Abstract. The pathogenesis of medulloblastoma (MB), the most common and aggressive brain tumor in children, is poorly understood. MB tumors respond to factors secreted by cerebellar Purkinje neurons, such as Sonic hedgehog (Shh) and Wingless-type MMTV integration site family member 3a (Wnt3a). Understanding the modulation of Shh/Wnt signaling is critical for the development of new MB treatments. Shh and Wnt3a induce MB cell proliferation and bind heparan sulfate glycosaminoglycan chains (HS-GaGs). HS-GaGs are components of syndecans, cell surface HS proteoglycans (HSPGs) which act as co-receptors for extracellular matrix-based ligands, and are targets of heparanase (hpSE). We hypothesized that extracellular hpSE activity modulates MB intracellular signaling of Shh/Wnt3a, involving syndecan-1 and -4 carboxy terminal-associated proteins and downstream targets. We compared the regulation of Shh/Wnt3a signaling subsequent to treatment with exogenous human active hpSE in MB lines possessing increased invasive abilities. We identified guanine nucleotide exchange factor-H1 (GEF-H1), a small GTPase guanine nucleotide exchange factor, as a new component of a syndecan signaling complex. Secondly, we demonstrated that HPSE modulated Shh/Wnt3-dependent expression and the intracellular distribution of GEF-H1, β-catenin and N-Myc. Thirdly, HPSE modulated Shh/Wnt3a-dependent gene expression of heparan sulfate proteoglycan (HSPG) and Gli transcription factors. Fourthly, pre-treatment with HPSE, alone or prior to Shh/Wnt3a exposure, altered small GTPase (Rac1/RhoA) activities differentially and promoted RhoA activation. Finally, the differential regulation of Rac1/RhoA activities by hpSE affected MB cell proliferation and invasion. Our results indicate that the hpSE/HSPG axis is implicated in critical MB cell signaling pathways with potential relevance for MB treatment.

Introduction

Medulloblastoma (MB) is the most common invasive brain tumor of childhood. It arises in the cerebellum, generally occurs during embryogenesis and is characterized by a massive expansion of cells from the external granular layer. These events are thought to originate in response to growth factor secretion from Purkinje neurons, such as Sonic hedgehog (Shh) and Wingless-type MMTV integration site family member 3a (Wnt3a) (1,2). An insidious feature of MB tumors is their metastatic propensity as evident from MB cell intravasation through the fourth ventricle into the cerebral spinal fluid, resulting in dissemination throughout the spinal cord (1,3,4). Despite advances in therapy, a significant number of patients succumb to the disease, and survivors often suffer permanent neurocognitive deficiencies (3-5).

Multiple signaling pathways have been associated with MB onset and growth, including Shh and Wnt. Shh is a secreted protein that binds to the transmembrane receptor Ptc1. Upon Shh binding, Ptc1 inhibition of Smoothened is relieved and Shh signal is transduced to the Gli family of transcription factors (6,7).

A key feature of both Shh and Wnt3a is their high-affinity binding to heparan sulfate glycosaminoglycan chains (HS-GAGs) (8). For example, treatment of cerebellar sections with bacterial heparinases abrogates Shh mitogenicity (9). HS-GAGs are components of HS proteoglycans (HSPGs), ubiquitous macro-molecules located in the extracellular matrix.

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Abbreviations: CT, carboxy terminal; DEPC, diethylpyrocarbonate; ECM, extracellular matrix; FBS, fetal bovine serum; GAG, glycosaminoglycan; GEF-H1, guanine nucleotide exchange factor-H1; GPC, glypican; GPl, glycosylphosphatidylinositol; GST, glutathione-S-transferase; GSTPD, GST pulldown; H2O, double-distilled water; HEPES, 4-(2-hydroxyethyl)-piperazine-1-ethanesulfonic acid, N-(2-hydroxyethyl)piperazine-N’-(2-ethanesulfonic acid); rhHPSE, recombinant human active heparanase (GS3); HS, heparan sulfate; HSPG, heparan sulfate proteoglycan; IPTG, isopropythiogalactoside; MALDI-TOF MS/MS, matrix-assisted laser desorption ionization time of flight tandem mass spectrometry; MB, medulloblastoma; MEM, minimal essential media; Shh, recombinant human Sonic hedgehog; Wnt3a, recombinant human Wingless-type MMTV integration site family member 3a; SDC, syndecan; SDS-PAGE, sodium dodecyl sulfate-polyacrylamide gel electrophoresis; Wnt, Wingless-type MMTV integration site

