Phosphorylation of Extracellular Proteins in Acinetobacter baumannii in Sessile Mode of Growth

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Acinetobacter baumannii is a problematic nosocomial pathogen owing to its increasing resistance to antibiotics and its great ability to survive in the hospital environment, which is linked to its capacity to form biofilms. Structural and functional investigations of post-translational modifications, such as phosphorylations, may lead to identification of candidates for therapeutic targets against this pathogen. Here, we present the first S/T/Y phosphosecretome of two A. baumannii strains, the reference strain ATCC 17978 and the virulent multi-drug resistant strain AB0057, cultured in two modes of growth (planktonic and biofilm) using TiO2 chromatography followed by high resolution mass spectrometry. In ATCC 17978, we detected a total of 137 (97 phosphoproteins) and 52 (33 phosphoproteins) phosphosites in biofilm and planktonic modes of growth, respectively. Similarly, in AB0057, 155 (119 phosphoproteins) and 102 (74 phosphoproteins) phosphosites in biofilm and planktonic modes of growth were identified, respectively. Both strains in the biofilm mode of growth showed a higher number of phosphosites and phosphoproteins compared to planktonic growth. Several phosphorylated sites are localized in key regions of proteins involved in either drug resistance (β-lactamases), adhesion to host tissues (pilins), or protein secretion (Hcp). Site-directed mutagenesis of the Hcp protein, essential for type VI secretion system-mediated interbacterial competition, showed that four of the modified residues are essential for type VI secretion system activity.

Keywords: phosphorylation, Acinetobacter baumannii, biofilm, extracellular proteins, proteomics, Hcp, T6SS and pil

INTRODUCTION

Acinetobacter baumannii is a Gram-negative nosocomial pathogen that mostly impacts patients in intensive care units and causes severe infections including pneumonia, bacteremia, endocarditis, skin and soft tissue infections, urinary tract infections, and meningitis (Bergogne-Bérézin and Towner, 1996; Dijkshoorn et al., 2007; Sengstock et al., 2010). This organism has been recently classified by the WHO as “Critical” (Priority 1, together with P. aeruginosa and Enterobacteriaceae) in the list of global priority antibiotic-resistant bacteria, for which the research and development of
new and effective antibiotic treatments are urgently required. Several factors support this ranking (Shrivastava et al., 2018). This rapidly emerging pathogen is problematic worldwide owing to the dramatic increase of infections caused by multidrug-resistant (MDR), extensively drug resistant and even pan-drug resistant, isolates (Goic-Barisic et al., 2017; Lim et al., 2018). A. baumannii also has the capacity to survive for long periods of time in hospital settings, due to its considerable ability to form biofilms (Antunes et al., 2014), enabling it to survive desiccation (Gayoso et al., 2014), oxidative stress (Soares et al., 2010), or antiseptic treatments (Peleg et al., 2008). Thus, A. baumannii has an outstanding ability to adapt to detrimental environmental conditions.

Post-translational modifications (PTMs) are an important and effective strategy that bacteria use to adapt to environmental conditions. Phosphorylation of proteins is an important signaling event in prokaryotes (Mijakovic et al., 2016) and plays important roles in bacterial virulence, adaptation, and resistance (Stock et al., 2000; Dworkin, 2015; Standish et al., 2016; Tierney and Rather, 2019; Xie et al., 2019; Lipa and Janczarek, 2020). The reversible and dynamic PTMs include the addition or removal of phosphoryl groups by kinases and phosphatases, respectively, and can alter protein function by modulation of protein conformation, subcellular localization and protein-protein interactions (Watanabe and Osada, 2012).

Proteomics is a strategy that allows the global characterization of modified proteins in a complex sample and has been used to identify serine/threonine/tyrosine phosphoproteomes for many bacterial species (Yagüe et al., 2019). One study that analyzed the phosphoproteomes of two A. baumannii strains during growth in intracellular medium showed that there was twice as many phosphorylated residues in the highly invasive multidrug-resistant clinical isolate Abh12O-A2 compared to those in the reference strain ATCC 17978, suggesting that phosphorylation plays a key role in the regulation of drug resistance mechanisms (Soares et al., 2014).

