Lamstatin – a novel inhibitor of lymphangiogenesis derived from collagen IV

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Abstract

The lymphatic system is essential for the maintenance of tissue homeostasis and immunity. Its dysfunction in disease (such as lymphangioleiomyomatosis) can lead to chylous effusions, oedema or dissemination of malignant cells. Collagen IV has six α chains, of which some of the non-collagenous-1 domains have endogenous anti-angiogenic properties, however, little is known about specific endogenous anti-lymphangiogenic characteristics. In this study we sought to investigate the expression levels of collagen IV non-collagenous-1 domains in lung tissue of patients with and without lymphangioleiomyomatosis to explore the hypothesis that a member of the collagen IV family, specifically the non-collagenous domain-1 of α5, which we named lamstatin, has anti-lymphangiogenic properties. Levels of lamstatin detected by immunohistochemistry were decreased in lungs of lymphangioleiomyomatosis patients. We produced recombinant lamstatin in an E.coli expression system and synthesized a 17-amino acid peptide from a theoretically identified, active region (CP17) and tested their effects in vitro and in vivo. Recombinant lamstatin and CP17 inhibited proliferation, migration and cord formation of human microvascular lung lymphatic endothelial cells, in vitro. Furthermore, lamstatin and CP17 decreased complexity and dysplasia of the tumour-associated lymphatic network in a lung adenocarcinoma xenograft mouse model. In this study we identified a novel, direct inhibitor of lymphangiogenesis, derived from collagen IV. This may prove useful for exploring new avenues of treatment for lymphangioleiomyomatosis and metastasis via the lymphatic system in general.

Keywords: Lymphangiogenesis • Lymphangioleiomyomatosis • collagen • type IV collagen alpha5 chain

Introduction

Lymphangiogenesis, the formation of new lymphatic vessels from pre-existing ones, has an essential role in the maintenance of tissue homeostasis and immunity. The recent elucidation of the role of the lymphatic network in the dissemination of malignant tumour cells [1] has highlighted the need for a greater understanding of the regulatory mechanisms involved. It is suggested that lymphangiogenesis is regulated by pro- and anti-lymphangiogenic factors maintaining a steady-state balance under normal conditions, however, in the presence of inflammation or disease, this balance is shifted towards pro-lymphangiogenic factors. Vascular endothelial growth factor (VEGF)-C and -D are the most prominent members of this group, but others such as basic fibroblast growth factor (bFGF) and angiopoietin (Ang)-1 have also been shown to induce lymphangiogenesis [2–5]. Vascular endothelial growth factor-C and -D bind to the tyrosine kinase receptor VEGFR-3 or Flt-4, which is highly expressed on lymphatic endothelial cells [6–10]. However, direct, tissue-derived inhibitors of lymphangiogenesis have received very little attention to date.
Collagen IV (Col IV) is an important component of the extracellular matrix (ECM), deposited as heterotrimers formed when any three chains of the six genetically different α chains (α1–α6) combine. Each α chain consists of a helical and a non-collagenous (NC)1 domain and all are present in the healthy lung [11]. Interestingly, the absence or decreased detection of members of the Col IV family, especially α5, is associated with higher levels of tumour cell invasion in adjacent tissue in breast, prostate, bronchoalveolar and colorectal cancer [12–16]. Some NC1 domains have been reported to have anti-angiogenic properties (e.g. α1, α2, α3) but others, such as α4, α5 and α6 have not yet been ascribed a function, especially in terms of lymphangiogenesis (reviewed in [17, 18]).

Lymphangioleiomyomatosis (LAM) is a rare (up to 5 per million) but progressive disease affecting predominantly women of childbearing age [19]. Lymphangioleiomyomatosis manifests itself either as part of the genetic disorder tuberous sclerosis (TSC) or occurs sporadically (~60% of cases), and demonstrates characteristics of metastatic disease [20–23]. The pulmonary component of LAM features alterations in the ECM, cystic lesions, as well as nodules consisting of and generated by abnormally proliferating and metastasizing smooth muscle-like cells (LAM cells) [24–26]. In addition, the number of lymphatic vessels is vastly increased in lungs from patients with LAM compared with lungs from individuals without LAM, and the increase of lymphangiogenesis in LAM patients is correlated with decreased survival rates [27]. Pro-lymphangiogenic factors, including VEGF-C (produced by LAM cells in vivo [27]), VEGF-D (elevated in vivo in patients with cystic fibrosis and those with bronchiectasis for collagen IV α3 [11]. Sections were deparaffinized and rehydrated through graded alcohol. Blocking serum (10% non-immune horse serum) was then added to the sections for 20 min. at room temperature. Without rinsing, either primary antibodies (collagen IV α1–6 NC1 a kind gift from Dr Sado at Shiget Medical Research Institute, Okayama, Japan (1 ng/ml) [16]) or isotype control antibody (Rat IgG, Jackson Immunoresearch, West Grove, PA, USA (1 ng/ml)) was added to the sections and incubated for 1 hr at room temperature. Sections were then washed with PBS and a goat anti-rat fluorescein isothiocyanate (FITC) [MP Biomedicals, Solon, OH, USA (1 ng/ml)] conjugated secondary antibody was added, and incubated for 30 min. at room temperature. Following a rinse with PBS, slides were mounted using Vectashield mounting medium (Vector Laboratories, Burlingame, CA, USA). Images were taken on an Olympus BX51 (Olympus Australia, Mt Waverley, Australia) fluorescence microscope and captured using Leica imaging software IM1000 (Leica, Wetzlar, Germany) as described previously [11]. The presence and intensity of staining for the collagen IV α1-6 chain NC1 domains in the images from four controls and eight LAM patients was scored by three independent observers, who were blinded to the diagnosis of the participants. Images were scored as follows: 0: Absence of stain; 1: Weak and discontinuous staining; 2: Weak and continuous staining; 3: Strong and continuous staining. Scores from the three observers were averaged and the standard error of the mean calculated.

