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Dual role of pericyte α6β1-integrin in tumour blood vessels

Louise E. Reynolds¹, Gabriela D’Amico¹*, Tanguy Lechertier¹*, Alexandros Papachristodoulou², José M. Muñoz-Félix¹, Adèle De Arcangelis³, Marianne Baker¹, Bryan Serrels⁴ and Kairbaan M. Hodivala-Dilke¹

¹Adhesion and Angiogenesis Laboratory, Centre for Tumour Biology, Barts Cancer Institute- A CRUK Centre of Excellence, Queen Mary University of London, Charterhouse Square, London, UK. EC1M 6BQ.

²Laboratory for Molecular Neuro-Oncology, Dept. of Neurology, University Hospital Zurich, Frauenklinikstrasse 26, CH-8091 Zurich, Switzerland.

³IGBMC UMR 7104, INSERM U964, Université de Strasbourg, BP. 10142, 1, Rue Laurent Fries, 67404 Illkirch Cedex, France.

⁴Cancer Research UK Edinburgh Centre, University of Edinburgh, Crewe Road South, Edinburgh, UK. EH4 2XR.

*These authors contributed equally
*Corresponding author: l.reynolds@qmul.ac.uk

Running title: Dual role for pericyte α6β1 in tumours
Abstract

The integrin α6β1 is a major laminin receptor, and formation of a laminin-rich basement membrane is a key feature in tumour blood vessel stabilisation and pericyte recruitment; processes that are important in the growth and maturation of tumour blood vessels. However, the role of pericyte α6β1-integrin in angiogenesis is largely unknown. We developed mice where the α6-integrin subunit is deleted in pericytes and examined tumour angiogenesis and growth. These mice had: i) reduced pericyte coverage of tumour blood vessels; ii) reduced tumour blood vessel stability; iii) increased blood vessel diameter; iv) enhanced leakiness and, v) abnormal blood vessel basement membrane architecture. Surprisingly, tumour growth, blood vessel density and metastasis were not altered. Analysis of retinas revealed that deletion of pericyte α6-integrin did not affect physiological angiogenesis. At the molecular level, we provide evidence that pericyte α6-integrin controls PDGFRβ expression and AKT/mTOR signalling. Together, we show that pericyte α6β1-integrin regulates tumour blood vessels by both controlling PDGFRβ and basement membrane architecture. These data establish a novel dual role for pericyte α6-integrin that affects the blood vessel phenotype during pathological angiogenesis.

Key words

Integrins; pericyte; tumour growth; angiogenesis
**Introduction**

Blood vessels comprise of endothelial cells supported by mural cells, also known as pericytes. Although the role of integrins in endothelial biology has been studied extensively (Avraamides et al., 2008, Germain et al., 2010), almost nothing is known about the role of pericyte integrins, including α6-integrin (Garmy-Susini et al., 2005, Liu and Leask, 2012). Integrins are transmembrane cell surface receptors that mediate cell-cell and cell-extracellular matrix (ECM) interactions. One of the major components of the vascular basement membrane is laminin whose predominant adhesive receptors include integrins α6β1 and α6β4 (Durbeej, 2010). Targeting endothelial integrins has proved to be relatively beneficial in treating cancer in preclinical studies (Yamada et al., 2006, Tabatabai et al., 2010) but has had limited success in clinical trials (Patel et al., 2001), therefore new targets, including pericytes are currently being examined.

Tumour blood vessels have many structural abnormalities including decreased endothelial barrier function, reduced pericyte recruitment and poor basement membrane organisation when compared with normal, quiescent vessels (Armulik et al., 2005). Studies have shown that pericyte recruitment and investment to blood vessels stimulates endothelial cell basement membrane (BM) deposition and organisation in vitro (Stratman et al., 2009). This is mediated mainly by secretion of endothelial PDGF-BB, attracting PDGFRβ-positive pericytes which adhere to the BM surrounding endothelial cells (Stratman et al., 2010, Abramsson et al., 2003). In vivo, mice lacking PDGF-BB/PDGFRβ signalling fail to adequately recruit pericytes to newly formed blood vessels resulting in severe perturbation of blood vessel stabilisation and maturation (Hellberg et al., 2010). Furthermore, interference with PDGF-BB/PDGFRβ signalling results in disruption of already established endothelial/pericyte associations and vessel destabilisation during retinal development (Benjamin et al., 1998). Whether pericyte α6-integrin might regulate tumour vessel stability was hitherto unknown.
In the present study, we examined the role of pericyte α6-integrin on tumour blood vessel function using a genetic ablation approach. Surprisingly, loss of pericyte α6-integrin did not affect tumour growth, angiogenesis or metastasis but did cause a decrease in pericyte association with tumour blood vessels and poor basement membrane organisation, with an associated increase in vessel leakage and instability. At the molecular level, we demonstrate a novel mechanism by which pericyte α6-integrin reduces PDGFRβ expression on pericytes and therefore diminishes responses to PDGF-BB. This, in turn, is the likely mechanism for aberrant pericyte investment of tumour blood vessels. Together, our data suggest that pericyte α6β1-integrin plays a dual role in regulating PDGFRβ expression and BM organisation that likely increases vessel leakage and instability.
Methods

Generation of mice

α6-integrin floxed mice (Bouvard et al., 2012, Germain et al., 2010) were bred with mice expressing Cre-recombinase under the control of the Platelet Derived Growth Factor Receptor β (PDGFRβ) promoter, PDGFRβCre (Foo et al., 2006) to generate pdgfrβcre-;α6fl/fl and pdgfrβcre+;α6fl/fl mice. mTmG mice were obtained from Cancer Research UK.

Immunostaining of tumour sections

Unless otherwise stated, frozen sections were fixed in ice-cold acetone for 10 min followed by permeabilisation with 0.5% NP-40 for 10 min. Sections were blocked for 45 min with 1.0% bovine serum albumin (BSA) and 0.1% Tween 20 in PBS. Primary antibodies were incubated overnight at 4°C followed by incubation with a fluorescently conjugated secondary antibody for 1 h at room temperature (1:1000; Invitrogen). The following primary antibodies were used: α6-integrin (Chemicon GoH3), NG2 (Millipore AB5320), endomucin (Santa Cruz) (all 1:100).

Quantitation of α6-integrin on pericytes

Quantification of α6-integrin expression on NG2-positive pericytes on tumour blood vessels from pdgfrβcre-;α6fl/fl and pdgfrβcre+;α6fl/fl mice was performed using ImageJ™ software. The mean pixel intensity of α6-integrin expression on NG2-positive pericytes was quantitated.

Immunostaining and quantitation of basement membrane

Immunostaining for laminin α4 chain (kind gift from Takako Sasaki) was performed as described previously (Sasaki et al., 2001). For laminin α5 chain, sections were fixed in 4% paraformaldehyde (PFA), washed twice with PBS, then blocked with 1% BSA for 30 min. Sections were incubated overnight at 4°C with primary antibody (1:400 dilution in blocking buffer) (kind gift from Jeffrey H. Miner (Pierce et al., 2000)), followed by several washes,
incubation with Alexa 488-conjugated anti-rabbit secondary antibody (1:1000; Invitrogen) and mounted.

For Collagen IV (Abcam) and Fibronectin (Abcam) staining, sections were fixed in 4% PFA, blocked with 3% NGS, 0.1% TritonX-100 (TX-100) in PBS for 30 min at room temperature. Primary antibody was diluted 1:200 in 1% normal goat serum (NGS), 0.1% TX-100 in PBS and incubated overnight at 4°C. Sections were washed three times with PBS, incubated with Alexa 488-conjugated anti-rabbit secondary antibody (Invitrogen) diluted 1:100 in 1% NGS TX-100, washed three times and mounted with Prolong Gold anti-fade with DAPI.