Very few global proteomic studies have characterized PTMs of extracellular proteins (Ouidir et al., 2014a; Gaviard et al., 2019). However, this analysis is very important as virulence factors are often secreted and the presence of PTMs, like phosphorylation, can impact their function. Recently, analysis of extracellular proteins produced by P. aeruginosa identified multiple PTMs on a range of secreted virulence factors (Gaviard et al., 2019). Moreover, the PTM distribution on these virulence factors appeared to be different between the intra- and extracellular-sourced proteins. Therefore, it is important to characterize the PTMs of proteins produced by A. baumannii as these modifications are likely to contribute to the adaptability, virulence, and persistence of the organism.

Here, we have investigated, for the first time, the serine, threonine and tyrosine phosphoproteome in the extracellular medium sourced from A. baumannii in a biofilm mode of growth. Two strains were investigated: the reference strain ATCC 17978 and the virulent multi-drug resistant strain AB0057, isolated from a patient with an A. baumannii bloodstream infection. Overall, a total of 109 and 146 phosphorylated extracellular proteins were identified for strains ATCC 17978 and AB0057, respectively. We also showed that one specific phosphorylation site was essential for the activity of the A. baumannii type VI secretion system (T6SS). This study provides the first global phospho-secretome profiling of A. baumannii and represents a promising starting point for further investigations on the biological role of phosphorylation in A. baumannii.

**MATERIALS AND METHODS**

**Strains and Growth Conditions**

For this study, two A. baumannii strains were used, ATCC 17978 (lacking the pAB3 plasmid, confirmed using PCR with previously described pAB3-specific primers) (Weber et al., 2015) and AB0057 (Hujer et al., 2006). Strains were grown overnight in Mueller Hinton Broth (MHB, Difco) at 37°C with shaking. Bacterial cultures were inoculated with approximately 1 × 10^7 Colony Forming Units (CFU)/mL from overnight cultures. Biofilm cultures were grown in 500 mL of MHB containing 18.75 g of sterile glass wool (Merck) (Crouzet et al., 2014, 2017) and incubated at 37°C for 24 h with slow shaking (90 rpm). In parallel, planktonic cultures were grown in the same conditions (without glass wool) and incubated at 37°C for 24 h with vigorous shaking (140 rpm). Bacteria were harvested by centrifugation (8,000 × g for 15 min at 4°C), and supernatants containing extracellular proteins stored for analysis. For each condition, bacterial cultures were grown in triplicate.

**Secreted Protein Extraction**

Each supernatant sample (500 mL) was filtered through a 0.22 µm membrane (GSWP 47 mm, Millipore) to remove residual bacteria. A protease/phosphatase inhibitor cocktail (50 µL, Halt® Protease and Phosphatase Inhibitor Single-Use Cocktail, EDTA-Free (100 ×), Thermo Scientific), and the histone deacetylase (HDAC) inhibitors nicotinamide (50 µL at 2 M, inhibitor of HDAC class III) and Trichostatin A (5 µL at 0.3 mM, inhibitor of HDAC classes I and II) (Sigma-Aldrich) were added to each sample. The secreted proteins were then concentrated to a volume of approximately 1 mL using Centricon plus-70 centrifugal filter units with a 10 kDa cutoff, according to the manufacturer’s protocol (Millipore). An ultracentrifugation step was applied to the concentrated samples (200,000 × g, 2 h, 4°C) to pellet outer membrane vesicles. The protein content of each supernatant representing extracellular proteins was then determined using the Bradford assay (Bio-Rad).

**Phosphopeptide Identification**

Samples were prepared as described previously (Ouidir et al., 2014a,b). Briefly, extracellular protein samples (50 µg) were subjected to a short electrophoresis separation in an SDS-PAGE stacking gel (7%) then protein bands were excised, and the protein reduced with DTT, alkylated with iodoacetamide, before overnight digestion with trypsin (2 µg per band). Phosphopeptide enrichment was performed using metal oxide affinity chromatography (MOAC) with titanium dioxide beads (TiO2, Carlo Erba) as previously described (Ouidir et al., 2014a,b). The enriched phosphopeptides were then analyzed.
using a LTQ-Orbitrap Elite mass spectrometer coupled to an Easy nLC II system or using Qxactive Plus coupled to an Ultimate 3000 (all from Thermo Scientific). Raw data files were processed using Proteome Discoverer 1.4 software (Thermo Scientific). Peak lists were searched using the MASCOT search software (Matrix Science) against the A. baumannii ATCC 17978 and A. baumannii AB0057 databases, containing 4097 and 4118 protein sequences, respectively. The false discovery rate (FDR) threshold for identifications was set at 1% (for proteins and peptides). For each identification, a peptide ion score higher than 14 for phosphorylation; a peptide rank of 1, a q-value and an expectation value below 0.05 was considered significant. To avoid biased automatic annotation, all phosphopeptide spectra were manually inspected. For the localization of the phosphosites, the probability results obtained from MASCOT and phosphoRS (Thermo Scientific) were used. MS proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the data set identifier. Our project accession in PRIDE is PXD027208.