**Material and methods**

**Immunohistochemistry**

We studied paraffin embedded tissue sections of bronchial rings from patients with LAM and sections derived from macroscopically normal tissue distant from tumours following cancer resections which served as controls, in addition to tissues from patients free of respiratory disease. All tissues were stained for the six collagen IV isoforms (NC1 domain-specific antibodies). An additional group of LAM patients was stained for collagen IV α1, α2 and α5. Bronchial rings from people with cystic fibrosis and bronchiectasis were also stained for collagen IV α3 and α5. We have previously described the staining of the controls used in this study [11]. The controls for staining collagen IV α1–α6 were derived from macroscopically normal tissue taken from a location distant from the tumour following cancer resections. In addition, tissue was also harvested from endobronchial biopsies or lungs removed at transplantation from patients free of respiratory disease. We have also previously described the staining of tissues from patients with cystic fibrosis and those with bronchiectasis for collagen IV α3 [11]. Sections were deparaffinized and rehydrated through graded alcohol. Blocking serum (10% non-immune horse serum) was then added to the sections for 20 min. at room temperature. Without rinsing, either primary antibodies (collagen IV α1–6 NC1 a kind gift from Dr Sado at Shiget Medical Research Institute, Okayama, Japan (1 ng/ml) [16]) or isotype control antibody (Rat IgG, Jackson Immunoresearch, West Grove, PA, USA (1 ng/ml)) was added to the sections and incubated for 1 hr at room temperature. Sections were then washed with PBS and a goat anti-rat fluorescein isothiocyanate (FITC) [MP Biomedicals, Solon, OH, USA (1 ng/ml)] conjugated secondary antibody was added, and incubated for 30 min. at room temperature. Following a rinse with PBS, slides were mounted using Vectashield mounting medium (Vector Laboratories, Burlingame, CA, USA). Images were taken on an Olympus BX51 (Olympus Australia, Mt Waverley, Australia) fluorescence microscope and captured using Leica imaging software IM1000 (Leica, Wetzlar, Germany) as described previously [11]. The presence and intensity of staining for the collagen IV α1-6 chain NC1 domains in the images from four controls and eight LAM patients was scored by three independent observers, who were blinded to the diagnosis of the participants. Images were scored as follows: 0: Absence of stain; 1: Weak and discontinuous staining; 2: Weak and continuous staining; 3: Strong and continuous staining. Scores from the three observers were averaged and the standard error of the mean calculated.

**Gene cloning, expression and purification of recombinant protein**

The DNA for the NC1 domain of human Col4α5 (lamstatin, chromosome Xq22.3) was extracted from primary human lung endothelial cells. Briefly, cells were extracted from blood vessels dissected from human donor lungs as described previously [11], expanded in tissue culture medium containing 10% FBS, 10 μg/ml endothelial cell growth supplement, 20 U/ml heparin and 2% antibiotics, and total RNA was extracted using the NucleoSpin RNA II kit according to the manufacturer’s instructions (Macherey Nagel, Düren, Germany). Total RNA was transcribed to cDNA using hexamer primers (New England Biolabs, Ipswich, MA, USA) and Superscript III (Invitrogen, Carlsbad, CA, USA). The MMP cleavage site prediction tool (http://www.dmbr.ugent.be/prx/bioit2-public/SitePrediction/index.php) was used to identify the MMP2 cleavage site at the beginning of the NC1 domain in the collagen IV α5a sequence, and primers that recognized the corresponding gene sequence were designed. The cDNA was then amplified with the following primers: 5′-TTCCATATGGATTTCTTATTACA-3′ (forward), 5′-CGGG ATCCCTATGCTCTCTCTGGGA-3′ (reverse) with restriction sites for NdeI (35 sec., annealing at 60°C for 30 sec. and elongation at 72°C for 60 sec. The ampiclon (675 bp) was eluted from a 1.5% Agarose gel (Amresco, Cochran Solon, OH, USA) using a QIAEX II gel extraction kit (Qiagen, Doncaster, VIC, Australia) and cloned into pcDNAs/FRT/TOP10 (Invitrogen) according to the manufacturer’s recommendations. The vector was transformed into TOPO10 E.coli (Invitrogen) and streaked on agar plates with ampicillin (100 μg/ml) (Sigma-Aldrich, St. Louis, MO, USA). Colonies were picked, expanded and the inserts within the isolated plasmids were subject to sequencing (Supamac, Sydney, Australia). Positive clones were selected and archived for later use.
Lamstatin was then subcloned into pET15b (via BamHI and NdeI) and transformed into BL21 (DE3) (Bioline, Sydney, NSW, Australia) for expression. E.coli were grown overnight, and then expansion cultures were started with an inoculum of OD 0.1 and grown until they reached OD 0.5. Expression was then induced with 11.2 mg/l of isopropyl 1-thio-β-β-galactopyranoside (IPTG; Sigma-Aldrich) for 4 hrs and cells were pelleted thereafter at 4°C at 4000 x g for 20 min. Pellets were collected and washed twice with buffer A and then resuspended in buffer A (7.9 g/l Tris–HCl, 1.46 g/l EDTA, pH 7.5). Cells were then sonicated on ice for 50 cycles (4 sec. at 60% of max. amplitude and 6 sec. pause). The suspension was pelleted at 15,000 x g for 20 min, before washing with solubilization buffer 1 (1% Triton X-100 and 180.2 g/l urea). The supernatant (15,000 x g, 20 min.) was removed and inclusion bodies were incubated with solubilization buffer 2 (354.4 g/l guanidine, 10.3 g/l NaHPO₃ and 1.58 g/l Tris–HCl, pH 5.5) for 2 hrs at RT. Insoluble debris was spun down and the lysate was either purified via a Nickel-sepharose column (AmershamPharmacia, GE Healthcare, Rydalmere, NSW, Australia) or directly processed by dilution and ultra filtration (Amicon Ultra15, 10 kD; Millipore, Billerica, MA, USA). Purified protein was analysed on PAGE for purity (Coomassie Blue staining) and stored at −80°C for later use. The protein concentration was measured by UV (280 nm; NanoDrop, Wilmington, DE, USA) and bicinchoninic acid assay (Sigma, Sydney, Australia). CP17 was obtained from AusPep (Tullamaine, Victoria, Australia) in HPLC grade purity.