Key words: heparanase, medulloblastoma, Sonic hedgehog, Wingless-type MMTV integration site family member 3a, guanine nucleotide exchange factor-H1, GTPase
and on the cell surface, consisting of a core protein and covalently attached HS-GAGs (10-14). Cell surface HSPGs, as members of the syndecan and glypican families, regulate cross-talk between tumor and host cells by controlling the location and activity of growth factors, acting as co-receptors. Their unique functions place HSPGs at the center of cell signaling integration (15).

HSPG expression characteristics have been linked to tumor cell metastasis (11-14). Notably, HSPG are targets of heparanase (HPSE), the dominant mammalian endoglycosidase (endo-β-D-glucuronidase) whose activity has been proven to be implicated in the angiogenic, tumorigenic and metastatic abilities of tumor cells (16,17). HPSE cleaves HS-GAGs at specific intrachain sites resulting in fragments (10-20 sugar subunits) which are biologically active, i.e., able to bind growth and angiogenic factors. Although Shh roles in MB have been demonstrated (2,18,19), mechanistic links between Shh deregulation and HPSE have yet to be investigated. Similarly, Wnt pathway activation involves the interaction of a Wnt ligand with Frizzled receptors, which leads to inhibition of the β-catenin regulatory complex. Modulation of Wnt signaling by HPSE has not been addressed, nor the mechanisms or functional links between Wnt activation and MB (20).

The invasive cell phenotype requires cytoskeletal dynamics driven by the small GTPases Rac and Rho (21,22). As tumor cells invade the surrounding tissue, they establish, abolish and/or relocate transient focal adhesions (22,23). Focal adhesion dynamics require Rac and Rho activities (21), and recent evidence supports roles of syndecan (SDC) cell surface HSPGs (24).

We hypothesized that HPSE downstream signaling events are crucial for driving MB cell proliferation and invasiveness, and that HPSE modulates HSPG-associated Shh/Wnt3a actions. To this end, we utilized human MB lines (D721 and D283) that we had previously characterized, establishing gradients of cell invasiveness and HPSE functionality (4,25). We identified GEF-H1 as a novel SDC-associated signal transduction protein in human MB cells. Of note, GEF-H1 is a Rho guanine nucleotide exchange factor with microtubule binding ability which also binds Rac1 as an inhibitor (26). We investigated GEF-H1, β-catenin and N-Myc protein expression and subcellular localization, in addition to cell surface HSPG and Gli 1-3 gene expression, by pre-treating MB cells with human active HPSE, followed by Shh or Wnt3a exposure. In this study, we demonstrate the roles of HSPG linking Shh/Wnt3a signaling to GEF-H1, Rac1/RhoA activities and MB cell proliferative and invasive events. Furthermore, we provide evidence that HPSE modifies MB cell responses to HS-binding Shh and Wnt3a.

Materials and methods

Cell culture and treatments. Human MB cell lines of low/high invasive abilities (D283 and D721) (4) were kindly provided at low-passage by Dr Darrell Bigner (Duke University, Durham, NC, USA) and cultured in improved minimal essential media (IMEM) with Richter's Zn²⁺ Option (Richter's modification), 2 mg/l L-glutamine, 2 mg/l L-proline, 50 µg/ml gentamicin sulfate (Gibco/Invitrogen, Carlsbad, CA, USA) supplemented with 20% fetal bovine serum (FBS), 100 U/ml penicillin and 100 µg/ml streptomycin (normal growth medium). The cultures were maintained at 37˚C in 5% CO₂. Due to the inherent difficulty in growing MB cells, special care was taken to grow cultures in flasks maintained in a canted position and at high cell density. HPSE treatments were for 12 h (200 ng/ml) using recombinant human active heparanase (rhHPSE) in IMEM with Richter's Zn²⁺ Option (Richter's modification), 2 mg/l L-glutamine, 2 mg/l L-proline, 50 µg/ml gentamicin sulfate supplemented with 10 mM HEPES (pH 6.5), 5% FBS, 100 U/ml penicillin and 100 µg/ml streptomycin. The cells were periodically tested for Mycoplasma negativity, and only low passage cells were used in the experiments.

