Mutation of Nogo-B Receptor, a Subunit of cis-Prenyltransferase, Causes a Congenital Disorder of Glycosylation

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SUMMARY

Dolichol is an obligate carrier of glycans for N-linked protein glycosylation, O-mannosylation, and GPI anchor biosynthesis. cis-prenyltransferase (cis-PTase) is the first enzyme committed to the synthesis of dolichol. However, the proteins responsible for mammalian cis-PTase activity have not been delineated. Here we show that Nogo-B receptor (NgBR) is a subunit required for dolichol synthesis in yeast, mice, and man. Moreover, we describe a family with a congenital disorder of glycosylation caused by a loss of function mutation in the conserved C terminus of NgBR-R290H and show that fibroblasts isolated from patients exhibit reduced dolichol profiles and enhanced accumulation of free cholesterol identically to fibroblasts from mice lacking NgBR. Mutation of NgBR-R290H in man and orthologs in yeast proves the importance of this evolutionarily conserved residue for mammalian cis-PTase activity and function. Thus, these data provide a genetic basis for the essential role of NgBR in dolichol synthesis and protein glycosylation.

INTRODUCTION

Nogo-B receptor (NgBR) was identified via expression cloning as a protein that interacts with the N terminus of Nogo-B, also called reticulon-4b (Miao et al., 2006). NgBR is a polytypic membrane protein, and its C-terminal domain shares significant homology with two gene products: (1) NUS1, a gene in yeast required for survival and N-glycosylation (Harrison et al., 2011; Yu et al., 2006) and (2) cis-prenyltransferases (cis-PTase), including genes in yeast (RER22 and SRT71), a human ortholog, (hCIT, also called dehydrodolichol diphosphate synthase [DHDDS]), and bacterial undecaprenyl pyrophosphate synthase (uPPS) (Sato et al., 1999; Schenk et al., 2001; Surmacz and Swiezewska, 2011). In lower organisms, single subunit cis-PTases such as UPPS catalyze the condensation reactions of isopentenyl pyrophosphate (IPP) with farnesy1 pyrophosphate (FPP) to synthesize linear poly-prenyl pyrophosphate with specific chain lengths. Polyprenyl pyrophosphate is dephosphorylated into polyprenol and then reduced by a polyprenyl reductase to produce dolichol (Cantagrel et al., 2010). In mammals, the relative contribution of Nus1/NgBR versus Rer2/Srt1/hCIT to cis-PTase activity and dolichol synthesis is unknown since loss of function of each grouping of genes results in reduced glycosylation.

Congenital disorders of glycosylation (CDG) are genetic diseases that represent an extremely broad spectrum of clinical presentations due to defects in several steps of protein glycosylation. Recently, there have been several reports of genetic defects in the dolichol biosynthetic pathway, such as mutations in DHDDS/hCIT and SRD5A3 (Cantagrel et al., 2010; Kasapkara et al., 2012; Zelinger et al., 2011; Züchner et al., 2011). DHDDS-CGD is associated with inherited retinitis pigmentosa, a disorder causing retinal degeneration, and DHDDS-CGD patients did not show the other typical CDG symptoms. SRD5A3-CGD affects the final step in dolichol synthesis. Its clinical features are typical for CDG type 1 glycosylation disorders, including psychomotor retardation, ocular malformations, cerebellar hypoplasia, skin lesions, and facial dysmorphism.

Here, we characterize the dolichol biosynthesis pathway in mice and yeast and demonstrate the necessity of both hCIT and NgBR for dolichol biosynthesis. In addition, we describe a unique congenital disorder of glycosylation caused by a mutation in NgBR, a conserved subunit of cis-PTase. Patients harboring a R290H mutation of NgBR have congenital scoliosis, profound psychomotor retardation, refractory epilepsy, and macular lesions showing retinitis pigmentosa. Thus, hCIT/NgBR heteromers are essential, conserved components of the machinery necessary for glycosylation reactions in mammals.
RESULTS AND DISCUSSION

