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A role for whey acidic protein four-disulphide-core 12 (WFDC12) in the regulation of the inflammatory response in the lung

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What is the key question?
Does WFDC12 possess innate defence properties similar to other WFDC proteins in the lung such as SLPI and elafin?

What is the bottom line?
We have functionally characterised a member of the WFDC family of proteins, WFDC12, and have shown it to possess antiprotease and anti-inflammatory activity and to be present at elevated levels in the respiratory tract of patients with acute respiratory distress syndrome (ARDS).

Why read on?
Given its properties we speculate that WFDC12 may play a role in protecting lung tissue during inflammation and infection.

Author Contribution
AMAG SW AS DMS DTM and NC performed the experiments; UH COK DP DFM PMN JSE and FTL provided biological samples; SH carried out analysis; AMAG SW and CCT designed experiments and drafted manuscript. All the authors reviewed the manuscript and approved its submission.
SECRETORY LEUKOCYTE PROTEASE INHIBITOR (SLPI) and elafin are members of the whey acidic protein (WAP), or WAP four-disulphide-core (WFDC), family of proteins and have multiple contributions to innate defence including inhibition of neutrophil serine proteases and inhibition of the inflammatory response to lipopolysaccharide (LPS). This study aimed to explore potential activities of WFDC12, a previously uncharacterised WFDC protein expressed in the lung. Recombinant expression and purification of WFDC12 were optimised in *Escherichia coli*. Recombinant WFDC12 inhibited cathepsin G, but not elastase or proteinase-3 activity. Monocytic cells pre-treated with rWFDC12 before LPS stimulation produced significantly lower levels of the pro-inflammatory cytokines IL-8 and MCP-1 compared to cells stimulated with LPS alone. Recombinant WFDC12 became conjugated to fibronectin in a transglutaminase-mediated reaction and retained antiprotease activity. *In vivo* WFDC12 expression was confirmed by immunostaining of human lung tissue sections. WFDC12 levels in human bronchoalveolar lavage fluid from healthy and lung-injured patients were quantitatively compared, showing WFDC12 to be elevated in both patients with acute respiratory distress syndrome and healthy subjects treated with LPS, relative to healthy controls. Together these results suggest a role for this lesser known WFDC protein in the regulation of lung inflammation.
INTRODUCTION

Members of the whey acidic protein (WAP)\textsuperscript{5}, or WAP four-disulphide-core (WFDC), family of human proteins are defined by the presence of one or multiple WAP domains – a 40-50 amino acid sequence containing eight conserved cysteine residues that fold to form four disulphide bonds[1]. The most studied members of this group of proteins are secretory leukocyte protease inhibitor (SLPI) and elafin, both of which contribute significantly to the innate immune response via their antiprotease, antimicrobial and anti-inflammatory activities. As a result, they have each been involved in clinical trials as potential therapeutic agents to aid the control of inflammation[2,3].

The small molecular weight proteins SLPI and elafin are found constitutively expressed in lung epithelia and other mucosal surfaces, demonstrating upregulation in response to a range of pro-inflammatory stimuli such as bacterial lipopolysaccharide (LPS), neutrophil elastase, IL-1β and TNF-α[4]. They are also produced by inflammatory cells such as neutrophils and macrophages[4].

SLPI and elafin each function as inhibitors of elastase, with elafin also inhibiting proteinase-3[4]. SLPI shows no inhibition of proteinase-3 but instead is active against cathepsin G, trypsin and chymotrypsin[4]. Both proteins are bactericidal against Gram negative and Gram positive species of bacteria[5,6], and display notable antifungal[4,6] and antiviral effects[7,8]. SLPI and elafin also play important immunomodulatory roles through their ability to reduce the host response to LPS. This is understood to occur by both extracellular and intracellular mechanisms[9–12].

