Palmitoylation of BMPR1a regulates neural stem cell fate

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Neural stem cells (NSCs) generate neurons and glial cells throughout embryonic and postnatal brain development. The role of S-palmitoylation (also referred to as S-acylation), a reversible posttranslational lipid modification of proteins, in regulating the fate and activity of NSCs remains largely unknown. We used an unbiased screening approach to identify proteins that are S-acylated in mouse NSCs and showed that bone morphogenetic protein receptor 1a (BMPR1a), a core mediator of BMP signaling, is palmitoylated. Genetic manipulation of S-acylated sites affects the localization and trafficking of BMPR1a and leads to altered BMP signaling. Strikingly, defective palmitoylation of BMPR1a modulates NSC function within the mouse brain, resulting in enhanced oligodendrogenesis. Thus, we identified a mechanism regulating the behavior of NSCs and provided the framework to characterize dynamic posttranslational lipid modifications of proteins in the context of NSC biology.

neural stem cell | neurogenesis | BMP receptor | palmitoylation | oligodendrogenesis

The activity and fate of neural stem cells (NSCs) in the developing and adult mammalian brain are tightly regulated by a plethora of cell intrinsic and extrinsic mechanisms ensuring that proper numbers and types of neuronal and glial cells are produced (1). By defining cellular and transcriptional programs that direct the fate of NSCs, substantial progress has been made to understand how NSCs are regulated within their endogenous niches (2–4). However, little is known about how posttranslational modifications (PTMs) of stem cell–expressed proteins affect NSC activity and subsequent fate choices. PTMs, such as phosphorylation, ubiquitination, sumoylation, acetylation, glycosylation, and lipidation among others, have been shown to affect the localization, activity, and stability of a large number of proteins, allowing for fast and partially irreversible modification of protein function (5–10). Thus, PTMs represent critical regulators of protein function in a variety of cell types, allowing for fast signal diversification and refined cellular responses to exogenous cues (11). Many proteins are modified by the attachment of lipid moieties such as myristoylation, prenylation, and palmitoylation (also referred to as S-acylation) that modulate protein function (5). Among all known lipid modifications, S-acylation represents the only known reversible form of lipid modification that has been shown to play a pivotal role in protein trafficking, stability, and function (5, 12). 16-Carbon palmitate residues are transferred onto cysteines of target proteins via palmitoyl-cysteine transferases (zDHHHC1–24) and can be dynamically removed by acyl-protein thioesterases (for example, APT1-2) (13). Indeed, recent data suggest that reversible palmitoylation is involved in a large variety of biological processes such as growth of Arabidopsis thaliana, melancorin signaling in the skin, synaptic function in the adult central nervous system, and many others (5, 14–16). Thus, over the last years large efforts have been undertaken to define the palmitoyl proteome in a variety of cell types and tissues (17).

Results

Identification of Palmitoylated Proteins in Mammalian NSCs. To identify a role for protein palmitoylation in mammalian stem cell function, we used an acyl-biotin exchange (ABE) assay to discover proteins that are S-acylated in NSCs isolated from the hippocampus of 8-wk-old mice (Fig. 1A). NSCs were lysed, and the protein samples were divided into 2 fractions. In the “hydroxylamine” (“HAM”) sample, the palmitate residue was cleaved off and exchanged with biotin. The “HAM condition served as a negative control. After the ABE reaction was completed, streptavidin beads were used to enrich for biotinylated proteins. Proteins enriched from “HAM and “HAM conditions were identified using mass spectrometry (MS) (Fig. 1B and SI Appendix, Fig. S1A). Proteins with a significant and at least 4-fold higher abundance in the “HAM sample were considered candidate proteins. With this approach, we identified 347 palmitoylated proteins in cultured NSCs (Fig. 1B), with an expected enrichment of identified proteins to be associated with membrane localization and function using Gene Ontology (GO) localization and molecular function analytic tools; T.W., K.B., D.G.-B., M.H., M.K.b.I., A.A., I.M., A.M., and D.K. performed research; P.P. contributed new reagents/authentic tools; T.W., K.B., D.G.-B., M.H., M.K.b.I., A.A., I.M., A.M., and D.K. analyzed data; and T.W. and S.J. wrote the paper.

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Data deposition: The mass spectrometry proteomics data have been deposited in the publicly accessible database ProteomeXchange Consortium via the Proteomics Identifications (PRIDE) partner repository with the dataset identifier PXD014355.

