Heparan sulfate antagonism alters bone morphogenetic protein signaling and receptor dynamics, suggesting a mechanism in hereditary multiple exostoses

Hereditary multiple exostoses (HME) is a pediatric disorder caused by heparan sulfate (HS) deficiency and is characterized by growth plate–associated osteochondromas. Previously, we found that osteochondroma formation in mouse models is preceded by ectopic bone morphogenetic protein (BMP) signaling in the perichondrium, but the mechanistic relationships between BMP signaling and HS deficiency remain unclear. Therefore, we used an HS antagonist (surfen) to investigate the effects of this HS interference on BMP signaling, ligand availability, cell-surface BMP receptor (BMPR) dynamics, and BMPR interactions in Ad-293 and C3H10T1/2 cells. As observed previously, the HS interference rapidly increased phosphorylated SMAD family member 1/5/8 levels. FACS analysis and immunoblots revealed that the cells possessed appreciable levels of endogenous cell-surface BMP2/4 that were unaffected by the HS antagonist, suggesting that BMP2/4 proteins remained surface-bound but became engaged in BMPR interactions and SMAD signaling. Indeed, surface mobility of SNAP-tagged BMPRII, measured by fluorescence recovery after photobleaching (FRAP), was modulated during the drug treatment. This suggested that the receptors had transitioned to lipid rafts acting as signaling centers, confirmed for BMPRII via ultracentrifugation to separate membrane subdomains.

In situ proximity ligation assays disclosed that the HS interference rapidly stimulates BMPRI–BMPRII interactions, measured by oligonucleotide-driven amplification signals. Our in vitro studies reveal that cell-associated HS controls BMP ligand availability and BMPR dynamics, interactions, and signaling, and largely restraints these processes. We propose that HS deficiency in HME may lead to extensive local BMP signaling and altered BMPR dynamics, triggering excessive cellular responses and osteochondroma formation.

Hereditary multiple exostoses (HME) is a congenital autosomal dominant disorder, also known as multiple osteochondromas, in which benign cartilage-capped bony tumors form along the border between the growth plate and perichondrium in long bones, ribs, vertebrae, pelvis, and cranial base in children and adolescents (1–4). Due to their location, large number, and size, the tumors (called exostoses or osteochondromas) can cause numerous health problems, including skeletal deformities, growth retardation, blood vessel and nerve impingement, early onset osteoarthritis, and chronic pain (5, 6). Most HME cases are caused by heterozygous loss-of-function mutations in the Golgi-associated and heparan sulfate (HS)-synthesizing enzyme EXT1 or EXT2 (7–10), resulting in a partial systemic HS deficiency (11, 12). It has long been assumed that osteochondroma formation is directly linked to HS loss, but a complete understanding of the nature of how this manifests into a pathogenic disorder is still lacking (13).

The HS chains are components of critical cell-surface and extracellular matrix–associated proteoglycans (HSPGs) that regulate numerous developmental and physiologic mechanisms and processes and in particular, the topography, range of action, and signaling activity of bone morphogenetic proteins (BMPs), fibroblast growth factors (FGFs), and other HS-binding signaling proteins (14–17). Indeed, we showed in previous studies that conditional Ext1/− ablation and ensuing severe decrease in HS levels caused ectopic canonical BMP signaling in the long-bone perichondrium in mouse models of HME (18). The induction of BMP signaling in the perichondrium was followed by a phenotypic switch in resident cells from mesenchymal/fibroblastic to chondrogenic and by formation of cartilaginous osteochondroma-like tissue masses over time. Our studies revealed for the first time that locally enhanced BMP signaling is a major culprit in osteochondroma induction and growth and that the tumors originate from perichondrium-associated stem and progenitor cells (13, 18). In very good agreement with these key findings, we showed in a more recent study

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This article contains Tables S1 and S2 and Figs. S1–S5.

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The abbreviations used are: HME, hereditary multiple exostoses; HS, heparan sulfate; HSGP, heparan sulfate proteoglycan; BMP, bone morphogenetic protein; rhBMP2, recombinant human BMP2; FGF, fibroblast growth factor; BMP, BMP receptor; FRAP, fluorescence recovery after photobleaching; pSMAD, phosphorylated SMAD; DRM, detergent-resistant membrane; DMEM, Dulbecco’s modified Eagle’s medium; HRP, horseradish peroxidase; BisTris, 2-[bis(2-hydroxyethyl)amino]-2-(hydroxymethyl)propane-1,3-diol; PLA, proximity ligation assay; ANOVA, analysis of variance; GAPDH, glyceraldehyde-3-phosphate dehydrogenase.

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that systemic administration of the BMP signaling antagonist LDN193189 markedly reduced osteochondroma formation in the HME mouse models (3), representing the first demonstration ever that osteochondroma formation is amenable to drug treatment. A study confirming our data has just been published (19). Together, the data indicated that a critical role of HS within developing and growing skeletal elements is to curb BMP action and signaling, possibly by limiting BMP availability and interactions with BMP receptors (BMPRs). Thus, aberrant function of these mechanisms resulting from decreases in HS levels can be pathogenic.

It is well established that cell-surface BMPRs are tetrameric complexes, each composed of two type I receptors (BMPRIa or BMPRIb) and two type II BMP receptors (BMPRII, ACVR2a, and ACVR2b) that transduce BMP action by mainly signaling via canonical phosphorylated SMAD1/5/8 proteins (20–23). Of particular relevance here are studies performed by Knaus and colleagues in which they analyzed and characterized the mechanisms of BMPR signaling in various types of cells in vitro (24–27). In particularly probing studies, they made use of combinations of high-resolution, live-cell imaging techniques and biochemical assays to investigate BMPR mobility, interactions, and signaling kinetics. They found that BMPRI and BMPRII have distinct mobility patterns under unstimulated conditions and that the highly mobile BMPRII population became immobilized and bound to BMPRI during rhBMP2 treatment. Data with C2C12 cells indicated that upon treatment with exogenous rhBMP2, the mobility of the BMPRII population was quickly reduced and the receptors were recruited into lipid rafts, where they oligomerized with the resident BMPRI population, eliciting canonical SMAD signaling (25).

