechanism(s) remain to be determined. It is very likely that multiple mechanisms occurring at different stages of development and/or in different tissues are involved in disease pathogenesis, and data are beginning to accumulate to support some of these mechanisms. Some of the teratogenic effects in these disorders likely result from a lack of cholesterol for incorporation into membranes. Methylated sterol intermediates, in particular, are unable to fulfill the role of cholesterol within membranes. They result in dramatic changes in fluidity within the lipid bilayers (59–61), and they are not incorporated well into lipid rafts (60,61). However, while there is some overlap in the phenotypes among all of the human and murine disorders, unique features of each would argue against cholesterol deficiency as the only mechanism.

Cholesterol levels in isolated Nsdhl<sup>fl/fl</sup>/Y GCPs were normal at P3, but became progressively lower than WT in P5 and P7 samples. We believe this reduction largely reflects depletion of maternal and fetal cholesterol supplied to the CNS prior to birth. The timing of cholesterol depletion in these cells corresponded closely to the appearance of defects in the EGL, with the earliest cerebellar phenotype detected in Nsdhl<sup>fl/fl</sup>/Y pups being decreased GCP proliferation at P5 that preceded increased apoptosis in these cells by approximately two days.

Cerebellar GCP proliferation is driven largely by SHH secreted by underlying Purkinje cells (29,30). NSDHL was not deficient in Purkinje cells of Nsdhl<sup>fl/fl</sup>/Y pups, and these cells appeared morphologically normal until P7. Thus, the proliferation defect is more likely to result from reduced response to SHH in the GCPs rather than impaired signaling from Purkinje cells. Indeed, isolated Nsdhl<sup>fl/fl</sup>/Y GCPs in vitro showed a markedly reduced response to exogenous SHH with regard to both proliferation and expression of SHH target genes, and addition of exogenous cholesterol restored this proliferative response to SHH to near wild-type levels (Fig. 5).

The hypothesis that reduced hedgehog signaling contributes to the pathology associated with genetic disorders of cholesterol biosynthesis was originally based on the fact that proper signaling by hedgehog ligands in vivo is, in part, dependent on their covalent modification by cholesterol (3). However, to date, no evidence has emerged to directly demonstrate that processing and modification of hedgehog ligands are affected in these disorders. Rather, Cooper et al. (62) demonstrated defects in the response of mutant cells to hedgehog signaling, likely at the level of SMO, in mouse embryonic fibroblasts (MEFs) derived from Dhcr7<sup>−/−</sup> and Sc5d<sup>−/−</sup> mice that are murine models of Smith-Lemli-Opitz syndrome and lathosterolosis, respectively. In addition, 7-dehydrocholesterol can substitute for cholesterol in SHH processing and activation (62). Furthermore, an active N-terminal hedgehog peptide, which is not modified by cholesterol, can be secreted and signal in recipient cells, although the signal may spread further extracellularly (63,64). We have also found normal processing of SHH in NSDHL-deficient MEFs, as well as in mutant Nsdhl<sup>−/−</sup> male embryos at E10.5 (F. Jiang and G.E. Herman, unpublished results).

A reduced response to hedgehog signaling in recipient cells in vivo has been demonstrated in NSDHL-deficient placentalas by...
our group (65), as well as in the rudolph (rud) mutant mouse that carries a hypomorphic allele of Hsd17b7 (66), encoding a 3β-ketosterol reductase that functions with NSDHL and a methyloxidase, SC4MOL, in the sterol C4 demethylation complex (see Fig. 1A). Rud embryos display severe CNS and skeletal defects, and do not survive past birth. Cultured MEFs derived from rud embryos show a reduced response to exogenous SHH in both proliferation and expression of Gli1 and Pcth1. In vivo, rud embryos display lower expression of a Ptc-LacZ reporter in the forebrain and developing long bones. PTC1H and SMO show normal translocation to the primary cilium of rud MEFs in response to SHH, and cholesterol levels in rud embryos are reported to be normal. Thus, the authors propose that the hedgehog-related defects result from impaired activation of SMO, most likely due to a deficiency of activating oxysterols or to inhibitory effects of accumulating sterol intermediates. Indeed, selected oxysterols can activate hedgehog signaling in vitro (67), and cause translocation of PTC1H and SMO to the cilium in the absence of ligand (68).