**Cells and media**

Human lung lymphatic endothelial cells (HMVEC-Ly) were purchased from Lonza (Basel, Switzerland) together with the EGM-2 MV BulletKit [composition: hEGF, Hydrocortisone, GA-1000 (gentamicin, Amphotericin-B), FBS (Foetal Bovine Serum), VEGF, hFGF-B, R3-IGF-1, Ascorbic Acid (Lonza)] for expansion. Human umbilical vein endothelial cells (HUVECs) were a kind gift from Dr Anthony Ashton at the Kolling Institute and Prof Jenny Gamble at the Centenary Institute, The University of Sydney. Human umbilical vein endothelial cells were cultured on gelatin-coated flasks in medium M199 containing sodium bicarbonate, non-essential amino acids, sodium pyruvate, 20% foetal bovine serum (FBS), 1% antibiotic–antimycotic mix, 50 μg/ml endothelial cell growth supplement (BD Bioscience, San Jose, CA, USA) and 50 μg/ml heparin (Sigma-Aldrich). Cells were used at passage 3–5. A549 cells originated from the ATCC (Manassas, Virginia, USA) and were grown in DMEM (Gibco, Invitrogen) with 10% foetal serum and standard antibiotic media. Primary human lung fibroblasts and airway smooth muscle cells were isolated and grown as previously described [11, 30–37]. The study was approved by the Ethics Review Committee of the South West Sydney Area Health Service, Royal Prince Alfred Hospital, St. Vincent Hospital and The University of Sydney human research ethics committee. Written informed consent was obtained from all participants.

**3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyl Tetrazolium bromide (MTT) cell viability assay**

Cells in medium were seeded at a density of 5000/well in a 96-well plate (Nunc, Thermo Fisher Scientific, Rockford, IL, USA) and 24 hrs later treated with lamstatin or CP17 as indicated. To ensure specificity of CP17, another collagen IV derived, blood vascular endothelium anti-proliferative peptide (T3) identified by Maeshima et al. [38, 39] was also tested. Cells were grown for 66 hrs after treatment, before MTT was added. After 6 hrs of incubation, MTT-formazan development was stopped with 10% SDS and 3.7 g/l HCl. OD was read at 570 nm (690 nm background reading) in a Spectramax M2 (Molecular Devices, Sunnyvale, CA, USA) spectrometer. Results were expressed as percentage of vehicle control. The vehicle for lamstatin was 292.2 ng/l EDTA/pH 3.5; that for the CP17 peptide was MilliQ water and for the T3 peptide we used 40% Acetonitrile with 0.1% trifluoroacetate.

**Trypan blue exclusion cell viability assay**

Cells were seeded and stimulated as described above. Instead of addition of MTT, cells were harvested by detaching them with 0.05% trypsin in 198 mg/l disodium EDTA for 5 min. and resuspended in fresh Hanks buffered saline solution. Twenty microlitres of cell suspension was then diluted 1:1 with trypan blue and loaded onto a counting chamber of a Nexcelom Cellometer™ AutoT4 (Nexcelom Bioscience, Lawrence, MA, USA). Samples were automatically counted and visually inspected for viable cells.

**Manual cell counts**

Human umbilical vein endothelial cells were seeded (1 X 10⁴ cells/cm²) in 12-well plates for 24 hrs before they were treated with lamstatin (2.5–10 μg/ml), vehicle (1 mM EDTA, pH 6.5) or CP17 (2.5–10 μg/ml) for 72 hrs. Cells were then rinsed with PBS, detached using trypsin and stained with trypan blue. Cells were then counted manually using a haemocytometer.

