In this work, we compared the profile of proteins secreted by planktonic and biofilm cultures of *Pseudomonas aeruginosa* using two-dimensional difference gel electrophoresis (2D-DiGE). This revealed that a novel metzincin protease, Mep72, was secreted during biofilm growth. Subsequent Western blotting and reverse transcription-PCR (RT-PCR) analyses demonstrated that Mep72 was expressed only during biofilm growth. Mep72 has a tridomain structure comprised of a metzincin protease-like domain and two tandem carbohydrate-binding domains. Unlike the only other metzincin (alkaline protease; AprA) in *P. aeruginosa*, Mep72 is secreted through the type II pathway and undergoes processing during export. During this processing, the metzincin domain is liberated from the carbohydrate-binding domains. This processing may be self-catalyzed, since purified Mep72 autodegraded *in vitro*. This autodegradation was retarded in the presence of alginate (an extracellular matrix component of many *P. aeruginosa* biofilms). The expression of full-length *mep72* in *Escherichia coli* was toxic. However, this toxicity could be alleviated by coexpression of *mep72* with the adjacent gene, *bamI*. Mep72 and BamI were found to form a protein-protein complex *in vitro*. 2D-DiGE revealed that the electrophoretic mobility of several discrete protein spots was altered in the biofilm secretome of an *mep72* mutant, including type III secretion proteins (PopD, PcrV, and ExoS) and a flagellum-associated protein (FlfD). Mep72 was found to bind directly to ExoS and PcrV and to affect the processing of these proteins in the biofilm secretome. We conclude that Mep72 is a secreted biofilm-specific regulator that affects the processing of a very specific subset of virulence factors.

*Pseudomonas aeruginosa* is an opportunistic human pathogen and a major cause of chronic infections in individuals with cystic fibrosis (CF). Chronic *P. aeruginosa* infections have long been associated with a biofilm mode of growth, characterized by the formation of sessile microbial communities and the production of exopolysaccharide (1–4). Such infections are particularly difficult to eliminate due to reduced immune clearance and their high tolerance to antibiotic treatment (5, 6). However, the biofilm phenotype remains poorly defined. A number of proteomic and transcriptomic analyses have been employed to investigate the lifestyle changes associated with the transition from a planktonic growth mode to a biofilm growth mode (7–12). In spite of this, there remains little consensus on what defines a biofilm (13).

*P. aeruginosa* is also a secretor. Secreted virulence factors are partly responsible for causing the extensive tissue damage associated with *P. aeruginosa* acute infections (14). Proteinaceous virulence factors are exported through the numerous secretion systems encoded by the *P. aeruginosa* genome (15). Many of these secreted proteins are hydrolytic enzymes which include the type I-secreted alkaline protease, AprA, and various type II-secreted proteases, such as elastase (LasB), staphylolysin (LasA), and PvdS-regulated protease (PrpL). Other virulence determinants are secreted through the type III secretion system (T3SS). The T3SS has been proposed to inject effector proteins directly into the host cell (16, 17). The expression of the T3SS often correlates with severe disease and increased mortality (18, 19).

A pair of two-component sensor systems have been identified that reciprocally regulate T3SS expression and biofilm formation (20, 21). As a result, biofilm cells often are thought of as being less virulent than planktonic cells, and the formation of biofilms is thought to represent a commitment toward chronic infection (22). However, biofilms also have been associated with acute infections (23), and chronic infections do not necessarily involve biofilm formation (24). Moreover, the expression of T3S proteins has been detected in biofilms grown under certain conditions, suggesting that biofilm formation and T3S are not always necessarily mutually exclusive phenotypes (11, 25).

To the best of our knowledge, only a few studies have investigated whether specific secreted proteins are associated with the biofilm growth mode. To date, most previous studies have focused on the production of secreted proteins by planktonic cultures (26–29). However, one study by Toyofuku and colleagues examined the secreted proteins that associated with the extracellular polysaccharide matrix. These workers demonstrated that outer membrane vesicles commonly associate with the biofilm matrix, implying that these vesicles are core constituents of *P. aeruginosa* biofilms (30). Another study showed that three extracellular proteases (AprA, LasB, and PrpL) were upregulated by Ca²⁺ in a mucoid *P. aeruginosa* strain (FRD1) grown under continuous-flow conditions (31). These proteases were upregulated concomi-
orarily with the alginate biosynthetic gene, algD, in the presence of high levels of Ca\(^{2+}\) in the culture medium. A study by Purschke and colleagues compared the secretomes of *P. aeruginosa* and *Candida albicans* in mixed biofilms (32). Here, they demonstrated that the diversity of *P. aeruginosa* proteins secreted in mixed-culture conditions was lower than in single-culture conditions. However, some secreted proteins (such as the virulence factor ToxA and hemophore HasAp) were uniquely expressed in mixed biofilms but were not detected in monocultures. Furthermore, proteinaceous factors, such as the adhesin CdaA, have been shown to be upregulated under biofilm conditions (33). CdaA is thought to contribute to biofilm structural integrity by either cross-linking Psl polysaccharide and/or by tethering Psl to cells.

Recently, Balyimez et al. reported that the transcription of two uncharacterized *P. aeruginosa* open reading frames (ORFs), PA2782 and PA2783, is under the control of the cyclic AMP (cAMP)-responsive transcriptional regulator, Vfr, and that *Escherichia coli* cells expressing PA2783 displayed proteolytic activity (cAMP)-responsive transcriptional regulator, Vfr, and that expression plasmid [pQE80(\[pMal-P2X\]](mep72)] was constructed (34). They subsequently renamed PA2783 as Mep72, after the cells expressing PA2783 displayed proteolytic activity. Mep72 is thought to be a protease expressed in biofilms specifically affects proteins exported by the type III secretion machinery.