The exact mechanism by which inactivation of Nsdhl disrupts hedgehog signaling has not been conclusively established. However, the lack of GCP proliferation or the lack of a Gli1 response to SHH in cultured mutant cells that is rescued by exogenous cholesterol (Figs 5 and 6) suggests that product deficiency is the primary abnormality. In addition, significant reduction of methylsterols using ketoconazole (Fig. 7A and D) did not restore the Gli1 response. Further evidence against accumulation of methylsterols being the primary mechanism is the lack of an effect of the 4-methylsterol T-MAS on Gli1 expression in cultured wild-type cells (Fig. 9). However, methylsterol accumulation is clearly associated with altered cell morphology and cell death and may be responsible for the massive apoptosis seen in the cerebellum in vivo. These abnormalities are also unlikely to be SHH-related. Although previous studies have shown that blocked hedgehog signaling can induce cell death in neuronal lineages in vitro model systems (69,70), conditional knockout of several genes involved in hedgehog signaling in the cerebellum causes reduced GCP proliferation without a subsequent increase in apoptosis (26,44). Moreover, apoptosis in Nsdhl<sup>-/-</sup>Y cerebella occurred throughout the inner granule layer (IGL) as well, where hedgehog signaling is not thought to be active.

We were also able to demonstrate differences in the response of cultured mutant GCPs to SHH versus the SMO agonist SAG. Ketoconazole could partially restore the Gli1 response of mutant cells treated with SAG, while it had no effect on the SHH-stimulated signal (Fig. 8). These data separate, at least partially, the cellular response to cholesterol depletion and methylsterol accumulation and are consistent with the model proposed by Stottmann et al. (66) for rud embryos of an effect at the level of SMO, possibly resulting from methylsterol accumulation.

Methylsterol accumulation has also been implicated in the pathogenesis of other cholesterol synthesis, as well as acquired, disorders. In CK syndrome, all of the affected males tested had normal sterol profiles, total cholesterol and steroid hormone levels, and lipoprotein profiles in plasma (10,19). However, when mutant lymphoblasts were cultured in lipid-depleted media, accumulation of 4-methylsters, similar to that seen in CHILD syndrome, was found. Furthermore, in the only affected male tested, the CSF cholesterol level was normal, while corresponding 4-methylsterol levels were increased (10). These data led the authors to speculate that accumulation of methylsters, rather than primary deficiency of cholesterol, caused CK syndrome. In the first patient described with autosomal recessive SC4MOL deficiency, another disorder with C4 methylsterol accumulation (11), the phenotype included severe psoriasiform dermatitis, arthropaligias, immune dysfunction, congenital cataracts, short stature, microcephaly, and ID. The serum cholesterol was low at 85 mg/dl (normal 140–176 mg/dl) with markedly elevated levels of 4,4'-dimethyl and 4α-monomethyl sterols detected in plasma, as well as in cultured skin fibroblasts grown in complete or lipid-depleted media. There was also increased proliferation of affected skin fibroblasts grown in lipid-depleted media that the authors attributed to the ‘meiosis-activating effects’ of the sterol metabolites that accumulate (55,56). They postulated that the hyperproliferative skin phenotype in disorders of the C4 demethylation complex results from elevations of these sterol intermediates. Finally, Astsaturov and colleagues identified SC4MOL and NSDHL in an siRNA screen for proteins that sensitize tumor cells to EGFR inhibitors, with the goal of improving cancer therapy (71). Depleting either enzyme in the presence of EGFR inhibitors led to increased apoptosis in several cancer cell lines, while other enzymes in the pathway did not produce similar effects. Depletion of the upstream enzyme CYP51A1 (see Fig. 1A) using siRNA or ketoconazole eliminated the effect, while added cholesterol did not. These authors also proposed that accumulation of 4-methylsterol is specifically responsible for this sensitization and further demonstrated that the loss of SC4MOL or NSDHL led to altered EGFR trafficking with higher degradation rates within the cell lines examined.

