A BMP7 Variant Inhibits Tumor Angiogenesis
In Vitro and In Vivo through Direct Modulation of Endothelial Cell Biology

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Abstract

Bone morphogenetic proteins (BMPs), members of the TGF-β superfamily, have numerous biological activities including control of growth, differentiation, and vascular development. Using an in vitro co-culture endothelial cord formation assay, we investigated the role of a BMP7 variant (BMP7v) in VEGF, bFGF, and tumor-driven angiogenesis. BMP7v treatment led to disruption of neo-endothelial cord formation and regression of existing VEGF and bFGF cords in vitro. Using a series of tumor cell models capable of driving angiogenesis in vitro, BMP7v treatment completely blocked cord formation. Pre-treatment of endothelial cells with BMP7v significantly reduced their cord forming ability, indicating a direct effect on endothelial cell function. BMP7v activated the canonical SMAD signaling pathway in endothelial cells but targeted gene knockdown using shRNA directed against SMAD4 suggests this pathway is not required to mediate the anti-angiogenic effect. Importantly, BMP7v selectively decreased ERK and AKT activation, significantly decreased endothelial cell migration and down-regulated expression of critical RTKs involved in VEGF and FGF angiogenic signaling, VEGFR2 and FGFR1 respectively. In an in vivo angiogenic plug assay that serves as a measurement of angiogenesis, BMP7v significantly decreased hemoglobin content indicating inhibition of neoangiogenesis. In addition, BMP7v significantly decreased angiogenesis in glioblastoma stem-like cell (GSLC) Matrigel plugs and significantly impaired in vivo growth of a GSLC xenograft with a concomitant reduction in microvessel density. These data support BMP7v as a potent anti-angiogenic molecule that is effective in the context of tumor angiogenesis.
Introduction

Bone morphogenetic proteins (BMPs) belong to the larger TGF-β family of extracellular ligands and are involved in a multitude of developmental and homeostatic processes [1]. BMPs bind to BMP type I and type II STK membrane receptors to transduce signals through SMADs 1/5/8 and SMAD 4, along with non-SMAD signaling pathways [2,3]. Recently the BMP family members have been implicated in tumor progression, most notably acting as suppressors of tumor-initiating precursor cells (TICs), also referred to as cancer stem cells (CSCs) [4–6]. In a wide range of tumor histologies, CSCs may give rise to the proliferating tumor mass as well as metastatic cells [7–9], with the importance of these cells in tumorigenesis further expanded to include their pivotal role in angiogenesis and vascular mimicry [10]. There have been a number of additional reports linking BMP family members and their cognate receptors with a role in angiogenesis [11]. BMP9 and BMP10 are high affinity ligands for the type I receptor ALK1, specifically expressed on endothelial cells and possibly playing a role in endothelial cell dynamics [12]. BMP4 induced human umbilical vein endothelial cell sprouting, migration and tube formation [13,14] along with paracrine effects on tumor vasculature in malignant melanoma [15]. These observations as well our previous results showing that a BMP7 variant (BMP7v) was not only able to reduce glioblastoma brain invasion but also associated angiogenesis [16] led us to further investigate the mechanism of this angiogenic inhibition.

We used an in vitro endothelial cord formation assay, a surrogate of angiogenesis, to investigate the role of BMP7v signaling in VEGF, bFGF, and tumor-driven angiogenesis [17,18]. To our knowledge BMP7v is the only systemically available member of this family reported to be efficacious in preclinical cancer models, having been engineered to address poor PK issues that plague the wild type protein [16], and therefore represents a significant step forward in its potential use in treating cancer. Unlike wild type BMP7, BMP7v was able to avoid inhibition by circulating endogenous inhibitors such as noggin, chordin and chordin-like 2 via reduced binding. BMP7v treatment led to disruption of neo-angiogenic cord formation in vitro (and a reduction in hemoglobin content in vivo) and regression of existing VEGF and bFGF established cords in vitro. Using a series of tumor cell models capable of secreting soluble pro-angiogenic factors and therefore driving angiogenesis in vitro, BMP7v treatment completely blocked cord formation. In addition, pre-treatment of endothelial cells with BMP7v led to a significant reduction in their cord forming ability, indicating a direct effect on endothelial cell function. BMP7v activated the canonical SMAD signaling pathway in endothelial cells but, surprisingly, SMAD signaling was not required to mediate the anti-angiogenic effect. BMP7v decreased endothelial cell migration and down regulated expression of the RTKs involved in VEGF and FGF angiogenic signaling, VEGFR2 and FGFR1 respectively, while selectively reducing ERK and AKT activation. As previously reported [16], BMP7v was able to significantly delay in vivo GSLC xenograft growth with a concomitant and significant reduction in CD31 positive endothelial cells and microvessel density (MVD). Consequently, BMP7v directly modulates key signaling components within angiogenic pathways to elicit an anti-angiogenic phenotype and thereby likely contributing to its anti-tumor effects [16].

Materials and Methods

Cell culture and BMPs

Tumor cell lines (U-87-MG, PC-3, 786–0, MDA-MB-231, SK-OV-3) were obtained from ATCC (Manassas, VA) in May 2004, July 2005, April 2010, February 2005, September 2003 respectively and grown according to ATCC guidelines, A-2780 were obtained from NCI in March 2005, LXFA-629 NSCL adenocarcinoma cells were obtained from Oncotest (Freiburg,
Germany) in July 2011, and were maintained in RPMI 1640 medium (Invitrogen, Grand Haven, NY) supplemented with 10% heat-inactivated FBS (Invitrogen) and 1% glutamine (Invitrogen). Tumor cells above were stored within a central cell bank that performs cell line characterizations. GSCLCs were isolated from primary tumors between December 2003 and November 2005 and grown as neurospheres as previously described (19). GSCLCs were characterized by flow cytometry and immunohistochemistry as previously described (19).