Targeted Disruption of NgBR Causes Early Embryonic Lethality In Vivo and Defective cis-PTase Activity and Cholesterol Levels in Isolated Fibroblasts

To examine the physiological significance of NgBR, we generated a conditional knockout mouse (Figures S1A–S1C available online). The NgBR knockout allele (NgBRΔα) was generated by crossing NgBR conditional allele (NgBRΔ1) with a progesterone Cre driver expressed in the male germ line (O’Gorman et al., 1997). Heterozygous NgBR mice (NgBRΔα/+) appeared normal, and intercrosses with NgBRΔαΔβ showed no viable homozygous mice (NgBRΔαΔβ) (Figure 1A). To determine when lethality occurred, timed pregnancies of NgBRΔα/α breeding were examined at embryonic day 6.5 (E6.5) and E7.5. No NgBRΔα/α embryos were identified at these time points indicating postimplantation embryonic lethality before E6.5 (Figures 1B and 1C). Next, we established mouse embryonic fibroblast (MEFs) cultured from NgBRΔα mice using an inducible Cre-loxP system. Reduced expression of NgBR in the tamoxifen inducible NgBR knockout (NgBR iKO) MEF cells was confirmed by PCR and western blotting for mRNA and protein levels, respectively (Figure S1D). NgBR iKO MEFs showed accumulation of free cholesterol as determined by filipin staining (Figure 1D), decreased cis-PTase activity in isolated membranes (Figure 1E), and mannose incorporation into protein (Figure 1F). Transduction of cells with lentiviral human NgBR rescues the increase in free cholesterol (Harrison et al., 2009) and the decrease in mannose incorporation (Figures 1E and 1F). Furthermore, we exposed cells to lovastatin, an inhibitor of 3-hydroxy-3-methylglutaryl-coenzyme A (HMG-CoA) reductase, the rate-limiting enzyme in the synthesis of isoprenoids, and measured cell viability using an MTT assay. NgBR iKO MEFs were unable to synthesize longer chain polyprenyl pyrophosphates. Expression of Nus1, Rap2, or mixtures of IVT Nus1 with Rap2 products in equal amounts (as shown by western blotting in the bottom panel) did not catalyze formation of polyprenols. Interestingly, only cotranslation of Nus1 with Rap2 and its orthologs in S. pombe or humans formed an active cis-PTase complex producing prenols of expected lengths (Figures 2C–2F, lane 5), indicating that both proteins are required for a functional enzyme (Figures 2D–2F). Collectively, the data support the heteromeric structure of mammalian and yeast cis-PTase and suggest that eukaryotic cis-PTase is assembled during translation since only cotranslation, but not mixing of the proteins, yields active enzyme. Taken together, these data provide a clear rationale for the role of RER2/NUS1 and related genes in dolichol biosynthesis and advance our understanding of this important pathway.

A Mutation on NgBR Causes Congenital Scoliosis, Severe Neurological Impairment, Refractory Epilepsy, Hearing Deficit, and Visual Impairment

Recently, exome sequencing of individual families with symptoms of a congenital disorder of glycosylation (CDG) has led to the discovery of mutations in DHDDS (hCIT) and SRD5A3, genes involved in the early steps of polyprenol synthesis (Cantagrel et al., 2010; Kasapkara et al., 2012; Zelinger et al., 2011; Züchner et al., 2011). DHDDS-CDG is associated with inherited retinitis pigmentosa, a disorder causing retinal degeneration, and SRD5A3-CDG patients exhibit psychomotor retardation, ocular malformations, cerebellar hypoplasia, skin lesions, and facial dysmorphism. In our clinic, a family of Roma origin (Figure 3A) composed of healthy, unrelated parents and four siblings was examined, and two siblings presented with congenital scoliosis, severe neurological impairment, refractory epilepsy, hearing deficit, and visual impairment with discrete bilateral macular lesions.
Figure 1. Characterization of NgBR Knockout Mouse Embryos and Fibroblasts

(A) Genotype obtained from the progeny of heterozygous mating. No NgBR<sup>D/D</sup> embryo was detected. 