The role of most other WFDC proteins remains largely unknown. Our focus here is on WFDC12 (also known as WAP2). Without the predicted signal peptide, the mature 88 amino acid protein has an estimated molecular weight of 9.7 kDa and an isoelectric point of 5.77[13]. It has a N-terminal WAP domain and a C-terminal region showing no significant homology with any currently recognised motifs[13]. The WFDC12 gene is found highly expressed in lung, skin, prostate and oesophageal tissue[13]. Whilst the murine homolog
protein SWAM2 has been shown to have antibacterial properties[14], there has been no data on the function of human WFDC12 reported in the literature.

The primary aim of this research was to investigate if WFDC12 harboured host defence properties akin to SLPI and elafin. Recombinant expression and purification of WFDC12 in *E. coli* were first optimised, followed by evaluation of the protein’s antiprotease, antibacterial and immunomodulatory action. Recombinant WFDC12 (rWFDC12) was found to inhibit activity of the serine protease cathepsin G. No antibacterial ability was observed, but rWFDC12 inhibited LPS-induced cytokine production in THP-1 monocytic cells. Similar to SLPI and elafin, rWFDC12 was found to be capable of binding the extracellular matrix protein, fibronectin, and retained its antiprotease activity. Finally we show WFDC12 expression in human lung tissue via immunohistochemistry, and quantify WFDC12 levels in supernatants from LPS-treated THP-1 cells and bronchoalveolar lavage fluid (BALF) from a range of patient cohorts.
METHODS

Recombinant WFDC12 expression and purification
The mature WFDC12 cDNA sequence was cloned by Gene Art (Life Technologies, Paisley, UK) into a pQE30 vector (Qiagen, West Sussex, UK) with a six histidine coding sequence 5’ to the cloning region. His-tagged rWFDC12 protein was expressed and purified as detailed in the Online Supplement.

Protease activity assays
Cathepsin G activity was measured using the fluorogenic substrate Suc-Ala-Ala-Pro-Phe-AMC (100 μM; PeptaNova, Sandhausen, Germany) as outlined in the Online Supplement. Neutrophil elastase and proteinase-3 activity were measured using MeO Suc-Ala-Ala-Pro-Val-AMC (50 μM; Enzo Life Sciences, Exeter, UK).

Radial diffusion assays
The type strain bacteria Pseudomonas aeruginosa ATCC 27853, Staphylococcus aureus ATCC 25923 and E. coli ATCC 25922 were routinely maintained at 37°C on Colombia Blood Agar plates with 5% sheep blood (Southern Group Laboratories, Northamptonshire, UK). Double layer radial diffusion assays were carried out as described previously[15]. Bacteria were washed and resuspended in 10 mM sodium phosphate (pH 7.4) buffer before incorporation into agarose underlay gels based on the same buffer. Full details are given in the Online Supplement.

Cell culture experiments and cytokine ELISAs
Experiments using THP-1 cells and measurement of cytokine release were carried out as outlined in the Online Supplement.

Cross-linking of rWFDC12 to fibronectin and evaluation of antiprotease activity of fibronectin-bound rWFDC12
Transglutamination reactions were based on previously described methods[16,17], with minor modifications using rWFDC12 and fibronectin (see Online Supplement).
**Immunostaining of human tissue from an ex vivo lung perfusion (EVLP) model**

Lungs harvested from potential donors which were found to be unsuitable for transplantation were obtained from IIAM (International Institute for Advancement of Medicine). Ethical approval for the use of these samples was obtained from the Queen’s University Belfast School of Medicine, Dentistry and Biomedical Science Research Ethics Committee (Reference Number 12.17). The lungs were perfused *ex vivo* based on previously described methods[18]. Briefly, the lungs were initially rewarmed over a period of 1 hr by perfusion of the pulmonary artery using DMEM (high glucose, 4.5 g/L) containing 5% albumin, which was contained in a reservoir surrounded by a heated water bath (38°C). The lungs were gently inflated with continuous positive airway pressure (10 cmH2O, 95% O2, 5% CO2). Fresh whole human blood (anticoagulated using acid citrate dextrose) was added to the perfusate once physiological temperatures were reached, and LPS (6 mg) was then introduced intrabronchially into a pre-selected lobe via a PE catheter (BD, 240 tubing). Experimentation ceased 4 hr after LPS instillation, upon which tissue samples for histology were taken from both non-LPS (control) and LPS-perfused lobes. Immunostaining methods for WFDC12 and selected cell markers (epithelial cells: pan cytokeratin, macrophages: CD68, neutrophils: neutrophil elastase) are detailed in the Online Supplement.