Because of its potency and multiple regulatory functions, BMP signaling needs to be highly regulated (28–30). As pointed out above, BMP family members all possess a high-affinity and specific HS-binding domain, and thus, it is likely that their interactions with HS chains and HSPGs represent an important mechanism of regulation of BMP biological action (14, 17). However, details remain unclear. Kuo et al. (31) analyzed the role of HS in the signaling activity of recombinant BMP2 and BMP4 in C2C12 and PC12 cell cultures. They found that when the cells were pretreated with heparitinase, their responses to exogenous BMPs and canonical signaling were diminished, accompanied by a reduction in BMPRI/II oligomerization, as revealed by protein cross-linking, immunoprecipitation, and fluorescence correlation microscopy. In related studies, Jiao et al. (32) and Manton et al. (33) observed that heparitinase treatment actually enhanced BMP signaling and osteogenic cell differentiation in response to exogenous BMPs. Similarly, we observed in mouse embryo limb mesenchymal cells in high-density micromass cultures that chondrogenic cell differentiation and canonical BMP signaling were greatly stimulated by treatment with heparitinase, heparanase, or the HS antagonist surfen, in the absence of exogenous BMPs (18, 34). Others found that recombinant BMP2 and BMP4, in which the HS-binding region was mutated and nonfunctional, exhibited higher activity in cultured cells and a broader and stronger action in vivo, as measured by Xenopus embryo ventralization assays (35, 36). Together, current evidence points to the overall conclusion that HS and HSPGs exert complex regulatory roles in BMP and BMPR function. They appear to be needed to capture and retain BMPs and can then exert positive or negative modulation of BMP signaling activity in distinct contexts and processes.

In the present study, we have interrogated these mechanisms in greater detail and specifically asked to what extent HS regulates mobility, dynamics, interactions, and signaling of BMPRIa and BMPRII members. To do so, we used live-cell imaging, fluorescence recovery after photobleaching (FRAP), FACS, and other biochemical tests in cultured cells and analyzed and measured those parameters before and after acute HS interference. For the latter, we used surfen to modulate HS function rather than content or structure. The data provide strong evidence that a main role of cellular HS is to limit BMP signaling and BMPR dynamics, suggesting that alterations of such basic restraining mechanisms during severe HS deficiency could be deleterious and promote cell misbehavior and disease progression.

Results

HS interference elicits increases in endogenous BMP activity and SMAD signaling

To assess the roles of HS in regulating BMP signaling in manners reflective of cellular homeostatic and dynamic mechanisms, we initially carried out studies on endogenously expressed ligands and receptors. Thus, we first determined the steady-state levels of gene expression of representative BMPs and BMPRs in Ad-293 and C3H/10T1/2. The cell lines used here are popular and have been used in numerous studies, including those dealing with pediatric skeletal diseases (37, 38). RT-PCR analysis showed that both cell lines displayed readily detectable levels of transcripts encoding Bmp2, Bmp4, BmprIa, and BmprII (Fig. SI, A and B) as well as those for the BMP early response target gene Id1 (Fig. 1, A and B). Both cell lines also exhibited appreciable basal levels of phosphorylated SMAD1/5/8 (pSMAD1/5/8) (Fig. 1, C and E, lane 1) that were quickly enhanced severalfold within 15 min of treatment with rhBMP2 and remained so for at least 1 h (Fig. 1, C, G, E, and I, lanes 2–6). The above data indicated that both cell lines had a basal level of BMP signaling and were able to adjust and increase it upon changes in ligand availability. To determine the extent to which HS regulated such basal levels of BMP signaling, the cells were treated with surfen and processed for analysis and quantification of the above parameters. In previous studies using analytical tools, including surface plasmon resonance, we and others showed that surfen competes with protein factor binding to HS and can dislodge BMPs prebound to HS (18, 39). In good agreement, we observed that surfen provoked a clear increase in pSMAD1/5/8 levels within 30 min of treatment in Ad-293 cells, amounting to over 4.0-fold by 1–2 h (Fig. 1, D and H, lanes 2–6). Similar responses occurred in C3H/10T1/2 cells but were slower (Fig. 1, F and J, lanes 2–6). Surfen treatment also elicited a strong stimulation of Id1 expression (Fig. 1, A and B). Notably and importantly, the drug-induced responses were largely prevented by co-treatment with recombinant noggin, a potent BMP antagonist (Fig. S2) (40). In addition, noggin treatment
decreased (i) basal Id1 expression and rhBMP2-induced Id1 overexpression (Fig. S2, A–C) and (ii) basal, rhBMP2-induced, and drug-induced pSMAD1/5/8 levels (Fig. S2, D–G). In sum, the above data indicate that basal levels of BMP signaling in Ad-293 and C3H/10T1/2 are readily increased following functional interference with HS. Because such responses were counteracted by noggin co-treatment, they probably reflected an increase in availability and activity of endogenous ligands at the cell surface.

To strengthen the latter conclusion, we directly investigated the possible presence of BMPs on the cell surface, using FACS. Ad-293 cells were lightly fixed with formaldehyde (and thus still impermeable), were incubated with primary rabbit BMP2/4 antibodies followed by secondary Alexa Fluor-488–labeled antibodies, and were finally subjected to FACS. Companion cells pretreated with surfen for 1 h were processed at the same time. Both cell populations produced strong signal patterns consisting of a prominent fluorescence peak at intensity 10^5 to 10^6 preceded by a small peak at about 10^4, probably representing background signal (Fig. 2, D and E). Relative fluorescence values for both populations were extremely strong (Fig. 2, F, G, H, I, J, and K).