Characterization of the banked cell lines was done last between December 2012 and August 2013 by a third-party vendor (RADIL, PCR profiling) for contamination of bacteria and viruses along with verification to be of human origin without mammalian interspecies contamination. The alleles for 9 different genetic markers were used to determine that the banked cells from ATCC and NCI matched the genetic profile that has been previously reported. Endothelial cord forming cells (ECFCs; Lonza, Walkersville, MD, January 2008) and adipose derived stem cells (ADSCs, Zen-Bio, Research Triangle Park, NC, March 2010) were maintained in EGM-2 MV media (Lonza) containing an additional 5% embryonic stem cell FBS (Invitrogen) for ECFCs. ADSCs and ECFCs were tested for their cord forming ability and pathogen tested as described above but no further method of authentication was done. ECFCs were grown in collagen-coated flasks; all other cells used tissue culture treated flasks in a humidified atmosphere at 37°C and 5% CO2. All cells were passaged for fewer than 4 months (ECFCs less than 7 passages, ADSCs less than 4 passages) before new cultures were initiated from frozen cells. BMP4, BMP9, and BMP10 were from R&D Systems (Minneapolis, MN); WT BMP7 and a modified version, BMP7v were manufactured as previously described [16].

**In vitro Cord Formation Assay**

ADSCs (Zen-Bio) were plated at 75,000 cells/well into 96-well HTS Transwell receiver plates (tumor-driven; Corning, Tewksbury, MA) or 50,000 cells/well (growth factor-driven) into 96-well black poly-D lysine coated plates (BD Biosciences, San Jose, CA), and tumor cells were plated at 25,000 cells/well in 96-well HTS Transwell plates (Corning) in co-culture MCDB-131 media (Invitrogen) supplemented with L-ascorbic acid 2-phosphate, dexamethasone, tobramycin, insulin (Sigma-Aldrich, St. Louis, MO), and cell prime r-transferrin AF (Millipore, Billerica, MA) for 24 hours. ADSC media was removed and 6,000 (tumor-driven) or 5,000 (growth factor-driven) ECFCs per well were over-seeded. Treatment with PBS or 2 nM BMP7v occurred 4 hours following ECFC plating, for 96 hours. Where applicable, 10 ng/ml VEGF or bFGF (Invitrogen) were added simultaneously with PBS or BMP7v. Cells were directly fixed for 10 min with 3.7% formaldehyde (Sigma Aldrich) followed by ice-cold 70% ethanol for 30 min at 25°C. Cells were rinsed once with PBS, blocked for 30 min with 1% BSA, and immunostained for 1 hour with antiserum directed against CD31 (R&D Systems, Cat# AF806, sheep polyclonal) diluted to 1 μg/ml in 1% BSA. Cells were washed 3 times with PBS and incubated for 1 hour with 5 μg/ml donkey α-sheep-Alexa-488 (Molecular Probes, Grand Island, NY), α-Smooth Muscle Actin Cy3 conjugate (1:200, Sigma-Aldrich, Cat# C6198, mouse monoclonal), and 200 ng/ml Hoechst 33342 (Molecular Probes) in 1% BSA, washed with PBS, then imaged using the cord formation algorithm on the Cellomics ArrayScan VTI at a magnification of 5X for neo-cord formation or 4X for established cords (Thermo Fisher Scientific, Pittsburgh, PA). For the established cord assay, ADSCs/ECFCs were treated with 10 ng/ml growth factor (VEGF or bFGF) for 96 hours prior to addition of PBS or 100 ng/ml BMP7v.

**Endothelial tube formation assay**

Human umbilical vein endothelial cells (HUVECs) (Lonza) were grown in EGM-2 MV media. HUVECs, pretreated with BMP4 (2 nM) and BMP7v (100 ng/ml) were plated (70,000 cells/
well) in Matrigel-coated 24 well plates (BD Biosciences), and incubated for 5 hours at 37°C. Tube formation was evaluated by phase-contrast microscopy, and cord length was measured using Image J analysis software (NIH).

**Immunofluorescence**

Cells were immunostained for CD31 (R&D Systems, Cat# AF806, sheep polyclonal), Ki67 (Millipore, Cat# AB9260, rabbit polyclonal), VEGFR2 (Cell Signaling Technology, Danvers, MA, Cat# 2479, rabbit monoclonal), FGFR1 (Cell Signaling Technology, Cat# 9740, rabbit monoclonal), or the In Situ Cell Death Detection Kit (Roche, Indianapolis, IN) per manufacturer’s recommendations, and 5 μg/ml goat α-rabbit-Alexa-647 (Invitrogen), or 5 μg/ml donkey α-sheep-Alexa-488 (Invitrogen) and 200 ng/ml Hoechst 33342, then imaged using the Target Activation algorithm on the VTI at an image magnification of 20X.

**Quantitative PCR analysis**

HUVECs or ECFCs (8000 cells/well) were plated in collagen I coated 96-well plates. Medium was removed 16 hours later, and BMP or vehicle was added in media with 100 μg/ml BSA. RNA was harvested 24 hours later via Qiagen RNeasy kit including the DNase treatment step per manufacturer’s instructions. RNA (500 ng) was converted to cDNA using Life Technologies cDNA archive kit per manufacturer’s instructions. Quantitative PCR analysis was performed with the following Applied Biosystems (Grand Island, NY) Taqman primer/probes: c-kit Hs00174029_m1, FGFR1 Hs00915142_m1, VEGFR2 Hs00911710, Kit ligand Hs00295067_s1, MCP-1 Hs00234140_m1, MMP-1 Hs00899658_m1, PECAM-1 Hs00169777_m1, PIGF Hs01119262_m1, β-actin 4333762T, GAPDH 4326317E, PGK1 5333765T, and PPIA 4333763T.

**ELISA analysis**

Cells were cultured as above for qPCR analysis for 72 hours. Cell culture supernatants were collected and debris was removed by centrifugation. Supernatants were applied undiluted to ELISAs (R&D Systems) per manufacturer’s instructions.

**ECFC migration**

Assays were performed as previously described [19]. Briefly, cells were plated in media containing 10% FBS on collagen I coated Oris Cell Migration plates containing seeding stoppers. Stoppers were removed 16 hours later, and treatments were added in Angiokit Optimised Growth Medium (Cell Systems Biology, Ontario). Medium was removed 24 hours later, and cells were fixed with Prefer (Anatech, Battle Creek, MI) for 30 minutes at RT, permeabilized with PBS containing 0.1% Triton X-100 for 15 minutes, washed twice with PBS and incubated with 50 μl of 15 μM propidium iodide (Sigma-Aldrich). Images were captured on the Acumen Explorer with a defined migration area.