The reduced migration of Nsdhl<sup>-/-</sup>/Y GCPs that was observed beginning around P5, is also unlikely to be related to abnormal hedgehog signaling since it has not been implicated in this process in GCPs. The tyrosine kinase receptor TrkB and its ligand, BDNF, are both expressed by GCPs; both are required for proper migration, among several other factors. Recent data suggest that, in rodent cortex and hippocampus, BDNF recruits TrkB to intracellular lipid rafts upon binding, while a small pool of TrkB in plasma membrane rafts is unchanged (72,73). Trafficking of TrkB into early endosomes, with subsequent activation of PI3K/AKT and Rac/cdc42, is necessary for the GCP chemotactic response to brain-derived neurotrophic factor (BDNF) (74). Given the recent finding by Sukhanova et al. (71) that NSDHL depletion alters trafficking of another receptor tyrosine kinase (EGFR), it is possible that a similar mechanism is causative here. Future studies examining the migration of control and mutant GCPs in vitro in the presence of a BDNF gradient could address this issue. It should be noted that, while disruption of several signaling pathways is possible, and, perhaps likely, we do not believe that there is a general disruption of all membrane signaling functions, since expression of Hes1, a Notch gene target involved in GCP proliferation in the cerebellum (75), was unaffected in cultured GCPs from mutant cells (not shown).

It is also possible that the migration abnormalities could arise from defects in the Bergmann glia. Postmitotic GCPs of the inner EGL first migrate tangentially, parallel to the pial surface, and then migrate radially inward along Bergmann glia processes to reach the IGL (25). During tangential migration, GCPs extend leading and trailing processes. For radial migration, a third process extends inward through the developing molecular layer, while the parallel fibers are retained and form synapses with Purkinje cell dendrites. Given the greatly reduced neurite outgrowth of Nsdhl<sup>-/-</sup>/Y GCPs during differentiation in vitro, it is possible that extension of processes from the mutant cells is likewise restricted in vivo, thus preventing normal migration. Neuronal growth cone motility is dependent on cholesterol-rich lipid rafts that are thought to organize signaling molecules that regulate adhesion and cytoskeletal dynamics during cell migration (76). Likewise, the Bergmann glial processes that appeared abnormal and fewer in number in Nsdhl<sup>-/-</sup>/Y P6 cerebella may provide a...
poor substrate for radial migration of GCPs. Migration of GCPs along Bergmann glia also involves the formation of gap junctions and intercellular signaling, that may be affected by disruption of normal membrane organization of lipid rafts in cholesterol-deficient cells (77,78).

In summary, we have demonstrated a novel, early postnatal lethal phenotype in a conditional allele of the cholesterol biosynthetic enzyme NSDHL. Our studies of the early postnatal cerebellum in vivo and in vitro demonstrate severe defects in GCP proliferation and hedgehog signaling that we believe are due primarily to cholesterol deficiency or depletion. In vitro studies in cultured GCPs also demonstrate a likely role for methylster accumulation in cell death, consistent with the work of others in cells (71). Further studies will be necessary to explore the pathogenesis of the migration defects. The Nsdhl<sup>fl/fl</sup> conditional allele should be extremely valuable to investigate perturbations in cholesterol homeostasis in other regions of the brain, as well as in adult and aging mice.

Materials and Methods

Nsdhl<sup>fl/fl</sup> allele

The Nsdhl<sup>fl/fl</sup> targeting vector was constructed by inserting three PCR products with engineered restriction sites into the vector pL451 that contains a neomycin selection cassette (Neo) flanked by Frt sites (22). The PCR products were amplified from BAC clone 44089, that includes the Nsdhl gene and was obtained from the RPC-22 BAC library of mouse 129S6/SvEvTac genomic DNA (79). First, an 824 bp PCR product that included Nsdhl exon 5 with flanking intronic sequence and an engineered LoxP site was inserted into the BamHI site downstream of the vector LoxP site. Second, a 1.7 kb fragment from intron 4 was inserted into the HindIII site upstream of the Neo cassette to provide a 5′ homology arm. Third, a 3.7 kb fragment that included exons 6 and 7 with flanking intronic sequence was inserted into the NotI site downstream of the floxed exon 5 fragment to provide a 3′ homology arm. Finally, a viral thymidine kinase (HSV-tk) expression cassette was inserted into the ClaI site upstream of the 5′ homology arm to provide negative selection against non-homologous recombination events. The wild type sequence of the Nsdhl exons and loxP sites in the final construct was verified by DNA sequencing. The construct was linearized with Sall and electroporated into TC1 (129/SvEvTac) ES cells by our institutional ES Cell Core laboratory. G418-resistant clones were isolated and replicated cultures were grown in the presence of ganciclovir to identify HSV-tk<sup>-</sup>-<span class="symbol" id="r1" class="r1">−</span> clones. Southern blot analysis of DNA from selected ES cells digested with Asp718 or BglII was used to identify clones with homologous integration of the Flx5 construct. Three properly targeted clones were electroporated with a plasmid (FLPo) that expresses a codon-optimized FLP recombinase (80) to remove the Neo cassette. Two Neo<sup>-</sup>-<span class="symbol" id="r1" class="r1">−</span> clones were injected into C57BL/6 blastocysts to generate chimeric founder animals. Three of the resulting male founders with a high percentage ES cell contribution transmitted the Nsdhl<sup>fl/fl</sup> allele to female offspring.