Results were analyzed using one-way ANOVA with Tukey’s multiple-comparison tests and are expressed as mean ± S.D. (error bars) for n = 3 separate experiments.
than those elicited by cells exposed to primary immune and secondary antibodies (Fig. 2F, 1° + 2° abs and 1° + 2° abs + Surfen). When Ad-293 cells were pretreated with trypsin for 15 min, fixed, and then reacted with primary immune and secondary antibodies, their fluorescence signal was drastically reduced (Fig. 2F, trypsin-treated conditions), indicating that BMP2/4 were surface-bound and susceptible to protease digestion. To further double-check the data, untreated cells and companion cells treated with surfen for 1 h were processed for immunoblot analysis of endogenous BMP2, using a rabbit monoclonal antibody. Indeed, both populations elicited a single prominent immunoreactive band of 15 kDa (Fig. 2G) under reducing conditions and present in similar relative amounts in both cell populations (Fig. 2H). Together, the above lines of experimentation indicate that Ad-293 cells do possess endogenous cell surface–bound BMP2 and/or BMP4. The apparent protein levels do not substantially change after acute drug treatment, suggesting that surfen did not dislodge the proteins away from the surface but probably made them available for BMPR interactions and signaling (see Fig. 1).

**BMPR mobility and association with lipid rafts change upon HS interference**

Given that BMP signaling and pSMAD1/5/8 levels increased readily after surfen treatment, we asked whether these responses were associated with, and probably due to, changes in BMPR mobility and clusterization into lipid rafts (41). For these analyses, we resorted to live-cell imaging during FRAP and created BMPR fusion protein expression constructs, using the SNAP-tag technology (42, 43). This approach has significant advantages over traditional methods, including the fact that the tag can be labeled with diverse nonpermeant fluorophores and is small, thus likely to interfere less, or not at all, with the fusion protein’s routing and destination through cellular compartments, including the endoplasmic reticulum, Golgi, and cell surface. Accordingly, Ad-293 cells were transfected with SNAP-BMPRIα or SNAP-BMPRII expression plasmids, and found that each cell population expressed the respective protein of appropriate molecular weight (Fig. S3, A–C). To demonstrate that the presence of the fusion proteins did not alter BMP signaling and may actually render the cells more responsive, Ad-293 cells were co-trans-
The significant decrease in BMPRII lateral mobility triggered by drug treatment above probably indicates that this receptor...
had been recruited to lipid raft domains well known to serve as centers for receptor oligomerization and signal transduction (41). To test this prediction, we determined the distribution of SNAP-BMPRII and SNAP-BMPRIa in detergent-resistant membranes (DRMs), which are mainly composed of lipid rafts (44, 45). Accordingly, Ad-293 cells expressing SNAP-BMPRII or SNAP-BMPRIa were treated with surfen or vehicle for 10–15 min and extracted with nonionic detergents, and the resulting cell extracts were processed for DRM fractionation by sucrose gradient ultracentrifugation (44, 45). Gradients were subdivided into consecutive fractions, and proteins in each fraction were separated and analyzed by gel electrophoresis and immunoblots. Caveolin-1 was used as marker of lipid rafts (41, 46). In vehicle-treated control cells, the receptor populations were mainly distributed over the cell surface (Fig. 6, A, E, I, and M). Microscopic inspection and image-based quantification of at least 30 cells in three separate experiments showed that control cells had an average of about 45 detectable dots/cell in Ad-293 cultures and about 30 detectable dots/cell in C3H/10T1/2 cultures (Fig. 7, Q and R). Strikingly and clearly, an acute treatment (10–15 min) with rhBMP2 or surfen led to a sharp increase in the number of fluorescent dots that was quite apparent and obvious by simple microscopic inspection (Fig. 7, B, F, J, and N (rhBMP2) and C, G, K, and O (surfen)) and was confirmed by quantification and statistical analyses (Fig. 7, Q and R). To demonstrate that these receptor interactions were in fact due to ligand availability, we preincubated rhBMP2 with noggin or surfen with noggin for 10 min at 37 °C. We then applied the rhBMP2/noggin and surfen/noggin mixtures (or noggin alone) to the cells for 10–15 min. A proximity ligation assay (PLA) was carried out, and indeed, we found that noggin prevented rhBMP2-induced (Fig. 55, B, E, and G) and drug-induced (Fig. 55, C, F, and G) BMPRIa and BMPRII interactions compared with BMP2 and surfen alone (Fig. 7). Companion cultures exposed to preimmune antibodies (same species, same dilution) and then processed for the entire proximity ligation procedure elicited no detectable signal (Fig. 7, D, H, L, and P), attesting to the specificity of the procedure.

Discussion

Like many other members of key signaling protein families (14, 17), BMPs have long been known to possess an HS-binding domain (36), but the importance and functional roles of the resulting HS-BMP interactions for BMPR action have remained poorly understood. The in vitro data we present here provide a series of compelling and interconnected observations leading to the overall conclusion that HS has a clear role in regulating the dynamic behavior of BMPRs and in modulating BMP signaling. Based on cellular responses detailed above, it appears that HS largely has a restraining influence on BMP activity and signaling and that its functional deficiency leads to increased signaling. Given that these responses are appreciable in the absence of exogenously provided BMPs, they clearly

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**Figure 5. HS interference does not significantly alter BMPRIa surface mobility.** A, representative SNAP-BMPRIa–expressing Ad-293 cell in which a region of interest was identified and is depicted by a yellow box. B–E, magnified images of the region of interest show prebleach at 0 s, bleach at 50 s, and recovery at 130 and 215 s. F–J, fluorescence intensity quantification of photobleached area and calculated half-time recovery (β1/2; Table 1). Treatment with rhBMP2 or surfen caused no major difference in β1/2 (G–I) or Fm (J) when compared with control SNAP-BMPRIa–expressing 293 cells. Results were analyzed using one-way ANOVA with Tukey’s multiple-comparison tests and are expressed as mean ± S.D. (error bars) for n = 17–30 cells/condition.