(B) Embryo resorption frequencies during postimplantation development. Resorption sites were apparent at E7.5 among ~25% decidua. 

(C) Representative decidua of E7.5 embryo resorption sites analyzed. Decidua were obtained from NgBR<sup>D/+</sup> breeding. Decidua with embryo contained normally developed E7.5 embryo (insert). Presumptive NgBR<sup>D/D</sup> decidua exhibit implanted site for embryo without evident embryonic material (arrowhead). 

(D) Filipin staining and quantitative representation for MEF. Filipin staining was performed 48 hr after Lenti-NgBR transduction into NgBR iKO MEF cells. U18666A was used as a positive control for inhibition of cholesterol trafficking. 

(E) Microsomal cis-PTase activity assay for NgBR iKO MEF. Enzyme activity was reduced by 83% in NgBR iKO MEF compare to control. 

(F) [2-<sup>3H</sup>]-mannose labeling of proteins in mouse embryonic fibroblasts. Tunicamycin (Tm) treatment was used as a control. 

(G) Statin sensitivity measured by MTT assay. Cell viability was determined by MTT assay after 16 hr exposure with various concentrations of lovastatin (1–80 μM). Cell viability was calculated by the following equation: MTT optical density value of treated sample / MTT OD value of nontreated sample. 

(H) RT-PCR for UPR pathway genes. Relative mRNA expression to control show increased expression. Data are representative of at least three experiments. *p < 0.05. Data are mean ± SE. See also Figure S1.
Proband II.3 was born at term with intrauterine growth retardation. Muscle hypotonia was present since birth, and congenital scoliosis and developmental delay were observed since early infancy. Tonic-clonic seizures, refractory epilepsy, and recurrent attacks of status epilepticus developed from the age of 11 months. Microcephaly (3rd centile), failure to thrive (<3rd centile), regression of psychomotor development, severe axial hypotonia and acral spasticity developed after discharge. Routine laboratory tests were unremarkable, and cholesterol level was within reference range. The boy died at the age of 29 months. Histopathological findings in autopsy tissue revealed nonspecific neuronal loss in brain cortex and cerebellum. Similarly to his brother, proband II.4 had generalized hypotonia, congenital scoliosis, and significant delay in motor milestones. Refractory epilepsy started at the age of 7 months, and he has been hospitalized several times with severe seizures. He lost any social interaction, and he displays no spontaneous movements. At the age of 4 years, he has microcephaly (0.6th centile), failure to thrive (<5th centile), and marked hypertrichosis. He has severe axial hypotonia, acral spasticity with preserved deep tendon reflexes, pseudobulbar palsy, and central visual and hearing impairment. MRI of the brain revealed severe cortical atrophy. A complete dilated fundus examination including color fundus photography was performed under general anesthesia. At the age of 31 months, except for an opacity located in the inferior half of the right cornea, there were bilaterally no other anterior segment abnormalities, and the vitreous was optically clear. There were no bone spicule pigmentations, but diffuse retinal pigment epithelium mottling could be observed bilaterally. Optic nerves appeared paler and retinal vessels narrower. Repeated examination at the age of 4 years documented a development of bilateral macular lesion showing foveal hyperautoflorescence (Figure 3B).

The exomes of parents and both affected probands were sequenced and searched for genetic variants in the internal exome database, the Exome Variant Server, and 1000 Genomes databases, and only four such variants were discovered; three are located in the autozygous region identified on chromosome 6 and one on chromosome 21 (Table S1). Corresponding genes were evaluated based on their potential contribution to the clinical phenotypes, and a homozygous missense mutation c.869G > A in the NUS1 (NM_138459) or NgBR was found. The
c.869G > A mutation was confirmed by Sanger sequencing, and the affected probands are homozygous for this mutation, whereas their parents and healthy siblings are heterozygous (Figure 3C). The mutation encodes for amino acid exchange p.Arg290His (R290H), which is located in the evolutionarily conserved C-terminal domain of NgBR (Figure 3D) and is predicted to affect protein function with a score of 0.00 according to SIFT and to be damaging using Polyphen. This mutation was not reported in dbSNP, 1000 Genomes, or the Exome variant server and was not listed in our internal exome database (>250 exomes). Targeted genotyping of genomic DNA from 255 individuals of Roma origin identified two additional heterozygous carriers of the c.869G > A mutation. Even though the identity and relation status of these two carriers is unknown, this finding suggests that the congenital disorder of glycosylation caused by a loss-of-function mutation of NgBR may be relatively frequent among the European Roma population.