Heparanase, HepIII, Shh and Wnt treatments. Preparations of highly active rhHPSE were kindly provided by Dr Israel Vlodavsky (The Bruce Rappaport Medical Faculty, Technion, Israel) (27). Heparanase activity was validated by utilizing a heparan sulfate-degrading enzyme assay kit (Takara Mirus, Madison, WI, USA). Following the mock or HPSE treatment, the cells were serum-starved for either 12 or 48 h, as indicated. The growth factor treatments were for 12 h with either recombinant human Shh (R&D Systems, Minneapolis, MN, USA) or recombinant human Wnt3a (R&D Systems) (100 ng/ml each). The removal of HS-GAGs of cell surface HSPG was accomplished by digestion with Heparitinase III (HepIII) from Flavobacterium heparinum (Sigma, St. Louis, MO, USA). The final HepIII concentration was 0.05 U/ml culture medium. The digestions were carried out for 1 h at 37˚C in 5% CO₂.

RT-PCR. Total RNA was isolated from the MB cells using the RNEasy Plus mini-kit (Qiagen, Valencia, CA, USA) according to the manufacturer's instructions. The RNA yield was determined using a Nanodrop ND1000 spectrophotometer (Nanodrop Products, Wilmington, DE, USA). To ensure a lack of genomic DNA contamination, 1 µg of total RNA was digested with DNase I (Invitrogen) prior to first-strand synthesis. Inactivated DNase I digestion (2 µl) was used as a template with a first-strand synthesis kit utilizing SuperScript II reverse transcriptase according to the manufacturer's instructions (Invitrogen). The first-strand synthesis reaction was then diluted 1:1 with H₂O to be utilized as a single-strand cDNA template. PCR amplification was performed in 20-µl reactions consisting of 1X AmpliTaq Gold buffer (Applied Biosystems, Foster City, CA, USA), 2 mM MgCl₂, 300 µM dNTP mix, 400 nM primer pair, 2 µl single strand cDNA template and 2 units AmpliTaq Gold Taq polymerase (Applied Biosystems). PCR conditions were: 94˚C, 2 min; 40 cycles of 94˚C, 20 sec; 58˚C, 15 sec; 72˚C, 42 sec; 72˚C, 30 sec. Gene accession numbers and DNA sequences for the oligonucleotide primer pairs utilized were: GAPDH, NM_002046.3, forward TTC CAC CCA TGG CAA ATT CC, reverse TGG CAG GTT TTT CTA GAC GG; HPSE, NM_006665.3, forward CTG GCA ATC TCA AGT CAA CC, reverse TCC TAA CCA GAC CCT CTT GC; SDC1, NM_001006946.1, forward TGA CTC TGA CAA CTT CTC CG, reverse TCT TCT TCA TGC GGT ACA GC; SDC2, NM_002998.3, forward AAC CAG AAA CTG AAC CTC GG, reverse TCT ACA TCC TCA TCA GCT CC; SDC3, NM_014654.3, forward AGT GAG AAC TTC GAG AGA CC, reverse TGT GTG GTC TCT TCT TCT GG; SDC4, NM_002999.2, forward TAG AAG GCC GAT
ACT TCT CC, reverse TCA CGC GTA GAA CTC ATT GG; GPC1, NM_002081.2, forward AGC CAT GAT TTT CAG GGA CC, reverse GGA TGC ATG TTT GGA AAA GC; GPC2 NM_152742.1, forward CTC GGT ATT CAG TTT TGC GG, reverse TTT CCC CAT AGA AGT CTC GC; GPC3, NM_004484.2, forward GCA AGT ATG TCT CCC TAA GG, reverse CTT GTA GCA GTG ACT TGG AAA CC; GPC4, NM_001448.2, forward CAA GCT GTT TGT CCT TCA CG, reverse AAC ATG GTG TCA CAG TCA CG; GPC5, U66033.1, forward GAC CAT TTC TTC AGG TTT GC, reverse GAA GTG CAG ATA GTC TGT GG; GPC6, NM_005708.2, forward TCC TCG TTT TTA TTT CAC CG, reverse AAG TGG TGC GAA CAA AAT GG; Gli1, NM_005269, forward ACC AAT CAG TAG CTA TGG CG, reverse TAT CAT CTT CCA AGA GTT CC; Gli2, NM_005270.4, forward ACA TCA ACA ACT CCC GAA GC, reverse TGT CCA GAT CCT CTC TGA GG; Gli3, NM_001668.5, forward TCT CCA TGA TCT CAG CAA CC, reverse AGA GTA GGT GAA GCT CAA GG. The PCR reactions were conducted in a Mastercycler ep gradient thermo-cycler (Eppendorf North America, Westbury, NY, USA).