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Fig. 1. Identification of S-acylated proteins in neural stem cells. (A) Schematic describing the enrichment strategy and mass spectrometry-based workflow used for the identification of S-acylated proteins. Mass spectrometry and a label-free analysis were used for identification and quantification of proteins. Protein lysates were divided into 2 fractions. In the “HAM + Biotin” condition, the palmitate was cleaved off and exchanged with a biotin. The “HAM - Biotin” condition controls for unspecific binding. Streptavidin beads were then used to enrich for biotinylated proteins, followed by mass spectrometry-based identification. NEM, N-ethylmaleimide. (B) ABE enrichment coupled with mass spectrometry–based analysis identified 347 potentially S-acylated proteins (blue). Proteins that were less than 4-fold or not significantly enriched between the “HAM” and “HAM” conditions are shown in black. (C) Probing identified proteins for GO. “Localization” and “Molecular function” show that a substantial fraction of candidate proteins is associated with membrane compartments (blue, enriched candidates; black, enriched and nonenriched proteins). (D) Comparison of candidates with the SwissPalm database for S-acylated mouse proteins. Of the identified candidate proteins, 21.39% had not been described previously as S-acylated (dark gray). (E) Enrichment and background intensities (blue) of previously confirmed S-acylated proteins identified in the NSC dataset. CANX, calnexin; FLOT1, flotillin-1; GRIA2, AMPA receptor 2. (F) Enrichment and background intensities (blue) of proteins validated in this study. NUP210, nucleoporin 210. Error bars represent mean ± SD. *P < 0.05, **P < 0.01, ***P < 0.001.

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analysis (Fig. 1C). Supporting the validity of the assay, we were able to identify previously validated S-acylated proteins (Fig. 1D and E). Remarkably, 21% of the identified candidate proteins had not previously been described as palmitoylated (Fig. 1D). Given that palmitoyl-proteome screening approaches may yield false-positive results (18), we used protein-specific ABE assays and an alternative method (metabolic labeling of palmitate residues [19]) to verify S-acylation of selected candidate proteins such as NUP210 and VAMP7 (vesicle-associated membrane protein 7) (Fig. 1F and SI Appendix, Figs. S1 B–H and S2A), thereby confirming the validity of the unbiased screening results.

**BMPR1a Is Palmitoylated at Multiple Cysteines by zDHHC20.** One of the candidate proteins we identified was the bone morphogenetic protein receptor 1a (BMPR1a), which together with other components mediates canonical and noncanonical BMP signaling (Fig. 1F) (20, 21). BMP signaling plays a pivotal role in brain development and NSC behavior and has been associated with a large number of related biological processes, among them cell growth, cell differentiation, and cancer (22, 23). We confirmed palmitoylation of BMPR1a by tagging the endogenous BMPr1a gene with GFP using CRISPR/Cas9 technology, allowing for efficient pulldown of BMPR1a in NSCs and subsequent ABE analysis (Fig. 2A and SI Appendix, Fig. S2B). Analogously, we confirmed palmitoylation of BMPR1a by tagging the endogenous BMPR1a with a hemagglutinin (HA) tag in mouse embryonic stem cells (ESCs) that were subsequently differentiated into NSCs and analyzed using metabolic labeling of palmitoylation sites within BMPR1a (SI Appendix, Fig. S2 C–F). Computational analyses suggested 5 cysteine residues to be palmitoylated within the BMPR1a protein sequence (24). However, using an acyl-PEG (methoxy(polyethylene glycol) exchange assay, allowing for determination of the number of S-acylated cysteines, we found that BMPR1a has 3 functional palmitoylation sites (cysteines C173, C175, and C180), which are located close to the transmembrane domain of the receptor (Fig. 2 B and C). Directed mutagenesis showed that S-acylation was absent when cysteine residues were exchanged into alanines from the overexpressed BMPR1a protein (Fig. 2 B and C and SI Appendix, Fig. S2 G and H). Importantly, exchange of palmitoylated cysteines into alanines did not affect binding to the BMP4 ligand, as shown by immunoprecipitation of BMPR1a followed by immunochemistry of if bound proteins (Fig. 2D). We also detected single spectral counts of the interaction partner BMPR2; however, they were not enough to draw any quantitative conclusions. Immunoprecipitated tdTomato samples were used to control for unspecific binding. Immunoprecipitation of overexpressed HA-tagged BMPR1a or BMPR1a C173/175A together with FLAG-tagged BMPR1a or BMPR2 in NSCs showed that BMPR1a C173/175A was still able to dimerize with BMPR1a and BMPR2 (SI Appendix, Fig. S3A). We next aimed to identify which of the 24 known palmitoyl-acyltransferases (zDHHC1–24) mediates palmitoylation of BMPR1a. We found zDHHC20 to colocalize with BMPR1a (Fig. 2E). Notably, we found overexpression of zDHHC20 to be sufficient to enhance S-acylation of BMPR1a (Fig. 2F), thus identifying a palmitoyl-acyltransferase that transfers palmitate residues onto BMPR1a.