**HS interference leads to increased interactions between BMPRIa and BMPRII**

The data above indicate that interference with HS promoted the recruitment of the BMPRII to lipid rafts, where they would presumably interact with resident BMPRIa to elicit an increase in BMP signaling (see Fig. 6). To obtain more direct evidence of changes in BMPRII/BMPRIa interactions upon HS interference, we carried out proximity ligation assays (47, 48). Ad-293 and C3H/10T1/2 cells were first treated with surfen, rhBMP2, or vehicle for 10–15 min and fixed. After blocking, the cells were incubated with oligonucleotide-containing antibodies to BMPRIa or BMPRII, rinsed, and incubated with a hybridization solution containing connector oligomers binding to their respective antibodies. This was followed by ligation and amplification in the presence of a 594 fluorophore, generating bright red fluorescence dots on the cell surface indicative of closely associated and interacting proteins (47, 48). Vehicle-treated cells displayed a basal number of red fluorescence dots that were uniformly distributed over the cell surface (Fig. 7, A, E, I, and M). Microscopic inspection and image-based quantification of at least 30 cells in three separate experiments showed that control cells had an average of about 45 detectable dots/cell in Ad-293 cultures and about 30 detectable dots/cell in C3H/10T1/2 cultures (Fig. 7, Q and R). Strikingly and clearly, an acute treatment (10–15 min) with rhBMP2 or surfen led to a sharp increase in the number of fluorescent dots that was quite apparent and obvious by simple microscopic inspection (Fig. 7, B, F, J, and N (rhBMP2) and C, G, K, and O (surfen)) and was confirmed by quantification and statistical analyses (Fig. 7, Q and R). To demonstrate that these receptor interactions were in fact due to ligand availability, we preincubated rhBMP2 with noggin or surfen with noggin for 10 min at 37 °C. We then applied the rhBMP2/noggin and surfen/noggin mixtures (or noggin alone) to the cells for 10–15 min. A proximity ligation assay (PLA) was carried out, and indeed, we found that noggin prevented rhBMP2-induced (Fig. 55, B, E, and G) and drug-induced (Fig. 55, C, F, and G) BMPRIa and BMPRII interactions compared with BMP2 and surfen alone (Fig. 7). Companion cultures exposed to preimmune antibodies (same species, same dilution) and then processed for the entire proximity ligation procedure elicited no detectable signal (Fig. 7, D, H, L, and P), attesting to the specificity of the procedure.

**Discussion**

Like many other members of key signaling protein families (14, 17), BMPs have long been known to possess an HS-binding domain (36), but the importance and functional roles of the resulting HS-BMP interactions for BMPR action have remained poorly understood. The in vitro data we present here provide a series of compelling and interconnected observations leading to the overall conclusion that HS has a clear role in regulating the dynamic behavior of BMPRs and in modulating BMP signaling. Based on cellular responses detailed above, it appears that HS largely has a restraining influence on BMP activity and signaling and that its functional deficiency leads to increased signaling. Given that these responses are appreciable in the absence of exogenously provided BMPs, they clearly
reflect mechanisms engaging resident endogenous ligands and receptors and, importantly, their native stoichiometries and dispositions. Admittedly, the signaling responses to drug treatment are slower than those elicited by exogenous rhBMP2 treatment and are also smaller in scale. However, robust responses to exogenous BMPs are likely to be unrealistic and above physiological levels, given that even at nanomolar rhBMP2 doses used here and in previous studies, such doses are bound to be vastly higher than those of endogenous ligands (probably including active and precursor forms). Together, our data suggest that the cell surface of Ad-293 and C3H/10T1/2 cells contains native BMPs interacting with HS and potentially available for action, becoming engaged in signaling upon modulation of HS restraining influence (Fig. 8).

Cell-surface receptors are highly dynamic entities, can move in and out of lipid rafts, and can establish functional interactions with their respective partners and appropriate ligands to signal (49, 50). The BMPRs conform to these general trends, and, as indicated above, different BMPRs examined so far have been found to have distinct cell-surface translocation dynamics and distribution. In cells such as C2C12, a large portion of the BMPRIIa population resides in lipid rafts and is rather static, whereas the BMPRII population is highly dynamic and seemingly distributed more broadly on the surface (20, 25, 26). The BMPRIIs quickly transition to lipid rafts upon acute treatment with rhBMP2 or rhBMP4. Our data with rhBMP2-treated Ad-293 and C3H/10T1/2 cells largely agree with those observations, implying that the differential dynamics and distribution of BMPRII versus BMPRIa may reflect fairly general cellular characteristics and that the recruitment of BMPRII to lipid rafts may set the overall degree of signaling. Our data, however, provide the first demonstration ever that a similar physical rearrangement of the receptors occurs upon functional interference with HS in the absence of exogenous rhBMPs. Because signaling eventually follows such receptor redistribution, the data imply that the dynamic response of the receptors to HS deficiency and their direct interactions were fruitful and productive and led to downstream effector action. It is important to note here that our data only explore one of the three type II BMP receptors; therefore, it is unknown whether ACVR2a or ACVR2b would behave in a similar manner to BMPRII. The data are in line with the possibility raised above that the Ad-293 and C3H/10T1/2 cells possess a reservoir of endogenous cell-surface BMPs that can be recruited and engaged in signaling action upon modulation of HS influence. This notion is supported by the fact that signaling in response to surfen was prevented by noggin co-treatment, strongly indicating that the endogenous ligands were accessible to, and blocked by, such an external protein antagonist. Our FACS data also are in line with that notion by showing that cell-surface BMP2/4 were accessi-