**Western blot analysis**

SDS (1%) whole-cell protein extracts were created, briefly sonicated, and protein was quantified using the Bradford method. Protein (25 μg) was subjected to electrophoresis on 4–20% pre-cast Tris-glycine gradient gels (Invitrogen), transferred to nitrocellulose (Invitrogen), blocked with 5% blotting grade blocker (Biorad, Hercules, CA) in Tris-buffered saline containing 0.1% tween (TBST), probed with primary antiserum, washed with TBST, and incubated with HRP-labelled secondary antibody. Membranes were washed with TBST, and signal was detected by ECL (Thermo Fisher Scientific). Antiserum directed against VEGFR2 (Cat# 2479,
rabbit monoclonal), FGFR1 (Cat# 9740, rabbit monoclonal), pERK1/2 (Thr202/Tyr204) (Cat# 4370, rabbit monoclonal), total ERK1/2 (Cat# 4695S, rabbit monoclonal), pSAPK/JNK (Thr183/Tyr185) (Cat# 9251, rabbit polyclonal), pSMAD1 (Ser463/465)/pSMAD5 (Ser463/465)/pSMAD8 (Ser426/428) (Cat# 9511, rabbit polyclonal), pCRAF (Ser338) (Cat#9427, rabbit monoclonal), pMEK1/2 (Ser217/221) (Cat# 9154, rabbit monoclonal), total SMAD4 (Cat# 9515, rabbit polyclonal) (all from Cell Signaling Technology), phospho-p38 MAPK (Thr180/Tyr182) (Epitomics, Burlingame, CA, Cat# 1229–1, rabbit monoclonal), and β-actin (Santa-Cruz Biotechnology, Santa Cruz, CA, Cat# sc-47778, mouse monoclonal) were diluted (1:1000) with 5% blotting grade blocker in TBST. Densitometry was performed using Image J analysis software.

**shRNA-mediated knockdown of SMAD4 in ECFCs**

ECFCs were transduced (MOI 9) with individual clone (non-target control, SCH202V) or equivalent amounts of five pooled clones of MISSION shRNA lentiviral SMAD4 transduction particles (TRCN0000010321, TRCN0000010322, TRCN0000010323, TRCN0000040028, TRCN0000040030) in growth media containing 8 μg/ml protamine sulfate for 72 hours prior to selection with 2.5 μg/ml puromycin (all from Sigma-Aldrich).

**In vivo Matrigel Plug Angiogenesis Assay**

ADSCs (2 x 10^6) and ECFCs (0.5 x 10^6) were mixed on ice with 200 μl growth factor reduced Matrigel (BD Biosciences) and injected into the flanks of athymic nude female mice (Harlan, Indianapolis, IN). Cells in Matrigel were mixed with 100 ng BMP7v or 100 ng BMP4, or mice were dosed twice daily by intraperitoneal injection of BMP4, BMP7, or twice daily by oral gavage with sunitinib (25 mg/kg) 4 hours prior to cell implantation. Five days post implant, implants were removed, flash frozen, and hemoglobin was quantified using the QuantiChrom Hemoglobin Assay Kit (Bioassay, Hayward, CA) as previously described [17].

**GSLC Xenografts**

Four-week-old male nude mice (HDS-athymic nude mice, Charles Rives, Milan, Italy) were used for the *in vivo* model of angiogenesis. The experiments with animals were approved by the Ethical Committee of the Catholic University School of Medicine, Rome. Green fluorescent protein (GFP) expressing GSLC1 and GSLC28 cells were harvested, washed twice, and resuspended in cold PBS at the concentration of 2 x 10^5 cells per ml. Cells in 100 μl of PBS were mixed with 100 μl of Matrigel (BD Biosciences) on ice, and the mixture was implanted by subcutaneous injection. Treatment with BMP7v (1 μg per day) [16] was initiated one week after grafting. BMP7v (1 μg in 100 μl of saline) was injected intraperitoneally once daily for 3 weeks (5 days per week; total 15 injections). Control mice were intraperitoneally injected with an equal volume of saline.

Implants were removed 4 weeks post-grafting, stored in 30% sucrose buffer overnight at 4°C, and cryotome sectioned at 20 μm. IHC was performed as previously described [16]. Immunodetection was performed using the avidin biotin complex peroxidase method (LSAB Dako, Golstrup) using freshly made dianinobenzidine as chromogen. Alternate sections were stained with H&E for morphological analysis. Immunofluorescence was performed at 4°C on free floating sections rinsed with PBS. All primary antibody solutions were prepared in PBS containing 0.3% Triton X-100, and incubated with the sections overnight. Sections were incubated using antisera directed against CD31 (1:50, Santa Cruz Biotechnology, Santa Cruz, CA, Cat sc-1506, goat polyclonal) or α-Smooth Muscle Actin (SMA; 1:200, Novus Biologicals, Cambridge, UK, Cat sc-5060-531, rabbit polyclonal). After 3 PBS washes, sections were
incubated for 1 hour at RT with Cy-3 conjugated anti-rabbit secondary antibody (1:400 Jackson Immuno Research Laboratories, West Grove, PA). Sections were mounted on slides with Vectashield mounting medium (Vector, Burlingame, CA). Images were obtained with a Zeiss LSM 510 META confocal laser scanning microscope (Zeiss, Milan). For quantitative analyses of angiogenesis, 10 separate fields were randomly selected/tissue section both in the periphery and in the center of the implants. “Periphery” = area lying between the subcutaneous fat and the outlying GFP-expressing GSLCs. “Center” = the area containing GFP-expressing GSLCs. Representative images were acquired using confocal microscopy and evaluated using the Zeiss AxioVision 4.4 dedicated software (Zeiss, Oberkochen, Germany).