Cloning. For affinity chromatography experiments, the pGEX-4T-3 vector was used (GE Healthcare, Piscataway, NJ, USA). RT-PCR was performed to amplify cDNA regions encoding the intracellular CT domains. Oligonucleotide primer pairs were designed for insertion into the BamHI/XhoI (New England Biolabs, Ipswich, MA, USA) restriction sites within the pGEX vector. The gene accession numbers and DNA sequences for the oligonucleotide primer pairs utilized were: SDC1, NM_001006946.1, GST SDCN1 CT forward TTT GGA TTC CGC ATG AAG AAG GAC GA; GST SDCN1 CT reverse TTT CTC GAG TCA GGC ATG GAA TTC CTC CTG; SDC4, NM_002999.2, GST SDCN4 CT forward TTT GTA TCC CGT ATG AAG GAG GAT GAA G; GST SDCN4 CT reverse TTT CTC GAG TCA CGC GTA GAA CTC ATT GG. The accuracy of these DNA sequences was confirmed by sequencing (SeqWright Inc., Houston, TX, USA).

Mass spectrometry. Protein bands excised from the SDS-PAGE were identified by matrix-assisted laser desorption ionization time of flight tandem mass spectrometry (MALDI-TOF MS/Ms) from the proteomic core facility at Baylor College of Medicine. Briefly, unique bands were excised from GSTPD and identified binding proteins using MALDI-TOF MS/Ms. Protein digests were then spotted on a MALDI target plate, and the analyses were performed using an ABI/SCIEX 4700 Proteomics Analyzer and TOF/TOF mass spectrometer. Detected mono-isotopic peptide masses were then analyzed by MS-Fit (Protein Prospector), and proteins were identified using peptide mass fingerprinting.

Cell lysates. RIPA buffer (Sigma) containing 150 mM NaCl, 1% IGEPAL CA-630 (Sigma), 0.5% sodium deoxycholate 0.1% SDS, 50 mM Tris, pH 8.0, supplemented with the protease inhibitors 1 mM PMSP (Sigma) and Complete-mini (Roche, Mannheim, Germany) was used to generate whole cell lysates. Nuclear and cytoplasmic lysates were generated using the NE-PER kit (Thermo, Waltham, MA, USA) according to the manufacturer's instructions.

**GTPase activity assays.** Rac1 and RhoA small GTPase activity assays were performed using respective GLISA kits (BK124, BK125) according to the manufacturer's instructions (Cytoskeleton, Denver, CO, USA). Briefly, after determining the cell number to cell lysis buffer volume ratio that would reproducibly result in protein concentrations of 0.5 µg/µl, the experimental treatments were performed. Cell lysates were snap-frozen in liquid nitrogen to preserve GTP-bound GTPase. Bound active Rac1 was detected with a Rac1-specific antibody. Bound active RhoA was detected using a Rac1-specific antibody. The degree of Rac1 activation was determined by comparison to corresponding values obtained from lysates derived from untreated serum-starved cells. GTP-bound RhoA GTPase activity was similarly assayed (GLISA kits; Cytoskeleton). The experiments were replicated, data were analyzed and means/standard deviations were calculated.