Figure 6. Surfen promotes recruitment of BMPRII population to lipid raft domains. Vehicle-treated (A and B) and surfen-treated Ad-293 cells (C and D) expressing SNAP-BMPRII or SNAP-BMPRIa were homogenized and fractionated by sucrose gradient ultracentrifugation. Fractions were blotted with anti-SNAP or anti-caveolin 1 antibodies. A, in vehicle-treated cells, the BMPRII population is distributed in both lipid raft and nonlipid raft domains. C, in surfen-treated cells, there is a robust shift in BMPRII population to lipid raft domains, depicted in particular in fraction 1. B, in vehicle-treated cells, the BMPRIa population is mostly in the lipid raft domains. D, in surfen-treated cells, the majority of the BMPRIa population is seen in lipid raft domains, but there are a significant amount of receptors in the nonlipid raft domains compared with the control cells. E and F, quantification of receptor population in either DRM or non-DRM (*, p < 0.05). Results were analyzed using Student’s t test and are expressed as mean ± S.D. (error bars) for n = 3 separate experiments.
Figure 7. HS interference leads to increased BMPRIa and BMPRII interactions. Antibody-based in situ proximity ligation assays were used to analyze BMPRIa and BMPRII interactions in Ad-293 and C3H10T1/2 cells. Treatment with rhBMP2 (25 ng/ml) or surfen (5 μM) led to a striking increase in receptor-receptor interactions (depicted by red fluorescence dots) in both Ad-293 cells (B, C, F, and G) and C3H/10T1/2 cells (J, K, N, and O) compared with control cells (A, E, I, and M). Companion cells reacted with preimmune antibodies exhibited no signal (D, H, L, and P), attesting to the specificity of the analysis. Note that E–H and M–P are magnified images of representative fields from each culture shown in A–D and I–L. Scale bar, 50 μm. Q and R, scatterplots depicting quantification of fluorescent signal per cell in each condition (**, p < 0.01; ***, p < 0.001; ****, p < 0.0001). Results were analyzed using one-way ANOVA with Tukey’s multiple-comparison tests and are expressed as mean ± S.D. (error bars) for n = 17–30 cells/condition.

Figure 8. Schematic depicting distribution of BMPRs and HSPGs before or after treatment with rhBMP2 or surfen. A, in control cells, the BMPRIa population would be located predominantly in lipid raft domains (orange membrane section), whereas the BMPRII population would be more widely distributed over nonlipid raft domains. Endogenous BMPs would largely be bound to HS and limit interactions with BMPRs and signaling. B, treatment with rhBMP2 would cause a major and rapid redistribution of BMPRIIs to lipid rafts, interactions with resident BMPRIa population, and strong downstream signaling. Excess rhBMP2 would accumulate onto HSPGs. C, in surfen-treated cells, the drug would compete with HS-BMP binding; promote availability and engagement of endogenous BMPs; and stimulate BMPRII recruitment to lipid rafts, interactions with BMPRIa, and downstream signaling.
ble to their antibodies, elicited a highly specific fluorescence signal, and were susceptible to protease degradation in live cells. Because the FACS profiles did not change substantially after surfen treatment, the data do indicate that the treatments did not dislodge a significant amount of endogenous ligands from the surface, instead presumably making them available for interactions and signaling with resident BMPRs.

Cell-surface BMPR dynamics have been previously studied by FRAP as well as FRET and protein cross-linking and immuno-precipitation, but not by proximity ligation. A major advantage of this procedure is that it provides direct visualization of endogenous protein-protein complexes and their distribution and composition within individual cells, including between cytoplasmic proteins or transcription factors, such as c-Myc and Max (47, 48). Measurements and computation have indicated that the complexes reflect interacting proteins at an average distance of 10–20 nm from each other. In line with pSMAD1/5/8 levels reflective of basal BMP signaling (in the absence of exogenous BMPs), untreated control Ad-293 and C3H/10T1/2 cells displayed an appreciable number of BMPR-Ia-BMPRII complexes on their surface depicted by the red fluorescence amplification spots. Within a 10–15-min treatment with rhBMP2, the red spot number increased as sharply as it did after surfen treatment (again without exogenous rhBMP2). Because rhBMP2 treatment caused a higher increase in spot number and higher pSMAD1/5/8 levels compared with surfen treatment, the data raise the interesting possibility that there is a fairly direct correlation between the number of surface BMPR complexes and the degree and levels of BMP signaling, without a need to invoke changes in catalytic capacity of each complex depending on treatment type or experimental condition. As in the case of data from FRAP or FRET analyses (25–27), the increases in complex numbers revealed by the ligation assays occurred within minutes of surfen treatment, in line with the highly dynamic nature of cell-surface receptors in response to ligands becoming acutely available. We should consider the possibility, however, that HS may have additional influences on the physical behavior of BMPRs and their signaling activity. BMPRs do not possess a stereotypic HS-binding domain (14), but it is possible that they may establish links to HS and HSPGs via intermediaries or with the PG core proteins. For instance, the core proteins syndecan-1 and syndecan-2 were shown to interact with integrins and the tyrosine phosphatase receptor CD148 on the cell surface (51, 52). In addition, HS is an integral component of the FGF–FGF receptor signaling complexes and is required for FGF signaling (53, 54). In addition, surfen may have influenced other glycosaminoglycans (39). Considered together, our data and previous studies point to the overall notion that HS may often, if not always, be involved in regulating protein signaling by influencing ligand availability and cell-surface receptor dynamics and function, although in distinct and even opposing manners and depending on receptor types, signaling nature, and cell types and context.