Fluorescence staining was visualised using the Axioplan microscope (Zeiss). Images were captured using Axiovision Rel. 4.0 software. The Axiovision software linear measuring tool was used to analyse the spread (in μm) of basement membrane surrounding blood vessels.

**Pericyte association**

For analysis of pericyte coverage, tumour sections were double immunostained for endomucin and NG2 (for details see Immunostaining of tumour sections). Pericyte coverage was quantified by counting the total number of endomucin positive blood vessels across whole tumour sections followed by the numbers of blood vessels positive for both endomucin and NG2. The percentage (%) of blood vessels with associated pericytes was calculated.

**Tumour blood vessel leakage**

To analyse tumour blood vessel leakage, Hoechst dye (4 μg/ml; Sigma, H33258) was used. Hoechst dyes diffuse quickly from vessels and binds to the DNA of cells surrounding blood vessels, allowing for quantitation of areas of uptake by perivascular tumour cells and hence blood vessel leakage (Janssen et al., 2005). Briefly, mice were injected sequentially with 100 μl PE-CD31 (Biolegend; to stain functional blood vessels) followed 9 min later with 100 μl Hoechst (4 μg/ml; Sigma, H33258) via the tail vein and 1 min later culled. Tumours were excised and snap frozen. 8-10 fields at x20 magnification were analysed by ImageJ™. For
quantitation, blood vessel leakage was calculated by counting the numbers of Hoechst positive nuclei surrounding PE-CD31 positive tumour areas. Results are shown in arbitrary units (AU).

**Tumour growth and angiogenesis**

The syngeneic mouse tumour cell lines B16F0 (melanoma, derived from C57BL6) and Lewis Lung Carcinoma (LLC) (both from ATCC) were used in subcutaneous tumour growth experiments. 1x10^6 B16F0 cells or 0.5x10^6 LLC cells, resuspended in 100 μl of phosphate buffered saline (PBS), were injected subcutaneously into the flank of 12-14 week old pdgfrβcre+;α6fl/fl and littermate control mice (pdgfrβcre-;α6fl/fl mice). Tumour growth was measured every 2 days using calipers. After 14 days, animals were culled, tumours excised and either fixed in 4% formaldehyde in PBS overnight, or snap-frozen in liquid nitrogen, for subsequent immunohistochemical analysis.

**Blood vessel density**

Size-matched tumours from pdgfrβcre-;α6fl/fl and pdgfrβcre+;α6fl/fl mice were snap-frozen and bisected, and cryosections made. Frozen sections were fixed in 100% acetone at −20°C, rehydrated in PBS for 10 min, and then blocked (PBS, 1% BSA, 0.1% Tween-20) for 45 min at room temperature. After a 5 min wash in PBS, sections were incubated with 1:100 anti-endomucin in blocking buffer for 45 min at room temperature. After three 5 min washes in PBS, sections were incubated with Alexa 488-anti-rat secondary antibody (Invitrogen) diluted in blocking buffer. After three 5 min washes in PBS, sections were washed briefly with distilled water before being mounted. The number of endomucin-positive blood vessels present across the entire area of each mid-line tumour section from size and age-matched tumours was counted and divided by the area of the section to determine tumour blood vessel density.
**Tumour blood vessel diameter**

The diameter of endomucin-positive blood vessels were quantified using the Axiovision™ software linear measuring tool.

**Perfused blood vessel analysis**

For analysis of the % of perfused, functional tumour vessels, 100 μl PE-CD31 antibody (Biolegend, London, UK) was injected via the tail vein 10 minutes (mins) prior to culling mice. Tumours were dissected immediately, snap frozen and sectioned. Frozen sections were then immunostained for endomucin as described above. To calculate the % number of functional vessels, the number of PE-CD31-positive blood vessels was divided by the total number of endomucin-positive blood vessels.

**Tumour metastasis**

0.5 x 10⁶ LLC tumour cells were injected subcutaneously and tumours were allowed to reach a size of 100 mm³ before surgical resection. The mice were monitored for up to 3 weeks, after which the metastatic burden quantified by counting numbers of surface lung metastases. Lungs were then fixed and sections stained with H&E for further analysis. The numbers of surface metastases was counted immediately after fixation, to give the total number of lung metastases per mouse. To examine internal metastases, H&E stained sections were analysed and the area of individual metastases was measured using Axiovision™ software, to give the internal metastatic nodule area.

**Primary endothelial cell, fibroblast and pericyte isolation**

Primary mouse endothelial cells were isolated from lungs and maintained as described previously (Reynolds and Hodivala-Dilke, 2006). Briefly, pdgfrβcre--;α6fl/fl and pdgfrβcre+;α6fl/fl mouse lungs were minced, collagenase digested (Type I, Gibco), strained through a 70 μm cell strainer (BD Falcon) and the resulting cell suspension plated on flasks
coated with a mixture of 0.1% gelatin (Sigma), 10µg/ml fibronectin (Millipore) and 30µg/ml rat tail collagen (Sigma). Endothelial cells were purified by a single negative (FCγ sort-RII/III; Pharmingen) and two positive cell sorts (ICAM-2; Pharmingen), using anti-rat IgG-conjugated magnetic beads (Dynal). During preparation of primary endothelial cells, lung fibroblasts were isolated from the non-endothelial cell population that was generated during the first positive sort. For all cell types, passaging occurred when cells reached 70% confluency. Cells were trypsinised, centrifuged, washed with PBS and replated on pre-coated flasks for endothelial cells and pericytes and non-coated flasks for fibroblasts. Fibroblasts were cultured in DMEM + 10% FCS to passage 4, Endothelial cells in MLEC (Ham’s F-12, DMEM (low glucose), 10% FCS, heparin and endothelial mitogen (Generon)) to passage 4-5. Pericytes were isolated from mouse brains as described previously (Tigges et al., 2012) and cultured in Pericyte medium (ScienCell) to passage 9.

**FACS analysis**

Primary mouse brain pericytes isolated from pdgfrβ<sup>cre-;α6fl/fl</sup> and pdgfrβ<sup>cre+;α6fl/fl</sup> mice were incubated with an anti-α6-integrin subunit antibody (GoH3; Abcam) to determine expression levels, for 30 min at 4°C. This was followed by incubation, for 30 min at 4°C with an appropriate FITC-conjugated secondary antibody. Unstained cells were used as a control. For characterisation of primary mouse brain pericytes, cells were washed with PBS and trypsinised at 37°C. The cell suspensions were washed in medium containing serum and centrifuged at 1, 200 rpm for 3 min. Cells were washed with cold FACS buffer (1% BSA/PBS) and fixed with 4% formaldehyde for 10 min at room temp. Cells were washed with FACs buffer and the cell suspensions were incubated with the following primary antibodies: PE-anti CD31 (Biolegend), PE-anti Mac1 (CD11b) (Biolegend), PE-anti GFAP (BD Biosciences), APC-anti PDGFRβ (Biolegend), PE-Cy7-anti CD146 (Biolegend) for 30 min. Cells were then washed three times in sample buffer and resuspended in a final volume
of 400 ml. As a control, unstained cells were FACS sorted. Primary mouse lung endothelial cells were incubated with PE-anti CD31 (Biolegend).

**Western blot analysis**

Primary lung endothelial cells, lung fibroblasts and brain pericytes isolated from \textit{pdgfr\textbeta{}cre-}\textbackslash{}\textit{a6fl/fl} and \textit{pdgfr\textbeta{}cre+};\textit{a6fl/fl} mice were grown to 70-80\% confluency then lysed in RIPA buffer. 15-30 \mu{}g protein was run on 8\% polyacrylamide gels then transferred to nitrocellulose membranes. Membranes were probed with primary antibody overnight at 4°C. All antibodies (ERK, AKT, \alpha{}6-integrin) for the signalling studies were purchased from Cell Signaling and used at a 1:1000 dilution. \alpha{}3-integrin antibody was purchased from Millipore (AB1920). The anti-HSC70 antibody, used as a loading control, was from Santa Cruz and used at 1:5000 dilution. For PDGF-BB stimulation, pericytes were serum starved for 6 h in Optimem with 0.5\% FCS, then stimulated with PDGF-BB (30 ng/ml; Peprotech, UK) for 0, 30 s, 60 s, 5 min, 15 min, 30 min before lysis. Densitometric readings of band intensities were obtained using the ImageJ™ software.