Mice

Mice homozygous for the Nsdhl<sup>fl/fl</sup> allele (Nsdhl<sup>fl/fl;1.1Hm</sup>) were maintained on a mixed B6;129-Nsdhl<sup>fl/fl</sup> background and also backcrossed onto C57BL/6. GFAP-cre [FVB-Tg(GFAP-cre)25Mes/J, stock #004600] and Gli<sup>1lacZ</sup> (Gli<sup>sensAl</sup>/J, stock #008211) mice were obtained from The Jackson Laboratory (Bar Harbor, ME). GFAP-cre animals were maintained as heterozygotes on an FVB background by brother-sister mating. GFAP-cre heterozygous males were crossed to Nsdhl<sup>fl/fl;Nsdhl<sup>fl/fl</sup></sup> females to produce the Nsdhl<sup>fl/fl</sup>/GFAP-cre and Nsdhl<sup>fl/fl;Y</sup> males that were used for experiments. Gli<sup>1lacZ</sup> animals were backcrossed to C57BL/6 and maintained as heterozygotes. GFAP-cre males were crossed to Gli<sup>1lacZ</sup> females to generate GFAP-cre/Gli<sup>1lacZ</sup> compound heterozygous males that were then crossed to Nsdhl<sup>fl/fl</sup>/Nsdhl<sup>fl/fl</sup> females to produce Nsdhl<sup>fl/fl</sup>/GFAP-cre/Gli<sup>1lacZ</sup> and Nsdhl<sup>fl/fl</sup>/Gli<sup>1lacZ</sup> mice that were used for Gli<sup>1lacZ</sup> reporter expression analysis (see below). Sox2-cre mice were obtained from the Jackson Laboratory (stock #014094) and were maintained in the Research Institute ES Cell Core. For studies described here, the Sox2-cre males were on a mixed background (immunohistochemistry) or at N5 generation of a backcross onto C57BL/6 (Western blotting). Genotyping of mice was performed by PCR on DNA from toe clips using primers and PCR conditions as recommended by the Jackson Laboratory or as shown in Supplementary Material, Table S1. The housing and handling of mice conformed to protocols approved by the Institutional Animal Care and Use Committee of the Research Institute at Nationwide Children’s Hospital.

Antibodies

Primary antibodies used for immunoblotting and immunohistochemistry were: NSDHL (rabbit polyclonal, 50), Brdu (rat monoclonal, AbD Serotec, Raleigh NC), GFAP (mouse monoclonal, Thermo-Fisher Scientific, Fremont CA.), TUJ1 (mouse monoclonal, Covance, Emeryville, CA), phosphohistone H3 (PHH3) (rabbit polyclonal, Cell Signaling, Danvers, MA) calbindin (rabbit polyclonal, Millipore, Temecula CA), TAG-1 (Developmental Hybridoma Bank, Iowa City, IO) and β-tubulin (rabbit polyclonal, Thermo Scientific, Fremont, CA). Secondary antibodies were: HRP-linked goat anti-rabbit IgG (Cell Signaling), DyLight 488-conjugated donkey anti-mouse IgG, DyLight 549-conjugated donkey anti-rat IgG, and DyLight 549-conjugated donkey anti-rabbit IgG (Jackson ImmunoResearch, West Grove, PA).