**Cord formation assay**

Matrigel (BD Bioscience) was prepared in 48-well plates according to the manufacturer’s recommendations. HMVEC-Lys were seeded at a concentration of 2 X 10⁴ cells/well and allowed to attach for 1 hr. The cells were then grown in complete EGM-2 MV in the presence or absence of lamstatin or CP17 (both 1.25, 2.5, 5, 10 μg/ml) or vehicle (292.2 ng/l EDTA/pH 3.5). Effects were observed 24 hrs after addition of lamstatin or CP17. Human umbilical vein endothelial cells were seeded on to gelretex (Invitrogen) coated 24-well plates for 2 hrs before being treated with lamstatin (10 μg/ml), vehicle or CP17 (10 μg/ml) for up to 18 hrs at 37°C, 5% CO₂. Images were taken using a Ti EIIp microscope (Nikon) with a constant magnification factor of 100× and the panorama setting on an Olympus CAMEA C-4000 camera (Olympus, Hamburg, Germany). The number of cords was counted from a 416 px x 333 px (=138.5 px²) area divided into nine segments of equal size, the cords per field were counted and the average number of cords per 1000 px² was calculated.

**Migration assay**

Subconfluent HMVEC-Ly was harvested after a 4 hrs starving period in EBM-2 MV (no bullet kit supplement, but 0.1% BSA) using non-enzyme detachment solution (Treviron, Gaithersburg, MD, USA), washed and the concentration adjusted to 4.0 X 10⁶ cells/ml in EBM-2 MV (0.1% BSA). A 24-well insert (BD Falcon™ FluoroBlok, BioCoat for Endothelial Migration, BD Bioscience) was brought to room temperature and the top well was filled with 250 μl of cell suspension with either vehicle control or 2.5, 5 or
10 μg/ml lamstatin or CP17. Cells were incubated for 1 hr at 37°C, before EBM-2 MV (complete), with or without treatment (see above), was added to the bottom well. The presence of the inhibitor in the top and bottom wells ensured a constant concentration over the course of the experiment. Cells were allowed 22 hrs to migrate before the filter was washed with HBSS twice and stained with Calcein (4 μg/ml in HBSS, Invitrogen Molecular Probes) for 90 min. Fluorescence was read in a Spectramax M2 (Molecular Devices, Sunnyvale, CA, USA) at 488 nm excitation and 520 nm emission. Experiments were repeated three times.

**Mouse model of angio- and lymphangiogenesis**

All animal experiments were approved by the provincial state office of southern Finland and carried out in accordance with institutional guidelines. NOD/SCID/gamma mice were injected in the ears with LNM35 AAV-EGFP-expressing tumour cells suspended in growth factor reduced Matrigel® with or without 10 or 100 μg/ml lamstatin, 10 or 100 μg/ml CP17 or respective vehicle (292.2 ng/ml EDTA pH 3.5). After 12 days of tumour growth, animals were killed and ears were collected. Whole mount staining of the ears was performed as described earlier [40]. Lymphatic vessels were stained with LYVE-1 (AlexaFluor 647; Invitrogen, Carlsbad, CA, USA), blood vessels were stained with ECAM-1 (BD Bioscience clone MEC13.3) and AlexaFluor 594. LNM35 tumour cells were detected using their green fluorescent protein (EGFP) expression. Confocal imaging (constant thickness and scanning intervals) enabled visualization of tumour-associated lymphatic networks. Analysis of the mean intensity (this value represents the mean intensity of the pixels inside the image boundaries) of LYVE-1 staining was performed with QuantiOne (Bio-Rad, Hercules, CA, USA). The number of branching points or loops was assessed as described by Shayan et al. [41]. In brief, confocal images of lymphatic vessels were loaded into Image J (www.sbweb.nih.gov/ij/) and a 5 × 5 grid was overlaid. Branches were defined as two clearly distinguishable vessels that separate out without rejoining. Loops were defined as small, circular vessel structures, which had to be in the same focal plane. Branches or loops were counted separately in each image. One image, created from one stack of 64 confocal images from a single location in the experimental ear was analysed per animal and mean values were calculated for each treatment group. For the lamstatin experiments we studied three animals in which no tumour cells were present, 11 which received vehicle alone, eight which received 100 μg/ml lamstatin, and nine which received 100 μg/ml lamstatin. Similarly, for experiments in which the effect of CP17 was examined we studied three animals in which no tumour cells were present, 11 which received vehicle alone, eight which received 10 μg/ml CP17 and six which received 100 μg/ml CP17. After 12 days tumour sizes were measured in all animals treated with either vehicle (292.2 ng/ml EDTA, pH 3.5), 10 or 100 μg/ml lamstatin. Tumour was assumed to be ellipsoid in shape and thickness, and width and length were measured using a micrometre calliper. Volume was calculated according to the formula for ellipsoid bodies and plotted in mm³.

**Statistical analysis**

Values were considered to be significantly different if P < 0.05. All calculations were performed with GraphPad Prism 5 (Macintosh version, GraphPad Software, San Diego, CA, USA). Non-parametric analysis of variance (ANOVA) with repeated measures was used (Kruskal–Wallis) with Dunn’s or Bonferroni’s post-tests or paired Student’s t-tests where appropriate.