**PDGFR\textbeta{} immunostaining of cells**

Immunostaining of brain pericytes, isolated from \textit{pdgfr\textbeta{}cre-};\textit{a6fl/fl} and \textit{pdgfr\textbeta{}cre+};\textit{a6fl/fl} mice for PDGFR\textbeta{} (28E1, Cell Signaling), was performed according to the manufacturers’ protocol.

**Aortic ring assay**

Thoracic aortas were isolated from \textit{pdgfr\textbeta{}cre-};\textit{a6fl/fl} and \textit{pdgfr\textbeta{}cre+};\textit{a6fl/fl} 8-10 week old mice and prepared for culture as described previously (Baker et al., 2012). Where indicated, culture medium was supplemented with PDGF-BB at 30 ng/ml (Peprotech, London, UK). PBS was used as a control. Rings were fed every 3 days with fresh medium with or without
PDFG-BB (30 ng/ml). Sprouting microvessels were counted after 9 days in culture, fixed and stained to identify endothelial cells and pericytes, as described previously (Baker et al., 2012).

**Scratch wound assays**

Primary mouse brain pericytes from pdgfr<sup>cre</sup>;α<sup>6</sup>fl/fl and pdgfr<sup>cre+</sup>;α<sup>6</sup>fl/fl mice were grown to confluency in Pericyte medium (ScienCell, CA. Cat no. 0010) in 6-well plates coated with 0.1% gelatin and fibronectin. Cells were serum starved overnight in Optimem containing 0.5% FCS. The following day, the monolayers were scratched horizontally and vertically through the centre of each well. The cells were either stimulated with PDGF-BB (30 ng/ml) or not and cell migration monitored over a 72 h period. At each timepoint, 12 photographs were taken of each scratch and the wound width was measured using ImageJ™ software. Results were normalised to time 0.

**Proliferation assay**

Primary WT and α<sup>6</sup>-null brain pericyte proliferation was assessed using the CellTiter 96® Aqueous One Solution Reagent (Promega), according to the manufacturer’s instructions. Plates were read at a wavelength of 490 nm, with absorbance measured relative to blank wells containing reagent only. Plates were coated with 0.1% gelatin and fibronectin prior to seeding the pericytes.

**Characterising transgenic mice**

The primers for the Cre PCR were forward primer: 5'-GCCGCATTACCGTGCTGCAAGA-3'; reverse primer: 5'-GTGGCAGATGGCGGGCAACACCATT-3. The reaction generates a fragment of approximately 1000 bp. The primers for α<sup>6</sup>-floxed PCR were forward primer: 5'-AGAAGGTGATGTTACCC-3' and reverse primer: 5'-AATGTAACTAGCATTCAAGT-3.
The PCR reaction generates a 154 bp fragment for the α6-floxed allele and a 120 bp fragment for the wild-type allele (described in Germain et al.).

**Whole-mount immunofluorescence on retinas**

Postnatal day (P) 9 eyes were fixed in 4% PFA overnight at 4°C. Retinas were dissected in PBS and, after 5 washes in PBS, incubated in blocking solution (1% FBS (Sigma), 3% BSA (Sigma), 0.5% Triton X-100 (Sigma), 0.01% Na deoxycholate (Sigma), 0.02% Na Azide (Sigma) in PBS, pH7.4) for 2 h at room temperature. Retinas were incubated for overnight at 4°C with primary antibodies: rabbit polyclonal anti-NG2 (Millipore, #AB5320, dilution 1:500) in blocking solution:PBS (1:1) overnight at 4°C. Retinas were then washed several times in PBS and incubated with secondary antibody AlexaFluor 488 and AlexaFluor 548 conjugates (Life Technologies), all diluted 1:300, overnight at 4°C. After several washes in PBS, retinas were post-fixed in 1% PFA and re-blocked in 1%BSA/0.5% Triton X-100 in PBS for 1 hr at room temperature. After 2 rinses in PBlec (0.1 nM CaCl2, 0.1 MgCl2 mM, 0.1 MnCl2 mM, and 1% Triton X-100 in PBS; pH6.8), retinas were incubated with biotinylated Isolectin B4 (B1205, VectorLabs, dilution 1:12.5) in PBlec overnight at 4°C. After several washes in PBS, retinas were incubated with Streptavidin Alexa Fluor 647 conjugate diluted 1:100 in 0.5%BSA/0.3% Triton X-100 in PBS overnight at 4°C. After several washes in PBS, retinas were incubated with Hoechst dye as a nuclear counterstain (Life Technologies), washed and mounted with ProLong Gold (Molecular Probes). Fluorescently labelled samples were imaged with a confocal microscope (Carl Zeiss LSM 710, Carl Zeiss) in multichannel mode. Three-dimensional projections were digitally reconstructed from confocal z-stacks using the LSM Zen 2009 software and Image J open source image processing software (version:2.0.0-rc-43/1.51d).
**Adhesion assay**

Adhesion assays were performed as previously described (Germain et al., 2010), using brain pericytes isolated from pdgfrβcre-; α6fl/fl and pdgfrβcre+; α6fl/fl mice. Adhesion of pericytes is presented relative to adhesion of pericytes to fibronectin for the same genotype.

**Immunostaining of immune cells, fibroblasts and endothelial cells**

Immunostaining of immune cells from B16F0 tumour sections was performed as described previously (Reynolds et al., 2010).

Primary mouse lung fibroblasts and endothelial cells were fixed with 4% PFA for 10 min, washed twice and blocked with 5% NGS in PBS for 30 min at room temperature. Primary antibodies to vimentin (1:100; Cell Signaling), NG2 (1:100; MAB 5385, Millipore), endomucin (1:100; V7C7, Santa Cruz) and α-sma (clone 1A4, Sigma) were incubated for 1 h at room temperature, washed three times with PBS, and incubated with the relevant secondary antibody for 45 min at room temperature.

**Reverse phase protein array (RPPA) Nitrocellulose slides**

Cell lysates were prepared using ice cold RIPA buffer. After normalization, to adjust protein concentrations, triplicate spots of each lysate were deposited onto16-pad Avid Nitrocellulose slides (Grace Bio) under conditions of constant 70% humidity using an Aushon 2470 Array platform (Aushon BioSystems). After printing and washing steps, the arrays were blocked by incubation in Superblock (Thermo Scientific #37535) for 10 min. The protein array chips were subsequently incubated for 1 h with primary antibody followed by repeat blocking with Superblock and 30 min incubation with anti-rabbit Dylight-800 conjugated secondary antibody (Cell Signaling, cat.no.5151).

Following secondary antibody incubation and subsequent wash steps the immune-stained arrays were imaged using an Innopsys 710IR scanner (Innopsys France). Microarray images were obtained at the highest gain without saturation of fluorescent signal detection.
analysis was performed using Mapix software (Innopsys France) to calculate the relative fluorescence intensity (RFI) value for each sample. An estimate of total protein printed per feature on the array was determined by staining an array slide with fastgreen protein stain. Readout values for all antibodies tested are expressed as a ratio of total protein loaded and are presented as the mean of technical replicates.

**Statistical analysis**

Statistical significance was calculated using Student's t-test. \( p < 0.05 \) was considered statistically significant.