GSLC tumor efficacy study

Athymic nude mice (4–6 weeks; Charles River) were implanted subcutaneously with 5 x 10^5 GSLC1 cells as described above [10,20]. Mice were kept under pathogen-free conditions in positive-pressure cabinets (Tecniplast Gazzada, Varese). Tumor diameter was measured using calipers and calculated as the mean value between the shortest and the longest diameters. Treatment with BMP7v (1 and 0.1 μg) or an equal volume of saline (controls) was initiated when the xenografts reached 9–13 mm in mean diameter and was administered by intraperitoneal injection every other day for 3 weekly cycles. Mice were sacrificed three weeks post-last dose, tumors excised; formalin fixed, paraffin-embedded sections (4 μm thick) were mounted on positive-charged glass slides. For antigen retrieval, deparaffinised and rehydrated sections were treated with citric acid buffer (pH 6.0) or with EDTA (pH 8.0), 2 x 5 minutes at 750 W, followed by inhibition of endogenous peroxidase with 3% H2O2 for 5 minutes. Sections were incubated with primary antibody and visualized using the avidin–biotin–peroxidase complex per manufacturer’s instructions. 3,3’-Diaminobenzidine was used as the enzyme substrate to observe specific antibody localization and Mayer hematoxylin was the nuclear counterstain. Samples were stained at least twice with high reproducibility. The histological pattern of tumor xenografts was assessed on H&E stained sections. The proliferative potential was assessed by Ki67 immunostaining (Dako, M7240, clone MIB-1, mouse monoclonal). Mitotic index (MI) was determined as the percentage of positive cell nuclei relative to the total number of cells in 20 high power fields. The expression of nestin, GFAP, and βIII tubulin, was assessed with anti-nestin (Santa Cruz Biotechnology, Cat# sc-23927, clone 10c2, mouse monoclonal), anti-GFAP (Millipore, Cat# AB5804, rabbit polyclonal), and anti-βIII tubulin (Millipore, Cat# CBL412X, clone TU-20, mouse monoclonal) antibodies. The expression of the endothelial phenotype was determined by CD31 (1:50, Santa Cruz Biotechnology, Santa Cruz, CA, Cat# sc-1506, goat polyclonal) immunostaining.

For CD31 and MVD quantitation, 5 μm thick paraffin-embedded sections were de-waxed, hydrated and heated with antigen retrieval. Sections were then treated with 3% H2O2 for 10 min, incubated with 10% human serum for 20 min and subsequently exposed to antibodies against mouse/human CD31 (Novocastra, Milan, NCL-CD31-1A10, mouse monoclonal) and human CD31 (Dako, clone JC70A, mouse monoclonal) or isotype-matched controls at appropriate dilutions using Antibody Diluent (Dako, S3022), at 4°C overnight. After washing, biotinylated secondary antibody against mouse immunoglobulin was applied for 30 min at RT (Dako, LSAB2 System–HRP, K0675). The slides were rinsed and incubated with streptavidin–biotin-peroxidase for 30 min. All sections were counterstained using AEC substrate and counter-stained with hematoxylin. MVD was determined by the number of microvessels positive for CD31 from images randomly selected from the tumor stroma. Positive vessels were counted from 3 fields and the average count/field was reported.
Statistical analysis
Statistical significance was assessed by a two-tailed Student t test with equal variance compared to data obtained for PBS controls (in vitro) or control groups (in vivo). Statistical significance was assigned to p values < 0.05.

Results
BMP7v reduced endothelial cord formation
BMP7v was engineered to address poor PK issues that plague the wild type protein [16], and therefore represents a significant step forward in its potential use in treating cancer. Unlike wild type BMP7, BMP7v was able to avoid inhibition by circulating endogenous inhibitors such as noggin, chordin and chordin-like 2 via reduced binding (S1 Fig), and for all of these reasons we chose to continue to evaluate this molecule preclinically. Previous results indicated an anti-angiogenic activity of BMP7v in intracerebral GSLC xenografts, possibly contributing to its anti-tumor activity [16]. Therefore, we further characterized the anti-angiogenic activity of BMP7v. An in vitro co-culture system containing a feeder layer of ADSCs, which are similar to mesenchymal stem cells, and ECFCs, a subtype of umbilical cord blood-derived endothelial cells which can form vascular networks, was used to analyze BMP effects on cord formation [17,18]. Cords were visualized by immunostaining for the endothelial cell-specific marker “cluster of differentiation 31” (CD31) for ECFCs, and α-smooth muscle actin (SMA) for ADSCs. ADSCs migrate near and around developing cords and differentiate into pericyte-like cells similar to what is seen in normally developing vasculature [21]. BMP7, BMP7v, BMP9, and BMP10 significantly (p < 0.001) reduced VEGF and bFGF-driven neo-angiogenesis (Fig 1A), while BMP4 only slightly enhanced cord formation compared to the PBS treated control (Fig 1A). BMP4, BMP7, BMP7v, BMP9, and BMP10 enhanced SMA staining, providing further support for their role in differentiation (with both BMP9 and BMP10 eliciting a profound response) (Fig 1A).

The anti-angiogenic effects of BMP7v observed in vitro prompted analysis of BMP7v effects on a neoangiogenesis Matrigel plug model that consists of the same ADSC/ECFC co-culture but can form functional blood vessels following co-implantation into the flank of a nude mouse [18,22]. Five days after implantation, the cells formed extensive networks of blood vessels whose functionality was assessed by measuring hemoglobin content. Both BMP7v (100 ng) and sunitinib (25 mg/kg) caused a significant reduction (p < 0.05) in hemoglobin content (Fig 1D).