NgBR R290H Mutation Triggers Defects in Cellular Cholesterol Trafficking and Dolichol Biosynthesis

To characterize the NgBR R290H mutation, fibroblasts were isolated from control and NgBR R290H patients and we examined the levels of NgBR mRNA, protein, and interaction of NgBR with hCIT (Figures S3A–S3C). We did not observe any significant differences in the migration of NgBR on SDS-PAGE or the levels of NgBR protein compared to WT (Figure S1B). This suggests that the translation and the subsequent processing of mutant NgBR protein were not altered by the presence of the mutation. NgBR was isolated as a protein that interacted with reticulon 4B, also called Nogo-B (Miao et al., 2006). Therefore, we examined whether Nogo-B levels and its interaction with NgBR were altered in carriers of the NgBR mutation. The levels of Nogo-B, its interaction with NgBR, and the localization of Nogo-B were not different (Figures S3D–S3F). Next, we assessed three aspects of NgBR function, free cholesterol levels, cis-PTase activity, and glycosylation. WT cells had little filipin-positive free cholesterol, whereas treatment with U18666A to induce a Niemann-Pick C (NPC) disease phenotype (Cenedella, 2009) increased free cholesterol (Figure 4A, quantified in the right panel). In contrast, NgBR R290H mutant cells exhibited increased accumulation of free cholesterol similar to cells where NgBR was silenced (Harrison et al., 2009). Additionally, cis-PTase activity (Figure 4B) and mannose incorporation into proteins (Figure 4C) was markedly lower in NgBR R290H fibroblasts compared to control. We also examined defective glycosylation of proteins in patient fibroblasts by western blotting for two known glycoproteins, LAMP-1 and ICAM-1 (He et al., 2012; Xiang et al., 2013). Both LAMP-1 and ICAM-1 were hypoglycosylated in the patient fibroblasts (Figure 4D). Thus, the NgBR R290H mutant is a loss-of-function mutation that affects cis-PTase function of NgBR without disrupting complex formation with hCIT or Nogo-B. The reduced cis-PTase activity in fibroblasts was manifested as altered dolichol profiles in the urine or serum as assessed by mass spectrometry of all carriers of
the R290H mutation (Figure S4), as recently described for patients harboring loss-of-function mutations in DHDDS (Wen et al., 2013).

Amino Acid at the Fourth Position from the C Terminus of NgBR Is a Functionally and Evolutionarily Conserved Residue

Alignment of NgBR orthologs from distantly related eukaryotic organisms reveals a high degree of conservation at the C terminus, with arginine or asparagine present at the fourth position from the C terminus (Figure 5A). To test the evolutionary conservation of this position, hCIT was expressed with NgBR or NgBR R290H in the nus1Δ, rer2Δ, srt1Δ triple knockout strain. Cells expressing the NgBR R290H allele have lower cis-PTase activity (Figure 5B), overall polyisoprenoids (Figure 5C), and dolichol levels as measured by MS (Figure 5D). In addition, we analyzed S. cerevisiae and S. pombe Nus1 mutants to determine the importance of the amino acid conservation at the fourth position from the C terminus in NgBR orthologs. S. cerevisiae Nus1 belongs to group of fungi and plants NgBR orthologs bearing asparagine instead of arginine, while S. pombe Nus1 encodes arginine at position 255 corresponding to the R290 in human NgBR. Therefore, we compared cis-PTase activity of the S. cerevisiae nus1Δ strain expressing wild-type Nus1, Nus1-N372H (mimicking NgBR R290H mutation), as well as Nus1-N372R. Also, we expressed wild-type SpNus1 or SpNus1-R255H in the nus1Δ fission yeast strain. Mutation of the same position in Nus1 in S. cerevisiae (Figures 5E and 5F) and S. pombe (Figures 5G and 5H) resulted in a similar loss of function. Interestingly, the N372R allele of Nus1 from S. cerevisiae affects only the chain length of the product (Figures 5F) but not the rate of incorporation of IPP (Figures 5E).