Western blotting. Cell lysates were quantified by standard spectrophotometric methods using either Protein Reagent (Bio-Rad, Hercules, CA, USA) or BCA Reagent (Thermo). Aliquots (20 µg of protein) of quantified cell lysates were resolved by 10-12% SDS-PAGE, transferred onto polyvinylidene difluoride (PVDF) membranes (Bio-Rad) in Tris-buffered saline (TBST) containing 5% non-fat dry milk and 0.05% Tween-20 (Sigma). Hybridizations were performed for 16 h at 4°C. The antibodies used were: SDC1 (clone B-A38, 854.070.000; Cell Sciences, Canton, MA, USA), SDC4 (ab24511; Abcam, Cambridge, MA, USA), GEF-H1 (55B6; Cell Signaling), N-Myc (clone BK124, BK125) according to the manufacturer's instructions (Cytoskeleton, Denver, CO, USA). Briefly, after determining the cell number to cell lysis buffer volume ratio that would reproducibly result in protein concentrations of 0.5 µg/µl, the experimental treatments were performed. Cell lysates were snap-frozen in liquid nitrogen to preserve GTP-bound GTPase. Bound active Rac1 was detected with a Rac1-specific antibody. Bound active RhoA GTPase activity was similarly assayed (GLISA kits; Cytoskeleton). The experiments were replicated, data were analyzed and means/standard deviations were calculated.

**Cell proliferation assays.** Cell proliferation was determined described (28). Briefly, GST-SDC1 or GST-SDC4 fusion constructs were transformed into the E. coli strain BL21 (Novagen, Madison, WI, USA). Expression was induced by the addition of IPTG to LB culture media. Induced fusion proteins from the bacteria were purified by lysing the cells using B-PER reagent (Thermo) and passing the cell lysates over columns containing glutathione beads (Thermo). GST fusion proteins were eluted from the columns with reduced glutathione. The purified GST fusion proteins (30 µg) were bound to glutathione bead columns, and 500 µg of whole cell lysates was passed over the columns. The columns were then washed three times, and the bound proteins were eluted with 2X Laemml but buffer with subsequent heating for 5 min at 95°C. Eluted proteins were detected by Coomassie blue. Bands of interest were excised from the gels and identified by mass spectrometry.

**GTP pulldowns.** This procedure was performed as previously described (28). Briefly, GST-SDC1 or GST-SDC4 fusion constructs were transformed into the E. coli strain BL21 (Novagen, Madison, WI, USA). Expression was induced by the addition of IPTG to LB culture media. Induced fusion proteins from the bacteria were purified by lysing the cells using B-PER reagent (Thermo) and passing the cell lysates over columns containing glutathione beads (Thermo). GST fusion proteins were eluted from the columns with reduced glutathione. The purified GST fusion proteins (30 µg) were bound to glutathione bead columns, and 500 µg of whole cell lysates was passed over the columns. The columns were then washed three times, and the bound proteins were eluted with 2X Laemmli buffer with subsequent heating for 5 min at 95°C. Eluted proteins were detected by Coomassie blue. Bands of interest were excised from the gels and identified by mass spectrometry.
10 µl AlamarBlue, and subsequently cultured for 4 h at 37°C. Proliferation was determined by measuring OD readings at 570 and 600 nm using a SpectraMax Plus384 spectrophotometer (Molecular Devices, Sunnyvale, CA, USA). Data were then analyzed for statistical significance. All assays were performed in triplicate.

**Invasion assays.** Transwell membrane inserts (Corning) were coated with Matrigel™ (BD Biosciences) at a 1:30 dilution in serum-free media overnight (16 h) before use. The inserts were washed with serum-free medium, and 1.5x10^5 cells/insert in IMEM + 0.1% BSA were placed on the top chambers. The bottom chambers contained IMEM medium with 20% FBS and N-formyl-Met-Leu-Phe (5 µM) as chemoattractant (Sigma). The assays were run for 12 h and then the inserts were removed, stained with 0.5% crystal violet solution (400 µl) (Sigma) for 10 min, washed and extracted with 200 µl of extraction solution (Cell Biolabs, San Diego, CA, USA) for 10 min. The OD values were read at 560 nm using 100 µl of extracted solution and statistically analyzed.