**Palmitoylation of BMPR1a Alters Its Function.** We next probed the functional relevance of BMPR1a palmitoylation by testing if acylation-deficient BMPR1a proteins are sufficient to rescue the complete loss of the function proliferation phenotype of BMPR1a in NSCs (26). Given their positioning within BMPR1a, we analyzed C173/175A and C180A exchanges separately. As expected, we found that CRISPR/Cas9-mediated deletion of BMPR1a reduced proliferation of NSCs in response to BMP4 exposure, as measured using 5-ethyl-2′-deoxyuridine (EdU) pulse labeling (SI Appendix, Fig. S3 B and C). Whereas wild-type and C180A overexpression normalized proliferation, the proliferation upon C173/175A overexpression was not rescued (SI Appendix, Fig. S3B). Consistent with this, we found that overexpression of C173/175A was not sufficient to transduce canonical BMP signaling, as measured by luciferase assays with BMP response elements (BREs) (SI Appendix, Fig. S3D) (27). Notably, overexpression of zDHHC20 was able to promote canonical BMP signaling in vitro (SI Appendix, Fig. S3E). This finding suggests that palmitoylation of cysteine residues C173/175 is required for canonical BMPR1a function, at least in vitro.

To analyze how palmitoylation of BMPR1a affects NSCs in vivo, we modified acylation sites within BMPR1a and aimed to generate mouse models harboring single (C180A) or double (C173/175A) mutants, again exchanging the palmitoylated cysteines with non-S-acylatable alanines. However, it turned out that C173/175A double mutants could not be obtained, most likely due to early embryonic lethality: out of 767 injected and transferred zygotes, we did not obtain a single viable double mutant mouse. This is consistent with the failure of C173/175A BMPR1a to rescue the full BMPR1a knockout in vitro (SI Appendix, Fig. S3F), as full knockouts of BMPR1a die at embryonic day E9.5 (28). However, we were able to generate C180A single mutants (Fig. 3A and B) and detected expression of BMPR1a using immunohistochemistry of BMPR1a in NSCs and the embryonic and adult brain (Fig. 3A and SI Appendix, Fig. S3F), as described before (26, 29). Thus, we next analyzed how palmitoylation of BMPR1a at position C180 affects its localization and function. Strikingly, we found that embryonic NSCs isolated from embryonic day 17.5 (E17.5) brains of C180A mutant mice showed reduced surface expression of the BMPR1a receptor (Fig. 3B) without affecting overall expression levels (Fig. 3C). In addition, fluorescence recovery after photobleaching (FRAP) analysis revealed an increase in the immobile fraction of overexpressed BMPR1a C180A compared to control BMPR1a (Fig. 3D). In FRAP experiments, the total recovery of fluorescence after photobleaching is influenced by mobile and immobile fractions of the BMP receptor. The mobile fraction undergoes exchange with the photobleached area, whereas the immobile fraction does not. Moreover, we found a robust reduction in endocytosis of BMPR1a in C180A mutant cells (Fig. 3E). Together, these findings indicate that palmitoylation of BMPR1a at position C180 is required for proper localization and mobility of the BMPR1a receptor (30).