Recently, altered ratios of plasma and urinary dolichols were observed in retinitis pigmentosa patients carrying the K42E mutation in DHDDS/hCIT (Wen et al., 2013). To compare the influence of NgBR R290H and hCIT K42E mutations on cis-PTase activity, we expressed hCIT or hCIT K42E with NgBR or NgBR R290H in the S. cerevisiae triple knockout strain and measured enzyme activity. Introduction of this mutant into the triple knockout strain expressing WT NgBR reduced steady-state cis-PTase activity to an extent similar to that of NgBR R290H expressed with WT hCIT (Figure 5I), and combining the mutations reduced activity, further demonstrating epistasis of the gene products.

NgBR and its orthologs are essential genes, and NgBR/hCIT heteromers are responsible for dolichol synthesis in mammalian cells (Figures 6A and 6B). Based on previous work, NgBR can interact with hCIT, NPC2, and Nogo-B (Figure 6B; numbered 1–3). The interaction with NPC2 was identified by an independent broad-based screening strategy (Harrison et al., 2009).
Genetic evidence for the importance of this interaction stems from data showing that NgBR knockout MEFs and patient fibroblasts harboring the R290H mutation exhibit increased free cholesterol levels. Since NPC2 is a soluble glycoprotein (Naur-еекiene et al., 2000) and glycosylation of NPC2 is important for its function (Chikh et al., 2004), it is feasible that in addition to a direct stabilizing effect of NgBR on NPC2, mutant NgBR can influence NPC2 glycosylation due to reduced cis-PTase activity contributing to this phenotype (Figure 6B). The interaction of NgBR with Nogo-B does not impact cis-PTase activity or cellular cholesterol content and may influence intracellular signaling pathways.

Little is known about the function of dolichol species in vivo besides its role as a glycan carrier, although in vitro evidence suggests that dolichol can modulate biophysical properties of membranes and serve as a cellular antioxidant (Surmacz and Swiezewska, 2011). Patients carrying a mutation in NgBR demonstrated altered ratios of dolichol in urine and in blood.
Although altered dolichol chain length ratios are important bio-markers in patients with mutations in hCIT/DHDDS and NgBR, alterations in dolichol chain length are unlikely to exert a domi-nant effect since lipid-linked oligosaccharides built on as few as 11 dolichol units seem to be efficient substrates in N-glycosylation reactions (Grabińska et al., 2010; Rush et al., 2010). How-ever, the overall lower dolichol content of cell membranes not only directly affects glycosylation but can impair membrane structure and, in turn, affect multiple cellular processes including sterol biosynthesis. In summary, the development of a knockout strain of mice, the establishment of a NgBR/hCIT reconstitution system in yeast, and the discovery of a highly conserved mutation in the NgBR mutation in humans will assist in the further characterization of the cellular functions of this essential polyisopo-ne lipid.

**EXPERIMENTAL PROCEDURES**

**Generation of NgBR Mouse Embryonic Fibroblasts**

NgBR<sup>fl/fl</sup> was crossed with NgBR<sup>+/-</sup>; R26CreER (Badea et al., 2003), and primary MEFs were prepared from E13.5 embryos. Each MEF line was derived from an individual embryo. Isolated MEFs were immortalized using an SV40-large-T-expressing retrovirus obtained from Genecopoeia (LP-SV40T-LV105-0205) according to the manufacturer’s protocol. Immortalized cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) with 1% penicillin/streptomycin containing 10% FBS. The genotypes of control MEFs and NgBR iKO MEFs used in this study are NgBR<sup>fl/+</sup> and NgBR<sup>+-/D</sup>; R26CreER. Both cell lines were treated with 1 μM 4-hydroxytamoxifen (Sigma) for more than 5 days to induce Cre recombination. mRNA or protein expression level was confirmed for each experiment. All experiments with NgBR<sup>fl/fl</sup> mice were approved by the Institutional Animal Care Use Committee at Yale School of Medicine.