**Statistics.** Data are represented as the mean ± standard deviation. Significance values were obtained by the Student’s paired t-test (P<0.05; P<0.01). P-values of <0.01 were considered highly significant.

**Results**

**Differential cell surface HSPG and GEF-H1 expression in medulloblastoma.** In previous studies, we demonstrated that heparanase expression of specified MB cell lines correlates with their invasive phenotype (25). We considered that SDCs, particularly SDC1/4, in addition to bearing HS-GAGs as substrates for HPSE enzymatic activity, may have intracellular CT signaling capabilities that have yet to be characterized. We hypothesized that HPSE modulates downstream signaling events that are dependent upon HS/HS-binding growth factor interactions. To determine the overall cell surface HSPG expression status in these MB cell lines, semi-quantitative RT-PCR was performed and gene expression levels for members of the SDC and glypican families were determined (12). We focused on SDC1/4, considering their established roles in cell invasion, cell adhesion and cytoskeletal rearrangement (13,29). Furthermore, SDC1 has been reported to affect Wnt responsiveness in other model systems (30), and SDC4 is a fundamental component of focal adhesions (31). Unique cell surface HSPG gene expression patterns were observed for each MB cell line analyzed (Fig. 1A). Notably, SDC1/4 gene profiling from D283 and D721 cells was detected under normal growth conditions (Fig. 1A). We combined SDC1/4, considering their established roles in cell invasion, cell adhesion and cytoskeletal rearrangement (13,29). Furthermore, SDC1 has been reported to affect Wnt responsiveness in other model systems (30), and SDC4 is a fundamental component of focal adhesions (31). Unique cell surface HSPG gene expression patterns were observed for each MB cell line analyzed (Fig. 1A). Notably, SDC1/4 gene profiling from D283 and D721 cells was detected under normal growth conditions (Fig. 1A).

Next, to identify SDC intracellular CT binding signal transduction components, GST fusion proteins were constructed containing the intracellular CT domains from human SDC1/4 and performed GST pulldown assays using cell lysates from D283 and D721 MB cell lines. Proteins from the GST pulldown eluates were then separated by SDS-PAGE. A single Coomassie blue-stained protein band of 110 kDa was excised preferentially from the D283 cell lysates compared to the D721 eluted proteins (Fig. 1B). This band was excised, subjected to trypsin digestion and the resulting peptides were identified by MALDI-TOF tandem mass spectrometry. This unique band was identified as human GEF-H1 (Fig. 1C). Furthermore, SDC1/4 and GEF-H1 protein expression was visualized in
Heparanase modulates GEF-H1, β-catenin and N-Myc subcellular distribution in medulloblastoma. Recent evidence demonstrated that GEF-H1 functions to facilitate RhoA activity in the cytoplasm at sites of focal adhesion turnover, such as in regions of membrane ruffling (32). Therefore, GEF-H1 localization may influence cell migration and cytoskeletal dynamics. To characterize endogenous GEF-H1 expression and subcellular distribution in response to treatment with Shh or Wnt3a, and subsequent to pre-treatment with exogenous rhHPSE, nuclear and cytoplasmic fractions were isolated. GEF-H1 antibody detected both phosphorylated and non-phosphorylated forms of GEF-H1 in these subcellular compartments (Fig. 2A). Exposure to rhHPSE altered the GEF-H1 expression and its subcellular redistribution. The MB cell lines possessed not only a differential GEF-H1 expression and subcellular localization by rhHPSE, but also dependency upon Shh or Wnt3a treatment (Fig. 2A).

Since the expression and subcellular distribution of β-catenin is a crucial component of adherens junctions and co-transcription in Wnt signaling (33), signal transduction cross-talk between Shh and Wnt3a may be involved in MB, as demonstrated in other systems (34). Exposure to HPSE altered β-catenin expression and subcellular localization; differential β-catenin protein expression and subcellular redistribution were detected following Shh or Wnt3a treatment (Fig. 2B). D283 cells displayed primarily nuclear β-catenin, while D721 cells exhibited robust cytoplasmic β-catenin content, increasing in response to Shh/Wnt3a (Fig. 2B). To note, the β-catenin content of D721 cells was drastically reduced subsequent to pre-treatment with active HPSE (Fig. 2B).