**Human BALF collection and measurement of WFDC12**

WFDC12 levels were evaluated in BALF samples obtained from a number of different clinical trials: Samples from patients within 48 hours of ARDS onset (n = 27) were taken from the HARP study[19]. LPS-treated healthy subjects (50 μg LPS, BALF at 6 hr post LPS-inhalation, n = 10) and healthy controls (n = 5) were taken from a separate study[20]. Additional healthy controls (n = 6) were taken from the ongoing ARENA study. Ethical approval for the use of samples from the HARP (ISRCTN70127774), LPS (ISRCTN21056528) and ARENA (NCT01659307) studies was granted by the local institution and the local research ethics committee (REC numbers 06/NIR02/77, 06/NIR02/91 and 12/NI/0082 respectively). A description of the bronchoalveolar lavage procedure, sample processing and quantification of WFDC12 levels by ELISA is outlined in the Online Supplement.
Statistical analysis

All data were analysed using GraphPad Prism 5.0 (GraphPad Software Inc., San Diego, CA) and are reported as mean ± SEM. Results are representative of at least n = 3 unless otherwise indicated. Means were compared by unpaired t-test or one-way analysis of variance (ANOVA) as appropriate with Tukey’s post-hoc test. Statistical significance is indicated as: * P < 0.05, ** P < 0.01, *** P < 0.001.
RESULTS

**Purification of rWFDC12** – The rWFDC12 used for the experiments in this study was purified under denaturing conditions as outlined in the methods section with a typical yield of approximately 8 mg per litre of culture. SDS-PAGE assessment showed the final product contained some multimers but was mostly rWFDC12 in monomeric form (Fig. 1A). The level of purity was reproducibly high. Identity of the purified protein was confirmed by 1D nLC-MS-MS (reverse phase chromatography).

**Antiprotease activity of rWFDC12** – Since SLPI and elafin are known to be potent inhibitors of neutrophil serine proteases in the lung, we tested rWFDC12 for similar activity. Cathepsin G protease activity (213 nM) was inhibited by rWFDC12 in a dose-dependent manner (Fig. 1B), with almost complete inhibition by the highest rWFDC12 amount tested (9.1 μM). No significant inhibition of elastase or proteinase-3 activity by rWFDC12 was observed (data not shown).

**Antibacterial assays** – The WFDC proteins SLPI and elafin are reported to demonstrate notable antibacterial effects and so we investigated the ability of rWFDC12 to inhibit the growth of the type strain bacteria *P. aeruginosa* ATCC 27853, *S. aureus* ATCC 25923 and *E. coli* ATCC 25922. Concentrations of up to 72.7 μM of protein were tested via radial diffusion assays but did not exhibit antibacterial effects (Fig. 2). The antimicrobial peptide cecropin A was used as a positive control.

**Effect of rWFDC12 on LPS challenge in monocytic cells** – The previously documented ability of SLPI and elafin to inhibit LPS-induced production of cytokines in monocyctic cells led us to investigate if WFDC12 had similar properties. THP-1 monocytic cells were pre-treated with rWFDC12 for 1 hr before stimulation with LPS for 16 hr. Measurement of secreted cytokines via ELISA showed that cells pre-treated with rWFDC12 had significantly lower levels of IL-8 (Fig. 3A) and MCP-1 production (Fig. 3B), but not IP-10 (data not shown).