**MATERIALS AND METHODS**

**Strains and growth conditions.** Bacterial strains and plasmids used in this study are listed in Table S1 in the supplemental material. *P. aeruginosa* cultures were grown at 37°C in AGSY medium (56 mM alanine, 17 mM K\(_2\)HPO\(_4\), 86 mM NaCl, 100 μM CaCl\(_2\), 10 mM MgSO\(_4\), 5 μM FeCl\(_2\), 7.5 μM MnCl\(_2\), 0.5% [vol/vol] glycerol, 3 g/liter yeast extract, pH 7). *P. aeruginosa* forms robust biofilms in AGSY medium, and transcriptomic/proteomic data are available (11,35). Planktonic cultures were grown in flasks with vigorous shaking and harvested after 3 h (late exponential phase; optical density at 600 nm [OD\(_{600}\)] = 1) or 9 h (stationary phase; OD\(_{600}\) = 9). Continuous-flow biofilms were grown in AGSY medium as previously described (35). Planktonic cells were harvested in early stationary phase. Biofilms were grown in a continuous flow system (35), and extracellular proteins were harvested for proteomic analysis after 24 h by collecting ca. 150 ml of the tube effluent directly into a flask packed in ice. Culture supernatants were clarified by centrifugation (8,000 × g) and filtration (0.2-μm filter; Millipore Stericup). Proteins were precipitated by addition of trichloroacetic acid (10%, wt/vol, final concentration) and harvested by centrifugation (100,000 × g, 4°C, 45 min). Pellets were washed with 50% ethanol and resuspended in ASB14 buffer (35). Samples were analyzed by two-dimensional difference gel electrophoresis (2D-DIGE), and statistical analysis was performed using BVA (biological variance analysis) and HCA (hierarchical cluster analysis) as previously described (35). Master numbers were assigned by the DeCyder software and refer to the position of the protein spot on the master gel. For the proteomic comparison of the mep72 mutant and wild-type secretome, a single biological sample was analyzed using dye-swapped technical replicates, and the modulated protein spots were identified as described above. For the comparison of the mep72 mutant and wild-type secretome, protein spots were selected for mass spectrometry analysis based on whether they were present in one sample and absent from the other, as opposed to protein spots that were modulated between samples. The samples were analyzed by 2D-DIGE, and statistical analysis was performed using biological variance analysis and hierarchical cluster analysis. Protein spots were excised and digested for liquid chromatography-tandem mass spectrometry (LC-MS/MS) analysis, and the output data were used to search the *Pseudomonas aeruginosa* database using MASCOT. Protein identifications were considered significant only if the MASCOT score was >80 and they contained one or more peptides with an E value of <0.01. Hits comprising more than one protein identification were discarded from further analysis.

**Western blot analysis.** The protein concentration in samples was quantified spectrophotometrically using Bradford reagent (Sigma). Equal amounts of protein from cell-associated and secreted fractions were separated on 12% sodium dodecyl sulfate-polyacrylamide gels, transferred to polyvinylidene fluoride membranes, and analyzed by Western blotting. Eox5 and PcV primary antibodies were a gift from Arne Rieetsch, Case Western Reserve University. Polyclonal anti-Mep72 primary antibody was prepared from rabbits that had been immunized with a peptide co-

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**Protein extraction.** Planktonic cell cultures were sedimented by centrifugation (3,200 × g, 30 min, 4°C). Biofilm effluent was collected on ice, and planktonic cells were removed by centrifugation. The average number of planktonic cells in the effluents before clarification was 2 × 10^7 CFU/ml (in comparison, planktonic cultures in stationary phase had an average cell density of 2 × 10^10 CFU/ml). The mean protein concentration in the 1-day biofilm effluents was 0.94 ± 0.30 μg/ml. In comparison, the planktonic stationary-phase (PS) samples contained 2.32 ± 0.79 μg/ml protein. Supernatants from the biofilm and planktonic cultures were passed through a 0.2-μm filter (Millipore), and protein was precipitated by the addition of solid trichloroacetic acid to a final concentration of 12.5% (wt/vol). Protein was left to precipitate overnight at 4°C and was collected by sedimentation (3,200 × g, 30 min, 4°C). Pellets were washed three times in 50% ethanol and redissolved in 1-[(3-cholamidopropyl)-dimethylammonio]-1-propanesulfonate (CHAPS) buffer (6 M urea, 2 M thiourea, 10 mM Tris, 4% CHAPS detergent, pH 8.3). Intracellular proteins (for Western blotting) were prepared by normalizing cell pellets to the same optical density (OD\(_{600}\)) using TE buffer (50 mM Tris, 4 mM EDTA, pH 8.3) and sonication on ice. Cell debris was removed by centrifugation (3,200 × g, 30 min, 4°C).