BMP7v acts directly on endothelial cells to prevent VEGF and FGF-driven neo cord formation and regress established cords
The ECFC/ADSC co-culture is a complex system requiring interactions of different cell types to form cords and the effects of the BMPs may be due to indirect effects mediated by the ADSCs. ADSCs and ECFCs were therefore individually assayed for response to BMPs using a simplified in vitro assay of culturing endothelial cells on basement membrane extract to form tubes [23]. HUVECs pre-treated for 48 or 72 hours with BMP7v prior to plating on Matrigel demonstrated a significant reduction (p < 0.001) in tube formation compared to PBS controls (Fig 1B) whereas BMP4 pre-treatment was largely ineffective. Additionally, mono-culture of HUVECs and ECFCs on tissue culture treated plastic were pre-treated with BMP7v and assayed for various angiogenic markers via qPCR and ELISA (S1 Table; all genes p < 0.05). Many pro-angiogenic genes were down regulated at the mRNA and protein level further suggesting a direct effect of BMP7v on ECFCs.
To investigate the effects of BMP7v on ECFCs alone in the co-culture system, ECFCs were pre-treated with BMP7v prior to cord formation analysis. ECFCs pre-treated with BMP7v followed by removal of BMP7v prior to cord formation resulted in a significant reduction in growth factor-driven cord formation \((p<0.001, \text{Fig 1C})\). Pre-treatment of ECFCs along with subsequent addition of BMP7v led to the greatest reduction in cord formation (Fig 1C).
Interestingly, cords that were established with growth factor prior to BMP7v treatment showed statistically significant regression ($p < 0.01$, Fig 2) suggesting BMP7v can also disrupt formed vessels. VEGF-driven established cords were less sensitive to BMP7v than FGF-driven established cords, which responded with a complete breakdown in cord structure while maintaining nidogen immunostaining, a well-defined marker for basement membrane (Fig 2A and 2B) [24]. These data suggest that BMP7v has a direct effect on ECFCs and their angiogenic function, and can uniquely inhibit establishment of newly formed cords, as well as regression of firmly established cords.

Fig 2. BMP7v reduced VEGF and bFGF established cords. (A) The ADSC/ECFC co-culture was unstimulated (basal) or stimulated with 10 ng/ml VEGF or bFGF for 96 hours prior to treatment with PBS or 100 ng/ml BMP7v for 72 hours and immunohistochemistry for CD31 (green), α-smooth muscle actin (red), and Hoechst 33342 to stain all nuclei (blue). Representative images (4X magnification) from three independent experiments are shown, and graphs represent mean connected tube area ± SEM after normalization to growth factor-induced cord values; asterisks denote statistically significant (***, $p<0.01$) differences compared to PBS controls. (B) The ADSC/ECFC co-culture was unstimulated (basal) or stimulated with 10 ng/ml bFGF for 96 hours prior to treatment with PBS or 100 ng/ml BMP7v for 72 hours and immunohistochemistry for CD31 (green), nidogen or collagen IV (red) and Hoechst 33342 to stain all nuclei (blue). Representative images (5X magnification) are shown.

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BMP7v does not alter proliferation or death of ECFCs but limits migration

An obvious potential anti-angiogenic mechanism of BMP7v involves regulation of ECFC growth, viability and/or migration. As determined through trypan blue exclusion, cell viability of the co-culture was not altered by BMP7v (data not shown) but migration of ECFCs alone was significantly, albeit weakly reduced \((p<0.05, \text{S2A Fig})\). ECFC proliferation (Ki67) and apoptosis (TUNEL) was not significantly affected by BMP7v (S2B Fig).

BMP7v reduced tumor cell-driven endothelial cord formation

To more closely represent tumor angiogenesis, BMP7v effects on tumor cell-driven cord formation were analysed. Tumor cells from a range of histologies releasing soluble proangiogenic factors were grown in a permeable transwell system with a co-culture of ECFCs and ADSCs, followed by analysis of cord formation. BMP7v significantly reduced \((p<0.001)\) cord formation induced by all tumor cell histologies tested, including GSLCs, suggesting a broad anti-angiogenic function of BMP7v \textit{in vitro} (Fig 3).

BMP7v activated SMAD signaling, down regulated MAPK family and AKT signaling, and reduced VEGFR2 and FGFR1 expression in endothelial cells

BMP7v activated the canonical SMAD signaling pathway in ECFCs and ADSCs as determined by SMAD phosphorylation (Fig 4A), which indicates functional BMP7 receptor expression. BMPs can also activate MAPK pathways; therefore, phosphorylation of p38 MAPK, ERK, and JNK was analysed. In ECFCs grown in ADSC conditioned media (since we could not directly assess this cell population in the context of the cord formation assay), BMP7v strongly reduced the level of pERK (Fig 4B). In addition, pAKT was significantly reduced (S3 Fig). The reduction was not a result of global signaling shutdown as p38 MAPK and SAPK/JNK signaling was largely unaffected (S3 Fig).

Therefore BMP7v was able to activate the canonical SMAD activation pathway while simultaneously and selectively down regulating signals emanating from growth (ERK) and survival (AKT) pathways. BMP7v strongly reduced protein expression of VEGFR2 and FGFR1 in ECFCs alone (Fig 4C) and in ECFCs (>80% reduction in protein expression) within the cord formation assay (Fig 4C and 4D). Previously, we reported on the shRNA knockdown of either VEGFR2 or FGFR1 in ECFCs leading to complete lack of response to either VEGF or FGF (18), respectively, and it is likely that BMP7v anti-angiogenic activity is at least partially a result of a SMAD-independent reduction in VEGFR2 and FGFR1 expression.

In order to determine if the anti-angiogenic effect of BMP7v is mediated through canonical SMAD signaling, stable shRNA knockdown in ECFCs of SMAD4, the obligatory nuclear shuttle protein for activated pSMAD1, 5, 8, was analysed for effects on cord formation. Surprisingly, shRNA knockdown of SMAD4 (>80% reduction in protein expression; Fig 5A) in ECFCs had little effect on VEGF and bFGF driven cord formation with or without BMP7v compared to shRNA knockdown of a non-targeting control (Fig 5B).

