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PV1 down-regulation *via* shRNA inhibits the growth of pancreatic adenocarcinoma xenografts

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Abstract

PV1 is an endothelial-specific protein with structural roles in the formation of diaphragms in endothelial cells of normal vessels. PV1 is also highly expressed on endothelial cells of many solid tumours. On the basis of *in vitro* data, PV1 is thought to actively participate in angiogenesis. To test whether or not PV1 has a function in tumour angiogenesis and in tumour growth *in vivo*, we have treated pancreatic tumour-bearing mice by single-dose intratumoural delivery of lentiviruses encoding for two different shRNAs targeting murine PV1. We find that PV1 down-regulation by shRNAs inhibits the growth of established tumours derived from two different human pancreatic adenocarcinoma cell lines (AsPC-1 and BxPC-3). The effect observed is because of down-regulation of PV1 in the tumour endothelial cells of host origin, PV1 being specifically expressed in tumour vascular endothelial cells and not in cancer or other stromal cells. There are no differences in vascular density of tumours treated or not with PV1 shRNA, and gain and loss of function of PV1 in endothelial cells does not modify either their proliferation or migration, suggesting that tumour angiogenesis is not impaired. Together, our data argue that down-regulation of PV1 in tumour endothelial cells results in the inhibition of tumour growth *via* a mechanism different from inhibiting angiogenesis.

Keywords: angiogenesis • fenestrae • vesiculo-vacuolar organelles • caveolae • pancreatic cancer • transendothelial channels • tumour microenvironment

Introduction

The vertebrate gene *Plasmalemmal Vesicle Associated Protein* (*PLVAP*) encodes for PV1, a 60kd, single span, type II integral membrane N-glycosylated protein [1–3]. PLVAP/PV1 expression is restricted to a subset of normal microvascular endothelial cells (ECs) and, in adult organisms, PV1 is not expressed in the ECs of all large vessels with the exception of the endocardium lining the heart chambers [1, 3–5]. In the ECs of normal capillaries of selected organs, PV1 is essential for the formation of the stomatal diaphragms of caveolae

and the diaphragms of fenestrae, transendothelial channels and vesiculo-vacuolar organelles [2, 4, 6–8]. Formation of the diaphragms seems to be the only cellular function of PV1 [9].

Endothelial diaphragms are present in ECs of tumour vessels undergoing angiogenesis (reviewed in [10]) and PV1 was reported to be highly expressed in the ECs of human solid tumours, pancreatic cancer included [11–14] [15]. PV1 expression on tumour ECs was known for more than two decades, through data obtained with the PAL-E and MECA32 monoclonal antibodies (mAb), shown recently to bind to human and mouse PV1 respectively [16, 17] (reviewed in [10]). *In vitro* data suggest that PV1 plays an active role in pathological angiogenesis by facilitating human EC tube formation in Matrigel [11, 12] and EC migration induced by angiogenic growth factors such as VEGF and HGF[11–13, 15, 18]. However, the usefulness of PV1 as a novel anti-cancer therapeutic target is unclear [12, 19].

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Besides its structural role and postulated roles in cell migration and angiogenesis, PV1 may also have a role in inflammation [10, 17]. PV1 blockade in mice using anti-PV1 antibodies results in obvious impairment of recruitment of neutrophils and macrophages at inflammation sites [17]. PV1 down-regulation by siRNA in TNF α -activated ECs *in vitro* inhibits diapedesis of leucocytes without affecting their adhesion to and rolling on activated ECs under flow [17].

Pancreatic ductal adenocarcinoma (PDAC) ranks fourth among cancers as a cause of death in the United States and Europe with a median survival of 6 months [20]. Five-year survival is <5% and is limited to stage I and II patients who can benefit from pancreas resection in combination with chemotherapy and radiotherapy [21]. Late stage (III and IV) unresectable PDAC patients have access only to palliative chemotherapy, yielding a median survival rate of 6–11 months [22, 23]. So far, all PDAC therapies are short-lived and associated with significant toxicities. Thus, pancreatic cancer patients are prime candidates for the benefit of synergistic adjuvant therapies to increase efficacy and/or manage toxicity.

To evaluate whether or not PV1 plays a role in angiogenesis in vivo and the potential of PV1 as a therapeutic target in PDAC treatment, we first tested its role in tumour growth in two different xenograft models of PDAC. We show that PV1 down-regulation by a single intratumoural delivery of PV1shRNA using lentiviruses results in reduced tumour growth in these two models. Because of the sequence mismatch between human and mouse PV1, we show that this effect is clearly the result of PV1 down-regulation in tumour stroma, which is of mouse origin. Moreover, in both tumours, PV1 is expressed only in ECs of tumour vessels and not expressed in tumour or stromal cells at protein or mRNA level. Taken together, these data argue that PV1 expression in tumour ECs is required for tumour growth in vivo. In addition, we show that deletion of PV1 in tumour ECs may not result in decreased tumour angiogenesis implicating additional mechanisms in PV1-promoted tumour growth.

Materials and methods

Materials

All general reagents were from Thermo-Fisher (Pittsburgh, PA, USA), unless otherwise stated.