**Ethical regulations**

All animals were used in accord with United Kingdom Home Office regulations (Home Office license number 70/7449). The in-house Ethics Committee at Queen Mary University of London has approved all experiments, using mice under the project license.
Results

Generation and characterisation of pdgfrβcre+;α6fl/fl mice

We have generated a new mouse model that enables us to study deletion of α6-integrin in pericytes. We bred α6-integrin floxed mice (Bouvard et al., 2012, Germain et al., 2010) with mice expressing Cre-recombinase under the control of the Platelet Derived Growth Factor Receptor β (PDGFRβ) promoter, PDGFRβCre (Foo et al., 2006), to generate pdgfrβcre-;α6fl/fl and pdgfrβcre+;α6fl/fl mice. Mice were born to pdgfrβcre+;α6fl/fl x pdgfrβcre-;α6fl/fl crosses at normal Mendelian ratios and male:female ratios with no obvious adverse phenotype (Fig. S1A-C). Mice were genotyped by PCR analysis (Fig. S1D). Histological analysis of H&E-stained sections of lung, heart, liver and spleen from pdgfrβcre-;α6fl/fl and pdgfrβcre+;α6fl/fl adult mice showed no apparent tissue defects (Fig. S1E). Furthermore, no apparent vascular abnormalities were observed in these tissues (Fig. S1F) or in the developing retina (Fig. S1G), suggesting that loss of PDGFRβ-driven α6 had no apparent effect on physiological angiogenesis. Finally, to confirm pericyte specific Cre expression in our mouse model, we crossed pdgfrβcre- and pdgfrβcre+ mice with the mTmG reporter mouse (Muzumdar et al., 2007). Analysis of tumour blood vessels from pdgfrβcre-;mTmG and pdgfrβcre+;mTmG mice showed that although tomato (mT) expression was observed in blood vessels in both pdgfrβcre-;mTmG and pdgfrβcre+;mTmG mice, GFP (mG) expression, was present in mouse tissue only after Cre excision, and only observed in pericytes in pdgfrβcre+;mTmG mice (Fig. S2A, B). As expected, α6-integrin is expressed on tumour endothelial cells, shown by co-expression of α6-integrin and the endothelial cell marker CD31 (Fig. S2C), suggesting that the deletion of α6-integrin in vivo was restricted to PDGFRβ-positive pericytes.

α6-integrin deficiency on pericytes impairs tumour blood vessel stabilisation

Very few studies have examined the role of pericyte integrins in tumour blood vessels in vivo (Garmy-Susini et al., 2005, Turner et al., 2014, Liu and Leask, 2012). To assess whether α6-
integrin is required for pericyte investment on blood vessels during tumour angiogenesis, \textit{pdgfr\textbeta cre-};\textalpha 6fl/fl and \textit{pdgfr\textbeta cre+};\textalpha 6fl/fl mice were injected with either syngeneic B16F0 melanoma or LLC tumour cells. We first established loss of \textalpha 6-integrin expression in pericytes in tumour blood vessels from \textit{pdgfr\textbeta cre+};\textalpha 6fl/fl mice. Tumour sections from \textit{pdgfr\textbeta cre-};\textalpha 6fl/fl and \textit{pdgfr\textbeta cre+};\textalpha 6fl/fl were double stained for NG2 (a pericyte marker) and \textalpha 6-integrin. Although \textalpha 6-integrin was expressed in NG2 positive pericytes in tumour blood vessels in \textit{pdgfr\textbeta cre-};\textalpha 6fl/fl control mice, \textalpha 6-integrin was undetectable in pericytes from \textit{pdgfr\textbeta cre+};\textalpha 6fl/fl mice (Fig. 1A) confirming that \textalpha 6-integrin expression is reduced on pericytes in tumours grown in \textit{pdgfr\textbeta cre+};\textalpha 6fl/fl mice. Quantification revealed a significant reduction in \textalpha 6-integrin expression in NG2-positive pericytes in tumour blood vessels from \textit{pdgfr\textbeta cre+};\textalpha 6fl/fl mice (Fig. 1B). Having confirmed deletion of \textalpha 6-integrin in pericytes in tumour blood vessels from \textit{pdgfr\textbeta cre+};\textalpha 6fl/fl mice \textit{in vivo}, we asked whether PDGFR\textbeta driven \textalpha 6-integrin deletion could affect pericyte association with tumour blood vessels. A significant reduction in the number of pericytes associated with blood vessels in B16F0 and LLC tumours grown in \textit{pdgfr\textbeta cre+};\textalpha 6fl/fl mice was observed (Fig. 1C). Thus, the absence of pericyte \textalpha 6-integrin corresponds with a reduction in pericyte association with blood vessels.

**Blood vessel leakage is enhanced in tumours from \textit{pdgfr\textbeta cre+};\textalpha 6fl/fl mice**

We next examined other possible biological consequences of the loss of pericyte \textalpha 6-integrin. We found that reduced pericyte investment correlated with a higher frequency of blood vessels with a large diameter (>100 \textmu m), in tumour blood vessels from \textit{pdgfr\textbeta cre+};\textalpha 6fl/fl mice (Fig. 1D). A known result of increased blood vessel diameter is poor functionality of blood vessels (Huang et al., 2010). We therefore examined tumour blood vessel leakage by measuring the relative level of perivascular Hoechst after intravenous injection of Hoechst dye. Blood vessel leakage was calculated by counting the numbers of Hoechst positive nuclei surrounding tumour blood vessels. Tumour blood vessels from \textit{pdgfr\textbeta cre+};\textalpha 6fl/fl mice
showed significantly more leakage of Hoechst dye than from pdgfrβcre-;α6fl/fl mouse tumours (Fig. 1E). Together, these data demonstrate that loss of pericyte α6-integrin is sufficient to reduce pericyte coverage of tumour blood vessels and increase vessel diameter and leakage.

**Blood vessel basement membrane organisation is altered in tumours grown in pdgfrβcre+;α6fl/fl mice**

Given that extracellular matrix (ECM) deposition is a crucial step in the maturation of tumour blood vessels, and requires the presence of both pericytes and endothelial cells, we examined whether the distribution of various BM matrices correlated with the observed decrease in pericyte tumour blood vessel association in pdgfrβcre+;α6fl/fl mice. Concomitant with reduced vascular stabilisation and increased leakiness in the pdgfrβcre+;α6fl/fl tumour blood vessels, a higher frequency of disorganised BM around tumour blood vessels was seen in pdgfrβcre+;α6fl/fl mice. Specifically, immunofluorescence analysis of collagen IV (Fig. 2A), fibronectin (Fig. 2B), laminins α5- (Fig. 2C) and α4-chains (Fig. 2D) in the BM around tumour blood vessels, revealed a “shorelining” pattern of BM components (DiPersio et al., 1997) in pdgfrβcre+;α6fl/fl mice. Quantification confirmed that the BM was wider with aberrant organisation, in blood vessels from pdgfrβcre+;α6fl/fl mice than from pdgfrβcre-;α6fl/fl mice (Fig. 2E). Adhesion assays, using mouse primary pericytes showed, as expected, that the absence of α6-integrin reduced significantly the adherence to Laminin 111 (Lm-1), but not fibronectin (Fig. S3A).

Since blood vessel stabilisation is a consequence not only of BM deposition but the association of supporting cells, ie. pericytes, these results show that pericyte investment and the formation of a normal BM is affected by deletion of pericyte α6-integrin, and likely contributes to the abnormal stability and leakage of tumour blood vessels in pdgfrβcre+;α6fl/fl mice.
Depletion of pericyte α6-integrin does not affect tumour growth or metastasis

Despite the dramatic changes in pericyte association, vessel leakage and BM architecture, no significant difference in tumour growth (Fig. 3A), blood vessel density (Fig. 3B), numbers of functional blood vessels (Fig. 3C), metastasis (Fig. 3D) or immune cell infiltrate (Fig. S3B) was observed between pdgfβcre-;α6fl/fl and pdgfβcre+;α6fl/fl mice. These data suggest that the changes in blood vessels observed in pdgfβcre+;α6fl/fl mice is not sufficient to affect tumour growth, angiogenesis or metastasis.