**Proteomic analysis.** Samples were harvested and prepared for proteomic analysis essentially as described by Mikkelsen et al. (35), with minor modifications. Briefly, for the growth-mode secretome comparison, four biological replicates were analyzed for each growth condition. Planktonic cells and biofilms of *P. aeruginosa* PAO1 were grown in AGSY medium as previously described (35). Planktonic cells were harvested in early stationary phase. Biofilms were grown in a continuous flow system (35), and extracellular proteins were harvested for proteomic analysis after 24 h by collecting ca. 150 ml of the tube effluent directly into a flask packed in ice. Culture supernatants were clarified by centrifugation (8,000 × g) and filtration (0.2-μm filter; Millipore Stericup). Proteins were precipitated by addition of trichloroacetic acid (10%, wt/vol, final concentration) and harvested by centrifugation (100,000 × g, 4°C, 45 min). Pellets were washed with 50% ethanol and resuspended in ASB14 buffer (35). Samples were analyzed by two-dimensional difference gel electrophoresis (2D-DIGE), and statistical analysis was performed using BVA (biological variance analysis) and HCA (hierarchical cluster analysis) as previously described (35). Master numbers were assigned by the DeCyder software and refer to the position of the protein spot on the master gel. For the proteomic comparison of the mep72 mutant and wild-type secretome, a single biological sample was analyzed using dye-swapped technical replicates, and the modulated protein spots were identified as described above. For the comparison of the mep72 mutant and wild-type secretome, protein spots were selected for mass spectrometry analysis based on whether they were present in one sample and absent from the other, as opposed to protein spots that were modulated between samples. The samples were analyzed by 2D-DIGE, and statistical analysis was performed using biological variance analysis and hierarchical cluster analysis. Protein spots were excised and digested for liquid chromatography-tandem mass spectrometry (LC-MS/MS) analysis, and the output data were used to search the *Pseudomonas aeruginosa* database using MASCOT. Protein identifications were considered significant only if the MASCOT score was >80 and they contained one or more peptides with an E value of <0.01. Hits comprising more than one protein identification were discarded from further analysis.
responding to residues CGIGYMGSDDKNSGR of the metzincin domain (residues 126 to 140) of Mep72. Immunization and antibody preparation was carried out by BioGenes (Germany). Anti-His primary antibody was purchased from Cell Signaling Technology. Enhanced chemiluminescence peroxidase-labeled anti-rabbit IgG antibodies (Sigma) were used as secondary antibodies. Blots were developed using Immobilon Western chemiluminescent horseradish peroxidase (HRP) substrate (Millipore).

**Protein refolding.** Inclusion bodies of His<sub>6</sub>-Mep72 were harvested and solubilized in 50 mM Tris-HCl (pH 8) containing 300 mM NaCl, 15 mM imidazole, and 8 M urea. The solubilized protein solution was clarified by centrifugation (9,000 × g, 30 min, 20°C) and loaded onto a 2-ml packed bed volume nickel-nitritriacetic acid (Ni-NTA) column equilibrated in the same buffer. The column was loaded and washed at room temperature. The column was washed with ca. 200 ml equilibration buffer. Elution was done by increasing the imidazole concentration to 300 mM. For refolding, the eluted protein was rapidly diluted into a 20-fold volumetric excess of 50 mM Tris-HCl (pH 8) containing 200 mM NaCl. Where indicated, the refolding buffer was supplemented with sugars or polymers. MBP-tagged Mep72 was purified in a similar way. Inclusion bodies were harvested and solubilized in 8 M urea. The sample was then rapidly diluted by squishing it into a larger volumetric excess of rapidly stirred refolding/column buffer (20 mM Tris-HCl [pH 7.4] containing 200 mM NaCl and 1 mM EDTA) at 4°C. After 60 min, the sample was clarified by centrifugation and the soluble supernatant was loaded directly onto an amylose affinity resin. The column was washed overnight, and the bound protein was eluted in column buffer containing 10 mM maltose. Eluted fractions were pooled and concentrated by ultrafiltration (10,000 molecular weight cutoff; Millipore) in a centrifugal concentrator.

**Immunoprecipitation (pulldown).** His<sub>6</sub>-BamI and MBP-Mep72 were expressed in *E. coli* DH5<sub>α</sub> from pQE80(bam1) and pMal-P2X (mep72), respectively, in 1 liter of LB. Cells were grown with vigorous shaking at 37°C to mid-log phase, after which protein expression was induced by the addition of IPTG to a final concentration of 0.3 mM. Cells subsequently were grown at 25°C overnight with shaking. Cells were sedimented by centrifugation (4,000 × g, 20 min, 4°C), and the pellets were resuspended in buffer A (50 mM Tris, 200 mM NaCl, pH 8.0). The cells were lysed by sonication on ice. Both recombinant proteins formed insoluble inclusion bodies. The inclusion bodies were collected by sedimentation (9,000 × g, 30 min, 4°C) and solubilized in buffer A containing 8 M urea. Insoluble material was removed by centrifugation (9,000 × g, 30 min), and the proteins were refolded by rapid injection into a 16-fold volumetric excess of rapidly stirred buffer A. The dilute bait protein suspensions then were loaded onto the appropriate affinity matrix (Ni-NTA–agarose [Qiagen] to capture the His-tagged BamI and amylose resin [New England BioLabs] to capture the MBP-tagged Mep72). The columns were then washed to remove unbound bait protein before applying the target protein solution. (As a control, the target protein solutions also were run through Ni-NTA or amylose columns to which no bait protein had been preadsorbed.) After washing the columns to remove unbound target proteins, the bound proteins were eluted with either 300 mM imidazole (for the Ni-NTA affinity matrix) or 10 mM maltose (for the amylose affinity matrix).

For the ExoS and PcrV pulldown, *P. aeruginosa* planktonic cultures (eight 25-ml cultures) were grown to stationary phase, and the culture supernatants were passed through a 0.2-μm filter (Millipore). Secreted proteins were concentrated in a 10-kDa cutoff centrifugal concentrator (Millipore) to a final volume of 1 ml (3,200 × g, 4°C). The concentrated protein was diluted 20-fold in buffer A before being loaded onto the affinity capture matrix.