**Results**

**Collagen IV α3 and α5 NC1 domains are reduced in LAM lung sections**

Paraffin embedded lung sections from LAM (n = 8) and control (n = 10) patients were immunohistochemically stained for all six isoforms of Coll IV. An additional 2 LAM patients were stained for collagen IV α1, α3 and α5. In Figure 1A, representative examples of LAM lung sections (top) are compared with well-defined control patients (which have been previously analysed for the expression of collagen IV α1–6 and published elsewhere in a separate study [11]) (middle). The isotype control is shown below the specific stains. Detection of tumstatin (the NC1 domain of collagen IV α3) or lamstatin (NC1 of collagen IV α5) staining was significantly reduced in the tissue from all LAM patients examined. However, these isoforms were present in the control patients. We did not identify any other Coll IV isoforms which were differentially stained.

Lymphangioleiomyomatosis (n = 8) and control (n = 4) sections were scored for the level of staining observed (Table 1). All LAM sections had reduced scores for the level of tumstatin observed (0.5 ± 0.16) as well as for lamstatin (0.56 ± 0.12) compared with the control sections. We have previously explored the role of the collagen IV α3 NC1 domain (tumstatin) in asthma [11], therefore in this study we chose to focus on the effects of the collagen IV α5 NC1 domain (lamstatin).

To test whether the absence of lamstatin was a feature of other chronic respiratory disorders, tissue sections from patients with cystic fibrosis (n = 4) and bronchiectasis (n = 1) were stained for col IV α3 and α5 and all were positive for tumstatin and lamstatin as shown in Figure 1B.

**Identification of the potentially active peptide in lamstatin**

We initially observed anti-proliferative activity of recombinant tumstatin (produced in our laboratory using a similar approach to that described here for lamstatin) and lamstatin on HMVEC-L-Ly (data not shown). To identify potentially active regions in lamstatin, we analysed the sequences of tumstatin and lamstatin for homology (using an approach similar to but not identical to that of Karagiannis et al. [17]) and a single and unique stretch of 17 aa with a molecular weight of 2022 Da was identified, which we termed as CP17. The aa sequence of CP17 is VCNFASRNDYSYWLSTP and it runs between aa 66 and 82 of CP17.

**Lamstatin and the consensus peptide CP17 reduce viability of proliferating HMVEC-L-Ly**

We treated adherent, subconfluent (approx 50%) and proliferating HMVEC-L-Ly cells from two healthy donors for 72 hrs with increasing concentrations of lamstatin or CP17. Lamstatin significantly reduced cell viability at concentrations of 5 and 10 μg/ml (maximal
reduction of ~25% for lamstatin) \((n = 5\) experimental repeats) (Fig. 2A). Conversely, lamstatin had no effect on the viability of primary lung fibroblasts \((n = 3\) non-LAM cell lines), primary airway smooth muscle cells \((n = 6\) non-LAM cell lines) and A549 epithelial cells \((n = 3\) experimental repeats) (Fig. 2A). CP17 produced a similar significant reduction in cell viability to lamstatin, at concentrations of 5 and 10 \(\mu g/ml\) \((n = 3\) experimental repeats) (Fig. 2B).

However, treatment with lamstatin had no effect on cell viability in growth factor-starved confluent HMVEC-LLys \((n = 6\) experimental repeats) (Fig. 2A). We also compared the effect of the previously published peptide T3 (one of the active regions of tumstatin) \([38]\) and found it had no effect on lymphatic endothelial cell proliferation \((n = 3\) experimental repeats) (data not shown).

To confirm that MTT cleavage is reduced as a consequence of decreased cell numbers, HMVEC-LLys were treated with vehicle or increasing concentrations of lamstatin and a manual trypan blue exclusion cell count was carried out. Lamstatin significantly reduced the total number of trypan blue negative cells over a period of 72 hrs (Fig. 2D) at concentrations of 2.5, 5 and 10 \(\mu g/ml\) \((n = 4\) experimental repeats).

### Lamstatin and CP17 reduce cord formation

To further characterize lamstatin’s anti-lymphangiogenic properties, we seeded HMVEC-LLy on Matrigel® and induced cord formation (an \textit{in vitro} equivalent of tube formation and a prerequisite for HMVEC-LLy neo-vascularization) with EBM-2 MV growth medium (Fig. 3A). Lamstatin significantly reduced the number of cords from 0.059 per 1000 px² to 0.028 per 1000 px² at a concentration of 5 \(\mu g/ml\) (~52% reduction, \(P < 0.05\)) and to 0.024 cords per 1000 px² at 10 \(\mu g/ml\) (~59% reduction, \(P < 0.05\)) \((n = 3\) experimental repeats) (Fig. 3B).