**Ability of rWFDC12 to bind fibronectin** – SLPI and elafin can each become cross-linked to extracellular matrix proteins, including fibronectin, in a transglutaminase catalysed reaction.
To examine if WFDC12 behaved in a similar manner, rWFDC12 was incubated with fibronectin and guinea pig liver transglutaminase for 2 hrs at 37°C. As observed by Western blot sample analysis, high molecular weight complexes (>250 kDa) appeared when rWFDC12 and fibronectin were incubated together in the presence of transglutaminase, but not when transglutaminase was absent (Fig. 4A). Accordingly, fibronectin was found depleted as visualised in a Colloidal blue stained gel (Fig. 4B). The heat-inactivated transglutaminase control reaction served to further certify that formation of the complexes was the result of specific enzymatic activity.

**Antiprotease activity of fibronectin bound rWFDC12** — In order to determine if fibronectin-conjugated rWFDC12 was still able to inhibit cathepsin G activity, rWFDC12 was added to fibronectin-coated wells of a 96-well plate, with/without transglutaminase. Unbound protein was washed away before addition of cathepsin G and fluorogenic substrate and the rate of substrate hydrolysis was monitored every 60 seconds for 1 hr. Wells containing transglutaminase and rWFDC12 showed approximately 26% less cathepsin G activity compared to wells with cathepsin G and transglutaminase in the absence of WFDC12 (Fig. 4C), indicating that immobilised rWFDC12 still functioned as an inhibitor. The level of inhibition was conceivably less than that observed with 5 μg of soluble rWFDC12 (Fig. 1B), since the cross-linking reaction is not wholly efficient. Wells containing rWFDC12 in the absence of transglutaminase showed no inhibition of cathepsin G activity (Fig. 4C), confirming specificity of the transglutaminase catalysed reaction.

**Immunostaining of WFDC12 in human tissue sections** — Human lungs rejected for transplant were perfused with LPS as outlined in the Methods section. Tissue samples from non-LPS (control) and LPS-perfused lobes were used for immunohistochemistry. Sections incubated with WFDC12 antibody showed positive staining (not detectable with isotype control) which correlated with epithelial, macrophage and neutrophil staining patterns (Fig. 5). WFDC12 expression was more apparent in tissue from LPS-perfused lung lobes due to increased infiltration of immune cells.

**Quantification of WFDC12 in THP-1 supernatants and human BALF samples** — The presence of a signal peptide suggested that WFDC12 could be secreted from cells; therefore we used
an indirect ELISA for measurement of WFDC12 in cell-free supernatants. THP-1 cells stimulated with LPS for 4 hr showed significantly increased WFDC12 secretion (Fig. 6A). Furthermore, we quantified WFDC12 levels in BALF from healthy patients and patients with lung injury. BALF from ARDS patients showed significantly increased WFDC12 relative to healthy controls (Fig. 6B). In addition, healthy volunteers administered LPS also showed significantly higher WFDC12 levels when compared to healthy controls (Fig. 6B).
DISCUSSION

SLPI and elafin are well characterised members of a family of proteins which each carry cysteine-rich motifs known as WFDC domains. Both SLPI and elafin have similar contributions to innate host defence via their inhibition of serine proteases, antimicrobial properties and dampening of the inflammatory response to LPS[4,21]. In order to investigate the role of WFDC12 in similar areas of host defence, we firstly optimised recombinant expression and purification of the protein, then tested it for a range of functions. We found antiprotease and immunomodulatory properties, but no antibacterial activity. Furthermore, the antiprotease action of rWFDC12 was preserved when it was immobilised to fibronectin.