**qRT-PCR.** All cells were harvested directly into RNAlater (Ambion), kept at 4°C overnight, and stored at −80°C until use. The culture–RNAlater mixture was thawed on ice, and a suitable amount of cells was sedimented by centrifugation. Cells were resuspended in water containing 1 mg lysozyme ml<sup>−1</sup> (Sigma) and incubated for 15 min at room temperature. RNA was extracted using an RNeasy Mini purification kit (Qiagen) by following the manufacturer’s instructions. The resulting RNA (200 ng) was used as a template for reverse transcription and conversion into cDNA using Superscript II reverse transcriptase (Invitrogen) by following the manufacturer’s instructions. Quantitative real-time reverse-transcription PCR (qRT-PCR) was performed on the cDNA using 1× SYBR green PCR master mix (Applied Biosystems) with 10 pmol of the appropriate primers (see Table S2 in the supplemental material). Amplification was carried out using an ABI PRISM 7300 real-time PCR system, and fluorescence data were processed using SDS software (ABI). Relative gene expression was obtained using 16S rRNA as the control, with mRNA/16S rRNA from each sample set equal to 1. Three biological replicates were examined for each growth condition.

**RESULTS**

**Biofilm and planktonic secretome comparison by 2D-DiGE.** We wanted to test directly whether biofilms of *P. aeruginosa* secrete an altered spectrum of proteins compared with planktonic cells. Continuous-flow biofilms were grown in silicon tubing as previously described (35), and secreted proteins were harvested. The secretome obtained from PAO1 biofilms grown for 1 day (1dB) was compared with that harvested from a stationary-phase planktonic culture of PAO1 (PS). The secretome samples were analyzed by quantitative 2D-DiGE (Fig. 1). One hundred sixteen spots were modulated in the biofilm samples (93 upregulated, 13 downregulated; *P* ≤ 0.01). A selection of the most highly modulated spots were excised from the gels and identified by LC-MS/MS and Mascot searching of the *P. aeruginosa* database (Table 1). Several spots corresponding to known virulence factors were upregulated in the biofilm secretome, including LasA, PasP, and PcrV. Additional upregulated spots corresponded to SpuE, AotJ, PA1342, and PotF5, all of which are predicted to be involved in polyamine uptake. Polyamines have been shown to play a role in the tolerance of *P. aeruginosa* to some antibiotics (36–38), although the mechanism underlying this currently is unknown. Other upregulated proteins included the ferric iron receptor...
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Known virulence factors containing proteins, integrins, and adhesins (42, 43). However, a transport of small molecules suspect is reminiscent of the conserved sequence motifs (the zinc-binding motif and Met-turn) characteristic of metzincins (46). The bacterial metzincins, also known as serralysins, often are associated with virulence. Indeed, the only other metzincin encoded by the PAO1 genome, Metz1, is known as serralysin, often is associated with virulence.

| Function and PA no. | Master no. | Gene name | Gene product | Level of abundance | Modulated ratio <sup>b</sup> (biofilm/PS) |
|---------------------|-----------|-----------|--------------|-------------------|-----------------------------------------|
| Transport of small molecules | PA0301 1022 | spaE | Polyamine transport protein | ↑ | 19.8 |
| | PA0487 969 | hitA | Ferric-iron-binding periplasmic protein HitA | ↑ | 17.5 |
| | PA2592 1019 | pff5 | Probable periplasmic spermidine/putrescine-binding protein | ↑ | 33.9 |
| | PA1493 960 | cysP | Sulfate-binding protein of ABC transporter | ↑ | 6.4 |
| | PA1493 1019 | aotJ | Arginine/ornithine binding protein AotJ | ↑ | 17.9 |
| | PA1342 1155 | ppcD | Probable lysine/arginine/ornithine-binding protein | ↑ | 9.9 |

Modulated ratio represents the fold change in the biofilm secretome compared to that in the planktonic secretome.

Summary of the identified protein spots modulated in the biofilm-grown culture secretome compared with their activity in the planktonic culture secretome.

An uncharacterized protein, PA5033, also was upregulated in the biofilm secretome. PA5033 is predicted to form a bicistronic operon with its adjacent gene, PA2782 (Fig. 2A). To test this, we harvested RNA from PAO1 biofilms. The RNA was converted to cDNA and used as a template to PCR amplify across the junction between PA2782 and mep72 using primers P1 and P2 (Fig. 2B). If the genes are transcribed as a single contiguous transcript, a 500-bp product would be anticipated, and this is what was observed (Fig. 2B). No 500-bp product was observed in the no-RT control. We obtained no evidence that PA2782 is cotranscribed with the upstream gene (PA2781) or that mep72 is cotranscribed with the downstream gene (PA2784) (data not shown).

Pfam domain analysis of Mep72 revealed that the protein is comprised of three domains: an N-terminal metalloprotease domain and two tandem C-terminal carbohydrates (Fig. 2C). The amino acid sequence of the N-terminal domain was analyzed using the sequence-structure homology recognition program FUGUE (44). This analysis suggested that Mep72 is structurally related to snake venom metalloproteases (adamalysins) and snake venom endothelial cell vascular apoptosis-inducing protein (VAP-1), which are members of the metzincin superfamily of zinc-dependent metalloproteases (45, 46). Further inspection of the primary structure of Mep72 revealed that it contained both of the conserved sequence motifs (the zinc-binding motif and Met-turn) characteristic of metzincins (46). The bacterial metzincins, also known as serralysins, are often associated with virulence. Indeed, the only other metzincin encoded by the PAO1
Expression of Mep72 is upregulated in continuous-flow biofilms. In order to determine the transcriptional expression profile of mep72, qRT-PCR was carried out on RNA harvested from biofilm cells (1dB) grown in a continuous-flow setup and from cultures of planktonic cells (Fig. 3A shows the data for exponentially growing planktonic cultures [PE]; essentially the same results were obtained for comparisons of stationary-phase planktonic cultures with biofilm cultures [data not shown]). mep72 transcript levels were 18-fold upregulated in 1dB cells compared with the level for planktonic cells. This is consistent with the notion that bamI and mep72 are functionally linked.