To support the notion that CP17 is the active region, the cord formation assay was repeated with concentrations of 1.25, 2.5, 5 and 10 \(\mu g/ml\) CP17. There were significant, comparable reductions in the number of cords after 24 hrs treatment with CP17 at concentrations of

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**Table 1** Levels of the six collagen IV \(\alpha\) chain NC1 domains observed in stained airway sections from lymphangioleiomyomatosis (LAM) and control individuals.

|       | \(\alpha1\) | \(\alpha2\) | \(\alpha3\) | \(\alpha4\) | \(\alpha5\) | \(\alpha6\) |
|-------|------------|------------|------------|------------|------------|------------|
| LAM (SEM) | 8         | 2.16 (±0.26) | 2.28 (±0.18) | 0.5 (±0.16) | 2.63 (±0.17) | 0.56 (±0.12) | 2.33 (±0.12) |
| Control (SEM) | 4         | 1.72 (±0.3) | 2.21 (±0.17) | 2.29 (±0.20) | 2.72 (±0.20) | 2.47 (±0.22) | 2.5 (±0.20) |

Scoring results obtained from three independent observers blinded to the patient diagnosis using the following scale: 0: Absence of stain; 1: Weak and discontinuous staining; 2: Weak and continuous staining; 3: Strong and continuous staining.
Lamstatin and CP17 are inhibitors of HMVEC-LLy migration

Effective induction of tissue lymphangiogenesis requires HMVEC-LLys to be able to migrate towards pro-lymphangiogenic gradients (e.g. of VEGF-C, VEGF-D, FGF-2 and other factors). We used full growth media (EBM-2 MV) as a chemoattractant, which is supplemented, with FGF-2 (see methods) amongst other agents. Human lung lymphatic endothelial cells exposed to increasing concentrations of either lamstatin or CP17 exhibited significantly reduced migration from 100% with no treatment to 25% with 10 μg/ml of lamstatin ($P < 0.01$) and from 100% to 13% for 10 μg/ml CP17 ($P < 0.01$) ($n = 3$ and three experimental repeats respectively) (Fig. 3C).
Lamstatin and CP17 inhibit HUVEC proliferation and cord formation \textit{in vitro}

We treated proliferating HUVECs for 72 hrs with lamstatin and manually counted using trypan blue exclusion \((n = 6\) experimental repeats) (data expressed as average number of cells/ml). Subconfluent HUVECs were treated for 72 hrs with lamstatin or CP17 and cell viability, measured with MTT, was expressed as percentage of vehicle control \((%\text{ VC}) (n = 6\) and \(n = 6\) experimental repeats). All data are expressed as mean \pm SE of the mean (SEM) and compared using one-way repeated measures ANOVA (Bonferroni correction for multiple comparisons) of various concentrations versus no treatment \(*P < 0.05\). HUVECs were seeded on geltrex-coated plates for 2 hrs before being treated with lamstatin \((10\ \mu\text{g/ml})\), vehicle or CP17 \((10\ \mu\text{g/ml})\) for up to 18 hrs. Cord formation in the presence of lamstatin \((n = 4\) experimental repeats) or CP17 \((n = 4\) experimental repeats) are expressed as mean \pm SE of the mean (SEM) of cords per 1000 square pixel \((px^2)\) and compared using paired student’s \(t\)-test, \(*P < 0.05\).

**Lamstatin and CP17 are potent inhibitors of tumour-induced lymphangiogenesis in a murine model of lymphatic dysplasia**

To address the hypothesis that lamstatin and its anti-lymphangiogenic component CP17 are inhibitors of lymphangiogenesis \textit{in vivo}, we used a LNM35 adenocarcinoma xenograft model which is known to have tumour-associated angiogenesis and lymphangiogenesis \cite{40}.
Lamstatin and CP17 inhibit lymphangiogenesis in a murine model of tumour-induced lymphangiogenesis. LNM35 tumour cells, expressing EGFP, were injected intradermally into ears of NOD/SCID/gamma mice with or without lamstatin or CP17 and Matrigel. No tumour = 200 μl Matrigel bolus only. Vehicle = tumour cells, Matrigel and vehicle (292.2 ng/l EDTA, pH 3.5). Lamst = Lamstatin either 10 or 100 μg/ml in a 200 μl bolus of LNM35 and Matrigel. CP17 = CP17 10 or 100 μg/ml in a 200 μl bolus of LNM35 and Matrigel. Images of representative staining for treatment of tumour-induced lymphangiogenesis with lamstatin and CP17 (lamstatin: no tumour n = 3, vehicle n = 11, 10 and 100 μg/ml lamstatin, n = 8 and n = 9 mice; CP17: no tumour n = 3, vehicle n = 11, 10 and 100 μg/ml CP17, n = 8 and n = 6 mice respectively). White layer = LYVE-1, lympatics; Green layer = LNM35 tumour; Red layer = PECAM-1 (CD31), blood vasculature. Scale bar represents 200 μm.