Antibodies

Rat anti-mouse CD31 (PECAM-1) clone MEC13.3, unlabeled or conjugated to APC, was from BioLegend (San Diego, CA, USA); Mouse anti- β -actin mAb (AC40) from Sigma-Aldrich (St. Louis, MO, USA); Biotinylated rabbit anti-mouse IgG from Vector Laboratories (Burlingame, CA, USA); HRP-conjugated goat anti-rat IgG-HRP, and goat anti-mouse IgG-HRP from Biodesign (Kennebunk, ME, USA); Goat antichicken IgG-Alexa 568 and Alexa 647, (Fab)2 fragment from Invitrogen (Eugene, OR, USA). Rat anti-mouse PV-1 mAb clone MECA-32 mAb [9] and chicken anti-human PV1C pAb [6] were as previously described.

Primary antibody labelling with fluorophores

Affinity purified primary antibodies, rat anti-mouse PV-1 mAb clone MECA-32 were labelled with Alexa 568 or 647 fluorophores (Molecular Probes; Invitrogen), as per manufacturer's instructions.

Cells and cell culture

Mouse lung endothelial cells (MLEC) isolated from lungs of wild-type mice or Cav1-knockout mice (MLEC-Cav1KO), and immortalized with polyoma middle T antigen, and flow sorted for CD31 and PV1 expression, are described before [9]. MLEC were cultured on plastic in MLEC growth medium consisting of a modified endothelial growth medium 2 (EGM2) (Lonza, Walkersville, MD, USA) formulation, from which the serum coming with the kit was omitted and replaced with heat-inactivated foetal bovine serum (FBS) (Hyclone, Logan, UT, USA) to 15% final concentration. The formulation was also supplemented with 100 µg/ml of each penicillin, streptomycin and glutamine (Invitrogen, Carlsbad, CA, USA), as described [9]. Bovine aortic (BAEC), human umbilical vein (HUVEC) and human lung microvascular (HLMVEC) endothelial cells were obtained from Lonza and were cultured in EGM2 or EGM-2MV medium, respectively. The MECA-32 hybridoma (developed by the laboratory of E. Butcher [24] at Stanford University) was obtained from the Developmental Studies Hybridoma Bank (Iowa City, IA. USA).

AsPC-1 and BxPC-3 human pancreatic cancer cell lines, obtained from ATCC (Manassas, VA, USA), were grown in RPMI (Mediatech Inc., Herndon, VA, USA), with 10% FBS (Omega Scientific, Tarzana, CA, USA), 100 U/ml penicillin and 100 μ g/ml streptomycin (complete medium).

PV1 siRNA sequences

PV1 siRNA-1, -2, -3, -4 (84094: PLVAP) were obtained from Dharmacon (Lafayette, CO, USA). The PV1 siRNA-5 duplex (ID85339; Ambion, Austin, TX, USA) [8] and Luciferase negative control siRNA [25] were previously reported.

PV1-knockdown effectiveness was evaluated in MLEC transfected with Oligofectamine (Invitrogen) in EBM2 culture medium lacking FBS and antibiotics, as per manufacturer's instructions. siRNAs were used at a final concentration of 5 nM and PV1 knockdown assessed 24, 72 and 96 h after transfection, using Western blotting with anti-PV-1 mAb MECA32. PV1-shRNA sequences are described in Table 1 and were designed as in [26].

Lentivirus production and EC infection

Oligonucleotides corresponding to the short hairpin RNA (shRNA) sequence of interest were annealed and cloned into the lentivirus vector, pLentiLox 3.7 (pll3.7) [26] (Addgene, Cambridge, MA, USA). Virus stocks were prepared

Table '	1 PV1-shRNA	sequences	used for	insertion	in lentiviruses
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Gene targeted	Strand	shRNA sequence
Luciferase	sense*	5'-T CTTACGCTGAGTACTTCGA TTCAAGAGATCGAAGTACTCAGCGTAAGTTTTTTC-3'
	antisense	5'-TCGAGAAAAAACTTACGCTGAGTACTTCGATCTCTTGAATCGAAGTACTCAGCGTAAGA-3'
PV1-1	sense	5'-T GCGAGCTGGAGGCGCGTAA TTCAAGAGATTACGCGCCTCCAGCTCGCTTTTTTC-3'
mouse 1080-1099	antisense	$5^\prime - TCGAGAAAAAAGCGAGCTGGAGGCGCGTAATCTCTTGAATTACGCGCCTCCAGCTCGCA-3^\prime$
PV1-2	sense	5'-T GAGGTGGATGTGCGCATTA TTCAAGAGATAATGCGCACATCCACCTCTTTTTC-3'
mouse 1124-1142	antisense	5'-TCGAGAAAAAAGAGGTGGATGTGCGCATTATCTCTTGAATAATGCGCACATCCACCTCA-3'
PV1-4	sense	5'-TAGGAGAAGTTCCAGGCGGATTCAAGAGATCCGCCTGGAACTTCTCCTTTTTTC-3'
mouse 690-708	antisense	5' TCGAGAAAAAAAGGAGAAGTTCCAGGCGGATCTCTTGAATCCGCCTGGAACTTCTCCTA-3'
PV1-5	sense	5'-T GGAATTCAAGAAAAGGATC TTCAAGAGAGATCCTTTTCTTGAATTCCTTTTTC-3'
mouse 1258-1276	antisense	5'-TCGAGAAAAAAGGAATTCAAGAAAAGGATCTCTCTTGAAGATCCTTTTCTTGAATTCCA-3'

*siRNA sequences used as template are in bold.

by co-transfecting pll3.7 with three packaging plasmids pMDLg/pRRE, CMV-VSVG and RSV-Rev, (Addgene) into 293T cells [25, 26].