To further verify that Mep72 expression is biofilm specific, secreted proteins from biofilm and planktonic cultures were extracted, and the expression of Mep72 was monitored by Western blotting with an antibody raised to the metzincin domain of Mep72 (Fig. 3B). Mep72 was undetectable in the secretome of aerobically grown PE and PS planktonic cultures but was abundant in continuous-flow biofilm supernatants. No Mep72 was detected in the biofilm secretome of an mep72 mutant (Fig. 3B). These transcriptomic and proteomic analyses indicate that Mep72 is a biofilm-associated gene product under these growth conditions.

Mep72 is secreted via the type II secretion system and is processed during export. Since Mep72 is secreted to the extracellular milieu and has a predicted N-terminal Sec secretion signal, it is likely to be exported via the type II general secretory pathway (GSP). Therefore, we predicted that Mep72 would serve as a substrate for the Xcp system, the main terminal branch of the GSP in *P. aeruginosa* (48). To test this, cell-associated and secreted proteins were harvested from biofilms of PAO1 and an ΔxcpP-Z mutant and compared by immunoblot analysis (Fig. 4). Comparison of the cell-associated fractions demonstrates that full-length Mep72 (65 kDa) is expressed in both strains. However, comparison of the secreted fraction reveals that while Mep72 is detectable in the biofilm secretome of PAO1, it is absent from the secretome of an ΔxcpP-Z mutant. This suggests that Mep72 is indeed secreted through the Xcp pathway. The data presented in Fig. 4 also show that Mep72 undergoes processing during or following its export. The cell-associated protein product corresponds to the predicted full-length isoform (metzincin domain plus tandem...
carbohydrate-binding domains) with a molecular mass of 65 kDa. However, the secreted protein product has a lower molecular mass (\( \sim 25 \) kDa) and, given its size and reactivity with the antibody (which was raised against a sequence of amino acids in the metzincin domain), most likely corresponds to the metzincin domain only. The same 25-kDa processed form was observed in the biofilm secretome of mutants containing disruptions in genes encoding LasA, LasB, AprA, and PrpL (data not shown), suggesting that the protein is processed by a minor protease or that it undergoes autoprocessing. Autoprocessing is not uncommon for members of the metzincin protease superfamily. Metzincin proteases often are translated as inactive precursor peptides which subsequently undergo an autoprocessing event that yields their enzymatically active, mature form (46).

**Ectopic expression of Mep72 in *E. coli* is toxic.** To further investigate the properties of Mep72, we recombinantly expressed the gene in *E. coli* DH5α. An N-terminal maltose-binding protein (MBP) with Sec leader peptide was fused to the mature portion (residues 21 to 599) of the mep72 ORF and expressed from an IPTG-inducible promoter. Upon induction with IPTG, a decrease in cell viability was observed (Fig. 5A). This reduction in cell viability suggests that recombinant Mep72 expression is toxic to the cell. We speculated that the product of the cotranscribed baml gene functions to counteract this effect. To test this, baml was cloned upstream of the MBP-Mep72 fusion protein. In this construct, both genes (baml and mep72) were expressed from the same IPTG-inducible promoter. Upon induction with IPTG, cell viability was restored to the level of the plasmid-only control (Fig. 5A). This suggests that Baml is essential for counteracting the toxic effect(s) of Mep72. This is reminiscent of the situation encountered with the alkaline protease, AprA, and its cognate inhibitor protein, AprI (49–51). Although the exact mechanism of AprI action remains unclear, it has been proposed that AprI binds to AprA, preventing premature protease activity until the protein has been secreted.

**Mep72 and Baml physically interact in vitro.** To determine whether the countertoxic effect of Baml on Mep72 was mediated through a physical interaction between the two proteins, a pull-down assay was carried out. Briefly, Baml containing an N-terminal hexahistidine tag (His6-Baml) was expressed in *E. coli*, refolded in vitro, and immobilized on an Ni-NTA affinity matrix. In a separate culture, MBP-Mep72 was expressed, refolded in vitro, and then passed through either a clean Ni-NTA affinity matrix (control resin) or through the Ni-NTA matrix to which His6-Baml had been attached. The columns were washed, following which the bound proteins were eluted and resolved by SDS-PAGE/Western blotting (Fig. 5B). MBP-Mep72 (105 kDa) bound to the column that was precharged with His6-Baml but did not bind the Ni-NTA affinity matrix in the absence of His6-Baml. This demonstrates that Baml and Mep72 do indeed physically interact in vitro and suggests that the countertoxic effect of Baml on Mep72 is mediated through formation of a protein-protein complex. It should be noted that since both the Baml and Mep72 protein preparations were necessarily obtained by refolding in vitro (these proteins partition into insoluble inclusion bodies when overexpressed in vivo), it is formally possible that only a small fraction of the proteins were in their physiologically active conformation. Therefore, the extent of coprecipitation observed...
In Fig. 5B, it is likely to underrepresent the probable interaction in vivo.

**Mep72 has autoproteolytic activity.** As with MBP-tagged Mep72, overexpression of His<sub>6</sub>-tagged Mep72 in liquid cultures of *E. coli* caused the culture to stop growing, and the protein partitioned into insoluble inclusion bodies. These inclusion bodies were harvested, washed, denatured in 8 M urea, and then refolded by rapid dilution to yield the native protein.