The above conclusions and putative scenarios are in agreement with the studies by Jiao et al. (32) and Manton et al. (33) indicating that BMP signaling and osteogenic cell differentiation increased in response to treatment with heparitinase and rhBMPs (55). They also agree with studies showing that recombinant BMP2 and BMP4 lacking a functional HS-binding region exhibited greater activity in cultured cells and a broader and stronger action in vivo as measured in Xenopus embryos (35, 36). Together with the data here and our previous work on limb bud chondrogenitor cells (18), current evidence does sustain the notion that HS largely limits BMP signaling in many/most contexts. However, this notion appears to be in sharp contrast with the observations by Kuo et al. (31) indicating that heparitinase pretreatment greatly diminished BMP signaling and BMPR interactions in PC12 and C2C12 cells in response to exogenous rhBMP2. One admittedly trivial explanation for these divergent observations is that different cell culture techniques, media, recombinant protein purity, and other factors may have influenced cell behavior and responses and, thus, experimental outcomes. A more interesting explanation is that the data are actually reconcilable, but we need to take into account whether exogenous BMPs were used and whether short- or long-term responses were analyzed. Thus, in the Kuo et al. study (31), the pretreatment with heparitinase may have decreased the ability of the cells to acutely “capture” the rhBMP2 added to the culture medium, leading to reduced pSMAD1/5/8 levels within the 1-h period of analysis compared with heparitinase-untreated cells. On the other hand, the positive effects of heparitinase treatment on osteogenic cell differentiation may reflect lower but prolonged responses to exogenous rhBMP2, eliciting beneficial differentiation effects long-term. In the case of chondrogenic differentiation we studied in the absence of exogenous BMPs, the beneficial effects of surfen or heparitinase would also reflect slow, low but sustained stimulation of BMP signaling. In sum, HS and HSPGs probably have multiple interconnected roles in the regulation of BMP and BMPR signaling. They appear to be needed to capture BMPs and retain/preserve them on the cell surface, making them readily available for signaling upon perturbation of the HS restraining function.

The observations here provide possible insights into the cellular pathogenesis of HME. It has long been known that the heterozygous loss-of-function mutations in Ext1 or Ext2 present in HME patients cause a systemic deficiency in HS levels of about 50% (11). This partial deficiency can in itself provoke certain physiologic abnormalities, including substandard lipid clearance and pancreas function (56, 57). However, it is not sufficient to cause formation of osteochondromas that underlie the more severe health problems in HME patients, including skeletal deformations, chronic pain, and even malignancy. In line with Knudson’s law of tumorogenesis (58), osteochondroma formation requires a “second hit,” such as loss of heterozygosity or other genetic changes that would lead to a steeper local drop in HS levels, as previous studies from our group and others have demonstrated (59–62). But how would such a steep drop in HS lead to osteochondroma formation? Because the tumors invariably form next to the growth plate of long bones and other skeletal elements in children and adolescents, we reasoned that perichondrium flanking the growth plates could be a key pathogenic player (18). Indeed, we showed that conditional ablation of both Ext1 alleles in mouse models of HME led to steep local loss of HS, changes in perichondrial cell phenotype, ectopic BMP signaling and chondrogenesis, and
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initiation of osteochondroma formation. Interestingly (and in line with the observations here), we observed the same chronological and phenotypic changes when WT mouse long-bone explants in organ culture were treated with surfen or heparitinase (18). Both treatments caused a prominent induction of ectopic BMP signaling in the perichondrium (as indicated by pSMAD1/5/8 levels) that was followed by ectopic chondrogenesis and osteochondroma-like tumor formation. These responses were absent in companion vehicle-treated controls. Thus, and as strongly indicated by the data in the present study, it is possible that under normal conditions, HS in perichondrium would restrain the activity of BMPs that are expressed in that tissue (63–65), preventing them from exerting their notorious pro-chondrogenic action and allowing the perichondrium to maintain its normal fibroblastic and mesenchymal character. A local steep drop in HS levels or function would, however, liberate BMPs, allow them to interact with their receptors, elicit canonical BMP signaling, and trigger ectopic chondrogenesis and osteochondroma initiation. Notably, the perichondrium possesses additional mechanisms that normally protect and sustain its mesenchymal character, including anti-chondrogenic factors and pathways such as FGFs and FGF signaling mediators ERK1/2 (40, 66). The steep local drop in HS leading to osteochondroma initiation would thus need to alter such mechanisms as well. Given that FGF signaling actually requires HS, it is possible that a severe local HS deficiency in HME could cause a decrease in FGF-dependent anti-chondrogenic action and a reciprocal increase in BMP-dependent pro-chondrogenic action, a combination of responses that could converge to promote ectopic chondrogenesis and osteochondroma formation. It is important to point out that these novel findings should be extended in cell types that are relevant to HME to fully understand the mechanisms of pathogenesis.

Experimental procedures

Construction of SNAP-BMPRIa and SNAP-BMPRII plasmids

Design and construction of these fusion protein vectors were based on previous studies (42, 43). BMPRIa and BMPRII cDNA clones and the pSNAPp vector were purchased from Origene and New England Biolabs, respectively. The pSNAPP vector has two multiple cloning sites that flank the N- and C-terminal ends of the SNAP protein. Thus, we designed primers to amplify the signal peptide with restriction enzymes Nhel and EcoRI and mature peptide with restriction enzymes BamHI and NotI of the receptors (Table S1). After electrophoresis, we cut and purified the inserts. To insert the signal peptide of the receptors into the pSNAPP vector, we first digested the vector with Nhel and EcoRI and then inserted the signal peptide via ligation using the Quick Ligation Kit according to the manufacturer’s protocol (New England Biolabs). After transformation, colonies were selected and subjected to PCR to identify positive colonies for the signal peptide. Those colonies were further amplified and purified. The newly constructed plasmids, RIA signal peptide–SNAP and RII signal peptide–SNAP, were cut with BamHI and NotI. The inserts of the mature peptide of the receptors were ligated and transformed. The transformed colonies were selected, subjected to colony PCR, and purified. In sum, for SNAP-BMPRIa, the SNAP-tag is positioned at the end of the signal peptide at the glutamine residue (position 22) and before the start of the mature peptide at the glycine residue (position 23). For SNAP-BMPRII, the SNAP-tag is positioned after the alanine residue (from position 25 to the end of the signal peptide) and before alanine (from position 26 to the start of the mature peptide). The final plasmids, referred to as SNAP-BMPRIa and SNAP-BMPRII, were submitted to our Nucleic Acid and Protein Core Facility to verify sequencing and construction.