**Bioinformatic Analysis of Modified Proteins**
Proteins were manually classified on the basis of predicted biological function using information gained from the annotated function, Uniprot2 and/or other available information (Resistome, VFDB experimentally demonstrated, Virulence Finder, E. coli main virulence genes, Integron and Macromolecular Systems) in the Genoscope databases (see text footnote 1; Vallenet et al., 2020). For hypothetical proteins, we used BLAST2Go 5.2.5 software, an automated functional annotation tool, for identification of (i) similar or potentially homologous sequences from NCBI database (NCBI-nr), (ii) protein domains and families from Interpro databases (EMBL-EBI), and (iii) Gene Ontology (biological process, molecular function). The Kyoto Encyclopedia of Genes and Genomes (KEGG) was used to map the pathways. STRING database (version 11.0) was used to investigate protein-protein network interactions with a high confidence score (0.7). WebLogo 2.8.2 and Icelogo4 were used with default parameters to visualize conserved patterns in sequence motifs (15 amino acids upstream and downstream of the phosphorylation site). Subcellular localization of the phosphoproteins, corresponding to the phosphopeptides identified, was predicted by PSORTb version 3.0.2 (Yu et al., 2010). To further characterize proteins, multiple bioinformatics tools were employed including SignalP 5.0 and TatP 1.0, to determine signal peptides of proteins predicted to be secreted by Sec and Tat translocons (Bendtsen et al., 2004, 2005b), LipoP2 to predict putative lipoproteins in the outer membrane by searching for lipoboxes (Juncker et al., 2003), and Secretome 2.0 and MatureP3 to predict secreted proteins (Bendtsen et al., 2005a; Orfanoudaki et al., 2017). MoonProt4 was used to find putative moonlighting proteins, defined as secreted proteins that have other known functions (Chen et al., 2018). BlastP5 was used to interrogate the Protein Data Bank (PDB) and structures to identify those with high sequence identity to the identified proteins. All figures representing three-dimensional structures were drawn with Pymol (Delano, 2002).

**Generation of an AB307-0294 hcp Mutant and Complementation Strain**
The AB307-0294 hcp mutant was generated using splice overlap extension (SOE) PCR and allelic exchange (Adams et al., 2008; Aranda et al., 2010; Tucker et al., 2014; see “Supplementary Materials and Methods” for full details). Briefly, a SOE PCR product consisting of a kanamycin cassette (containing neo), flanked by Flp recombinase Recognition Target (FRT) sites and regions representing upstream and downstream of hcp, was introduced into A. baumannii AB307-0294 via electroporation (Choi et al., 2006). Selected kanamycin resistant hcp mutants were provided with the plasmid pAT03 and grown in the presence of isopropyl-β-d-1-thiogalactopyranoside (IPTG) to induce Flp recombinase expression and excise the kanamycin cassette from the genome. The integrity of the markerless Δhcp mutant was confirmed by PCR and DNA sequencing. For complementation, an intact copy of hcp, including the native ribosome binding site, was PCR-amplified and cloned into the pBASE expression plasmid (Supplementary Table 2). The hcp complementation plasmid and empty pBASE were then used to separately transform the AB307-0294 hcp mutant, generating strains AL3844 (Δhcp[pBASE]) and AL3942 (Δhcp[hcp]).

**Mutation of Individual Hcp Phosphosites in A. baumannii Strain AB307-0294**
To assess the role of specific amino acids in Hcp function, the Δhcp mutant was separately provided with a range of plasmids, each expressing hcp with specific mutations to produce a single amino acid change in the protein (S18A, S18D, S31A, S41A, T43A, or S44A changes) compared to the wild-type Hcp. Each change to hcp was generated using PCR or SOE-PCR with the appropriately designed primers. PCR fragments were then cloned into pBASE (see “Supplementary Materials and Methods” for full details).