Purified Mep72 did not detectably proteolytically degrade general substrates, such as azocasein, gelatin, or elastin, even after prolonged incubation (data not shown). However, it did undergo rapid autodegradation. As soon as the His<sub>6</sub>-Mep72 was renatured, it began to autodigest (Fig. 6A). Autodigestion was essentially complete after 60 min, with little evidence of partially digested intermediates apparent. In contrast, MBP-tagged Mep72 remained stable for hours, suggesting that the presence of the MBP tag blocks autoproteolysis. To further test this possibility, we treated the MBP-tagged Mep72 with factor Xa, which cleaves at the junction between the two fused proteins. Once again, the liberated Mep72 rapidly autodigested (data not shown). Interestingly, the liberated MBP tag apparently was unaffected by the released Mep72, again indicating that the observed proteolysis is highly specific. Taken together, these data suggest that recombinant Mep72 is unstable due to autoproteolysis, at least under the conditions prevailing after refolding.

**Mep72 autoproteolysis is retarded by the presence of alginate.** Mep72 contains two C-terminal carbohydrate-binding domains, raising the possibility that Mep72 binds polysaccharides. Therefore, we wondered whether polysaccharides influence Mep72 autoproteolysis. To investigate this, purified His<sub>6</sub>-Mep72 was renatured in the presence of various carbohydrates (including monosaccharides, disaccharides, and polysaccharides), and the subsequent autoproteolysis was monitored by SDS-PAGE (Fig. 6B). Incorporation of alginate into the refolding buffer clearly

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**FIG 6** Refolding, autoproteolysis, and toxicity of Mep72. (A) Shown is a Western blot of Mep72 undergoing autoproteolysis. Denatured His<sub>6</sub>-Mep72 was refolded by rapidly diluting the urea denaturant. Samples were incubated for the indicated times before the addition of SDS sample buffer to stop any further reaction. The products were resolved by SDS-PAGE and analyzed by Western blotting. An Mep72 sample incubated for 60 min in 8 M urea (i.e., maintained in a denatured state) was used as a control. (B and C) Alginate retards Mep72 autoproteolysis. (B) Urea-denatured Mep72 was refolded as described for panel A, except that the indicated sugars or sugar polymers were included in the refolding mixture. Reactions were allowed to proceed for 20 min before quenching with SDS sample buffer and resolution by SDS-PAGE. The resulting Coomassie blue-stained gel is shown. The refolding buffers contained Tris-HCl–NaCl alone (Tris) or Tris-HCl–NaCl supplemented with 1% (wt/vol) alginate (Alg), glucose (Glu), arabinose (Ara), sucrose (Suc), maltose (Mal), polygalacturonic acid (PGA), or cellulose (Cel), as indicated. (C) Kinetics of Mep72 autoproteolysis with or without 1% (wt/vol) alginate. His<sub>6</sub>-Mep72 was refolded as outlined above. Aliquots of the refolded protein mixture were removed and quenched after a 2-, 10-, and 30-min reaction, as indicated. Shown is a Coomassie blue-stained gel of the products. (D) The metzincin domain and carbohydrate-binding domains of Mep72 are not themselves intrinsically toxic in the absence of BamI. Serial dilutions of *E. coli* containing plasmid-encoded, IPTG-inducible N-terminal MBP fusions to the metzincin- and carbohydrate (CHO)-binding domains of Mep72 were spotted onto LB-agar plates containing IPTG. The spotted dilutions were allowed to grow for 24 h before being photographed to record cell viability.
retarded the rate of autoproteolysis compared with the other sugars tested. These data indicate that the half-life of the protein can be affected by factors that presumably bind to the carbohydrate-binding domains. Further kinetic analyses confirmed that Mep72 autolysis is substantially delayed in the presence of 1% alginate (Fig. 6C).

As shown in Fig. 5A, high-level expression of MBP-tagged Mep72 is toxic to E. coli. However, we noted that prolonged (overnight) incubation of induced liquid cultures of E. coli expressing MBP-tagged mep72 sometimes led to a restoration in growth, presumably due to the outgrowth of less toxic spontaneous bypass mutants. Western analyses revealed that in most of these mutants, expression of Mep72 was abolished (data not shown). However, in one case (bypass mutant IP28), the mep72 ORF was found to contain a C→T transition, which led to the introduction of a stop codon in place of Gln137. This mutation truncates the protein downstream of the metzincin domain (residues 46 to 175) within the first carbohydrate-binding domain (CBM domain; residues 302 to 427). Western blotting confirmed that IP28 expressed a truncated, metzincin domain-containing protein. The location of this truncation was surprising, since we had assumed that the toxic effects of Mep72 expression in E. coli arose due to inappropriate proteolytic activity mediated by the metzincin domain, yet in IP28 the metzincin domain was intact. Therefore, we examined whether the ectopic expression of the metzincin domain alone or the tandem CBM domains was toxic to E. coli. However, the expression of neither construct was toxic (Fig. 6D).

| Function and PA no. | Master no. | Gene name | Gene product | Level of abundance | Fold modulation |
|---------------------|------------|-----------|--------------|-------------------|----------------|
| Spots displaying increased abundance in the wild-type (PAO1) secretome | | | | | |
| PA2783 | 1426 | mep72 | Metzincin protease Mep72 | ↑ | 55.8 |
| PA2783 | 1421 | mep72 | Metzincin protease Mep72 | ↑ | 21.6 |
| PA1706 | 1483 | pcrV | Type III secretion protein PcrV | ↑ | 7.9 |
| PA1094 | 810 | flID | Flagella capping protein FlID | ↑ | 7.6 |
| PA2783 | 880 | mep72 | Metzincin protease Mep72 | ↑ | 7.2 |