Although SLPI is composed of two WAP domains, its antiprotease activity has been mapped exclusively to just one of these, the C-terminal WAP domain[22]. Elafin’s antiprotease activity has been proven to reside in its single WAP domain[23]. Recent scrutiny of WAP domains from human WAP proteins challenged the previous supposition that all of these proteins are protease inhibitors[21]. Spacing of the eight conserved cysteine residues in the domain was identified as fundamental in the determination of inhibitory potential[22,23]. The inhibitory active sites of the SLPI and elafin domains lie between cysteines 2 and 3, which are separated by eight amino acids. Among the other aligned WAP domains, this space was sometimes just three residues, leading to the proposal that, upon folding, these proteins would not contain adequate space for an inhibitory active site[21]. However, the spacing within this area of WFDC12 is relatively large (seven residues), leading us to hypothesise that WFDC12 could also, like SLPI and elafin, harbour antiprotease properties. Indeed, our data confirms rWFDC12 as an inhibitor of the neutrophil serine protease cathepsin G. We speculate that the specificity of WFDC12 towards inhibition of cathepsin G, but not elastase or proteinase-3, is due to the presence of a phenylalanine residue in the putative inhibitory active site. This is a relatively large, hydrophobic amino acid which may not fit the smaller active sites of elastase or proteinase-3.

The antimicrobial activities of SLPI and elafin are associated with their cationic nature (net charges of +12 and +3 respectively) which facilitates them to disrupt the anionic bacterial
cell membrane[6]. However, antimicrobial peptides/proteins may also be anionic and function via alternative mechanisms[24]. Of particular interest to us, the mouse protein SWAM2 (homolog of WFDC12), is known to be anionic and anti-bacterial[14]. We therefore looked for similar function of WFDC12 (also anionic) against selected type strain bacteria. No apparent antibacterial effects were observed with the concentrations of rWFDC12 tested, (up to 72.7 μM), however we cannot rule out the possibility that WFDC12 may be effective against other species of bacteria.

Elafin, its precursor trappin-2, and SLPI have each been shown to become cross-linked to various extracellular matrix proteins such as fibronectin and elastin via a transglutaminase-mediated reaction[16,17]. They are also known to retain their antiprotease effects whilst immobilised[16,17], and we have demonstrated that rWFDC12 follows suit. We hereby build upon the previously outlined theory that trapping of protease inhibitors by extracellular matrix proteins serves to protect tissue from excessive degradation by neutrophil serine proteases released during inflammation[16,17]. The combination of increased exposure of extracellular matrix proteins and increased transglutaminase activity observed under inflammatory conditions is presumed to amplify conjugation of these inhibitors[16].

Elafin and trappin-2 each contain one or multiple Gly-Gln-Asp-Pro-Val-Lys consensus sequence motifs, a recognised substrate for the transglutaminase enzyme[16]. The SLPI molecule lacks this transglutamination consensus sequence, yet also readily conjugates with fibronectin and elastin in a similar reaction[17]. Mass spectrometry was used to identify the transglutaminase-reactive Gln and Lys residues within each of these three proteins[17]. WFDC12 also lacks the aforementioned motif, but contains several Gln and Lys residues which could be responsible for the fibronectin-conjugation we have shown here.

We have also demonstrated the anti-inflammatory nature of rWFDC12 in its ability to inhibit LPS-induced cytokine production. Although the effect does not appear as potent as the classical examples, SLPI and elafin, we acknowledge the limitations of our recombinant protein production in E. coli. Furthermore, since LPS can activate many signalling pathways within cells, WFDC12 may target one pathway exclusively and not others, thereby limiting the degree of inhibition observed in cytokine release. With regards to the mechanism of
inhibition, it is possible that WFDC12 is complexing with LPS or IL-8/MCP-1 to reduce the
signal detected. However, considering that SLPI and elafin can function via an effect on
intracellular signalling molecules such as IκBα/β[9,12], we would anticipate that WFDC12
also has intracellular effects on LPS signalling.

We have shown in vitro that WFDC12 secretion is significantly increased in LPS-treated THP-
1 cells. WFDC12 was detectable in human clinical samples by both immunohistochemistry of
lung tissue sections and analysis of BALF via ELISA. The levels of WFDC12 in ARDS BALF were
slightly lower than, but still comparable to, that of SLPI and elafin. WFDC12 was detected at
13.6 (± 1.2) ng/ml, equivalent to 1.24 (± 0.11) nM; we have previously reported nanomolar
levels of SLPI (approximately 8.75 nM) and elafin (approximately 5.11 nM) in BALF from
ARDS patients[25]. The significantly elevated levels of WFDC12 (relative to healthy controls)
found in BALF from ARDS patients and healthy volunteers post LPS-inhalation may represent
an upregulation and/or increased secretion of WFDC12 from epithelial and immune cells as
part of the acute response to inflammation. Taken together, the results from this study
suggest that WFDC12 may offer protective effects to the host via its antiprotease and anti-
inflammatory properties.