Histology and immunohistochemistry

Preparation of mouse tissues for histology, immunohistochemistry and hematoxylin/eosin staining was performed as previously described (50). Briefly, tissues were fixed in either Bouin’s fixative or 4% paraformaldehyde (PFA) in PBS by whole-body transcardial perfusion. Primary antibody dilutions were: NSDHL (1:1000), PHH3 (1:250), calbindin (1:3000), Brdu (1:1000), GFAP (1:250), TAG-1 (1:10), and TUJ1 (1:1000). All immunohistochemistry samples were counterstained with hematoxylin. For Brdu labeling in vivo, mice received an intraperitoneal injection of Brdu (100 μg/ml in PBS) at a dose of 1 μg/g body weight at designated times. For immunodetection of Brdu, brains fixed with Bouin’s fixative were embedded in paraffin and 5 μm sections were mounted on Fisherbrand Superfrost Plus slides (Fisher Scientific). Sections were deparaffinized and incubated in 3% hydrogen peroxide, 10% methanol in PBS for 30 min, followed by a 1 h incubation in 200 μg/ml pepsin A ( Worthington, Lakewood, NJ), 0.01N HCl in PBS. The samples were denatured in 2N HCl for 1 h at 37° C, followed by 10 min of incubation in 0.1 M sodium borate, pH 8.5. After rinsing in PBS, samples were incubated in blocking solution [2% normal goat serum in TBST (50 mM Tris, 150 mM NaCl, 0.25% Triton X-100) for 2 h. Sections were incubated overnight at 4°C with rat anti-BrdU antibody in blocking solution. After washing with PBS, samples were incubated for 4 h with
biotinylated anti-rat IgG secondary antibody, followed by HRP-conjugated streptavidin using a Vectastain Elite ABC kit (Vector Laboratories). Signal was detected using VECTOR NovaRED peroxidase substrate kit (Vector Laboratories, Burlingame, CA). For TUNEL assays, brains were fixed by perfusion with 4% PFA, and sections were assayed using the ApopTag Peroxidase In Situ Apoptosis Detection kit according to the manufacturer’s instructions (Millipore). Stained samples were photographed using SPOT RT Color digital cameras operated by SPOT version 3.5 software (Diagnostic Instruments, Sterling Heights, MI) mounted on a Nikon SMZ-10A dissecting microscope or a Nikon Eclipse E800 microscope. Areas of the EGL and IGL were measured using SPOT software. Quantification of PHH3-positive cells and BrdU-labeled cells was performed by manual counting on photomicrographs. For GCP migration analysis using BrdU pulse-labeling, the outer external granule layer (oEGL) on stained sections was defined, using SPOT software, as a zone 20 μm in width adjacent to the pial surface around the perimeter of the lobule. The inner granule layer was defined as the remaining area of the lobule, excluding the white matter at the center of the lobule.

**Lacz staining**

Glial1lacZ reporter expression analysis was performed on brains from Nsdhlfl/GFAP-cre/Glial1lacZ and Nsdhlfl/Gli1lacZ males as previously described (81). Briefly, brains were fixed in 0.2% glutaraldehyde, 2% formalin for 2 h following transcardial perfusion, infused with 30% sucrose and floating sections (60 μm) were cut from frozen tissue using a sliding microscope. Floating sections were incubated in X-gal staining solution for 4 h at room temperature, rinsed in PBS and fixed in 4% PFA. Stained samples were photographed using a Nikon SMZ-10A dissecting microscope as described above.

**Immunoblotting**

Total protein extracts were made by homogenizing pooled E9.5 control or mutant embryos in RIPA buffer with 1x Protease Inhibitor Cocktail (Sigma, St. Louis, MO). Immunoblots to detect NSDHL were performed as previously described (50). Approximately 5–20 μg of protein from each sample was resolved by SDS-PAGE. Blots were probed sequentially with anti-NSDHL and anti-β-tubulin primary antibodies at 1:2000 and 1:4000 dilution, respectively.

**Confocal microscopy**

Floating sections (40 μm) were cut from frozen brains that had been fixed with 4% PFA and infused with 30% sucrose. Sections were stained with anti-GFP primary antibody and DyLight 488-conjugated donkey anti-mouse IgG. Z-stack fluorescence images using 2 μm optical sections were acquired on a Zeiss LSM 510 Meta confocal microscope using LSM software.