**Detection of the T6SS Protein Hcp Using Western Immunoblotting**
Production and secretion of the Hcp protein in A. baumannii was assessed by subjecting whole-cell lysates (WCL) and culture supernatants to western immunoblotting using anti-Hcp polyclonal sera, as described previously (Fitzsimons et al., 2018) but with minor modifications (see “Supplementary Materials and Methods” for full details). The intensity of each

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1. [http://www.genoscope.cns.fr](http://www.genoscope.cns.fr)
2. [http://www.uniprot.org/](http://www.uniprot.org/)
3. [https://weblogo.berkeley.edu/](https://weblogo.berkeley.edu/)
4. [https://omics.ugent.be/icelogoserver/](https://omics.ugent.be/icelogoserver/)
5. [http://www.cbs.dtu.dk/services/SignalP/](http://www.cbs.dtu.dk/services/SignalP/)
6. [http://www.cbs.dtu.dk/services/TatP/](http://www.cbs.dtu.dk/services/TatP/)
7. [http://www.cbs.dtu.dk/services/LipoP/](http://www.cbs.dtu.dk/services/LipoP/)
8. [http://www.cbs.dtu.dk/services/SecretomeP/](http://www.cbs.dtu.dk/services/SecretomeP/)
9. [http://www.stepdb.eu/MatureP.php](http://www.stepdb.eu/MatureP.php)
10. [http://www.moonlightingproteins.org/](http://www.moonlightingproteins.org/)
11. [https://blast.ncbi.nlm.nih.gov/](https://blast.ncbi.nlm.nih.gov/)
immunoreactive band was quantified from the peak area of histograms generated using ImageJ software (Schneider et al., 2012). Significance of differences in Hcp production were assessed using one-way analysis of variance (ANOVA; GraphPad Prism, version 9) with Tukey's multiple-comparison post-test; a P value of <0.05 was accepted as statistically significant.

RESULTS AND DISCUSSION

The ability of *A. baumannii* to form biofilms significantly threatens human health as it enhances the ability of this pathogen to persist and survive in response to desiccation and various stresses and for prolonged periods in hospital environments. In order to identify phosphorylation events that occur during biofilm growth and determine those which may play important roles in bacterial virulence or resistance, we investigated the phospho-secretome of AB0057 in planktonic and biofilm modes of growth. We also assessed the @ @ phospho-secretome of ATCC 17978 pAB3− strain, known to have an active T6SS and to produce biofilms (Weber et al., 2015; Moon et al., 2017).

Phospho-Secretome Analysis of *A. baumannii* Strains ATCC 17978 and AB0057

Using a large-scale proteomic approach, we first assessed the phosphorylation of serine, threonine and tyrosine residues on extracellular proteins (S/T/Y phosphoproteomes) derived from biofilm and planktonic growth of *A. baumannii* reference strain ATCC 17978 and the virulent isolate AB0057. After bacterial culture, extracellular proteins were digested and phosphorylated peptides were enriched using titanium dioxide (TiO₂), as described previously (Ouidir et al., 2014a). Phosphopeptide identification, all data were manually checked to confirm the best mass spectrum interpretation and peptide sequence and localization of the PTM. If localization of the phosphorylation site was not possible then the putative modified residues were annotated in brackets.

For both strains, the overall number of phosphorylated sites was higher in peptides derived from the biofilm mode of growth than from the planktonic mode of growth. Indeed, 137 modified sites (corresponding to 124 unique peptides and 97 unique proteins) in ATCC 17978 and 155 modified sites (corresponding to 144 unique peptides and 119 unique proteins) in AB0057 were identified in biofilm, while 52 phosphosites (44 unique peptides and 33 unique proteins) in ATCC 17978 and 102 phosphosites (92 unique peptides and 74 unique proteins) in AB0057 were identified in planktonic mode (Figures 1A,B and Supplementary Tables 3–10). More phosphorylated sites were identified in the peptides derived from the virulent strain AB0057 than in peptides from the reference strain ATCC 17978, especially in the planktonic growth samples. Interestingly, less than half of the phosphosites identified in the peptides derived from planktonic grown cells were also present in peptides derived from biofilm grown cells (Figures 1C,D).

No significant difference was observed in the phosphorylation distribution onto S/T/Y (Supplementary Table 9). To identify conserved motifs, sequences within 15 amino acids upstream and downstream of the phosphorylated sites were examined for similarity. However, no consensus motif surrounding each of the phosphorylated sites was found.