An imbalance between endogenous proteases and their respective inhibitors is understood
to be a major contributing factor to the pathology of various inflammatory lung diseases
including ARDS and cystic fibrosis (CF)[26,27]. Antiproteases have therefore become a focal
point for development of new treatments for diseases such as these, and are yielding
promising results. For example, recent studies with α1-antitrypsin augmentation have
demonstrated a reduction in protease activity, neutrophil number and pro-inflammatory
cytokine levels[28,29]. SLPI is another attractive therapeutic candidate given its additional
antimicrobial/anti-inflammatory properties, and delivery of aerosolised SLPI showed
favourable results in CF trials[2]. However, deposition of the protein to poorly ventilated
areas of the CF lung was later shown to be not wholly efficient[30]. In addition, it is now
known that both SLPI and elafin are prone to proteolytic cleavage by the excessive levels of
proteases associated with dysregulated inflammation[25,31], perhaps limiting their full
therapeutic capacity. Therefore WFDC12 should ideally be examined for similar
susceptibility before investigation as a possible therapy.
Further exploration of WFDC12 properties is needed to determine its potential as a therapeutic agent in chronic inflammatory disorders of the lung. Given that elafin inhibits elastase and proteinase-3 but not cathepsin G, combination with WFDC12 would result in the targeting of all three neutrophil serine proteases and could prove more highly effective. Furthermore, delivery of WFDC12 as a fibronectin-conjugate may offer advantages of increased bioavailability and stability in an environment laden with proteases.
REFERENCES


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Competing interests: None

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The abbreviations used are: AMC, 7-amino-4-methylcoumarin; ARDS, acute respiratory distress syndrome; BALF, bronchoalveolar lavage fluid; CF, cystic fibrosis; EVLP, ex vivo lung perfusion; HRP, horseradish peroxidase; IL-8, interleukin-8; IP-10, interferon gamma-induced protein 10; LPS, lipopolysaccharide; MCP-1, monocyte chemotactic protein-1; MeOSuc, methoxy succiny|; NE, neutrophil elastase; SLPI, secretory leukocyte protease inhibitor; TNF-α, tumour necrosis factor alpha; WAP, Whey Acidic Protein; WFDC, WAP Four Disulphide Core
FIGURE LEGENDS

FIGURE 1. A. A typical rWFDC12 preparation. Induced E.coli M15 were harvested and lysed in a denaturing buffer. Cleared bacterial lysate was allowed to bind Ni-NTA agarose at 4°C for 1-2 hr, then poured into a column and washed extensively. The eluted rWFDC12 was refolded via a step-wise dialysis protocol before analysis by Coomassie stained gel and Western blot (polyclonal rabbit anti-WFDC12 antibody). B. WFDC12 inhibits the activity of the serine protease cathepsin G. Cathepsin G (213 nM) was incubated alone or with rWFDC12 for 15 min before addition of a fluorogenic substrate (Suc-Ala-Ala-Pro-Phe-AMC). SLPI was used as a positive control. Results are expressed as percentage of control (untreated cathepsin G).

FIGURE 2. Radial diffusion assay testing potential antimicrobial activity of WFDC12. Bacteria were incubated with a range of rWFDC12 concentrations. Cecropin A and water were used as positive and negative controls respectively.

FIGURE 3. Investigation of potential anti-inflammatory activity of WFDC12. THP-1 monocytic cells were incubated for 16 hr with media alone (control), LPS or rWFDC12; or pre-treated with rWFDC12 for 1 hr before stimulation with LPS for 16 hr. A. IL-8 and B. MCP-1 secretion were measured by ELISA. Statistical significance is indicated as: * P < 0.05, ** P < 0.01, *** P < 0.001.