α6-integrin controls pericyte PDGFRβ expression

In mice, genetic ablation of PDGF-BB and PDGFRβ leads to various vascular abnormalities associated with pericyte loss, including a loss of pericyte association to blood vessels (Abramsson et al., 2003). Since PDGF-BB is one of the major growth factors involved in pericyte investment to blood vessels (Abramsson et al., 2003, Furuhashi et al., 2004) we sought to investigate whether loss of α6-integrin affected pericyte responses to PDGF-BB. Initially, primary pericytes, fibroblasts and endothelial cells were isolated from pdgfβcre-;α6fl/fl and pdgfβcre+;α6fl/fl mice and characterised (Fig. S4A-C). Western blot analysis and FACS confirmed deletion of α6-integrin only in pericytes (Fig. 4A). This loss of α6-integrin was not compensated for by the overexpression of another laminin receptor, the α3-integrin subunit in α6-null pericytes (Fig. 4B). Further Western blot analysis showed that α6-integrin levels were normal in fibroblasts and endothelial cells isolated from pdgfβcre-;α6fl/fl and pdgfβcre+;α6fl/fl mice (Fig. 4C). The loss of α6-integrin in pericytes correlated with an approximately two-fold reduction in levels of PDGFRβ in α6-null pericytes, suggesting that α6-integrin was indeed regulating the expression of PDGFRβ protein (Fig. 4D). Additionally, PDGFRβ levels were significantly reduced in PDGFRβ immunostained α6-null pericytes compared with WT pericytes in vitro (Fig. 4E).
PDGF-BB responses are diminished in α6-null pericytes

Since changes in receptor expression levels do not always reflect changes in downstream signalling of the corresponding tyrosine kinase receptor (Platta and Stenmark, 2011), we next sought to confirm whether the decreased expression of PDGFRβ correlated with diminished responses to PDGF-BB stimulation in α6-null pericytes. We showed that PDGF-BB stimulated ex vivo microvessel sprouting was reduced in aortic rings isolated from pdgfrβcre+;α6fl/fl mice compared with PDGF-BB stimulated sprouting from pdgfrβcre-;α6fl/fl mouse aortic rings (Fig. 5A). In this assay the vessel sprouts become surrounded by pericytes which proliferate and migrate along the endothelium (Nicosia, 2009). We were unable to analyse pericyte coverage in pdgfrβcre+;α6fl/fl aortic rings due to the complete lack of sprouts that grew in response to PDGF-BB. In response to PDGF-BB, α6-null pericyte migration and proliferation was significantly reduced when compared with WT controls (Fig. 5B, C). We next examined the effect of α6-deficiency on PDGF-BB stimulated downstream signalling in pericytes. Interaction of PDGFRβ with PDGF-BB activates several signalling pathways, including MAP Kinase (ERK) and PI3K/AKT. Western blot analysis showed that, in α6-null pericytes, PDGF-BB mediated stimulation of ERK and AKT pathways were both reduced significantly (Fig. 5D, E). Since integrins and PDGFR can activate many downstream signalling pathways, we performed a non-candidate proteomic study using Reverse Phase Protein Array (RPPA) analysis, to identify other possible pathways that may be affected by the absence of α6-integrin on pericytes. We found that components of the AKT/mTOR signaling pathway (AKT, P70 S6kinase, mTOR, pS6 ribosomal protein and 4E-BP1) were significantly downregulated in α6-null pericytes (Fig. S4D). This pathway is activated downstream of integrins and is known to be regulated by PDGFR expression and function (Zhang et al., 2007). These results suggest that absence of α6-integrin results in a reduction of PDGFRβ levels, significantly reducing pericyte responses to PDGF-BB.

Collectively, these results demonstrate a novel role for pericyte α6-integrin in both the
regulation of PDGFRβ levels that was previously unknown, and in BM organisation. This dual mechanism confers a blood vessel phenotype that affects only the primary tumour.
**Discussion**

The role of pericyte α6-integrin expression has not been addressed previously. We have now shown that, the combined effects of the down-regulation of PDGFRβ and the changes in pericyte adhesion, upon deletion of α6β1-integrin, induces destabilisation of tumour blood vessels. Genetic ablation of pericyte α6β1-integrin correlates with reduced pericyte investment to tumour blood vessels, changes in BM architecture and reduced PDGFRβ expression levels and PDGF-BB-mediated downstream signalling. These correspond with increased blood vessel leakage without affecting tumour growth or metastasis. Together, our results suggest a dual function of α6-integrin on pericytes and here we will discuss both the effect of reduced PDGFRβ levels and decreased pericyte adhesion in turn.

Very few studies have linked pericyte integrins with growth factor receptor regulation. Currently, α5β1 integrin has been shown to regulate signalling through PDGFRβ, in vascular smooth muscle cells (Veevers-Lowe et al., 2011), and inactivation of PDGF-BB signalling can decrease α1β1 integrin levels (Hosaka et al., 2013). In another study, NG2 depletion in pericytes was shown to reduce β1-integrin-mediated signalling (You et al., 2014). Although limited, these studies suggest possible cross-talk between pericyte integrins and growth factor receptors; a mechanism that has been shown previously between endothelial cell integrins and growth factor receptors (da Silva et al., 2010, Germain et al., 2010, Reynolds et al., 2002). In parallel, *in vitro* studies using fibroblasts have highlighted the ability of integrins to enhance PDGF-dependent responses (DeMali et al., 1999, Sundberg and Rubin, 1996). Here we show for the first time that α6β1-integrin acts as a regulator of PDGFRβ – controlling its expression and signalling upon stimulation with PDGF-BB, in pericytes. Our data indicate that depletion of α6-integrin on pericytes leads to a significant reduction in the levels of PDGFRβ. In turn, this leads to a downregulation in the activation of the MAPKinase and AKT signalling pathways, which are both known to be critical for cell migration and proliferation (Lemmon and Schlessinger, 2010). Our observation of reduced pericyte
PDGFRβ levels, signalling and resulting inhibited responses to PDGF-BB, in α6-integrin deficient pericytes, may explain the tumour blood vessel phenotypes we observe in the pdgfrβcre+;α6fl/fl mice; including reduced pericyte blood vessel investment and increased detachment of pericytes to endothelial cells, since these functions have been reported to be mediated by PDGF-BB (Abramsson et al., 2003, Hellberg et al., 2010). It has been well documented that tumours transplanted into PDGF-B retention motif-deficient (pdgf-bret/ret) mice have an approximately 50% reduction in numbers of pericytes that poorly associate with the blood vessel wall and result in leaky vessels (Abramsson et al., 2003). It is noteworthy that although we observe a similar leaky blood vessel phenotype to that reported in tumour growth in pdgf-bret/ret mice, our results indicate that α6β1 integrin is a regulator of pericyte function rather than numbers, as seen in the pdgf-bret/ret mice (Lindblom et al., 2003).

One surprising observation was that despite such striking blood vessel defects in the pdgfrβcre+;α6fl/fl mouse tumours, including increased vessel leakage, we did not observe any changes in tumour growth, angiogenesis or lung metastasis (Cooke et al., 2012, Zang et al., 2015). Our work is in line with previous studies showing that early ablation of NG2+ cells or PDGFβR+ pericyte depletion does not necessarily affect tumour growth and metastasis (Keskin et al., 2015) suggesting that increased leakage alone is not sufficient to enhance metastasis. Indeed, increased vascular leakage has been shown to be insufficient for metastasis per se (Thurston et al., 1999). We hypothesise that despite the observed decrease in blood vessel pericyte coverage in pdgfrβcre+;α6fl/fl mice, the remaining pericytes attached to the endothelial cells provide enough survival factors, for example VEGF and Ang-1, to allow ECs to survive, allowing for normal tumour angiogenesis and tumour growth. Studies have shown that decreased PC coverage can lead to regression of blood vessels but the magnitude is tumour specific and does not necessarily retard tumour growth (Sennino et al., 2007). Indeed, treatment of RipTag2 tumours with anti-PDGFRβ antibody reduces pericyte numbers and enlarges blood vessels but does not reduce tumour vascular density.
(Song et al., 2005). Although not within the scope of this study, we believe that the phenotype observed in our pdgfrβcre+;α6fl/fl mice may help to improve chemotherapy efficacy to primary tumours, due to the leaky vessel defect.