Mouse lung endothelial cells were incubated (4 hrs) with lentiviruses encoding for PV1shRNA or controls (empty virus, shLuc) at 1 and 10 MOI, with or without 8 μ g/ml polybrene in infection medium (EBM-2), followed by incubation (24–48 hrs) with MLEC growth medium; infectivity was measured by flow cytometry using GFP expression as readout. The ability of pll3.7-based lentiviruses to infect AsPC-1 and BxPC-3 tumour cells was already demonstrated [25].

PV1-shRNA validation – immunofluorescence on cells and confocal microscopy

Mouse lung endothelial cells were infected with lentiviruses encoding for shLuc, shPV1-1 and shPV1-5 at an MOI = 0.1 and allowed to express shRNA for 48 hrs. Cells were labelled live (30 min., 10°C, EBM2 + 1% BSA) with fluorescent anti-PV1 (1.5 μ g/mI), rinsed (3×, RT) in PBS containing calcium and magnesium (PBS-CM) and chased (37°C, MLEC growth medium), rinsed (3×, RT) in PBS-CM, fixed (10 min., RT) in 4% paraformaldehyde in PBS-CM, rinsed again in PBS-CM, mounted in PermaFluor and examined by confocal fluorescence microscopy using a Zeiss 510 Meta confocal system (Carl Zeiss, Thornwood, NY, USA). Image stacks were acquired with the pinhole set at 1 Airy unit and transformed through the maximum-intensity projection function to obtain global images of the cells using ImageJ software (NIH, Bethesda, MD, USA).

Flow cytometry

Labelled cells were analysed using a FacsCalibur or a CANTO flow cytometer controlled by either CellQuest or DIVA software respectively (BD Biosciences, San Diego, CA, USA). Data analysis was carried out using FlowJo 9.4 software (Tree Star, Ashland, OR, USA). Each experiment had four replicates per time-point and was repeated three

times. Median fluorescence from at least 5000 live cells was calculated in each sample. Statistical significance was calculated using Student's *t*-test.

Generation of stable MLEC cell lines expressing PV1shRNA

Mouse lung endothelial cells were infected with a low (0.1–0.5) MOI of lentiviruses encoding for PV1shRNA or controls (empty virus, shLuc) in presence of 8 μ g/ml polybrene in infection medium, allowed to express the shRNAs for 48 hrs when the cells were sorted based on GFP expression. Transduction efficiency in these conditions was ~1–5% and purposely designed to limit the number of multiple lentiviral integrations in the genome. The cell populations obtained were labelled MLEC-shPV1-1, -2, -4, -5, MLEC-shLuc MLEC-EV (empty vector).

RNA isolation

Total RNA was isolated using RNAeasy kit (Qiagen, Valencia, CA, USA) and quality controlled using Bioanalyzer (Agilent/Stratagene, Santa Clara, CA, USA) and NanoDrop (Thermo-Fisher).

Real-time quantitative PCR

Quantitative real-time PCR was performed as before [9], using the Gene Expression Assays mouse PV1 (Mm00453379_m1), mouse Actin B (Mm00607939_s1), human PV1 (Hs00229941_m1), human Actin B (Hs03023880_g1) (Applied Biosystems, Foster City, CA, USA). Reactions were performed in triplicate, utilizing cDNA corresponding to 10 ng RNA. Data were analysed on the basis of the relative expression method with the formula $2^{-\Delta\Delta}CT$, where $\Delta\Delta C_T = \Delta C_T$ (sample) $-\Delta C_T$ (calibrator = average C_T values of all samples), and ΔC_T is the C_T of the house-keeping gene [beta-Actin] subtracted from the C_T of the target gene.

Evaluation of cell-surface PV1 levels by flow cytometry

Adherent MLEC stably expressing different shRNAs were labelled (30 min., 4°C) live with 1.5 μ g/ml MECA-32-Alexa 647 mAb in MLEC growth medium, rinsed (3 \times , RT) in PBS and non-enzymatically detached using EDTA (Cell Dissociation Solution; Sigma-Aldrich). Cells were then mixed with an equal volume of 1% BSA in PBS, and kept on ice in the dark until examined using flow cytometry.

Western blotting

Equal amounts (20 μ g/lane) of MLEC proteins were immunoblotted with MECA-32 and mouse anti- β -actin monoclonal antibody (clone AC40), as described. [9]. Signal quantization by densitometry on TIFF files was carried out using GelEval v1.35 software (FrogDance, Dundee, UK).