Spots displaying decreased abundance in the wild-type (PAO1) secretome

PA1709 | 1401 | popD | Type III secretion protein PopD | ↓ | 7.1 |
| PA3841 | 757 | exoS | Type III secretion effector ExoS | ↓ | 5.1 |
| PA3841/PA0044 | 751 | exoS-exoT | Type III secretion effector ExoS/ExoT | ↓ | 3.8 |

*Summary of the identified protein spots modulated in the PAO1 secretome compared to their activity in the mep72 mutant secretome. While LC-MS/MS analysis identified spot 757 as ExoS, spot 751 was identified as either ExoS or ExoT. Given that ExoS and ExoT share 74% amino acid sequence homology, many of the peptide sequences retrieved following trypsin digestion are almost identical. In these circumstances, ExoS and ExoT cannot be distinguished from each other.

a ↑ and ↓ arrows indicate whether the spot was more or less abundant in the PAO1 secretome than in the mep72 mutant secretome.

Flagella and type III secretion factors are modulated in a mep72 mutant. Given the results described above, we surmised that Mep72 is not a general protease and that its target is highly specific. To test whether specific pseudomonal proteins are affected by Mep72 activity, we used 2D-DiGE to compare the proteins secreted by a biofilm of PAO1 with those secreted by a biofilm of the mep72 mutant. The logic here was that since proteolytic processing often gives rise to a change in the electrophoretic mobility of target proteins on 2D gels, spot shifts between the wild type and mep72 mutant indicate that the corresponding proteins are targets of Mep72 activity. Consistent with the notion that Mep72 is not a general protease, the secretome of the mep72 mutant overall was very similar to that of the wild type. However, a small number of discrete changes in spot intensity were clearly observed in the 2D gel electrophoretogram (see Fig. S1 in the supplemental material).

Analysis of the spots more abundant in the PAO1 secretome (Table 2) revealed three isoforms of Mep72, one with a high molecular weight (spot 880) and two more-abundant spots with lower molecular weights (spots 1421 and 1426). The peptide sequences of the two lower-molecular-weight isoforms mapped onto the metzincin domain only, while the peptide sequences of the higher-molecular-weight isoform mapped onto the metzincin domain and the first carbohydrate-binding domain of the protein. These data are consistent with our earlier observation that Mep72 is likely to be processed during or shortly after secretion. Additional protein spots modulated in the PAO1 secretome included the type III secretion translocation factor PcrV and the flagellum-capping protein FlID.

Analysis of the protein spots that were more abundant in the mep72 mutant secretome revealed additional type III secretion factors. These included the T3SS translocation pore component PopD and the effector proteins ExoS and, possibly, also ExoT (ExoS and ExoT share very high levels of amino acid sequence similarity). These data suggest that Mep72 targets certain secreted T3S and flagellar proteins. Therefore, we postulated that there is a direct physical interaction between Mep72 and the T3SS proteins (see below).
our subsequent analyses on ExoS. ExoS displayed decreased apparent stability in the mep72 mutant secretome, as evidenced by the appearance of multiple lower-molecular-weight bands. Unlike ExoS, which was found only in the culture supernatant, PcrV was identified by Western analyses in the culture supernatant and in the matrix surrounding the biofilm cells. In both of these fractions, PcrV migrated as a single band in the 2D gel, with a lower apparent molecular weight. These data indicate that, contrary to expectation, Mep72 protects these targets against proteolytic processing/degradation.

**Mep72 and T3S factors interact in vitro**. A pulldown experiment was conducted to determine if there was any physical interaction between Mep72 and the T3S factors. MBP-Mep72 was overexpressed in E. coli, refolded in vitro, and then immobilized on an amylose affinity matrix. Secreted proteins from a stationary-phase culture of PA01 were allowed to bind, and the matrix was washed. Bound proteins were eluted, separated by SDS-PAGE, and probed by immunoblot analysis with antibodies raised against PcrV or ExoS (Fig. 7B). PcrV and ExoS coeluted with MBP-Mep72, suggesting a direct physical interaction between these proteins. As noted previously, since Mep72 was necessarily refolded in vitro for this experiment, these results are likely to underrepresent the interaction(s) occurring in vivo.

**DISCUSSION**

In the current study, we used global proteomic analysis to directly compare the secretome of biofilms and planktonic cell cultures. Through this analysis, we identified a novel metzincin protease (Mep72) whose expression appears to be highly biofilm specific under these growth conditions. Further characterization of Mep72 revealed that it is secreted through the main terminal branch of the type II general secretory pathway (Xcp system) and is processed during (or possibly shortly after) export. We also found that in the absence of coexpressed BamI, heterologous mep72 expression is toxic to E. coli, and that Mep72 and BamI form a protein-protein complex. mep72 expression in biofilms was found to alter the electrophoretic mobility of a discrete subset of secreted proteins, especially those associated with T3S. Mep72 subsequently was shown to physically capture PcrV and ExoS in vitro.

Recently, Balyimez et al. presented a preliminary report showing that in wild-type PA01, the PA2782-PA2783 genes are under the control of the CRP homologue Vfr (34). However, their studies were carried out using planktonic cultures, where PA2783 expression is, in our hands, very low. Consistent with this, when they examined the expression of PA2783 in the UWGC mutant bank strain, lacZ::phoA fusion also was extremely low. Balyimez et al. also found that when PA2783 is expressed in E. coli, the culture produces a zone of clearing when spotted onto skim-milk (protease) bioassay plates. However, we find this difficult to reconcile with the fact that E. coli lacks the main terminal branch of the general secretory pathway and lacks any mechanism by which to secrete proteases into the extracellular milieu. Instead, and given our results showing that the expression of PA2783 is toxic to E. coli, we suspect that the zone of clearing observed by these workers arose due to the release of intracellular proteases as a result of autolysis. Consistent with this, these workers failed to identify any secreted protease activity in concentrated supernatants from E. coli cultures overexpressing PA2783, although they did identify a low level of protease activity associated with the outer membrane fraction. Based on these findings, Balyimez et al. concluded that PA2783 is a membrane-bound 65-kDa endopeptidase and named it Mep72 (34).