Protein analysis in cell cultures

Ad-293 cell line, a derivative of the commonly used HEK 293 cell line with improved adherence, was purchased from Agilent Technologies, and the C3H/10T1/2 cell line was purchased from ATCC. Cells were grown in monolayer and transfected with SNAP-BMPRIa and/or SNAP-BMPRII or left untransfected. Transfection was carried out using Fugene 6 (Promega) reagent according to the manufacturer’s protocol. Briefly, cells grown in 6-well plates for protein analysis were transfected with 1 μg of plasmid and 3 μl of Fugene 6 reagent per well in 10% FBS/DMEM and incubated for 24–48 h. Following transfection, the medium was changed to 0.1% BSA/DMEM, and the next day, the cells were treated with vehicle (control), rhBMP2 (25 ng/ml; Gemini Bioproducts), surfen (5 μM; Open Chemical Repository: NSC 12155), noggin (50 ng/ml; R&D Systems), rhBMP2 (25 ng/ml) plus noggin (50 ng/ml), and surfen (5 μM) plus noggin (50 ng/ml) in 0.1% BSA/DMEM for 15 min, 30 min, 1 h, 2 h, and 6 h. Doses used were based on previous studies (18, 67). Cultures were lysed in 1× radioimmune precipitation assay buffer with protease and phosphatase inhibitors, samples were centrifuged at 13,200 rpm at 4 °C, and supernatants were collected. Protein concentration for each sample was determined using the MicroBCA protein assay kit (Thermo Scientific) according to the manufacturer’s protocol. Total cellular proteins (30 μg/lane) were electrophoresed on 4–12% NuPAGE BisTris gels (Life Technologies) and transferred to polyvinylidene difluoride membranes (Millipore). Membranes were blocked in 5% BSA, 1× Tris-buffered saline/Tween 20 (TBST) and incubated overnight at 4 °C with pSMAD1/5/8 (1:1000; Cell Signaling). Membranes were washed in 1× TBST and incubated with anti-rabbit HRP-linked antibody (1:2000; Cell Signaling) for 1 h at room temperature. Antigen-antibody complexes were detected with the SuperSignal® West Dura Extended Duration Substrate (Thermo Scientific) chemiluminescent detection system using the ImageQuant LAS 4000 luminescent image analyzer (GE Healthcare). Membranes were reblotted with SMAD1 (1:1000; Cell Signaling) for normalization. For loading control, membranes were blotted with GAPDH antibodies (1:1000; Santa Cruz Biotechnology). To detect the mature form of BMP2, membranes were blocked in 5% nonfat dried milk, 1× TBST and incubated overnight at 4 °C with anti-BMP2 (1:1000; Abcam) and GAPDH (1:1000; Santa Cruz Biotechnology). After washing, the membranes were incubated with secondary antibodies, anti-rabbit HRP-linked antibody, and anti-mouse HRP-linked antibody. Primary and secondary antibodies were diluted in 5% nonfat dried milk, 1× TBST. Antigen-antibody complexes were detected as mentioned above. ImageJ was used to determine band intensities.
**Gene expression analysis**

Total RNA was isolated from control, rhBMP2-treated, surfen-treated, noggin-treated, rhBMP2 plus noggin-treated, and surfen plus noggin-treated transfected or nontransfected Ad-293 and C3H/10T1/12 cultured cells from 6-well plates (67), using TRIzol reagent (catalog no. 15596-026, Life Technologies) according to the manufacturer’s protocol. RNA was quantified by Nanodrop. One microgram of total RNA was reverse-transcribed using the Verso cDNA kit (catalog no. AB1435/A, Thermo Scientific). Quantitative real-time PCR was carried out using SYBR Green PCR Master Mix (Applied Biosystems) in an Applied Biosystems 7500 machine according to the manufacturer’s protocol. *Gapdh* was used as the endogenous control, and relative expression was calculated using the △△CT method. Real-time PCR was performed using GoTaq DNA polymerase (Promega) in a ProFlex PCR system (Applied Biosystems) in an Applied Biosystems 7500 machine according to the manufacturer’s protocol.

**FACS**

Ad-293 cells were grown in DMEM containing 10% FBS in 6-well plates. The cells were treated with DMSO or surfen (5 μM) for 1 h at 37 °C in DMEM containing 0.1% BSA. Following treatment, the cells were washed with 1× PBS and removed from the plates by scraping in 1× PBS, 2 mM EDTA or by trypsin. The cells were washed with 1× PBS and fixed with 2% buffered formalin for 15 min on ice. The cells were washed with 1× PBS and blocked by incubation in 1× PBS, 1% BSA (PBSB) for 10 min on ice. Aliquots of 106 cells (100 μl) were incubated with control mouse IgG (Cell Signaling, catalog no. 5415) or anti-BMP2/4 antibody (H-1; Santa Cruz Biotechnology) in 1× PBSB on ice for 2 h. The cells were washed and incubated with an Alexa 488 goat anti-mouse IgG (Jackson ImmunoResearch) for 1 h on ice. The cells were washed with 1× PBS and analyzed on a BD Accuri flow cytometer, located in the Flow Cytometry Core Laboratory at Children’s Hospital of Philadelphia. The data were analyzed using CFlow Plus software. The Accuri C6 recorded 10,000 events and then the Ad-293 cell population. The established gate was used in subsequent readings (10,000 events), and the mean FL1-A values of the gated cells were graphed.