As well as pericyte α6-integrin deficiency affecting PDGFRβ levels and upstream signalling, we also show that deletion of α6-integrin affects the ECM adhesion properties of pericytes in vitro and in vivo. The BM that surrounds blood vessels is necessary for vessel integrity, stability and maturation. Recent studies have highlighted the importance of normal endothelial cell-pericyte interactions for proper BM organisation (Davis et al., 2013, Stratman et al., 2009, Baluk et al., 2003). When this interaction is de-stabilised, BM organisation is affected, resulting in decreased vessel integrity (Stratman et al., 2009, Davis et al., 2013). Therefore, it is possible that the phenotype we observe in the microvessel BM of pdgfrβcre+;α6fl/fl mice may be due, at least in part, to a loss of pericyte adhesion to endothelial cell basement membrane. Similar phenotypes have been shown in other studies, for example: inactivation of the β1-integrin subunit in mural cells leads to failure of these cells to associate with the subendothelial BM (Abraham et al., 2008); genetic ablation of a related laminin receptor, α3-integrin results in an epidermal BM defect, in which components of the BM show a disorganised expression pattern (DiPersio et al., 1997, Georges-Labouesse et al., 1996, Has et al., 2012) very similar to that observed in tumour blood vessels of α6-integrin pericyte deficient mice. Foxf2 (forkhead transcription factor, specifically expressed by pericytes) deficiency in brain pericytes leads to significantly reduced PDGFRβ and αvβ8 integrin levels with thinning of the vascular basal lamina, resulting in a leaky blood brain barrier (Reyahi et al., 2015); inactivation of PDGF-BB signalling decreases α1β1 integrin levels and impairs pericyte adhesion to extracellular matrix components of blood vessels (Hosaka et al., 2013).

Together these studies all support a common notion that crosstalk between pericyte integrins with PDGFRβ signalling can affect vascular BM organisation and vessel function. It is also
conceivable that microvessel BM disorganisation may be a contributing cause of decreased supporting cell coverage in the tumour blood vessels in pdgfrkcre+;α6fl/fl mice. For example, it has been shown that deletion of laminin-α4 chain in mice causes impaired vessel growth due to reduced pericyte recruitment to blood vessels (Abrass et al., 2010). In vitro, pericyte recruitment during tube formation is necessary to stimulate endothelial BM formation (Stratman et al., 2009). Overall, our data suggest that the absence of pericyte α6-integrin leads to i) the reduced investment of pericytes to tumour microvessels possibly due to reduced PDGFRβ levels, and ii) this is associated with poor vessel BM architecture leading to vascular leakage.

Our study provides new insights into the regulation of tumour blood vessels by pericyte α6-integrin which points towards an important role in the regulation of tumour vessel leakage.
Acknowledgments

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Author contributions

L.E.R and K.M.H-D designed the experiments; L.E.R performed the experiments. T.L performed image analysis and mTmG studies. A.P. performed immune cell immunostaining and quantification. G. D performed confocal microscopy and retinal angiogenesis studies. J.M.M-F and L.E.R performed the FACs analysis. A.DA provided the $\alpha_6$-integrin floxed mice. M.B undertook the initial tumour experiments. B.S performed the RPPA analysis. L.E.R and K.M.H.-D wrote the paper.

Conflict of interest

The authors declare no conflict of interest.

Abbreviations

BM, basement membrane; ECM, extracellular matrix; LLC, Lewis Lung Carcinoma
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Figure Legends

Figure 1. Reduced pericyte investment and increased vessel leakage in pdgfrβcre+;α6fl/fl mice.

(A) B16F0 tumour sections from pdgfrβcre--;α6fl/fl and pdgfrβcre+;α6fl/fl mice were double-immunostained for NG2 (red) and α6-integrin (green) expression to determine in vivo expression of α6-integrin in pericytes. α6-integrin was observed in NG2-positive pericytes in tumour blood vessels from pdgfrβcre--;α6fl/fl mice. In contrast, α6-integrin expression was significantly reduced in pericytes in tumour blood vessels from pdgfrβcre+;α6fl/fl mice. Magnified regions show α6-integrin positive pericytes (to give a yellow signal) on pdgfrβcre--;α6fl/fl blood vessels but α6-integrin negative pericytes on pdgfrβcre+;α6fl/fl blood vessels. Arrows, α6/NG2 co-expression; arrowheads, NG2 expression alone. (B) Quantitation of α6-integrin expression in NG2-positive pericytes from tumour blood vessels in pdgfrβcre--;α6fl/fl in pdgfrβcre+;α6fl/fl mice. Bar chart represents mean pixel intensity of α6-integrin expression in NG2-positive cells+SEM; n=9-10 tumours per genotype. (C) Pericyte association with blood vessels. The % of blood vessels with associated NG2-positive pericytes in B16F0 and LLC tumours grown in pdgfrβcre--;α6fl/fl mice was reduced significantly compared with pdgfrβcre--;α6fl/fl mice. Scatter graphs represent % blood vessels that are NG2-positive in B16F0 and LLC tumours+SEM; n= 4-6 mice/group. Representative images of tumour sections stained with the pericyte marker, NG2 (green) and endomucin (red). Arrows, endomucin positive staining; arrowheads, NG2 positive staining. (D) Blood vessel diameter. The frequency of vessels with a diameter ≥100 μm was greater in both B16F0 and LLC tumours grown in pdgfrβcre+;α6fl/fl mice when compared with pdgfrβcre--;α6fl/fl mice. Bar charts represent the % of blood vessels ≥100 μm diameter+SEM. (E) Blood vessel leakage. Mice were injected via the tail vein with Hoechst dye and tumour sections were analysed for blood vessel leakage by measuring the numbers of tumour cells that had taken up Hoechst. Blood vessels in the tumours grown in pdgfrβcre+;α6fl/fl mice showed...
significantly more leakage than blood vessels in tumours grown in \textit{pdgfrβcre-};\textit{α6fl/fl} mice. Bar chart shows relative Hoechst leakage+SEM; \textit{n}= 6 tumours/genotype. \textbf{Dotted lines}, Hoechst leakage. \textit{*p <0.05, **p <0.005, ***p <0.0009}. Scale bar in \textit{A}= 50 μm, \textit{C}= 100 μm and \textit{E}= 200 μm.

\textbf{Figure 2. Tumours grown in \textit{pdgfrβcre+};\textit{α6fl/fl} mice have aberrant BM organisation around blood vessels.}

LLC tumours were stained with antibodies to the extracellular matrix proteins (\textit{A}) collagen IV, (\textit{B}) fibronectin, (\textit{C}) laminin α5 and (\textit{D}) laminin α4 chains. For all matrices, disorganisation of basement membrane with a “shoreline” pattern was observed more frequently around blood vessels in tumours grown in \textit{pdgfrβcre+};\textit{α6fl/fl} mice when compared with \textit{pdgfrβcre-};\textit{α6fl/fl} mice. \textbf{(E)} The width of the BM surrounding blood vessels was analysed. Boxes show magnified regions of BM. \textbf{Brackets}, identify representative BM widths. \textit{n}= 8-10 sections/genotype; \textit{**p}<0.005. Scale bars = 50 μm.