The majority of proteins were phosphorylated at only one site (Figure 1E). In strain AB0057, elongation factor Tu (AB57_0362) had the highest number of phosphorylations [six phosphorylations on T17, (T26/T27), Y77, Y88, T94, and T109]. In strain ATCC 17978, five phosphorylated sites were identified in the T6SS tube protein Hcp (S18, S31, S41, T43, and S44) but these additions were only present on extracellular Hcp isolated from biofilm cultures (Supplementary Table 8).

Initial bioinformatic analyses indicated that the majority of the modified proteins were predicted to be cytoplasmic, with only 3% predicted to be secreted (Supplementary Tables 7, 8). However, it is well known that many predicted cytoplasmic proteins can also be secreted, e.g., DnaK (Karched et al., 2019), EF-Tu (Chiu et al., 2017), and GroEL (Jeffery C.J., 2018). Such multifunctional proteins are also known as moonlighting proteins and may be involved in adhesion to the host or in microbial pathogenesis (Hagemann et al., 2017; Harvey et al., 2019; Jeffery C.J., 2019; N’Diaye et al., 2019). Nine phosphoproteins in ATCC 17978 and 10 phosphoproteins in AB0057 were identified as putative moonlighting proteins with adhesion or binding activities (Supplementary Tables 7, 8). The sequences of all phosphorylated proteins were analyzed using a number of bioinformatic tools. In total, 23 and 29 proteins with signal peptides were identified in ATCC 17978 and AB0057, respectively, and are therefore predicted to be secreted by the Sec pathway. Using SecretomeP, 36 proteins in ATCC 17978 and 35 proteins in AB0057 were predicted to be secreted via a non-classical secretion pathway (Kang and Zhang, 2020). MatureP identified 29 proteins in ATCC 17978 and 46 proteins in AB0057 secreted by the Sec secretory system. Furthermore, 12 lipoproteins were identified (five in ATCC 17978 and seven in AB0057). Taking into consideration the data from all of these analyses, the final number of predicted secreted proteins (estimated to be 3% using initial bioinformatic analysis) was determined to be 38%.

The phosphorylated proteins identified were classified according to the classical functional categories (Figures 1F,G). The majority of planktonic and biofilm derived extracellular proteins were classified into the amino acid metabolism group and the unknown function group, some of which in the latter group may potentially contribute to virulence by currently unidentified mechanisms. Interestingly, compared to ATCC 17978 planktonic growth, there were more ATCC 17978 biofilm-derived extracellular proteins classified into bacterial secretion system, iron metabolism, oxidative phosphorylation and folding and maturation functional groups (Figure 1F). For the clinical isolate AB0057, compared to planktonic growth, there were more biofilm-derived extracellular proteins classified into the adaptation protein functional group (Figure 1G). Furthermore, differences in the distribution of the identified proteins into functional groups were clearly noted between the two strains.
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Figure 1 | Numbers of phosphosites (A) and phosphoproteins (B) identified in A. baumannii ATCC 17978 and AB0057 strains in planktonic and biofilm modes of growth. (C,D) Phosphosite distribution between planktonic and biofilm modes of growth for ATCC 17978 and AB0057. (E) Number of phosphosites per protein in planktonic and biofilm modes of growth for ATCC 17978 and AB0057, respectively. (F,G) Functional categories of the phosphorylated proteins in planktonic and biofilm modes of growth for ATCC 17978 and AB0057, respectively. Hatched red: ATCC 17978 in planktonic. Hatched blue: ATCC 17978 in biofilm. Red: AB0057 in planktonic. Blue: AB0057 in biofilm.

for the secreted proteins in the following categories: DNA/RNA metabolism, transcriptional regulation, and drug resistance.