**Filipin Staining**

Filipin staining was performed as previously described (Harrison et al., 2009). In brief, cells were fixed in 4% paraformaldehyde for 10 min and permeabilized in 0.1% Triton X-100 for 5 min. Cells were then incubated with a 50 μg/ml concentration of filipin (Sigma, F4767) for 1 hr. As a positive control for induction of cholesterol accumulation, cells were treated for 8 hr with 1 μM U18666A (EMD Biosciences). Relative intensity of filipin staining was quantified by calculating average pixel intensity using Adobe Photoshop according to the following equation: average filipin intensity = total intensity above low threshold / number of pixels above low threshold (Pipalia et al., 2006).

**Microsomal cis-PTase Activity Assay**

For mammalian cells, crude microsomes were prepared as described before (Rush et al., 2010) with minor modification. cis-PTase activity in mammalian

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**Figure 6. Functions of the NgBR/hCIT Complex in Cellular Metabolism**

(A) The NgBR and hCIT complex promotes cis-PTase activity. NgBR/hCIT catalyzes the condensation of isopentenyl pyrophosphate with farnesyl pyrophosphate to generate a polyprenol pyrophosphate. Polyprenol diphosphate is dephosphorylated by unidentified phosphatase and reduced by polyprenol reductase (SRD5A3) to form dolichol. Finally, dolichol is phosphorylated by dolichol kinase prior to the synthesis of dolichol-linked sugars required for glycosylation pathways.

(B) (1) NgBR and hCIT assembly is essential for cis-PTase activity generating polyprenol pyrophosphate on the cytoplasmic leaflet of the ER membrane. Polyprenol pyrophosphate serves as an intermediate in synthesis of dolichol-linked saccharides. Dolichol-pyrophosphate tetradesaccharide (LLO) is indispensible for protein N-glycosylation reactions. Dolichol-phosphate mannose (DolPMan) is also involved in O-mannosylation, GPI-anchor synthesis, and C-mannosylation. (2) NgBR influences cholesterol trafficking by directly interacting with NPC2 and indirectly via modifying NPC2 N-glycosylation. (3) The interaction between Nogo-B and NgBR does not influence glycosylation or cholesterol trafficking, and the function of this interaction remains to be clarified.
cells was assayed as described before (Harrison et al., 2011) with minor modification. For S. cerevisiae and S. pombe, membrane fraction was prepared as described before (Szkopinska et al., 1997). cis-PTase assay using yeast membranes was performed as described (Szkopinska et al., 1997) with minor modifications. For a detailed description, please see the Supplemental Experimental Procedures.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures, three figures, and one table and can be found with this article online at http://dx.doi.org/10.1016/j.cmet.2014.06.016.

**AUTHOR CONTRIBUTIONS**

E.J.P. and K.A.G. contributed equally to all aspects of this paper: E.J.P. characterized NgBR deficient mice and performed all mammalian cell based studies and cloning/plasmid construction, and K.A.G. developed yeast strains and cis-PTase characterization in vivo and in vitro. Both authors contributed to the writing and editing of the manuscript. Z.G. and R.W. contributed to MS analysis of urinary and serum dolichol levels and the writing of the manuscript. V.S., H. Hartmannová, and K.H. performed genotyping, linkage analysis, homozygosity mapping, and exome sequencing. V.B., J.S., N.O., H. Hansíková, and H. Hůlková contributed to acquisition of clinical specimens and phenotypic characterization of patients. T.H. and J.Z. were responsible for clinical and diagnostic assessment of affected patients. L.J. performed high-resolution imaging of Nogo-B in patient cells. W.C.S., E.J.P., K.A.G., and S.K. were responsible for concept development and preparation of the manuscript. S.K. was responsible for overseeing the genetic aspects of the study, and W.C.S. was responsible for overall integration and execution of the scientific approaches.

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