**Palmitoylation of C180 Affects Noncanonical BMP Signaling.** To investigate the effects on BMP signaling in C180A mutant cells, we next analyzed signaling activity in proliferating and differentiating BMP4-stimulated cells isolated from C180A mutant mice and controls. We found that stimulation with BMP4 successfully promoted canonical BMP signaling in C180A-derived cells and control cells, as measured by levels of phosphorylated SMAD1/5 (SI Appendix, Fig. S3 G and H). Corroborating these results, we also found BMP4-dependent induction of inhibitor of differentiation 1-4 (Id1-4) gene transcripts at comparable levels in C180A compared to control cells as measured using digital droplet qPCR (SI Appendix, Fig. S3 I and J). This is consistent with the finding that overexpression of C180A does not prohibit canonical BMP signaling as measured using a BRE-dependent luciferase assay (SI Appendix, Fig. S3D). However, the C180A mutation caused a decrease in noncanonical, active extracellular signal-regulated kinase 1/2 (ERK1/2) in nonstimulated and BMP4-stimulated NSCs derived from C180A mice and controls (Fig. 3F) and an increase in active ERK1/2 during in vitro differentiation of C180A-derived cells (SI Appendix, Fig. S3K), which is known to promote oligodendrocyte differentiation (31). These findings indicate that palmitoylation-mediated localization and intracellular trafficking of BMPR1a at site C180 affects noncanonical, ERK-dependent BMP signaling (30).
C180 S-Acylation of BMPR1a Alters Fate of NSCs In Vivo. To probe potential effects of altered BMPR1a palmitoylation on NSC activity and fate, we isolated embryonic NSCs at E17.5 and analyzed their functional properties. We found that C180A mutation caused enhanced NSC proliferation in vitro, as measured by EdU pulse labeling (Fig. 4A). Furthermore we found increased levels of key markers of oligodendrocytic differentiation and maturation, OLIG2 (oligodendrocyte transcription factor) and NG2 (neural/glial antigen 2), after 1 and 4 d of in vitro differentiation by Western blotting (Fig. 4B and C and SI Appendix, Fig. S4A). Analysis by immunocytochemistry against NG2 confirmed that more NG2+ cells were present after 4 d of in vitro differentiation at the expense of a reduced number of GFAP+ (gliarial fibrillary acidic protein) cells (Fig. 4D), indicating that acylation-deficient C180A cells show an altered NSC behavior with increased oligodendrogenesis. Thus, our findings give direct evidence that BMPR1a S-acylation is an essential modulator of NSC activity and fate choices in vitro.
oligodendrogenesis, we analyzed the generation of late embryonic/early postnatal oligodendrocytic cells. Therefore, we injected E17.5 C180A and control mice with the thymidine analog BrdU and analyzed the number and fate of BrdU-labeled cells in postnatal brains at postnatal day 7 (P7). Corroborating the in vitro data, we detected an increased density of OLIG2\(^+\) cells in the corpus callosum and an increase in the BrdU\(^+\)/OLIG2\(^+\) fraction of BrdU\(^+\) cells in the cortex of P7 C180A mutant mice compared to controls (Fig. 4E and F). Thus, the observed increased oligodendrogenesis appears to be based on increased differentiation toward the oligodendrocytic lineage and an increase in proliferation of oligodendrocyte precursor cells.

Strikingly, we also found an increased density of OLIG2\(^+\), NG2, and APC\(^+\) (adenomatous polyposis coli) cells in the cortex (Fig. 5 A, C, and D), corpus callosum (Fig. 5 E and F), and hippocampus of adult mice (SI Appendix, Fig. S4B), confirming and extending the phenotype we had observed in the P7 postnatal brain (Fig. 4E). Interestingly, we found no differences in the number of NeuN-expressing neurons in the cortex (Fig. 5B) or in NSC activity (SI Appendix, Fig. S4C) in the neurogenic niche of the adult hippocampus that were not permissive for oligodendrogenesis (32) in C180A mutant mice compared to controls, suggesting that the palmitoylation-dependent function of BMPR1a requires the physiological oligodendroglial fate potential of endogenous NSCs.

Fig. 3. Palmitoylation at C180 is important for BMPR1a function. (A) Schematic of genomic changes introduced into the Bmpr1a C180A transgenic knock-in mouse. The genetic sequence coding for cysteine 180 of BMPR1a was altered to encode for an alanine. BMPR1a is expressed (green) at E17.5 in the ventricular zone and colocalizes with the stem cell–associated intermediate filament NESTIN (red), as assessed by immunohistochemistry. VZ, ventricular zone; IZ, intermediate zone; CP, cortical plate. (B) BMPR1a cell surface expression is reduced in Bmpr1a C180A–mutated NSCs (gray), as measured by cell surface protein biotinylation. The cell surface protein \(\beta\)-dystroglycan (\(\beta\)-DG) was used for normalization and to compensate for different reaction efficiencies. GAPDH and BMPR1a in the input and unbound fraction revealed a comparable loading and efficient enrichment of BMPR1a. IP, immunoprecipitation; IN, input fraction; UB, unbound fraction. (C) BMPR1a expression is unchanged in NSCs extracted from control (white) and Bmpr1a C180A (gray) embryos (E17.5). (D) The mobile fraction of overexpressed BMPR1a C180A is decreased compared to BMPR1a. Shown are FRAP experiments using BMPR1a-mNeonGreen-3HA (control) and BMPR1a-C180A-mNeonGreen-3HA. Note the reduced mobile fraction in C180A BMPR1a. (E) The rate of BMPR1a endocytosis, as measured by reversible biotinylation, is reduced in NSCs extracted from Bmpr1a C180A (gray) embryos compared to controls (white). The cell surface protein \(\beta\)-DG undergoing regular endocytosis was used as a loading control to normalize between sample-dependent differences in reaction efficiencies. (F) Noncanonical BMP signaling is affected in the Bmpr1a C180A knock-in NSCs (gray) compared to control cells (white), indicated by reduced active ERK 1/2 in knock-in cells. (Scale bars: 100 \(\mu\)m.) Cont, control. Error bars represent mean ± SD. *\(P < 0.05\), **\(P < 0.01\), ***\(P < 0.001\).
Discussion