Pancreatic tumour xenograft model

Female athymic mice (Nu/Nu, Charles River) were injected subcutaneously into the dorsal flank area with 1 \times 10⁶ of either ASPC-1 or B×PC-3 cells. For each cell type, the mice were divided randomly into four equal groups of EIGHT mice to be left untreated or injected with shPV1-1-LV, shPV1-5-LV or shLuc-LV. Once tumours reached a volume of 50 mm³ (8–10 days after injection of the cells), they were injected with 4.10⁷ viral particles in 50 µl of OptiMem (Invitrogen). Tumour diameters were measured every 3 days. Tumour volumes were calculated as $\pi/$ 4 × width × height × length of the tumour. Experiments were terminated when the tumour diameter reached 15 mm following the procedures approved by the Dartmouth College IACUC.

Statistics on tumour growth

Data were analysed using ANOVA and Tukey HSD test for parametric data, or the Kruskall and Wallis test for non-parametric data using the Dunn-Benferroni test for multiple comparisons (VassarStats website). P < 0.05 was taken as the level of significance.

Colocalization of PV1 and CD31 in tumour samples by confocal microscopy

AsPC-1 and BxPC-3 tumours were induced as described and allowed to grow for 21 days. One hr before harvesting, mice were injected *via* the tail vein with 10 mg/kg Dark Red Fluorescent (660/680) FluoSpheres[®] in PBS (Molecular Probes/Invitrogen, Cat# F8789) to label the intravascular space. The mice were killed with CO₂, tumours harvested unfixed, embedded in optimal cutting temperature compound (OCT). Co-immunofluorescence experiments on methanol fixed 10 μ m thick sections were performed with 1.5 μ g/ml of Alexa 647-coupled MECA-32 mAb and Alexa488-coupled rat anti-CD31 MEC13.3 mAb and examined by confocal

fluorescence microscopy using a Zeiss 510 Meta confocal system. The controls consisted of non-immune rat IgG2a (clone 2A3) labelled with Alexa 647 or 488. Choice of fluorophores ensured lack of spectral overlap between the labels to virtually eliminate co-localization artefacts.

Tumour vascular area density

Tumours processed as described above were stained with a rat anti-CD31 mAb (clone MEC13.3) (BD Biosciences) using a Vectastain ABC Peroxidase kit (Vector Laboratories). Sections were examined using an Olympus BX-61 upright microscope equipped with an Olympus DP70 camera.

For quantitative morphometric analysis of CD31 staining in tumour samples, 10 random fields/slide at $20 \times$ magnification were quantified for DAB signal using Image-Pro Plus program (Version 7.0; Media Cybernetics, Silver Spring, MD, USA). Three tumours per group were analysed. Vascular area density [27, 28] was calculated as the ratio of CD31 positive pixels to the total area of the image, as described before [25].

Cell proliferation assays

To assess doubling time, cells were seeded in six well plates in triplicate at a density of 10^4 cells/well and the doubling time was calculated over 5 days. Every 24 hrs, cell numbers were determined by counting the cells using a haemocytometer. The experiments were repeated twice.

To assess cell proliferation based on DNA content, we employed an established assay [29, 30]. Briefly, MLEC were seeded at 1000 cell/well in 96-well plates (n = 8 wells per each time-point per genotype), cultured in full growth medium and collected every day for 7 days. Each day the wells were briefly rinsed in PBS, and the cells frozen until the last time-point was collected. Cells were lysed by osmotic shock with low (1×) strength SSC and then incubated with 1 mg/ml Hoechst 33258 (Enzo Life Sciences, Farmingdale, NY, USA). The signal was quantitated by spectrophotometry using a Synergy HT plate reader (BioTek, Winooski, VT, USA).

Adenovirus constructs and transduction

Adenoviral AdEasy-1 system (Agilent/Stratagene) was used to generate adenoviruses expressing human PV1-3xHA (AdPV1HA). Briefly, the sequence coding for human PV1-3xHA was PCR amplified, inserted into the multiple cloning site of the pShuttle vector and the resulting construct was recombined with pAdEasy-1 vector by co-transfection in the BJ5183 *E. coli* strain (Agilent/Stratagene). Positive clones were linearized with *Pac I* restriction enzyme and transfected into 293 cells for Adenovirus production using the CsCl₂ density gradient method or the VivaPure AdenoPACK kits (VivaScience, Sartorius, Germany).

BAECs and HUVECs were transduced (4–6 hrs, 37° C) with adenoviruses at an MOI of 10 PFUs/cell for AdGFP or 50–200 viral particles/cell for AdNull and AdPV1HA in EBM2 supplemented with 2% FBS, cells were rinsed twice in EGM2 and incubated (24 hrs, 37° C) in growth medium. The cells were routinely used 24 hrs after infection. Empty adenovirus (AdNull) or adenovirus encoding GFP (AdGFP) at the same MOI were used as controls.

Expression of PV1-HA was determined in HUVEC or BAEC transduced in 100 mm dishes with an adenoviral MOI of 1. Twenty-four hrs

after transduction, the cells were processed for Western blotting, as described above for MLEC, using 25 μ g of total protein per well and using chicken anti-human PV1C pAb.