**GCP culture**

Granule cell precursors were isolated from individual cerebella at P4 and cultured as previously described (37) except that Percoll gradient separation and preplating were omitted. Briefly, freshly dissected cerebella were dissociated using the Papain Disassociation System Kit according to the manufacturer’s instructions (Worthington Biochemical, Lakewood, NJ). GCPs were plated at a density of 2.5 × 10⁵ cells/cm² on poly-D-lysine coated 6-well or 12-well tissue culture plates (Corning, Corning, NY) in 3 or 1 ml respectively of serum-free culture medium (SFM), consisting of Neurobasal-A medium (Life Technologies, Carlsbad, CA) supplemented with GlutaMAX I (Life Technologies), 250 μM KCl, 50 I.U./ml penicillin, 50 μg/ml streptomycin and B-27 supplement (Life Technologies) and cultured at 37°C, 5% CO₂ for 48 h. Where indicated, SFM was supplemented with 15 μg/ml cholesterol (Sigma) from a 15 mg/ml stock solution in 95% ethanol, 1 μg/ml recombinant mouse sonic hedgehog N-terminus (SHH-N) (R&D Systems, Minneapolis, MN) from a 100 μg/ml stock solution in 1% BSA in PBS, 5 μg/ml human LDL (Biomedical Technologies, Stoughton, MA) from a 5 mg/ml stock solution, 2 μM ketoconazole (Santa Cruz Biotechnology, Santa Cruz, CA) from a 1 mm stock in DMso, and/or 100 nm smoothed agonist, SAG (EMD Millipore, Billerica, MA) from a 100 nm stock in DMso. Equal volumes of vehicle alone were added to all untreated wells.

For BrdU labeling of GCPs in vitro, BrdU was added to each well to a final concentration of 10 μg/ml during the last 4 h of incubation. The cells were fixed with 2% PFA and immunostaining was performed as described above, except that BrdU primary antibody incubation was followed by an 8 h incubation with Dylight 549-conjugated donkey anti-rat IgG secondary antibody. Coverslips were mounted on slides with ProLong Gold antifade reagent with DAPI (Life Technologies). Quantitation of BrdU labeling was performed by manually counting the number of BrdU-positive cells from a total of at least 1500 total cells from five random fields on each coverslip. Results represent the combined data from three independent experiments.

In preliminary experiments using cultured GCPs, we occasionally noted highly variable or discrepant results. In subsequent experiments, we saved a small portion of the freshly isolated GCPs and performed RT-PCR on isolated RNA using primers that distinguished between the Nsdhfl allele, that produces WT message, and the recombinant Nsdhfl allele. In approximately 5% of Nsdhfl/Y samples analyzed (6/124 samples examined), the Nsdhfl allele constituted ≥5% of the PCR product, indicating incomplete cre-mediated excision. The amount of WT mRNA present correlated with the discrepant experimental results. For data presented here, any mutant samples in which WT product was detected were excluded from the analyses. This included in vitro proliferation experiments, sterol analyses, qPCR studies, and studies examining the effects of ketoconazole, SAG, or T-MAS on cultured GCPs.

For treatment of cell cultures, T-MAS (4,4-dimethyl-cholest-8(9),24-dien-3ß-ol) (Avanti Polar Lipids, Alabaster, AL) was complexed with low fatty acid bovine serum albumin (CellMaxx bovine albumin, catalog #199899, MP Biochemicals, Solon, OH) as previously described (82). Pure T-MAS was stored in dessicated vine albumin, catalog #199899, MP Biochemicals, Solon, OH) as previously described (8). Total protein extracts were made by homogenizing pooled E9.5 control or mutant embryos in RIPA buffer with 1x Protease Inhibitor Cocktail (Sigma, St. Louis, MO). Immunoblots to detect NSDHL and β-tubulin were performed as previously described (50).