BMP7v reduced angiogenesis in GSLC xenografts

Subcutaneous grafting of GSLCs as Matrigel implants in immunosuppressed mice provides a well-suited model to study early tumor angiogenesis \textit{in vivo} [25]. Four weeks after grafting, the implant appeared as a gelatinous mass with small vessels travelling across the surface (S4A Fig). The center of the implant was populated by clusters of tumor cells which strongly expressed nestin, a neuronal stem cell marker and, to a lesser degree, GFAP, an astrocyte differentiation
Immunofluorescence of anti-CD31 revealed that the implant was crossed at its periphery by vessels arising from the host, which sprouted tiny branches towards the GFP-expressing tumor cell clusters (S4C Fig). These vessels showed a well-defined wall with endothelial cells packed in the typical cobblestone pattern with α-SMA staining cells, likely to be pericytes, attached (and denoting a relatively established vasculature as compared to the central structures). Conversely, the vascular channels that sprouted amongst the clusters of central tumor cells were

Fig 3. BMP7v reduced tumor-driven cord formation. ADSC/ECFC co-cultures with permeable transwells containing media (no cells-basal) or indicated tumor cells (GSCLC1, GSCLC28, A-2780, SK-OV-3, LXFA-629, MDA-MB-231, PC-3, U-87-MG, and 786–0) were treated with PBS or 100 ng/ml BMP7v for 96 hours prior to immunohistochemistry for CD31 (green), α-smooth muscle actin, (red), and Hoechst 33342 to stain all nuclei (blue). Representative images (5X magnification) are shown. All tumor cell lines treated with BMP7v displayed statistically significant (***, p<0.001) differences in connected tube area compared to PBS controls.

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irregular in shape and their walls often appeared discontinuous. Systemic administration of BMP7v resulted in a dose-dependent, tumor growth delay (p values <0.01 and <0.05 for 1 and 0.1 μg doses, respectively) similar to what was previously reported for orthotopic GSLC tumors [16]. Quantitative image analysis of the vessels revealed that BMP7v exerted an inhibitory effect on the vascularization of the implants (Fig 6A), which was more pronounced in the center (immature vessels) than the periphery (established vessels). In BMP7v treated implants, the number of neovessels in the center of both GSLC1 and GSLC28 implants was significantly reduced (p<0.05) compared with controls (S2 Table and Fig 6B). In the central region, both the total and relative vessel area was significantly lower (p<0.05) than controls (S2 Table and Fig 6B). Though there was not a significant difference in the number of peripheral vessels between control and

Fig 4. BMP7v reduced VEGFR2 and FGFR1 expression in endothelial cells. (A) Whole cell protein extracts were isolated following 15 or 60 minute PBS or 100 ng/ml BMP7v treatment and subjected to Western blot analysis using antiserum directed against phospho-SMAD1,5,8 (pSMAD1,5,8), phospho-ERK1/2 (pERK1/2), ERK1/2, and β-actin as a loading control. Results shown are representative of three independent experiments. (B) ECFCs were treated with PBS or 2nM BMP7v or BMP4 in ADSC conditioned defined co-culture media for the times indicated. Whole cell protein extracts were isolated following treatment and subjected to Western blot analysis using antiserum directed against phospho-CRAF (pCRAF), phospho-MEK1/2 (pMEK1/2), phospho-ERK1/2 (pERK1/2), and β-actin as a loading control. (C) Whole cell protein extracts were isolated from ECFCs following 72 hours of PBS or 100 ng/ml BMP7v treatment and subjected to Western blot analysis using antiserum directed against VEGFR2, FGFR1, and β-actin as a loading control. Graph represents mean densitometry ± SEM from three independent experiments, and asterisks denote statistically significant (***, p<0.001) differences compared to PBS controls. (D) The ADSC/ECFC co-culture was unstimulated (basal) or stimulated with 10 ng/ml VEGF or bFGF for 96 hours prior to treatment with PBS or 100 ng/ml BMP7v for 72 hours prior to immunohistochemistry for VEGFR2 (VEGF) or FGFR1 (bFGF) (red) and Hoechst 33342 to stain all nuclei (blue). Representative images (5X magnification) are shown. Upon BMP7v treatment, mean staining intensity for VEGFR2 and FGFR1 was statistically decreased (***, p<0.001) compared to PBS controls.

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BMP7v treated mice, their area did differ significantly ($p < 0.05$) due to the smaller vessel surface following BMP7v treatment (S2 Table and Fig 6B). It is worth noting that in both control and BMP7v treated mice, the vessels at the implant periphery contained α-SMA expressing cells, which were absent in the newly formed vessels of the central region. Control tumors displayed a typical histological pattern of GSLCs grown subcutaneously (S5 Fig). Prominent features included extensive areas of necrosis, arrangement of tumor cells in perinecrotic palisades, and remarkable angiogenesis. The tumor cells were rounded or polygonal in shape conferring an epithelial appearance to the tumor parenchyma. The proliferation
index (Ki67 immunostaining) was 18–20%. Nestin was expressed at high levels by virtually all tumor cells while ~30–40% of the cells expressed GFAP and βIII tubulin was weakly present in ~5–8% of the cells (S6 Fig). In xenografts treated with BMP7v (1 μg), confluence of cystic spaces and reduction in angiogenesis were the most dramatic changes. Clusters of tumor cells with elongated morphology were surrounded by fibrous tissue in many regions of the tumor. Nestin was expressed in <50% of the tumor cells with GFAP expressed throughout. There was a dramatic increase in GFAP and βIII tubulin expression in tumor cells (40–50%), most of which displayed an elongated morphology with a concomitant reduction in Ki67 expression (4–5%). To determine if BMP7v inhibits the GSLC-derived angiogenesis, we measured the human MVD in BMP7v treated or control tumors. After confirming the species specificity of
the antibodies (data not shown), BMP7v treatment clearly resulted in a statistically significant reduction in MVD following quantitation of human or both mouse and human CD31 (Fig 6D and S6 Fig).