Endothelial cell migration assays

For scratch-wound assay, cells were grown to confluence in triplicate in six well plates, the monolayer was scratched in five different places using a pipette tip and cell debris removed by rinsing the cells with medium. Images of the cells were acquired at 2, 4, 8, 12 and 20 hrs using a Leica DMIL inverted microscope equipped with a $10\times$ phase contrast objective and a QImaging 12-bit CCD camera, and analysed with ImageJ. Fifteen images per well were acquired at each time-point. Cell migration was quantified by determining the area of the wound at noted time-points at predetermined sites.

Electric cell-substrate impedance sensing (ECIS) automated cellmigration assay was performed on an ECIS system (Applied BioPhysics) equipped with a "wound healing module" and using 8-chambered goldplated electrode arrays (8W10E). Cells (5×10^4 /well) were plated on sterile 8W10E electrodes pre-coated with 0.1% porcine gelatin (collagen I), grown to full confluence and transduced with adenoviruses. Immediately after adenoviral transduction, the electrode arrays were mounted on the ECIS system within an incubator (37° C, 5% CO₂) and connected to its recorder device. The cell monolayer was wounded by electric pulses, the medium changed and the transendothelial electrical resistance, an index of endothelial cell barrier function, was measured in real time and recorded for 50 hrs in 5-min. intervals.

Results

PV1 down-regulation by lentiviral-encoded shRNA

Small interfering siRNA sequences directed against mouse PV1, previously reported [6, 8] or designed *de novo*, were transfected in MLEC and PV1 knockdown determined by immunoblotting. As all the PV1 siRNAs were able to down-regulate PV1 (data not shown), these sequences were used to design the shRNA sequence to be introduced into the pll3.7 vector for lentivirus (LV) generation. The lentiviral vector also encodes for the green fluorescent protein (GFP) for ease of identification of transduced cells. The siRNA and shRNA sequences used are shown in Table 1.

We first determined that all LVs generated were able to infect mouse ECs at low MOI, as shown by flow cytometric quantification of GFP-expressing cells (Fig. 1A) or using fluorescence confocal microscopy (Fig. 1B).

Next, the effectiveness of PV1 down-regulation in MLEC stably transduced with shRNA-LVs targeting PV1 or luciferase (shLuc-LV) was tested by multiple complementary approaches. We chose to study long-term, stable expression of shRNAs, as it is more relevant to the *in vivo* experimental approach. According to real-time PCR (not shown) and Western blotting results (Fig. 1C), total cellular levels of PV1 were decreased by more than 70% for all the

PV1-shRNA-expressing cell lines. As PV1 function occurs at the cell surface [9], we determined the effect of different shRNAs on plasma membrane levels of PV1 protein. We show using confocal microscopy (Fig. 1B) and flow cytometry (Fig. 1E) that all mouse PV1-shRNA-LVs efficiently silenced PV1 protein expression at the cell surface by more than 80% compared to non-infected or shLuc-LV-transduced MLEC. The lack of effect of shLuc control LV on PV1 protein levels is shown in Figure 1E. The efficiency of shLuc-LV control virus to decrease luciferase activity in 293 cells constitutively expressing luciferase was previously validated [25]. The two most efficient mouse shRNA-LVs, namely shPV1-1 and shPV1-5, were used in all subsequent experiments.

Intratumoural delivery of PV1-shRNA inhibits the growth of established pancreatic tumours

To determine whether the LV-mediated PV1 down-regulation is able to inhibit PDAC growth, we employed a model system consisting of heterotopic tumours derived from two different human PDAC cell lines (AsPC-1 and BxPC-3) grown subcutaneously in nude mice.

Following subcutaneous injections of AsPC-1 or BxPC-3 cells, tumour growth was monitored and mice with established tumours (volume of 50 mm³) were enrolled in four different groups, one to be left untreated and three treated with one injection either of shPV1-1-LV, of shPV1-5-LV or shLuc-LV. Tumour growth was determined over a period of several weeks. A statistically significant difference in tumour volumes between shLuc-LV-injected tumours and shPV1-LVinjected tumours was seen as early as 24 and 33 days following viral injection of AsPC-1- and BxPC-3-derived tumours respectively. The subsequent growth of all shPV1-LV-injected tumours remained significantly attenuated. By the end of the experiment, AsPC-1 tumours that were infected with shPV1-1-LV exhibited a 35% (P < 0.05) decrease in volume and 65% (P < 0.05) with shPV1-5 (Fig. 2A) in comparison with tumours infected with shLuc-LV or control tumours left untreated. No statistically significant differences were detected between the growth of tumours infected with shLuc-LV and control tumours left untreated. With respect to BxPC-3, shPV1-1 inhibited tumour growth by 57% (P < 0.01) and shPV1-5 one of 49% (P < 0.01) (Fig. 2B). Taken together, these data demonstrate that PV1 down-regulation by two different shRNAs are able to inhibit pancreatic tumour growth in two different human-PDAC xenograft models.