\textbf{Figure 3. Tumour growth, angiogenesis and metastasis are not affected in \textit{pdgfrβcre+};\textit{α6fl/fl} mice.}

\textbf{(A)} Subcutaneous B16F0 and LLC tumour growth was similar in both \textit{pdgfrβcre+};\textit{α6fl/fl} mice and \textit{pdgfrβcre-};\textit{α6fl/fl} mice. Bar charts represent mean tumour volumes±SEM; \textit{n}=20-30 mice/genotype. \textbf{(B)} Endomucin staining of midline sections of age- and size-matched B16F0 and LLC tumours showed no significant differences in blood vessel density between \textit{pdgfrβcre+};\textit{α6fl/fl} mice and \textit{pdgfrβcre-};\textit{α6fl/fl} mice. Blood vessel density is given as the number of blood vessels/mm$^2$ of midline tumour section. Representative images of endomucin stained tumour blood vessels are shown. Bar chart represents mean tumour blood vessel density of size-matched tumours±SEM; \textit{n}= 6 tumour sections/genotype, \textit{ns}=no significant difference. \textbf{(C)} The number of perfused tumour blood vessels was assessed by tail vein injection of PE-CD31 antibody before tumour excision in \textit{pdgfrβcre+};\textit{α6fl/fl} mice and
pdgfrβcre-;α6fl/fl control mice and comparing the numbers of CD31-positive vessels with numbers of endomucin stained vessels. B16F0 and LLC tumours grown in pdgfrβcre+;α6fl/fl mice had a similar number of functional PE-CD31 endomucin expressing tumour blood vessels compared to pdgfrβcre-;α6fl/fl littermate control mice. Bar charts represent the percentage of PE-CD31 perfused vessels over total number of endomucin positive blood vessels per midline tumour section from pdgfrβcre+;α6fl/fl mice and pdgfrβcre-;α6fl/fl mice + SEM; n=10 tumour sections per genotype, ns=no significant difference. Representative images of endomucin stained, PE-CD31 positive perfused blood vessels from B16F0 tumours are shown. (D) Metastasis is not affected in pdgfrβcre+;α6fl/fl mice. Subcutaneous LLC tumours were resected when they reached approximately 100 mm³. At 3 weeks post-resection mice were culled and lungs removed to assess metastasis. Lungs were fixed and the number of surface metastases was counted. There were no significant differences in the total number of surface metastases between pdgfrβcre+;α6fl/fl mice and pdgfrβcre-;α6fl/fl mice (bar chart). Lungs were then sectioned and stained with H&E. Measurement of individual metastatic nodule areas showed no difference between genotypes (scatter graph). Representative high power images of H&E stained lung sections show areas of metastasis (dotted line) (upper images) as well as low power images of metastatic lung tissue (arrows) (lower images). Scale bar in B= 100 μm, C= 50 μm, D= 50 μm (upper panel), 5000 μm (lower panel).

Figure 4. Pericyte α6-integrin regulates PDGFRβ expression

(A) Pericytes were isolated from pdgfrβcre-;α6fl/fl and pdgfrβcre+;α6fl/fl mice, and Western blot analysis performed to assess α6-integrin deletion. α6 deletion was only observed in pericytes from pdgfrβcre+;α6fl/fl mice. FACS analysis also confirmed loss of α6-integrin surface levels in pericytes isolated from pdgfrβcre+;α6fl/fl mice compared with pericytes isolated from pdgfrβcre-;α6fl/fl mice (line graph and equivalent data dot plot). Bar charts represent densitometric readings of Western blots, corrected for loading. N=3 separate
lysates/genotype. (B) Western blot analysis confirmed that WT and α6-null pericytes have similar levels of α3-integrin, a second laminin receptor, suggesting that there was no compensation of α3-integrin in the absence of α6-integrin. N=3 separate lysates/genotype. (C) Fibroblasts and endothelial cells were isolated from pdgfrβcre−;α6fl/fl and pdgfrβcre++;α6fl/fl mice and Western blot analysis performed to assess α6-integrin levels. α6-integrin levels were not affected in either cell type isolated from either pdgfrβcre−;α6fl/fl and pdgfrβcre++;α6fl/fl mice. (D) Western blot analysis revealed PDGFRβ protein levels were significantly reduced in primary α6-null pericytes. Bar chart shows densitometric values of PDGFRβ levels, corrected for loading. HSC70 was used as a loading control; n=3 separate lysates/genotype; *p <0.05; ns=no significant difference. (E) Immunostaining of PDGFRβ in α6-null pericytes in culture was significantly reduced compared with WT pericytes. Scale bar in E = 50 μm.

Figure 5. Responses to PDGF-BB are reduced in α6-null pericytes

(A) Aortic rings were isolated from pdgfrβcre−;α6fl/fl and pdgfrβcre++;α6fl/fl mice and treated with either PBS or PDGF-BB (30 ng/ml) for up to 10 days. PDGF-BB stimulation increased vessel sprouting in pdgfrβcre−;α6fl/fl but not pdgfrβcre++;α6fl/fl aortic rings. Bar charts show average number of aortic sprouts/genotype+SEM; n=40-50 rings/genotype in triplicate. (B) Confluent monolayers of primary WT and α6-null pericytes were wounded manually, after 24 h serum starvation, then stimulated, or not, with PDGF-BB (30 ng/ml). Wound width was quantified up to 48 h. Wound closure was significantly reduced in α6-null pericytes in response to PDGF-BB. Bar charts show wound width in the absence (left panel) or presence (right panel) of PDGF-BB, normalised to time 0; n=6/genotype/time point in triplicate. (C) Proliferation of primary WT and α6-null pericytes in the presence of Optimem or PDGF-BB was measured. α6-null pericytes proliferated significantly less in the presence of PDGF-BB compared with WT pericytes. Graphs represent relative proliferation; n=3
biological repeats. (D) Western blot analysis of phosphorylated-ERK1/2, total ERK1/2, phosphorylated-AKT and total AKT, from WT and α6-null pericyte lysates after serum starvation and stimulation with PDGF-BB for 0, 30 s, 60 s, 5 min, 15 min, 30 min. Downstream signalling responses to PDGF-BB in α6-null pericytes were reduced significantly. Individual cropped blots are representative of samples run on the same gel under identical experimental conditions. HSC70 acted as the loading control. (E) Graphs show densitometric readings of fold change of ratios p-AKT/totalAKT to time 0; fold change of ratios p-ERK1/2/totalERK1/2 to time 0±s.e.m. N=3 independent experiments. *p <0.05, **p <0.005.
SUPPLEMENTARY INFORMATION

Dual role of pericyte α6β1-integrin in tumour blood vessels

Louise E. Reynolds¹, Gabriela D’Amico-Lago¹, Tanguy Lechertier¹*, Alexandros Papachristodoulou², José M. Muñoz-Félix¹, Adèle De Arcangelis³, Marianne Baker⁴, Bryan Serrels⁴ and Kairbaan M. Hodivala-Dilke¹

¹Adhesion and Angiogenesis Laboratory, Centre for Tumour Biology, Barts Cancer Institute- A CRUK Centre of Excellence, Queen Mary University of London, Charterhouse Square, London, UK. EC1M 6BQ.

²Laboratory for Molecular Neuro-Oncology, Dept. of Neurology, University Hospital Zurich, Frauenklinikstrasse 26, CH-8091 Zurich, Switzerland.

³IGBMC, UMR 7104, INSERM U964, Université de Strasbourg, BP. 10142, 1, Rue Laurent Fries, 67404 Illkirch Cedex, France.

⁴Cancer Research UK Edinburgh Centre, University of Edinburgh, Crewe Road South, Edinburgh, UK. EH4 2XR.

*These authors contributed equally

⁹Corresponding author: l.reynolds@qmul.ac.uk
## Supplementary Figure 1. Characterisation of pdgfrβcre-;α6fl/fl and pdgfrβcre+;α6fl/fl mice and morphological analysis of tissue and postnatal retinal angiogenesis.