Our data suggest that Mep72 is in fact a truly secreted protein and that it exhibits minimal endopeptidase activity on generic substrates, such as milk protein. Furthermore, the protein undergoes autoprocessing during export to yield a ca. 25-kDa mature protein.

As with many other metzincins, we found that Mep72 undergoes extensive processing before yielding the mature extracellular form. In this regard, Mep72 shares similar genomic and biochemical properties with its sibling metzincin, AprA. Like AprA, Mep72 is coexpressed with an inhibitor protein, BamI, which presumably suppresses the toxic activity of the metzincin until the latter has been secreted (49, 50). However, unlike AprI (the inhibitor of AprA), which has a well-defined tertiary structure, BamI contains several regions that are predicted to be natively unfolded (data not shown). Moreover, although AprA has been shown to cleave generic protein substrates, such as gelatin and casein (52), we found no evidence that Mep72 is proteolytically active against these. The two metzincins also differ in their mode of export; while AprA has its own dedicated type I secretion system, Mep72 is secreted through the Xcp-encoded type II secretion pathway.

Only a small number of proteins in the mep72 mutant secretome displayed altered mobility on 2D gels, but these included components of the T3S and motility apparatus. It has not escaped our notice that the T3S and flagellar apparatus are evolutionarily related and that the proteins which pass through these assem-
blages are among the few secreted proteins which bypass the periplasm completely during export. One interesting, albeit speculative, possibility is that as they pass through the periplasm, most *P. aeruginosa* secreted proteins become modified in some way that protects them from the action of potent (and in many cases, relatively nonspecific) cosecreted proteases, such as LasB. From an anthropocentric viewpoint, without such a mechanism, it may be counterproductive to secrete exoproteins into a preexisting, self-made proteolytic soup. Such protective posttranslational modifications may account for the “charge trains” commonly seen in 2D gels of *P. aeruginosa* secretome samples, a possibility that we are currently investigating. Presumably, given that the T3S and flagellar proteins bypass the periplasm, they do not receive any such protective modification. This may explain why these proteins in particular are targeted by Mep72, which, counterintuitively, appears to protect T3S proteins (PcrV and ExoS) from proteolysis (Fig. 7A). How Mep72 recognizes its target proteins is not known. Presumably, Mep72 recognizes specific sequence motifs in these targets. Alternatively, it may be that the absence of a protective periplasm-conferred modification (that is normally present on other secreted proteins) promotes Mep72 binding. Posttranslational modification of extracellular proteins is not in itself unusual; as noted by Nouwens and colleagues, a plethora of secreted proteins undergo posttranslational modification in *P. aeruginosa*, leading to the formation of multiple distinct spots of the same protein on 2D gels (27). Indeed, LasB is represented by as many as 11 discrete spots (or processed isoforms), and the LasA protease is known to be processed by other secreted endopeptidases, including LasB (53, 54). It is possible that Mep72 contributes to this complex network of processing by protecting its substrates from proteolysis by other secreted proteases.

Mep72 was shown to influence the electrophoretic mobility of a number of structural components of the T3SS translocation complex (PcrV and PopD). Here, we demonstrate a physical interaction between Mep72 and PcrV by an *in vitro* pulldown assay. However, it remains unclear whether Mep72 associates with assembled but surface-detached (broken) needle complexes or whether PcrV and PopD are truly secreted proteins. This is still an area of active research; for example, a study by Akopyan and colleagues demonstrated that translocators and effectors of the *Yersinia* T3SS localized to the surface of bacterial cells before target cell contact (55).

Purified Mep72 underwent rapid and essentially complete autodigestion following refolding in *vitro*. This feature may be physiologically relevant (e.g., serving to self-limit the accumulation of the protein in culture supernatants), or it may be an artifact of the refolding procedure; loosely folded or partially folded intermediates may be more susceptible to proteolysis by small amounts of correctly folded protease. One possible missing refolding factor was zinc ions. However, the addition of supplementary zinc to the refolding buffer had no impact on the autolytic kinetics of the refolded His-tagged protein (data not shown). Another possible refolding factor is BamI. This protein clearly confers protection upon cells expressing Mep72 (Fig. 5A), although its precise mode of action is uncertain. It remains unclear whether this protein simply binds to Mep72 and inhibits its protease activity (as in the AprI-AprA interaction) or whether this binding simply prevents Mep72 from recognizing its target proteins. These are open questions that we are currently investigating further.

Curiously, a nucleotide substitution that led to the introduction of a stop codon 5’ of the carbohydrate-binding domains obviated toxicity, as revealed by a bypass screen. Additionally, expression of neither the metzincin domain nor carbohydrate-binding domains alone resulted in toxicity. This suggests that all 3 domains must be present for toxicity to be manifest. One possibility is that all three domains are required for proper folding of the protein.

In summary, we have demonstrated that the secretome of planktonic cell cultures is very different from that of biofilms. This work led to the discovery and subsequent characterization of Mep72, a novel metzincin. Current work is aimed at better understanding the precise mechanism by which Mep72 affects the T3S proteins and the biochemical consequences of this processing. We are also investigating what regulates *baml-mep72* expression in biofilms.

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