PV1 is expressed specifically in the EC of tumour xenografts and is not expressed in tumour cells

The sequence mismatches between mouse PV1 shRNAs and human PV1 strongly argued that tumour growth inhibition occurred *via* PV1 down-regulation in the tumour microenvironment (*i.e.* ECs, stromal cells and tumour-infiltrating leucocytes), which is of murine origin. To strengthen this observation, we defined the precise cellular type(s) expressing PV1 in the AsPC-1- and BxPC-3-derived tumours. Although PV1 is expressed in tumour ECs in many solid tumours



Fig. 1 PV1 knockdown by lentiviral shRNA. (**A**) Flow cytometry analysis demonstrating the ability of each of the generated LVs to infect mouse ECs, as demonstrated by GFP expression. The cells were incubated with a LV dose of \sim 0.1 MOI. The GFP-positive cells were used for sorting to generate stable cell lines expressing the various shRNAs. (**B**) Laser confocal microscopy demonstrating the down-regulation of PV1 signal in cells transduced with low MOI of shPV1-1-LV (top panels) or shPV1-5-LV (bottom panels). PV1 signal was detected with the anti-PV1 MECA-32 mAb directly coupled to a fluorophore (*red*). Transduced cells were identified by their expression of GFP (green). The images represent maximum-intensity projections of confocal stacks encompassing all the cell thickness, acquired with a $60 \times$ objective and no zoom. Non-transduced cells serve as an internal control for normal PV1 levels. Bars 20 µm. (**C**) Total cellular-PV1 protein levels in MLEC stably transduced with anti-PV1 and anti-actin antibodies, as described in Methods. (**D**) Quantization of PV1 level on the surface of mouse ECs transduced with LVs encoding for shLuc, shPV1-1, -2, -4 and -5. The data represent median fluorescence (n = 4, P < 0.01, error bars -stdev). (**E**) Examples of flow cytometry data showing PV1 levels on the surface of MLEC parental (*orange trace*), MLEC-shLuc (*green trace*), MLEC-shPV1-1 (*blue trace*), MLEC-shLuc (*red trace*).



including pancreatic cancer, there is no previous information regarding PV1 expression in AsPC-1- and BxPC-3-derived tumours. To address this, the vascular space of tumour-bearing mice was labelled with fluorescent microspheres, tumours harvested and sections stained with anti-PV1 antibodies directly labelled to fluorophores. We used the MECA-32 mAb that recognizes only mouse PV1 [24] [16]. and the anti-hPV1C pAb that specifically recognizes human PV1 [6]. In agreement with data published for other tumours [31–34], mouse-PV1 expression was detected solely in the pancreatic tumour ECs, as shown by its co-localization with the fluorescent microspheres as luminal markers of the functional blood vessels (Fig. 3A and B) and EC markers PECAM-1/CD31 (Fig. 3A and B), ICAM-2/CD102



Fig. 3 In both AsPC-1 and BxPc-3 pancreatic tumours PV1 is expressed only in tumour endothelial cells (**A**–**B**) Confocal microscopy of frozen sections from AsPC-1 pancreatic tumours grown in nude mice, with anti-PV1 (red) and anti-CD31 (green) and antibodies directly coupled to fluorophores. The vascular lumen (blue) was labelled by the introduction of dark red microspheres in the circulation. In the merged field, white means triple co-localization. Last panel in each shows nuclei stained with DAPI. (**A**) is a representative field showing vessels inside the tumour whereas (**B**) shows vessels at the tumour periphery. Magnification $20 \times$. Bars – 20μ m. (**C**) Immunohistochemistry of serial frozen sections from human pancreatic xenograft tumours grown in nude mice, with anti-PV1 (PV1) and anti-CD31 (CD31) antibodies. The nuclei are counter stained with haematoxylin. Bars – 20μ m. (**D**) PV1-mRNA levels in HLMVEC and human pancreatic tumour cell lines AsPC-1 and BxPC-3 measured by real-time quantitative PCR. The data were obtained from triplicate samples and normalized to β -actin levels ($\Delta\Delta$ Ct). Bars – SEM (**E**) Confocal microscopy of frozen sections from AsPC-1 pancreatic tumours grown in nude mice, stained with anti-human PV1 (red) showing lack of expression of PV1 by human cancer cells. In the same specimen, mouse PV1 (blue) is present and colocalizes with the vascular marker CD102 (green), both detected by antibodies directly coupled to fluorophores. Magnification $20 \times$. Bars – 10μ m.

(Fig. 3E), endoglin/CD105 and CD34 (data not shown). Similar results were obtained by staining of serial sections with anti-PV1 mAb MECA-32 and anti-CD31 (Fig. 3C). We also found that both AsPC-1 and BxPC-3 cells in culture lack expression of PV1 mRNA as shown by real-time PCR (Fig. 3D). Positive controls for real-time PCR were human lung microvascular endothelial cells (HLMVEC) and HLMVEC treated with phorbol myristate acetate, expressing low and high levels of PV1 respectively [6]. As a control for differences between cancer cells in culture and tumours *in vivo*, tumour sections were also probed with human PV1 antibodies. No expression of human PV1 was detected in the AsPC-1 or BxPC-3 tumour sections.

Thus, in both AsPC-1 and BxPC-3 tumours, PV1 is specifically expressed in the tumour ECs and not in the tumour cells *per se*, in good agreement with data in the literature obtained in other tumours.