Since a burst of N-Myc expression has been associated with proliferation of external granular layer neurons of the developing cerebellum and MB (1), human MB model system was analyzed for N-Myc expression. Treatment with exogenous active HPSE may modulate the MB cell response to Shh or Wnt3a treatments. Cytoplasmic and nuclear fractions were probed for N-Myc protein expression, and N-Myc modulation by Shh or Wnt3a was distinctly observed (Fig. 2C). No N-Myc expression was detected in D283. Only exposure with Shh or Wnt3a induced N-Myc in the D721 cells. To note, HPSE abrogated N-Myc expression in the MB cell lines (Fig. 2C). The purity of the nuclear fractions was confirmed by probing with fibrillarin (Fig. 2D).

Heparanase modulates Gli gene expression in medulloblastoma. Regulation of Gli transcription factor genes is a key downstream read-out of activation of the Shh signal transduction pathway (1). Since Gli genes are activating (Gli1) or repressing (Gli2/Gli3) (35), the expression of the Gli genes subsequent to HPSE exposure was analyzed, followed by treatment with Shh or Wnt3a. Gli from the D283 cells were expressed under control conditions and HSPE treatment reduced Gli2/3 gene expression (Fig. 3). Treatment of D283 cells with Shh also resulted in Gli2/3 gene expression (Fig. 3). However, subsequent to treatment with exogenous HPSE, as without growth factor treatment, Gli2/3 gene expression was reduced (Fig. 3). The Gli gene expression from the poorly invasive D283 cells was modulated by treatment with...
Heparanase modulates Shh and Wnt-induced medulloblastoma cell invasion. It is known that the invasive phenotype requires a dynamic cytoskeleton (38). Considering that cytoskeletal dynamics rely upon the activity of Rac1 and RhoA small GTPases (38), and that HPSE regulates Rac1 and RhoA activities, MB cells of varying invasive characteristics were pre-exposed (4) with HPSE, followed by Shh or Wnt3a treatment.

HPSE pre-treatment significantly increased the invasiveness of the D283 MB cells (P<0.05, P<0.01) in the conditions tested (Fig. 5). Conversely, highly invasive D271 cells experienced less invasiveness in response to HPSE. Treatment with Shh or Wnt3a subsequent to HPSE did not restore D271 cell invasiveness (Fig. 5). Taken together, these data indicate the differential roles of HPSE in regulating MB cell invasion in highly invasive vs. non-invasive cell lines.

Discussion

Interest concerning the roles of the HSPGs in MB is growing and has become a subject of continued investigations. In this study, we characterized the relevance of the HPSE/HSPG axis on Shh and Wnt3a signaling in a human MB cell system consisting of two lines with distinct tumorigenicities. We provide first-time evidence demonstrating i) a differential cell surface HSPG expression, notably of SDC1 and 4; ii) the identification of a guanine nucleotide exchange factor, GEF-H1, associated with SDC CT domain; iii) the importance of GEF-H1 in HSPG-modulated Shh and Wnt3a signaling affecting Rac1/RhoA activities and cell invasiveness; iv) the fact that active HPSE significantly affects Shh and Wnt3a signaling and alters the expression and localization of key downstream effectors, namely β-catenin and the Gli family of transcription factors.

Cell surface SDC1 and 4 are central in cell invasion and focal adhesion, and small GTPase activity is critical to cell motility (11,13). We provided evidence that a small GTPase regulatory protein, GEF-H1, is differentially expressed under normal (serum-containing) growth conditions among MB lines previously characterized in regards to HPSE functionality towards invasiveness (4,25). Recently, Sanz-Moreno et al indicated that the interchangeable plasticity of Rac1 vs. RhoA activity is important in tumor cell migration (38). GEF-H1 seems to be uniquely situated to participate in such interplay, since it binds and regulates both Rac1 and RhoA (39,40), and thus can be considered a key regulator of tumor cell migration.
and phenotypic plasticity. This is of relevance as it refutes the expectation that a global Rhoa preference throughout a cell is due to the presence of more GEF-H1 protein.