(A) *Pdgfrβcre-;α6fl/fl* mice crossed to *pdgfrβcre+;α6fl/fl* mice produce litters that had a similar...
number of $\text{pdgf}^{\beta}\text{cre}^{-};\alpha 6^{fl/\text{fl}}$ and $\text{pdgf}^{\beta}\text{cre}^{+};\alpha 6^{fl/\text{fl}}$ mice in predicted Mendelian ratios, (B) and similar ratios of $\text{pdgf}^{\beta}\text{cre}^{+};\alpha 6^{fl/\text{fl}}$ males and females, identified at weaning. (C) Mice were weighed at 3 months old. No differences were observed in the weights of sex-matched mice between the two genotypes. (D) All mice were analysed by PCR genotyping and shows a product identifying $\text{pdgf}^{\beta}\text{cre}^{+}$ PCR at approx. 1000 bp. $\alpha 6$-integrin-floxed PCR shows products identifying homozygous $\alpha 6$-integrin floxed ($\alpha 6^{fl/\text{fl}}$) (154 bp), WT non-floxed ($\alpha 6$-integrin wt) (120 bp), and heterozygous ($\alpha 6$-integrin fl/wt) mice (154 bp and 120 bp). (E) Representative H&E stained sections of lung, heart, liver and spleen from 12 week old $\text{pdgf}^{\beta}\text{cre}^{-};\alpha 6^{fl/\text{fl}}$ and $\text{pdgf}^{\beta}\text{cre}^{+};\alpha 6^{fl/\text{fl}}$ mice. No gross morphological defects were observed. (F) Representative high power images of blood vessels in lung, heart and liver from $\text{pdgf}^{\beta}\text{cre}^{-};\alpha 6^{fl/\text{fl}}$ and $\text{pdgf}^{\beta}\text{cre}^{+};\alpha 6^{fl/\text{fl}}$ mice showed no obvious morphological blood vascular defects between the genotypes. (G) Confocal microscopy images show 3D Z-stack reconstructions of flat whole-mount immunofluorescent stained retinas from $\text{pdgf}^{\beta}\text{cre}^{-};\alpha 6^{fl/\text{fl}}$ and $\text{pdgf}^{\beta}\text{cre}^{+};\alpha 6^{fl/\text{fl}}$ mice at postnatal day (P) 9. Arterial, vein and capillary vessels in the superficial retinal plexus are visualised with Isolectin B4 (IB4, green). Pericytes are identified with NG2 (red); n=4 retinas/genotype. Scale bar in E=20 μm; F=100 μm; G=50 μm.
Supplementary Figure 2. mTmG reporter activity in pericytes after PDGFRβCre excision. mTmG mice are used as a reporter of Cre activity in Cre+ mice. We generated pdgfrβcre−;mTmG and pdgfrβcre+;mTmG mice. (A) Schematic representation of the mTmG reporter effect in pdgfrβcre+ mice. In non-Cre expressing cells mT (red) signal is observed in endothelial cells (EC). In Cre expressing cells the mT transgene is excised and the mG
(green) signal is observed in pericytes (PC). (B) B16F0 tumour cells were injected subcutaneously into \textit{pdgfr\textsuperscript{cre-};mTmG} and \textit{pdgfr\textsuperscript{cre+};mTmG} mice and tumours excised 12 days post inoculation. Tumour blood vessels were examined for expression of both membrane-targeted tandem dimer Tomato (mT) (red), seen in all host tissue and membrane-targeted green fluorescent protein (GFP) (mG) (green). GFP expression was observed only in \textit{pdgfr\textsuperscript{cre+};mTmG} mice, as expected and almost exclusively in PCs surrounding blood vessels (\textit{white box}, magnified region). (C) Tumour blood vessels immunostained with antibodies to \textit{\alpha6-integrin} and CD31 revealed co-expression of the 2 markers, showing that \textit{\alpha6-integrin} expression is not affected on endothelial cells \textit{in vivo}. Scale bars in B, C =50 \textmu m.
Supplementary Figure 3. Adhesion of wild-type and α6-null pericytes to Laminin-1 and immune cell infiltration in tumours.

(A) 5x10^4 wild-type (WT) and α6-null mouse brain pericytes were allowed to adhere to Laminin-1 (Lm-1; 10μg/ml) and Fibronectin (Fn; 10μg/ml) coated 96-well plates for 1 h at 37°C. α6-null pericytes do not bind effectively to the Lm-1 matrix compared with WT.
pericytes. No difference in Fn adhesion was observed between WT and α6-null pericytes. Bar chart represents % adhesion of pericytes to Lm-1 relative to adhesion to Fn for the same genotype. n=3 biological repeats; **p<0.01. Sections from tumours grown in pdgfrβcre-;α6fl/fl and pdgfrβcre+;α6fl/fl mice were stained with antibodies to inflammatory cell markers (B) CD8 (T cells), (C) CD45 cells and (D) F480 (macrophages). No difference in inflammatory cell infiltration was observed between tumours from either genotype. Bar charts represent % immune cell infiltration; n=6 tumours/genotype; ns= not significantly different. Scale bar =100μm.

Supplementary Figure 4 Reynolds et al
Supplementary Figure 4. Characterisation of primary mouse brain pericytes, primary mouse lung endothelial cells and fibroblasts and Reverse Phase Protein Array (RPPA) analysis reveals the mTOR signalling pathway is attenuated in α6-null mouse brain pericytes. (A) Primary mouse brain pericytes were characterised using the following markers, by FACS: PDGFRβ, CD146, CD31 (endothelial marker), glial fibrillary acidic protein.
Figure 1 Reynolds et al

A

B

C

Pericyte association (% blood vessels with pericytes)

B16F0

***

Pdgfr/cre-; α6fl/fl
Pdgfr/cre+; α6fl/fl

LLC

Pdgfr/cre-; α6fl/fl
Pdgfr/cre+; α6fl/fl

D

Tumour blood vessel diameter > 100μm (%)

B16F0

**

LLC

E

Tumour blood vessel leakage (AU)

B16F0

Pdgfr/cre-; α6fl/fl
Pdgfr/cre+; α6fl/fl
Figure 3 Reynolds et al

A

B16F0

LLC

Tumour volume (mm³)

Days

0 2 4 6 8 10 12

0 200 400 600 800

B

B16F0

LLC

Blood vessels (mm²)

Pdgfβ/cre-ακ6fl/fl

Pdgfβ/cre+ακ6fl/fl

ns

ns

Endomucin

C

B16F0

LLC

Perfused vessels (% PE-CD31 positive vessels)

Pdgfβ/cre-ακ6fl/fl

Pdgfβ/cre+ακ6fl/fl

ns

ns

Endomucin stained

PE-CD31 Ab perfused

Double-positive perfused blood vessels

D

Total number of lung metastasis per mouse

Pdgfβ/cre-ακ6fl/fl

Pdgfβ/cre+ακ6fl/fl

ns

ns

Individual metastatic nodule area (mm²)

Pdgfβ/cre-ακ6fl/fl

Pdgfβ/cre+ακ6fl/fl

High power

Low power
(GFAP) and Mac1 (fibroblast markers). Red graph = unstained cells, blue graph = stained cells. The results are shown as a histogram (left) and dot plot (right). *pdgfrβcre-;α6fl/fl* mouse brain pericytes were also immunostained for NG2 (pericyte marker) and visualised by epifluorescence microscopy. (B) Primary mouse lung endothelial cells were stained for CD31 followed by FACS analysis and immunostained for endomucin and vimentin (expressed by ECs) and α-sma (fibroblast marker). (C) Primary mouse lung fibroblasts were immunostained for vimentin (positive marker for fibroblasts), NG2 and endomucin (negative markers for fibroblasts). Rat IgG was used as a negative control. (D) RPPA analysis of protein extracted from WT and α6-null pericytes showed a significant reduction in many proteins in α6-null pericytes. These included components of the mTOR signalling pathway – AKT, p70S6kinase, mTOR, pS6 ribosomal protein and 4E-BP1; this pathway is important for several cell functions including growth, migration, proliferation and angiogenesis and is known to be activated by integrins.