**Discussion**

In addition to its direct tumor cell effects in preventing the growth of GSLCs, our previous work on BMP7v suggested an anti-angiogenic role for this agent [16]. Here we show that BMP7v is an anti-angiogenic molecule *in vitro* and *in vivo* and acts directly on endothelial cells to mediate this effect. BMP7v is the only reported systemically available BMP in the context of cancer therapy and, when combined with a marked reduction in binding to endogenous BMP inhibitors such as chordin, chordin-like 2 and noggin represents a potentially significant advance in the field. BMP7v treatment of ECFC and HUVEC mono cultures induced multiple changes in gene and protein expression, in particular pro-angiogenesis molecules (including VEGFR2 and PlGF). In addition, when HUVECs were pretreated with BMP7v and cultured in Matrigel, there was a large reduction in the number of endothelial tubes formed. Furthermore, cord area was decreased by BMP7v when ECFCs were pretreated prior to co-culture with ADSCs. An examination of exogenous factors revealed that BMP7v inhibited cord formation in both VEGF and bFGF stimulated co-cultures, two most prominent stimulators of neovascularization [26,27]. Additionally, BMP7v inhibited tumor-driven cord formation across a range of human tumor cell models demonstrating that BMP7v is effective in a complex “physiological system.” Our previous results indicated that BMP7v did not alter expression of secreted proangiogenic factors from tumor cells [16], and BMP7v did not alter VEGF or bFGF secretion from the co-culture (data not shown). These data indicate that BMP7v may directly alter the ability of endothelial cells to form cords rather than affect the synthesis and/or release of soluble proangiogenic factors. In total, our *in vitro* data support a direct role for BMP7v modulating endothelial cell angiogenesis.

At least some of the anti-angiogenic effects of BMP7v may be explained by decreases in VEGFR2 and FGFR1 levels in ECFCs (60% and 80%, respectively). Given the large number of gene changes observed in the mono cultures, there are likely a host of additional genes exclusively affected by BMP7v, further contributing to the anti-angiogenic effect. For example, expression of proangiogenic signaling molecules such as cKIT and KIT ligand, as well as PlGF are significantly impaired by BMP7v treatment. The BMP7v-driven anti-angiogenic effect is in contrast to current anti-VEGF therapies that block a single growth factor pathway and therefore have a more focused and defined phenotypic effect that falls short of regressing established cords [28,29]. Unlike anti-VEGF therapies, BMP7v effectively prevented neovascularization and regressed growth factor-dependent established cords *in vitro*, though the latter effect was much stronger for FGF-dependent cords than VEGF-dependent ones. This raises the intriguing possibility that BMP7v may be an effective anti-angiogenic agent in tumors that shift away from VEGF-dependent angiogenesis to a more FGF-dependent state [30].

To understand the more detailed mechanism by which BMP7v affected vessels, we investigated a number of additional BMP-dependent functions. Upon binding type I and type II receptor homodimers, BMPs activate SMADs 1, 5, and 8, which then associate with SMAD4, leading to nuclear translocation and gene expression [31]. While BMP7v potently stimulated phosphorylation of SMADs 1, 5, and 8 in ECFCs, significant knockdown of SMAD4 still resulted in inhibition of cord formation. Therefore, either SMAD signaling was not required for the anti-angiogenic effect of BMP7v, or a very small amount of SMAD4 expression was sufficient to mediate signal. It is known that BMPs can exert their actions via non-SMAD pathways [2,3]. For example, BMP7 stimulation of chemotaxis of monocytoic cells occurred in a
SMAD4-independent fashion [32]. Another pathway crucial to angiogenesis, RAS/ERK [33], was inhibited by BMP7v in growth factor stimulated ECFCs and it is reported that ERK and SMAD pathways can work together to regulate angiogenesis [33,34]. In fact there are numerous reports of a positive role for the ERK pathway in directly controlling angiogenesis, strengthening the likelihood that BMP7v down regulation of ERK activity, coupled with a significant reduction in proangiogenic receptor expression, significantly contributed to its antiangiogenic response [35,36]. Activation of the SMAD pathway and down-regulation of ERK pathways by BMP7v may therefore be critical in inducing an anti-angiogenic phenotype. Furthermore, there are additional signaling proteins that are modulated by BMPs, including p38 MAPK, JNK, STATs, and PI3K/mTOR, but these pathways appear to be largely unaffected by BMP7v treatment [37–40].

Clearly, BMP7v did not block in vitro angiogenesis via an apoptotic or antiproliferative response. While there was a slight reduction in ECFC migration upon BMP7v treatment, it is unlikely that this effect is responsible for the potent anti-angiogenic effect. Despite the lack of overt apoptotic or anti-proliferative markers, cords in vitro were clearly eliminated following BMP7v treatment, as evidenced through persistent nidogen and collagen IV immunostaining (reflecting the maintenance of a "basement membrane" [41]) in the absence of coincident CD31 immunostaining. One possibility is that BMP7v caused the ECFCs to differentiate into a cell type incapable of supporting cord formation (endoMT) [42] much like BMP4 which was shown to convert endothelial cells into stem-like cells [43]. Our attempts to demonstrate that BMP7v induced a similar differentiation event in ECFCs were unsuccessful (data not shown). An additional possibility is that BMP7v may have induced programmed necrosis in endothelial cells, a potentially important mechanism for tissue remodeling [44].

Importantly, BMP7v inhibited vessel formation in two different in vivo models; in ADSC/ECFC Matrigel plugs as measured by hemoglobin content and in a GSLC xenograft as evidenced by a reduction in neo-formed host vessel area and number. We previously reported that BMP7v in part exerted its anti-tumoral effect on GSLC tumors through an inhibition of angiogenesis [16] and our current results support this hypothesis. Interestingly, the effect of BMP7v was strongest within the tumor center, where vascularization was irregular and α-SMA-staining pericytes were not found, compared to vessels in the periphery which clearly formed more established, pericyte-associated structures [28,45]. Given that BMP7v down regulated VEGFR2 and FGFR1, and that VEGF and bFGF are key players in new vessel formation, it is not surprising that the immature, central vessels would be more sensitive to BMP7v. While BMP7v did not affect vessel number in the periphery, there was a significant change in vessel area. Therefore, BMP7v did have an effect on the more established peripheral vessels and raises the intriguing possibility that BMP7v is inducing what Jain described as "vascular normalization", as opposed to the outright regression seen in the implant center [46].