PV1 down-regulation does not affect vascular density of tumours

Next, we investigated whether the tumour growth inhibition induced by PV1 down-regulation is because of inhibition of tumour angiogenesis. Morphometric analyses of tumour sections stained with antibod-



Fig. 4 Down-regulation of PV1 does not decrease tumour vascular density (**A**) Immunohistochemistry with anti-CD31 antibodies of AsPC-1 (left panels) or BxPC-3 (right panels) tumours untreated (parental) or treated with shLuc-LVs (shLuc), shPV1-1-LV (shPV1-1) and shPV1-5-LV (shPV1-5). Bars $-20 \ \mu m$. (**B**) Morphometric analysis of vascular density in AsPC-1 and BxPC-3 tumours [shLuc-LVs (Luc), shPV1-1-LV (sh1) and shPV1-5-LV (sh5)]. Error bars - SEM.

ies against vascular markers such as CD31 were performed to determine the vascular area density, which is a measure of angiogenesis [35]. Representative images of CD31 staining are shown in Figure 4A. Quantitation data show that there are no statistically significant differences in vascular area density between the control and PV1 shRNAs-treated AsPC-1- and BxPC-3-derived tumours (Fig. 4B).

This result raises doubts that tumour growth inhibition was the result of decreased angiogenesis in this model system suggesting that endothelial PV1 is necessary for tumour growth *via* other mechanisms than promoting angiogenesis.

PV1 down-regulation does not modify endothelial cell proliferation and migration *in vitro*

Angiogenesis, defined as the formation of new vessels from pre-existing ones is based on the ability of ECs to proliferate and migrate [36] [37]. To strengthen the above observations we have next investigated whether the modulation of PV1 expression level influences any of these EC cellular functions *in vitro*. To determine the effect of PV1 gain of function, we used ECs that do not express PV1 (such as HU-VEC and BAEC) either not transduced or transduced with AdPV1HA, AdNull or AdGFP. To determine the effect of PV1 loss of function we have compared MLEC, which constitutively express PV1 and with MLEC stably expressing different shRNAs.

De novo expression of PV1 does not modify proliferation rate of HUVEC (Fig. 5A) or BAEC (not shown), as determined using an assay measuring DNA content. Controls consisted of non-transduced cells and cells transduced with the same MOI of AdGFP or AdNull. Flow cy-tometric analyses of PV1 and GFP expression shows that the cells are efficiently transduced by the adenovirus (Fig. 5B). Reciprocally, in MLEC that constitutively express PV1, PV1 down-regulation by transduction with lentiviruses encoding PV1 shRNAs does not modify PV1 proliferation, as measured by population doubling time (Fig. 5C) or DNA content (Fig. 5D).

Cell migration was assessed using wound assays by either the classical scratch method or by ECIS (electric cell impedance sensing). By the two methods, we found no statistically significant differences between ECs expressing PV1 and those that did not (Fig. 5E and F). Taken together, these data argued that either PV1 overexpression or PV1 down-regulation did not modify EC proliferation and migration.

Discussion

PV1 is an endothelial protein with important roles in formation of diaphragms of caveolae, fenestrae, transendothelial channels and, possibly, vesiculo-vacuolar organelles. The diaphragms are structures with unknown functions. PV1 is specifically expressed on normal ECs in a subset of vascular beds. PV1 is also specifically expressed in ECs of many human and mouse tumours, pancreatic cancer included. The strong expression of PV1 in the tumour ECs raises the question as to its function, if any, especially related to tumour initiation and progression.



Fig. 5 PV1 levels do not modify EC proliferation and migration (**A**) Proliferation of HUVEC transduced with adenoviruses encoding for PV1 (AdPV1) or GFP (AdGFP) measured with the DNA content assay. Data were obtained from n = 8 sample replicates per condition. Error bars – SEM. (**B**) Flow cytometric analysis of PV1 (*left*) and GFP expression (*right*) in HUVEC transduced with AdPV1HA (AdPV1) or AdGFP (AdGFP). Control cells were either non-transduced or AdNull transduced HUVEC. Representative samples (of n = 4) are shown in each panel. Control cells (*grey histogram*), AdGFP-transduced cells (*dashed line histogram*), AdPV1HA-transduced cells (*solid line histogram*). (**C**) Average doubling time of MLEC parental or stably expressing shLuc (Luc), shPV1-1 (sh1) or shPV1-5 (sh5). Data were obtained from triplicate samples. Error bars – SEM. (**D**) Proliferation of MLEC parental or stably expressing shLuc (Luc), shPV1-1 (sh1) or shPV1-5 (sh5) measured with the DNA content assay. Data were obtained from n = 8 samples per condition. Error bars – SEM. (**E**–**F**) Migration of HUVEC transduced with adenoviruses encoding for PV1 (AdPV1) or GFP (AdGFP), as measured by the monolayer wound assay at 8 hrs after monolayer wounding (**E**) or by the ECIS method (**F**). For the latter, the cells were plated onto electrodes at near confluence and the monolayer allowed to mature for 44 hrs when the cells over the electrodes were eliminated with electric pulses and the monolayer recovery was monitored for 11 hrs. The traces are representative samples. Data were obtained from n = 4 sample replicates per condition. Error bars in **E**) – STDEV.