**Sterol analysis**

GCPs were isolated from cerebella at P3, P5 and P7 by papain dissociation as described above. The cells were rinsed by suspending and pelleting two times in ice-cold PBS. Cell pellets were frozen and stored at −80°C. The yield of GCPs from each cerebellum was 2–5 × 10⁵ cells, depending on the age of the animals. For analysis of cultured cells, GCPs were isolated from P4 cerebella and cultured in SFM with the indicated treatments for 48 h as described above. The cells were rinsed two times with ice-cold
PBS and frozen at −80°C in the tissue culture plate. Cell pellets or frozen cells in 6-well plates were solubilized in 1.0 ml of 0.05% SDS. An aliquot of the lysate was used for protein measurement (BCA™ Protein Assay Kit, Pierce, Rockford, IL) and another for measurement of sterols. Sterols were extracted from the lysate by Folch extraction with chloroform/methanol (2:1), after internal standard [epicoprostanol, EPIC, Sigma C-2882 or cholestane (Steraloids, Newport, RI, C3300-000)] was added. The organic phase was removed and evaporated under nitrogen at 40°C. Sterols were then saponified by the addition of ethanol/KOH and incubation at 37°C for 1 h. Sterols were extracted twice with hexane. The combined hexane extracts were evaporated to dryness and derivatized with N,O-bis(trimethylsilyl)trifluoroacetamide (BSTFA, Sigma 33027) at 80°C for 30 min. Concentrations of the trimethylsilyl ethers of sterols were measured according to standard methods developed by Kelley (83) using gas chromatography with a ZB1701 column (Phenomenex, Torrance, CA; 30 m, 0.25 mm ID, 0.25 mm film thickness) and FID. In brief, cholesterol (Sigma C-8667) or desmosterol (Steraloids C3151-000) and EPIC or cholestane internal standards were used to generate calibration curves for cholesterol or desmosterol quantification, respectively. Replicate calibration curves demonstrated satisfactory linearity over the range 0.2–10 mg and 20–800 ng for cholesterol using 1 µg cholestane internal standard and 20–800 ng for cholesterol and desmosterol using 100 ng EPIC internal standard (with R² values > 0.995). Concentrations of other sterols were calculated by comparison of their peak area to that of internal standard assuming a 1:1 weight ratio via GC-FID sterol detection (83). Using this technique within-run precision for replicate methyl sterol analysis (n = 3) in representative P3 and P5 samples was determined to be <21% relative standard deviation (RSD). Sterol peaks were identified by comparison against authentic standards for desmosterol, lanosterol, and the methylsterol T-MAS. Derivatization of lanosterol and T-MAS were not complete under the conditions used (see Supplemental Material, Fig. S5C and data not shown). Quantitation of T-MAS in Figure 9 was based on GC-FID peak area detected for derivatized plus underderivatized T-MAS. For other methylsterols, peak identification was by reanalysis of samples using GC-mass spectrometry and mass spectrum matching against National Institutes of Standards and Technology (NIST) data in the Wiley 7th edition library with a quality score of 90 obtained for 4α-methylcholesta-8(9),24-dien-3β-ol and 92 for 4α-methylcholesta-8(9),24-dien-3β-ol on isolated GCPs from P7 (see Supplementary Material, Fig. S5D).

qPCR
Total RNA was extracted from 2 × 10⁶ cultured GCPs or whole cerebella using Trizol reagent (Life Technologies), followed by clean-up on RNeasy Mini Columns (Qiagen, Valencia, CA). RNA quality was assessed using a Bioanalyzer (Agilent, Santa Clara, CA) and all samples had a RIN >8. CDNA was synthesized from 1 µg of RNA for each sample using the SuperScript III First-strand Synthesis kit (Life Technologies) according to the manufacturer’s instructions. The volume of each CDNA sample was brought up to 100 µl with water and stored in aliquots at −80°C. Real-time PCR reactions were performed on 2 µl of CDNA using Power SYBR Green PCR Master Mix (Applied Biosystems, Foster City, CA) with 0.4 µM of each primer. Primer sequences are listed in Supplementary Material, Table S1. Amplification was carried out in MicroAmp optical 96-well plates (Applied Biosystems) on a 7500 Real-time PCR System instrument (Applied Biosystems). The amplification program was 50°C for 2 min, 95°C for 10 min, 40 cycles of 95°C for 15 s and 60°C for 1 min, followed by a melt curve stage. The relative quantity of CDNA for each gene was calculated using the ΔΔCT method (84), with Gapdh used as the reference gene in all samples. Primers for Sc4mol and Hmgcr were previously validated for qPCR (85). The amplification efficiency of additional primer pairs used in this study was measured by generating a standard curve from a dilution series of cDNA derived from WT GCPs. The R² values of the standard curves for all primer pairs were ≥ 0.992. The amplification efficiency (E = 10(−ΔΔCtΔn)) ranged from 95 to 98% for all primer pairs except Cnd1 (92%) and Cnd2 (87%). The ΔΔCt values were calculated using 95% as the efficiency. Melt curve analysis of products from each primer pair generated a single peak, indicating a unique amplicon for each. The results represent the combined data from at least three independent RNA samples, each assayed in triplicate.

Statistical analysis
For time course experiments in Figures 2–4, the data were analyzed by two way ANOVA using the R statistical environment (http://www.R-project.org), and P-values for pairwise differences in least square means were calculated using the lsmeans package (http://CRAN.R-project.org/package=lsmeans). For all other experiments, statistical differences between samples were determined by pair-wise comparison of the means of WT versus mutant sample data using two-tailed Student’s t-test (SPSS version 19), with P < 0.05 considered significant. All bar graph values are the mean ± SEM.

Supplementary Material
Supplementary Material is available at HMG online.

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