Animals were given a single administration of lamstatin or CP17 together with the LNM35 AAV-EGFP tumour cells and subsequent tumour and associated lymphatic and blood vessel network development were observed by confocal microscopy (lamstatin experiment no tumour n = 3, vehicle n = 11, 10 and 100 μg/ml lamstatin n = 8 and n = 9, CP17 experiment no tumour n = 3, vehicle n = 11, 10 and 100 μg/ml n = 8 and n = 6 mice respectively) (Fig. 5). Lamstatin produced an overall significant reduction in the degree of lymphatic vascularization, as indicated by the decrease in overall LYVE-1 staining by ~60% (P < 0.05) (Fig. 6A). Lamstatin at a concentration of 10 μg/ml reduced the mean intensity of LYVE-1 staining from 11.3 intensity units (IU) (vehicle) to 4.3 IU (P < 0.05) and at 100 μg/ml to 4.6 IU (P < 0.05). More specifically, in the presence of lamstatin the lymphatic network dysplasia was reduced and the vessels appearance was less disorganized and more like that seen in the non-tumour tissue (Fig. 5, top row: lamstatin 10 compared with lamstatin 100 or vehicle and no tumour control). The tumour cells (LNM35 AAV-EGFP) (Fig. 5, second row), visualized in green with EGFP expression, clustered together to form a mass, whereas those of the lymphatic or blood lineages were more organized, lining themselves up into vessel structures. In the matrigel plugs containing tumour cells, treatment with lamstatin or CP17 had no effect on the tumour cells. However, lamstatin at the highest concentration (100 μg/ml) significantly reduced both the number of loops (P < 0.05) (Fig. 6B) and the number of branches (P < 0.05) (Fig. 6C) (indications of tumour-induced lymphangiogenesis) to levels observed in no-tumour, control animals. Consistent with our hypotheses, CP17 also proved to be a potent inhibitor in this model (Fig. 5). CP17 at the highest concentration (100 μg/ml) significantly reversed the increased numbers of loops (P < 0.05; Fig. 6B) and branches (P < 0.05; Fig. 6C) to levels observed in healthy animals.

In contrast, lamstatin had no significant effect on the fluorescence intensity (IU) of the blood vessels at either 10 or 100 μg/ml (PECAM-1, red staining in Fig. 5 and analysis in Fig. 6D). CP17 also did not change the mean fluorescence intensity (IU, PECAM-1, red staining in Fig. 5) of the blood vessels at either 10 and 100 μg/ml, compared with the vehicle control (Fig. 6D). To exclude the possibility that lamstatin indirectly inhibited lymphangiogenesis by reducing the initial tumour mass, cell viability of LNM35 tumour cells in vitro (via the MTT assay) was measured. No significant effect of lamstatin on tumour cell viability was detected (data not shown). To support this result, we measured the in vivo tumour sizes of treated animals and no effect of either 10 or 100 μg/ml lamstatin treatment was detected when we compared matrigel plugs with tumour + vehicle and those with tumour + lamstatin (Fig. 6E).

**Discussion**

In this study we report that the Col IV a5 NC1 (which we have named lamstatin) is significantly reduced in lung tissue sections of LAM patients and has novel anti-lymphangiogenic properties. In vitro, lamstatin and the 17 aa functional peptide (CP17) reduced migration of lymphatic endothelial cells. Furthermore, in vivo, in a murine model of tumour-induced lymphangiogenesis, lamstatin and CP17 dramatically reduced lymphangiogenesis. Thus, we identify lamstatin as a direct inhibitor of lymphangiogenesis, the absence of which may profoundly influence lymphangiogenesis and lymphatic endothelial cell survival.

Reports of endogenous lymphangiogenic inhibitors are few in number. Recently, Albuquerque and colleagues identified a soluble VEGFR-2 monomer, able to capture and deplete VEGF-C, which is
responsible for the lack of lymph vessels observed in the cornea of mice and possibly human beings [42]. The angiostatic proteins 16K human prolactin and collagen IV a2 non-collagenous-1 (NC1) domain (canstatin) have also now been described as inhibitors of lymphangiogenesis [43, 44]. Shao and Xie showed that endostatin, a proteolytic fragment of the NC1 domain of Col XVIII directly reduced the viability, in vitro, of pig thoracic duct lymphatic endothelial cells. In addition, endostatin directly interfered with the migration of lymphatic endothelial cells in an in vitro assay [45] and inhibited lymphatic tumour growth and lymphangiogenesis in a Lewis lung carcinoma xenograft [46]. In contrast, Brideau and colleagues showed an indirect anti-lymphangiogenic effect of endostatin in vivo [47]. Endostatin decreased in vitro and in vivo migration of tumour-associated mast cells, a substantial source of pro-lymphangiogenic VEGF-C. The effects of endostatin, 16K human prolactin and canstatin were not limited to the lymphatic system as they also