Functions of Selected Phosphorylated Proteins
Type VI Secretion System: Phosphorylation of Hcp
The T6SS is a multiprotein apparatus that delivers protein effectors into both prokaryotic and eukaryotic cells (Pukatzki et al., 2009; Ho et al., 2017). In various Gram-negative species it plays a critical role in biofilm formation (Aschtgen et al., 2008) and as a defensive weapon in inter-bacterial competition and/or attack of host cells (Schwarz et al., 2010; Russell et al., 2011; Basler et al., 2013; Carruthers et al., 2013). In the A. baumannii strains AB0057 and ATCC 17978, the genes encoding the main structural components of the T6SS apparatus are encoded within one, well conserved, locus composed of 18 and 22 genes, respectively. Furthermore, three distinct VgrG loci are located elsewhere on the genome. Each VgrG locus encodes the VgrG tip protein, and a cognate effector protein and a cognate immunity protein. Hcp proteins form the needle of the T6SS via the stacked formation of multiple donut-shaped Hcp hexamers. In some bacterial species Hcp can act as a specialized effector or as a transporter and/or chaperone of T6SS cargo effectors (Ruiz et al., 2015). The presence of Hcp in the cell culture supernatant is a molecular marker of a functional T6SS (Mougous et al., 2006; Pukatzki et al., 2009; Weber et al., 2013; Corcionivoschi et al., 2015). Hcp proteins (AB0057_1481 and ABYAL1534) were detected in both planktonic and biofilm modes of growth, suggesting a functional T6SS is produced by both strains under both growth conditions.
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FIGURE 2 | Hcp phosphorylation sites: (A) Crystal structure of the A. baumannii Hcp (PDB 4W64) with specific phosphosites colored (S18, light green; S31, red; S41, yellow; T43, blue; S44, purple) and the Hcp hexameric donut structure viewed from the side (left) and bottom (right). (B) Alignment of Hcp protein from A. baumannii ATCC 17978 (A.b_ABYAL1534) and P. aeruginosa (P.a_PA0085). Completely invariant residues are indicated with an asterisk (*), highly conserved regions are marked with a colon (:), and semi-conserved residues are dotted (.). Identified peptides and phosphosites in our study are colored and designated by arrows.

(Supplementary Table 10). In agreement with the literature (Weber et al., 2015), Hcp is secreted by ATCC 17978 lacking the plasmid pAB3 (encoding a TetR repressor) at greater amounts when compared to secretion by strain AB0057. Interestingly, the five phosphosites (S18, S31, S41, T43, and S44) on Hcp were only detected in biofilm mode of growth for ATCC 17978 (Supplementary Tables 6, 8). Examination of the predicted 3D model of the A. baumannii Hcp hexamer indicates that these phosphosites are positioned at the top and the bottom of the donut-shaped structure (Figure 2).

To test the importance of the Hcp phosphosites for T6SS function, we mutated selected sites in a recombinant version of Hcp derived from A. baumannii strain AB307-0294 (100% amino acid identity with ATCC 17978 and AB0057 Hcp). Individual amino acid substitutions of the identified Hcp phosphosites were performed with alanine, a non-phosphorylatable amino acid. Each modified hcp gene was cloned into the expression plasmid pBASE and introduced into an A. baumannii AB307-0294 hcp mutant, and T6SS function in each strain assessed via the immunodetection of Hcp secretion. An S18A amino acid substitution in Hcp abolished Hcp secretion via T6SS activity as no Hcp was detected in the supernatant (Figure 3). Similarly, an S18D phosphomimetic substitution in Hcp also resulted in no Hcp secretion. In contrast, Hcp proteins with S31A, S41A, T43A, or a S44A substitution were functional as T6SS-secreted Hcp was detected in the supernatant, though levels of secreted Hcp were significantly reduced for Hcp proteins with S31A, S41A, and T43A substitutions, compared to mutant provided with wild-type Hcp or Hcp S44A. In all strains provided functional copy of wild-type or mutated hcp, Hcp protein was detected in the whole cell lysate, indicating that each substitution had no effect on the production of Hcp (Figure 3). The TssM protein is an essential component of the T6SS membrane-stabilizing structure. The ΔtssM strain was used as an additional negative control in the immunoblot as this mutant cannot secrete Hcp due to lack of a functional T6SS (Fitzsimons et al., 2018).