On the basis of its specific location, PV1 was hypothesized to play an active role in tumour angiogenesis, which is sustained by the published *in vitro* data where PV1 down-regulation by siRNA was shown to block the formation of capillary tubes by human ECs in Matrigel and migration towards tumour-secreted angiogenic growth factors [12]. These *in vitro* data, supporting a role of PV1 in EC migration, have not yet been validated *in vivo* prompting us to ask this question in a heterotopic xenograft tumour model.

As a model system, we have chosen heterotopic human tumour xenografts in athymic nude mice treated with LV encoded PV1 shRNA delivered by intratumoural injection. Tumours were derived from AsPC-1 and BxPC-3 cells, two established human PDAC cell lines isolated from metastatic and primary PDAC tumours, respectively. The model is simple to execute and the intratumoural injection of lentiviruses expressing various shRNAs was previously validated to be an effective means in down-regulating target molecules in this model [25]. Because the tumour microenvironment (i.e. tumour ECs, stromal cells and tumour-infiltrating leucocytes) is all derived from the mouse host, the model also allows the study of the down-regulation of PV1 expressed in the stroma *versus* tumour cells using speciesspecific shRNAs.

The salient finding of our work is the efficacy of PV1 down-regulation by shRNA in inhibiting the growth of pre-existing tumours in both PDAC models. This study demonstrates, for the first time, the potential therapeutic effect of targeting PV1 for the treatment of PDAC. As shown before, no inhibitory effect of the empty virus or shLuc-LV was observed when compared to the untreated tumours, demonstrating that tumour inhibition is not because of non-specific effects of the LVs or the shRNA. Moreover, this also shows that shR-NAs do not inhibit angiogenesis non-specifically, as was suggested by other studies with targeted and non-targeted siRNAs acting *via* toll-like receptor 3 (TLR3) [38, 39]. The specificity we observe could be because of the fact that in our model, shRNAs are expressed inside the LV-infected cells and as such, may not engage TLR3. Finally, the inhibition of tumour growth by two different PV1 shRNAs in two different models rules out (or makes highly unlikely) off-target effects of individual shRNAs.

The LVs infect all cell types present in the tumour [25], therefore it was important to demonstrate the specificity of PV1 location in these tumours. Staining tumour sections with PV1 and various EC markers used to detect tumour vessels, we show that PV1 is specifically expressed in ECs and not tumour cells, as shown for many other tumours before [10–12, 15]. We verified that tumour cells lack PV1 at both protein and mRNA level. Most importantly, the shRNA sequences used do not target human PV1. These data demonstrate that down-regulation of endothelial PV1 is responsible for the anti-tumour effect, opening the possibility to use PV1 as a therapeutic target in a broad range of cancers, where PV1 is expressed in the vessels.

The effect of PV1 down-regulation on tumour growth can be reasonably explained by the effect of either inhibition of tumour angiogenesis resulting in decreased ratio of tumour vascular volume to tumour tissue volume and/or impaired function of the tumour vessels. We find that the vascular area counts inside and at the periphery of the tumour were similar in both shPV1 and control-treated tumours, suggesting that angiogenesis may not be impaired in the PDAC models used. Although the value of density counts in measuring whether angiogenesis is inhibited or not has limits, as discussed in the past by Hlatky, et al., [40], these conclusions are supported by the lack of effect of PV1 gain and loss of function on both EC proliferation and migration in vitro. Moreover, analysis of PV1-knockout mice demonstrates that PV1 is not required for both vasculogenesis and angiogenesis during mouse development, as these mice have normally formed vessels (D. Tse, S. Deharvengt, C. McGarry, D. S. Longnecker, C. Carriere and R. V. Stan, unpublished data).

Several laboratories have demonstrated PV1 to be expressed specifically in tumour ECs and we have recently published data strongly suggesting that, at cellular level, the function of PV1 in ECs is to form diaphragms [9]. These diaphragms are associated with four different EC structures, of which caveolae [41] and vesiculo-vacuolar organelles [42] were previously postulated as regulating tumour growth. Additional experiments will be needed to dissect which diaphragm on which organelle is important.

At the vascular physiology function level, to date, PV1 was implicated in the maintenance of basal permeability in normal vessels (Tse, et al., 2012, submitted) and in the recruitment of leucocytes into inflammation sites, by unknown mechanisms [17]. Tumour vessels are fenestrated in most solid cancers and are quite disorganized and leaky: the significance of this increased leakiness for tumour growth is under debate [43]. If we are to speculate, the deletion of PV1 will presumably increase the permeability of tumour vessels by deletion of the diaphragms of fenestrae, and possibly those of VVOs. Presently, it is not obvious what consequences this increased leakiness of already leaky vessels will have on tumour growth. Moreover, data in the literature argues that decreasing tumour vascular permeability inhibits tumour growth [44]. Finally, PV1 promotion of diapedesis of leucocytes in inflammation sites [17] raises the possibility of PV1 being important for recruitment of tumour-infiltrating leucocytes with roles in promoting tumour growth [45], which is currently under study in immunocompetent models of cancer.

Finally, by far the most exciting, is the translational potential of PV1 function to be harnessed in devising new drugs against cancer. Here too, experiments employing disease-relevant genetically engineered mouse models will be needed to further explore the role of PV1 in tumour growth.

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Conflict of interest

The authors declare no conflict of interest.

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