Fig. 6 Lamstatin and CP17 inhibit lymphangiogenesis in a murine model of tumour-induced lymphangiogenesis. LNM35 tumour cells, expressing EGFP, were injected intradermally into the ears of NOD/SCID/gamma mice with or without lamstatin (white bars) or CP17 (black bars) and Matrigel. (A) Mean intensity values of LYVE-1 fluorescence staining in lamstatin and CP17 treated animals (Lamstatin vehicle n = 11, 10 and 100 µg/ml lamstatin n = 8 and n = 9, CP17 vehicle n = 11, 10 and 100 µg/ml CP17 n = 8 and n = 6, mice respectively). *P < 0.05 one-way ANOVA (Kruskal–Wallis, with Dunn’s multiple comparisons corrections) treatment versus vehicle. (B) Assessment of loops in tumour ear model of lamstatin-treated animals (no tumour n = 3, vehicle n = 11, 10 and 100 µg/ml lamstatin, n = 8 and n = 9 mice respectively) and CP17-treated animals (no tumour n = 3, vehicle n = 11, 10 and 100 µg/ml CP17, n = 8 and n = 6 mice respectively). (C) Morphological analysis of branching points of lymphatic vessels in whole mount staining of LNM35 tumours with lamstatin (no tumour n = 3, vehicle n = 11, 10 and 100 µg/ml lamstatin, n = 8 and n = 9 mice respectively) or CP17 (no tumour n = 3, vehicle n = 11, 10 and 100 µg/ml CP17, n = 8 and n = 6 mice respectively) treatment; (D) Percentage of mean PECAM-1-positive blood vessel area (% mean PECAM-1 PVA) of animals treated with vehicle, lamstatin (vehicle (292.2 ng/l EDTA, pH 3.5) n = 11, 10 and 100 µg/ml lamstatin n = 8 and n = 9 mice respectively) or CP17 (vehicle (MilliQ water) n = 11, 10 and 100 µg/ml CP17 n = 8 and n = 6 mice respectively). All data presented as mean ± SEM, one-way ANOVA (Kruskal–Wallis, with Dunn’s multiple comparisons corrections) treatment versus no treatment, *P < 0.05. (E) Lamstatin has no effect on tumour size after 12 days. After 12 days tumour sizes were measured in animals treated with either vehicle (292.2 ng/l EDTA, pH 3.5), 10 or 100 µg/ml lamstatin. Data are expressed as mean ± SEM. Vehicle n = 11, 10 and 100 µg/ml lamstatin n = 8 and n = 9 mice.
decreased new blood vessel formation [47]. We did not identify significant effects of lamstatin on new blood vessel formation in vivo, however, in vitro inhibition of HUVEC proliferation and cord formation was evident to some extent. The reason for the disparity in our in vivo and in vitro findings may reflect the differences in endothelial cell types examined—HUVECS versus murine tumour-associated endothelial cells. Tumour endothelial cells are abnormal [48, 49], compared with non-diseased cells, and thus their response to lamstatin and CP17 may be different.

The absence or decreased levels of histochemical detection for members of the Col IV family, especially Col IV α5 as we have reported here in LAM, are associated with higher levels of invasion in breast, prostate, bronchoalveolar and colorectal cancer [12, 13, 50–52]. In addition, increased lymphangiogenesis is commonly found in those cancers (explicitly breast and prostate) and results in lymph node metastasis and invasion [14, 53–56]. It is therefore conceivable that lamstatin functions as a general inhibitor of lymphangiogenesis in the human body, and depletion could lead to the formation of new lymphatic vessels constituting a pro-lymphangiogenic environment and facilitating invasiveness and metastasis.

Lamstatin, and the consensus peptide CP17, had inhibitory effects on the individual mechanisms underlying lymphangiogenesis in vitro. Both molecules reduced the viability of proliferating but not quiescent HMVEC-LLy cells. This reduced cell viability was the result of a reduction in the number of viable cells, potentially due to the induction of cell death or through the inhibition of cell progression through the cell cycle. Conversely, lamstatin had no effect on the viability of primary lung fibroblasts, primary airway smooth muscle cells and αS5 epithelial cells, indicating that, within the cell types we tested, the effect is specific for HMVEC-LLy cells. Lamstatin and CP17 reduced HMVEC-LLy cord formation, an in vitro equivalent of tube formation, and a prerequisite for HMVEC-LLy neo-vascularization. In addition, lamstatin and CP17 inhibited the capacity of HMVEC-LLys to migrate towards a pro-lymphangiogenic gradient. This suggests that lamstatin interferes with several crucial lymphangiogenic properties of HMVEC-LLy. Furthermore, CP17 harbours a functional site of 17 or fewer amino acids which may be responsible for the effects seen with lamstatin, as these properties are also observed with the peptide.

Lamstatin and CP17 were effective at reducing lymphangiogenesis in our in vivo model. We found that lamstatin inhibited lymphangiogenesis, but had little or no effect on blood vascularization (angiogenesis) or tumour cell proliferation. The observed effects of lamstatin in this model were restricted to the dysplastic lymphatic network. We can exclude the possibility that lamstatin indirectly inhibited lymphangiogenesis via reducing the initial tumour mass because lamstatin did not alter the viability of the LNM35 tumour cells in vitro nor was tumour size in vivo related to the treatment in vivo. CP17 also proved to be a potent inhibitor of all of the indices of lymphangiogenesis measured in this model. These findings confirm our in vitro data and strongly suggest that lamstatin and its functional component CP17 are inhibitors of lymphangiogenesis in vitro and in vivo, even in the highly pro-lymphangiogenic environment such as that exists in the vicinity of a tumour.

Although LAM is not yet considered to be a malignant disease, LAM cells are highly mobile and metastasize, and it is suggested that dissemination of LAM cell clusters occurs via the lymphatic system [57] which is increased in patients with LAM [27, 57, 58]. In this study, lamstatin was undetectable in the airway tissue sections of all eight LAM patients we tested. Therefore, the absence of lamstatin in LAM lung tissue may facilitate the dissemination process because lymphatic vessels can easily be formed in the absence of an endogenous anti-lymphangiogenic factor.

In summary, we identified a novel inhibitor of lymphangiogenesis absent in lung tissue sections from patients with LAM. Lamstatin and CP17 decrease proliferation and migration of HMVEC-LLy in vitro and decrease tumour-induced lymphatic neo-vascularization in an in vivo model. Lamstatin may offer insights into the pathophysiology of LAM and may also provide a novel avenue for the treatment of diseases which are associated with excessive lymphangiogenesis, such as metastatic cancer.

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Conflict of interest

The authors confirm that there are no conflicts of interests.

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