FIGURE 4. WFDC12 binds fibronectin via a transglutaminase mediated reaction. Recombinant WFDC12 (5 µg) was incubated with fibronectin (20 µg) and guinea pig liver transglutaminase (1 mU) in 200 mM Tris HCl pH 7.5, 20 mM CaCl₂ and 0.1 mM DTT for 2 hr at 37°C. Reactions were terminated upon addition of sample treatment buffer without reducing agent, boiled and subsequently analysed by SDS-PAGE. A. Samples were separated in both 7.5% and 15% gels, then assessed via Western blot with a polyclonal rabbit anti-WFDC12 antibody. The 7.5% gel (upper panel) was used to view cross-linked protein (>250 kDa) whilst unbound rWFDC12 was viewed on the 15% gel (lower panel). B. Samples were separated in a 7.5% gel and stained with Colloidal blue to allow visualisation of fibronectin
and rWFDC12-fibronectin complexes. C. WFDC12 complexed to fibronectin inhibits cathepsin G activity. Recombinant WFDC12 (4.5 μM) or buffer alone was incubated with/without guinea pig liver transglutaminase (1 mU) in fibronectin coated wells (20 μg/well) of a 96-well plate for 2 hr at 37°C. Unbound rWFDC12 was washed away before addition of cathepsin G (213 nM) and subsequent incubation for 15 min at 37°C. Fluorogenic substrate was then added and relative fluorescence recorded every minute for 1 hr. W = rWFDC12; F = fibronectin; Tg = transglutaminase; Hi = heat inactivated.

**FIGURE 5.** Expression of WFDC12 and cell specific markers in human lung tissue. Lungs deemed unsuitable for transplantation were perfused *ex vivo*, with LPS introduced to a selected lobe for 4 hr. Tissue samples from control and LPS-perfused lobes were formalin-fixed and paraffin-embedded for immunohistochemistry. Tissue sections were stained with rabbit anti-WFDC12, mouse anti-pan cytokeratin (epithelial cell marker), mouse anti-CD68 (macrophage marker) and mouse anti-neutrophil elastase (neutrophil marker). Representative images (n = 2) were taken at X 40 objective lens magnification. Brown stained cells indicate the localisation of positive staining. Scale bars shown represent 50 μm.

**FIGURE 6.** Quantification of WFDC12 in THP-1 supernatants and human BALF samples. A. THP-1 monocytes were stimulated with LPS for 4 hr and WFDC12 secretion measured by ELISA. B. WFDC12 was measured by ELISA in BALF samples from the following patient groups: Healthy controls (n = 11), healthy subjects sampled 6 hr following LPS inhalation (n = 10), and patients within 48 hr of ARDS onset (n = 27). Statistical significance is indicated as: * $P < 0.05$, ** $P < 0.001$. 
Figure 1

A  Coomassie stained gel  Western blot

37 kDa  trimer  15 kDa  monomer
25 kDa  dimer
20 kDa
15 kDa

B  Cathespin G activity (% of control)

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Figure 2

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Cecropin A (25 µM)
Figure 3

A

B

IL-8 (pg/ml)

MCP-1 (pg/ml)

0 455 0 22.7 45.5 90.9 455 nM WFDC12

+ LPS

+ LPS
Figure 4

A

WFDC12-fibronectin complexes →

WFDC12 →

- W W W W W

F F F F F

B

WFDC12-fibronectin complexes →

Fibronectin →

- W W W W W

F F F F F

C

Cathepsin G (% inhibition)

WFDC12 +Tg  WFDC12 -Tg
Figure 6

A

B

WFDC12 pg/ml

Control       LPS

Healthy      LPS      ARDS

WFDC12 pg/ml

0  5000  10000  15000  20000

0  1000  2000  3000  4000

*  ***