In P. aeruginosa PAO1, Hcp protein not only forms the T6SS tube but also associates with effectors, protects them from proteolysis, and likely traffics with effectors during transport (Silverman et al., 2013). Specific amino acids in Hcp, such as S31 or L28, can inhibit or affect the ability of the protein to secret the associated effector. Thus, T6SS effectors form a unique network of interactions with Hcp. In Francisella tularensis, the importance of the phosphorylation of the T6SS sheath protein IgB was recently highlighted (Ziveri et al., 2019). A Y139 mutation in IgB was able to abrogate the T6SS assembly regardless of the substituted residue (Y139A, Y139D, or Y139E). The phosphorylation status of Y139 therefore modulates the dynamics of T6SS assembly. Our results indicate that the phosphorylation of Hcp amino acid S18 seems to play a pivotal role in A. baumannii Hcp secretion via an active T6SS. These results highlight the importance of the characterized phosphorylated residues, especially S18, on T6SS activity in A. baumannii.
Pili are assembled via a chaperone-usher pathway (CUP) that involves a periplasmic chaperone and an outer membrane usher for secretion to the cell surface (Tomaras et al., 2003; Zavyalov et al., 2010; Busch and Waksman, 2012; Garnett et al., 2012; Bao et al., 2013; Berry et al., 2014; Werneburg and Thanassi, 2018). One of the most well-studied type 1 pili is the Csu system (Tomaras et al., 2003; Pakharukova et al., 2018). The Csu pilus is made of four subunits, CsuA/B (the major pilin subunit), CsuA, CsuB, and the tip subunit CsuE, which uses three distal hydrophobic loops to interact with hydrophobic substrates (Pakharukova et al., 2018). The CsuC and CsuD proteins are the chaperone and the usher components of this system, respectively (Pakharukova et al., 2018). Analysis of A. baumannii strains ATCC 17978 and AB0057 detected phosphorylations on the major subunit CsuA/B (Supplementary Tables 7, 8 and Supplementary Figure 1) within the N-terminus region known to be involved in the interaction with the chaperone and in subunit polymerization. Indeed, assembly of each pilus starts in the periplasm with stabilization of a subunit by the chaperone. This occurs via a donor strand complementation process, where the β-strand G1 of the chaperone induces an immunoglobulin-like fold of the subunit (Choudhury et al., 1999; Sauer et al., 1999). Following folding, a donor β-strand exchange mechanism occurs where the G1 β-strand is displaced by the N-terminal extension (donor strand Gd) of an incoming subunit (Remaut et al., 2006; Allen et al., 2012). The phosphorylation of S36 (in AB0057) and S38 (in ATCC 17978) was detected in the CsuA/B subunit proteins produced by both strains. These residues are located in the region immediately following the end of the Gd β-strand (Supplementary Figure 1). It was previously shown that mutation of the P5 residue (I37A mutant) leads to a drastic decrease of CsuA/B polymerization (Remaut et al., 2006; Pakharukova et al., 2015). Our finding in A. baumannii suggests phosphorylations could decrease the accommodation of the Gd β-strand into the groove of a neighboring subunit and thus impact polymerization. Other phosphorylations were identified in the subunit A strand (T39, T42, and S46), that can be hydrogen bonded to the A1 β-strand of the CsuC chaperone during the formation of a high-energy folding intermediate and before pilus polymerization and translocation (Pakharukova et al., 2015). Overall, these modifications could regulate CsuA/B interactions with the CsuC chaperone and contribute to the stabilization of its soluble form.

In addition, phosphorylations were also detected on the main subunits (FimA-like) of two other CUP pili in strain AB0057. The genes for these CU systems are located in two loci, each encoding four proteins (Supplementary Table 7) as follows: the FimA-like main pilin (AB57_1747 and AB57_2007 have 30 and 28% identity, respectively, with the E. coli FimA), the PapD chaperone (AB57_1746 and AB57_2006), the PapC outer membrane usher (AB57_1745 and AB57_2005) and the tip adhesion pilin (AB57_1744 and AB57_2004). These CUP pili were shown to be involved in biofilm formation on solid supports and in bacterial attachment to A549 epithelial cells (Rumbo-Feal et al., 2013; Álvarez-Fraga et al., 2016). As shown by the sequence alignment of these subunits with FimA proteins from different species (Supplementary Figure 2), one phosphorylation is commonly localized on the B° β-strand of both subunits, before the second cysteine necessary for the formation of a disulfide bond, and this residue is conserved in all the aligned pili subunits (Puorger et al., 2011; Zyla et al., 2010).
Interestingly, the B’’ β-strand (T45-C58) corresponds to a sequence required for targeting FimA monomers to mitochondria (Sukumaran et al., 2010; Zyla et al., 2019). Indeed, previous studies demonstrated that soluble FimA monomers from invasive bacterial pathogens (E. coli K1, Salmonella or Shigella) may target the VDAC-hexokinase complex at mitochondria to delay host cell apoptosis during the early phase of infection (Sukumaran et al., 2010). As A. baumannii is able to invade epithelial cells via a zipper-like mechanism and persist in these cells (Choi et al., 2005, 2008; Parra-Millán et al., 2018), FimA-like subunits may possess a similar function and phosphorylation could be an important event to modulate pili subunit-mitochondria interactions.