**FRAP**

Ad-293 cells were grown in 35-mm glass-bottom well dishes coated with poly-L-lysine (MatTek) and transfected with 1 μg/35-mm dish of SNAP-BMPRIα or SNAP-BMPRII for 24 h using Fugene 6 transfection reagent. After 24 h, the cells were incubated with 2 μM of SNAP-tag 488 (New England Biolabs) for 30 min at 37 °C. Cells were then washed three times with complete medium and imaged in FluoroBrite™ DMEM medium (Thermo Fisher Scientific). Cells were stimulated with vehicle (1× PBS or DMEM), rhBMP2 (50 ng/ml), surfen (5 μM), noggin (50 ng/ml), rhBMP2 plus noggin, or surfen plus noggin. FRAP was performed on a laser-scanning confocal microscope (Olympus FV-1000) with an IX81 inverted microscope equipped with dual scanners for simultaneous imaging and stimulation bleaching. A 40× oil immersion objective lens with an aperture of 1.30 (Olympus UPlanFL) was used for imaging. Additionally, we used an argon ion laser and HeNe laser for imaging and a 405 diode laser for bleaching under the control of Fluoview software. 35-mm glass bottom well dishes were placed in a circular holder connected to a heating system so that all FRAP assays were carried out at 37 °C. A small circular region of the membrane was bleached for 200 ms at 50% laser power. Images were taken approximately every 3 s over a period of 250 s. A series of 15 prebleached images were taken, followed by the bleaching and recovery. The images were analyzed by Fiji (ImageJ) and corrected and normalized by FRAP Analyzer software. The normalized values were plotted nonlinear regression curve fit and one-phase association (GraphPad Prism version 6) to calculate the half-time recovery (τ1/2) and mobile fraction (Fm).

**DRM isolation**

Ad-293 cells were grown in 100-mm dishes and transfected with 8 μg of SNAP-BMPRIα or SNAP-BMPRII plasmid and 18 μl of Fugene 6 reagent per 100-mm dish. After 48 h, cells were treated with vehicle (1× PBS or DMSO) or surfen (5 μM) for 10 min at 37 °C. DRM isolation was carried out according to previous studies (44, 45). Briefly, cells were washed with ice-cold 1× PBS and ice-cold 1× Tris-HCl, NaCl, and EDTA (TNE). Lysates were collected in 1× TNE and centrifuged for 5 min at 380 × g at 4 °C. The pellets were resuspended in 275 μl of 1× TNE plus protease inhibitors and homogenized via a 25-gauge needle. Following homogenization, 250 μl of 2% Triton X-100 was added to 250 μl of the cell homogenate and placed on ice for 30 min. For the step sucrose gradient, 1.25 ml of 56% sucrose was added to 250 μl of 35% sucrose in 1× TNE plus protease inhibitors and homogenized via a 25-gauge needle. Following homogenization, 250 μl of 2% Triton X-100 was added to 250 μl of the cell homogenate and placed on ice for 30 min. For the step sucrose gradient, 1.25 ml of 56% sucrose was added to the sample for a final percentage of 40% sucrose. The solution was transferred to a 5-ml centrifuge tube (Beckman Coulter). The sample was overlaid with 2.75 ml of 35% sucrose and then 0.25 ml of 5% sucrose. The samples were centrifuged for 18 h at 39,000 rpm at 4 °C using an SW55 Ti rotor. The next day, 15 fractions of 300 μl each were collected per condition. The samples were electrophoresed on 4–12% NuPAGE BisTris gels and transferred to polyvinylidene difluoride membranes. Membranes were blocked in TBST and incubated overnight at 4 °C with anti-SNAP-tag (1:1000; New England Biolabs) and caveolin-1 (1:1000; Cell Signaling). Membranes were washed in 1× TBST and incubated with anti-rabbit HRP-linked antibody (1:2000; Cell Signaling) for 1 h at room temperature. Antigen-antibody complexes were detected with the SuperSignal® West Dura Extended Duration Substrate chemiluminescent detection system.

**PLA**

This procedure was carried out based on established methods (47, 48). Ad-293 cells and C2H/10T1/2 cells were grown on poly-L-lysine–coated coverslips in 12-well plates. Cells were treated with vehicle (1× PBS/DMEM), rhBMP2 (50 ng/ml) or surfen (5 μM) for 10 min at 37 °C. For noggin experiments, rhBMP2 (25 ng/ml) or surfen (5 μM) were preincubated with noggin (50 ng/ml) for 10 min at 37 °C before the addition to the cell culture medium. Cells were then washed with 1× PBS and fixed in 2% paraformaldehyde. After fixation, cells were rinsed in 1× PBS and blocked in 1% BSA, 1× PBS for 30 min at room
temperature. Cells were incubated with the following antibodies overnight at 4 °C: rabbit anti-BMPRIa (1:75; Origene) and mouse anti-BMPRII (1:250; Thermo Fisher Scientific). Mouse IgG (Cell Signaling) and rabbit IgG (Cell Signaling) were used for negative controls. The next day, we used the Duolink In situ Red Starter PLA Kit Mouse/Rabbit (Sigma) according to the manufacturer’s protocol. Briefly, the cells were washed with 1× PBS and incubated with anti-rabbit plus probe and anti-mouse minus probe solution for 1 h at 37 °C. Next, cells were incubated in ligation solution for 30 min at 37 °C. Following ligation, the signal was amplified via polymerase by incubating the cells in amplification solution for 100 min at 37 °C. Cells were washed with wash buffer and allowed to dry for 10 min in the dark. Coverslips were inverted and mounted onto glass slides using Duolink In Situ Mounting Medium with DAPI. Slides were imaged on a Nikon Eclipse TE2000-U microscope using a ×20/0.75 objective and a ×40/0.95 dry objective (Nikon Plan Apo). Images were captured using an Evolution QEI monochrome camera (Media Cybernetics) and NIS Elements BR version 3.2 software. Color was added to images, and blue and red channels were merged through the ImageJ program. To quantify the fluorescent red amplification dots/cell, images were made binary under an RGB threshold, and “Particle Analysis” was utilized through ImageJ.

**Statistical analysis**

Results were analyzed using GraphPad Prism version 6 software. A one-way analysis of variance (ANOVA) with Tukey’s multiple-comparison test or Student’s t test was used to identify the differences. The threshold of significance for all tests was set as p < 0.05.

**Author contributions**—C. M. and M. P. designed the experiments. C. M. performed most of the experiments. P. C. B. and E. Y. designed and performed FACS analyses. H. T. provided technical expertise in conducting FRAP assays and data interpretation and assisted in manuscript revision. C. M., P. C. B., and M. P. analyzed the data. C. M. and M. P. wrote and revised the manuscript.

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