We addressed this issue by comparing Rhoa activities between the D283 and D721 cells. We observed a correlation between HPSE-induced GEF-H1 expression (Fig. 2A) and decreased Rac1, but increased Rhoa activity in the D283 cells.
cells, a situation one would anticipate in association with GEF-H1 modulation of Rac1 and RhoA activities. We identified increased Rac1 and RhoA activities in HPSE-pre-treated D283 cells decreasing GEF-H1 expression in response to such treatment (Fig. 2A). This indicates that D283 may rely upon an alternate (or additional) Rho GEF (Fig. 5B) (32).

Similarly, and considering that the β-catenin status in MB tumors is clinically relevant (20), we observed primarily nuclear β-catenin in our least invasive D283 cells. By contrast, the more aggressive D283 cells had increased cytoplasmic localization of β-catenin augmented without the presence of heparanase. Furthermore, the Shh/Wnt treatment increased cytoplasmic β-catenin, and exposure to active HPSE abrogated β-catenin expression and localization in these cells (Fig. 2B).

Highly proliferative D283 cells did not express nuclear N-Myc constitutively or in response to growth factor stimulation (Fig. 2C). The less proliferative D721 cells proved to be the more growth factor-inducible and this finding correlated well with N-Myc expression (Fig. 2C).

Stimulation by Shh and/or Wnt has been demonstrated in other systems to induce Gli gene expression (41). The expression pattern of the three Gli genes has been termed the ‘Gli code’ as the resulting phenotypic outcomes of active or repressive can be predicated by the combination of Gli gene expression (35). We demonstrated that the less invasive D283 cells were Gli repressive under conditions without exogenous active HPSE. Highly invasive D283 cells were Gli inert, regardless of HPSE treatment. Furthermore, we showed that when a MB cell responds to Shh or Wnt3a by Gli gene expression, subsequent exposure to exogenous active HPSE will likely result in repressing Gli gene expression. Our data also indicated that D283 MB cells demonstrate Wnt/Gli cross-talk (Fig. 4). HPSE pre-treatment inhibited Rac1 activity, but facilitated RhoA activity in poorly invasive D283 cells. Thus, the ratios of Rac1 and RhoA activity in these cells can be modulated to favor RhoA GTPase activity, in response to exogenous active HPSE, which represses the Rac1 activity while concomitantly increasing GEF-H1 protein expression. This is particularly valid when GEF-H1 is highly present in the cytoplasm (Fig. 2A). To note, the greatest increase in Rac1 GTPase activity was in the D721 cells, and this response was Shh-dependent. Although it has been shown that Shh down-regulates MB cell growth when cells are explanted into culture (42), our study is still relevant to gain insights into the cellular physiology of MB. For example, we previously demonstrated modulation of GEF-H1 induced signaling by HPSE in brain metastatic settings (43). However, it remains to be demonstrated whether or not GEF-H1 expression is consistently associated with decreased invasiveness in other tissue types. HPSE pre-treatment augmented Rac1/RhoA small GTPase activity, which is indicative of increased cytoskeletal dynamics.

Furthermore, we demonstrated that HPSE pre-treatment increased the invasiveness in a RhoA-dependent manner under conditions of increased GEF-H1 expression which augmented the invasiveness of D283. Conversely, we found that the highly invasive D283 cells did not show reduced Rac1 activity to accompany the increased RhoA activity in response to HPSE pre-treatment and decreased GEF-H1 expression while increasing RhoA activity. Therefore, an alternate Rho GEF may facilitate the increased RhoA activity in these cells. Along with reduced GEF-H1 expression associated with HPSE pre-treatment, D281 also demonstrated reduced invasiveness associated with increased RhoA activity.

In summary, we demonstrated a correlation between cell surface HSPG-induced mechanisms and MB cell invasiveness linking SDCs to GEF-H1 and Rac1/RhoA activities, the involvement in Shh/Wnt signaling pathways and their modulation by the presence of exogenous active heparanase. We also demonstrated roles for heparanase in regulating MB cell proliferation and adhesion in addition to its established heparan sulfate chain degrading activity. These findings provide impetus in further deciphering the HPSE/HSPG axis to affect gene regulation in MB and to generate synergies towards the discovery of novel markers for MB clinical intervention.

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