We showed that a large number of proteins are palmitoylated in mammalian NSCs. Thus, we provided a palmitoylation proteome resource that will be helpful in discovering mechanisms of NSC control in the embryonic and adult mammalian brain. We focused on BMPR1a, newly discovered to be palmitoylated in NSCs, and showed that BMPR1a is modified by zDHHC20–mediated S-acylation of 3 cysteine residues that affect BMP signaling in vitro and in vivo. Thus, we provided direct in vivo evidence of NSC-associated palmitoylation of a target protein that affects the fate of mammalian NSCs. Future studies will aim to probe if enhanced oligodendrogenesis in C180A mice may also cause enhanced remyelination in the context of demyelinating lesions, which may open novel avenues to enhance myelin repair in the adult brain (33).

The identification of palmitoylation of BMPR1a may have broad consequences for our understanding of health and disease given that BMP signaling has been implicated in a large number of biological processes ranging from stem cell–associated differentiation to cancer growth (23, 34). Whether BMPR1a S-acylation is affected in disease states will need to be determined. Notably, palmitoylation of BMPR1a is not specific to NSCs, as we also found acylation of the receptor in pluripotent ESCs (37).
Appendix, Fig. S2 C–F). Unfortunately, exchange of cysteines C173/175 into alanines may be nonviable in mice, hindering in vivo analyses at this time. However, the findings that palmitoylation of C173/175 seems to be required for proper embryonic development further support the notion that palmitoylation of BMPR1a is of substantial functional relevance and that C173/175 S-acylation is required for canonical BMP signaling, whereas S-acylation of C180 affects noncanonical BMP signal transduction. Future studies will need to use either conditional or in vitro approaches to exchange S-acylated cysteines on both alleles, which
may allow more in-depth functional analyses of the C173/175 sites. As we showed, in vivo manipulation of palmitoylation of BMPR1α seems to selectively affect BMP signaling, resulting in enhanced generation of oligodendrocytes, and our in vitro data suggest that this is based on altered noncanonical BMP signaling.

BMP receptors are present at the cell surface as hetero- or homeric complexes (35), and while BMPR1α mainly associates with detergent-resistant membrane (DRM) fractions, BMPR2 was reported to localize to DRM and non-DRM compartments (36). However, it remains poorly understood how shifts between canonical and noncanonical BMP signaling are regulated (37–39). A prerequisite for SMAD-independent signaling is the presence of BMPR1 in cholesterol-rich microdomains regulating lateral mobility of type I receptors, which has been described to be crucial for regulating noncanonical BMP signaling without affecting canonical signaling function (36, 40). Thus, future studies are necessary to address the impact of BMPR1α S-acylation on its segregation to specific membrane fractions, its mobility, and the formation of receptor complexes.

We used a screening approach to identify palmitoylated proteins in mammalian NSCs. We showed that BMPR1α is S-acylated and that palmitoylation affects the functionality of the BMP signaling pathway that is critically involved in a variety of cell biological processes (Fig. 5G). Manipulating palmitoylation of BMP signaling components may represent an approach to targeting BMP function in health and disease.

Materials and Methods

Stem Cell Cultures. Primary adult mouse NSCs were kept in the presence of EGF (20 ng/ml), FGF-2 (20 ng/ml), and heparin (5 mg/ml) in a monolayer culture at 37 °C and 5% CO2. DMEM/F12 GlutaMax supplemented with antibiotics (Anti-Anti) and N2 was used as a standard medium. For some culture conditions, an overnight BMP4 treatment was performed as described earlier (41). For the generation of E17.5 cortical NSC cultures, pregnant conditions, an overnight BMP4 treatment was performed as described earlier (41). For the generation of E17.5 cortical NSC cultures, pregnant

Acetyl-Biotin Exchange Assay and Mass Spectrometric Analysis. The protocol was performed as previously described (16). Please refer to the SI Appendix for a more detailed description. Samples were adjusted to the same volume and concentration (5 mL and 2.2 mg/ml) by the addition of LB. Sample preparation, liquid chromatography (LC)–MS/MS analysis, and data analysis were performed as described previously (41) with some minor changes: raw files were processed with Progenesis QI for proteomics (Nonlinear Dynamics) and Gapdh, Actin, and Tubulin were used for normalization between samples. These proteins bind due to their high abundance to the beads in a palmitoyl-
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