BMP7v administered at or above doses of 0.1 μg reduced the growth rate of GSLC xenografts. This effect depended on a continuous supply of BMP7v because tumor size increased upon BMP7v discontinuation. However, the apparent tumor regrowth that followed discontinuation of treatment may partly be ascribed to the development of cysts and not entirely to tumor regrowth. The development of cysts was associated with the occurrence of fibrous septa and sclerotic areas in the context of the tumor parenchyma, strongly suggesting that regression had occurred. The occurrence of regressive changes was paralleled by a reduction in the density of microvessels and by astrocytic and neuronal differentiation. Therefore, intraperitoneal administration of BMP7v inhibited the growth of subcutaneous GSLC xenografts and resulted from a reduction in cell proliferation, occurrence of regressive phenomena, inhibition of tumor angiogenesis and enhancement of neural cell differentiation.
In summary, BMP7v sets into motion changes in endothelial cell function that ultimately lead to a reduction in organized vessels and endothelial cell markers. In addition, BMP7v has been shown to play a role in promoting cancer stem cell differentiation as well as a significant reduction in GSC growth in vivo [16]. Therefore, a dual effect of BMP7v on both tumor stem cell differentiation and the invading endothelium may result in a potentially novel anti-tumor therapy.

Supporting Information

S1 Fig. Analysis of wild-type BMP7 and BMP7v binding inhibitors. Hep3B2 cells stably transfected with a hepcidin promoter luciferase construct (Hep3B2_HepPro_luc) were used for BMP inhibitor experiments. Hep3B2_HepPro_luc cells were plated at 30,000 cells per well in a tissue culture treated 96 well plate in DMEM (Hyclone) 5% FBS (Gibco) supplemented with Non-essential amino acids (Hyclone) and 200 μg/ml gentecin (Hyclone) for 24 hours. Cells were then starved in OMEM + 0.2% BSA for 5 hours. Cells were treated with a mixture of BMP and inhibitors in OMEM (Gibco) + 0.2%BSA (Gibco) for 18 hours then developed for luciferase activity utilizing Luciferase reporter Gene Assay Kit (Roche). BMPs were added at concentrations in the linear range of the assay (BMP-7s were used at 1 nM and 100 nm). Inhibitors were added at 100X molar excess of the BMP. BMP-7s were generated at Eli Lilly and Company.

S2 Fig. BMP7v reduced endothelial cell migration. (A) ECFCs were plated into Oris cell migration plates with the stopper in place. Following 24 hours, stoppers were removed and PBS or 1.6 nM BMP7v or BMP4 was added and cells were allowed to migrate for 24 hours prior to fixation and staining with propidium iodide (scratch area image represents stopper in place for entire experiment just prior to fixation). Cells were imaged on the Acumen Explorer and cell number within the scratch area (green box; blue box designates non-scratch area) was analysed. Representative images are shown, numbers represent percent decrease in cell number in the scratch area compared to the PBS control ± SEM from three independent experiments, and asterisks denote statistically significant (*, p<0.05) differences compared to PBS controls. (B) The ADSC/ECFC co-culture was stimulated with 10 ng/ml VEGF or bFGF simultaneously with PBS or 100 ng/ml BMP7v for 72 hours prior to immunohistochemistry for Ki67 or TUNEL (red) and Hoechst 33342 to stain all nuclei (blue). Representative images (5X magnification) are shown, graphs represent mean percent responders ± SEM from three independent experiments.

S3 Fig. Effect of BMP7v on various signalling pathways. ECFCs were treated with PBS or 2nM BMP7v or BMP4 in ADSC conditioned defined co-culture media for the times indicated. Whole cell protein extracts were isolated following treatment and subjected to Western blot analysis using antiserum directed against phospho-CRAF (pCRAF), phospho-MEK1/2 (pMEK1/2), phospho-p38 MAPK (p-p38), pERK1/2 (pERK1/2), phospho-SAPK/JNK (pJNK), and β-actin as a loading control. Note, for the convenience of the reader, some of the blots of Fig 4 are repeated in this figure.

S4 Fig. BMP7v reduced angiogenesis in GSLC xenografts. (A) Appearance of the Matrigel implant upon microsurgical inspection at 4 weeks post grafting. (B) Histological pattern of Matrigel implant at low (a, H&E; scale bar, 2,500 μm) and higher magnification with details of peripheral (b) and central regions of implant (c) (H&E; scale bars, 80 μm). Anti-nestin
immunostaining (d; scale bar, 30 μm). Fluorescence microscopy showing clusters of GFP positive GSLCs and their GFAP expression (GFAP red in e; scale bar, 30 μm). (C) Quantitative assessment of vascularity in Matrigel implants after anti-CD31 immunostaining (red, anti-CD-31; green, GFP; blue, DAPI) at the periphery (upper panel; scale bar, 150 μm) and center of implant (lower panel; scale bar, 80 μm).

S5 Fig. BMP7v reduced proliferation and induced differentiation of GSLCs in vivo. Three weeks after the last treatment, control tumors (CNTR) or tumors treated with BMP7v (1 μg or 0.1 μg) were assessed for morphology by H&E staining. In addition, tumor cell proliferation was analysed by Ki67, and tumor cell differentiation was assessed by nestin, GFAP, and βIII tubulin (βIII Tub) immunohistochemistry.

S6 Fig. BMP7v treatment reduced microvessel density (MVD) in tumor xenografts. (A) Immunohistochemistry was performed on vehicle (CNTR) or BMP7v treated (0.1 μg or 1 μg) tumors three weeks following treatment using a human-specific CD31 antibody. (B) Graph represents MVD assessed from both human and mouse CD31 immunostaining, and asterisks denote statistically significant (**, p<0.05; ***, p<0.001) differences compared to vehicle controls (CNTR).

S1 Table. Summary of fold-change in gene expression upon BMP7v treatment. Quantitative PCR analysis was performed, and data was normalized to the geometric mean of at least three housekeeping genes. ND, not determined. Changes in red denote p<0.05.

S2 Table. Summary of results from quantitative assessment of angiogenesis in Matrigel implants.

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Author Contributions
Conceived and designed the experiments: CMT JM RP RD SR LRV LS. Performed the experiments: CMT JM RP EV LW WB QGD LM SG LML GS AB MLC MT. Analyzed the data: CMT JM RP EV LW QGD LM SG LML GS AB MLC MT. Wrote the paper: CMT JM RP SR LS.

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