The tip pilins of both CUP systems, AB57_2004 and AB57_1744, shared 36 and 35% identity, respectively, with the adhesin MrkD1P of Klebsiella pneumoniae (PDB code 5W12).
Drug Resistance: Phosphorylation of β-Lactamases

The AB0057 A. baumannii strain is a multidrug resistant clinical isolate; it is particularly resistant to β-lactams including carbapenems due to the presence of several β-lactamases (Hujer et al., 2006). We detected several phosphorylations on different β-lactamases produced by AB0057 (Supplementary Table 7). We observed four phosphosites [(Y63/Y66/Y67), S88, S90 and (Y223/Y227)] on ADC (Acinetobacter-derived cephalosporinase) proteins, AB57_0009 and AB57_2796. As these two proteins share 97.68% identity, it is not possible to precisely assign all of the phosphorylation sites to one or the other protein. However, phosphosites S88 and S90 were unambiguously assigned to the AB57_0009 protein. Residue S88 corresponds to the catalytic serine of the enzyme, making a covalent bond with the antibiotic before hydrolysis of the β-lactam, and S90 is oriented toward the small helix H6” that follows the Ω loop, shown to be extended of five amino acids in the AB57 protein compared to other members of the class C β-lactamases. The phosphoY sites were located in the Ω loop [(Y223/Y227)] and the β-strand [(Y63/Y66/Y67)] (Figure 4), those in the latter being involved in the structural stability (Caselli et al., 2018). The Ω loop, localized at the entrance of the enzyme active site, is involved in the catalytic cycle of the β-lactam hydrolysis and most of identified mutations in this loop alter the susceptibility of the bacterium to β-lactam antibiotics.

For the β-lactamase AB57_0551, we identified one phosphosite [S79/T80] corresponding to the catalytic serine residue S79. This protein shares 99% amino acid identity with the class-D β-lactamase OXA-23 whose structure has been solved in complex with a molecule of meropenem linked covalently to the catalytic serine (PDB code 4JF4) (Harper et al., 2018).

In agreement with these results, a link between phosphorylation and the modulation of β-lactamase activity was recently reported. Lai et al. (2016) showed the importance of the S90 phosphosite in the catalytic motif S88VS90K of AmpC (ADC) from A. baumannii SK17, corresponding to the phosphoS90 found in AB57_0009. The phosphorylatable S90A mutant had higher β-lactamase activity and increased resistance to imipenem, in contrast to the phosphomimetic S90D and S88D mutants which were more sensitive to imipenem. Thus, phosphorylation in the catalytic site region could repress enzyme activity. It should be also noted that phosphorylation of S90 (AB57_0009) and [S79/T80] (AB57_0551) were exclusively detected in extracellular proteins derived from planktonic growth. Taken together, these data suggest that during planktonic lifestyle, A. baumannii AB0057 would be more sensitive to antibiotics such as carbapenems.

Secondary Functions: Phosphorylation of Moonlighting Proteins

Moonlighting proteins are proteins that can have two or more distinct biochemical functions (Jeffery C., 2019). They increase the complexity of bacterial regulation and the specific function of each depends on cellular localization, cellular ligands, and/or substrates. In this current study, 39 moonlighting phosphoproteins were identified, including DnaK, EF-Tu, GroEL, and peroxiredoxin (Supplementary Tables 7, 8). Thirteen of these proteins had been assigned a second role in adhesion (to plasminogen, eukaryotic cells, fibrinectin), virulence or modulation of host immune response (Supplementary Table 11; Das et al., 2010; Henderson and Martin, 2013; Jeffery C., 2018). EF-Tu functions at the ribosomal level, but in P. aeruginosa it is also exported and binds to plasminogen (Kunert et al., 2007) and in M. pneumonia it has been shown to bind to fibronectin (Dallo et al., 2002). In Bacillus cereus, EF-Tu can be recruited onto the bacterial surface and act as an environmental sensor to detect the principal skin neutrophili, substance P (N’Diyaye et al., 2019). A. baumannii EF-Tu from both ATCC 17978 and AB0057 contained many phosphorylated sites. As all of the moonlighting proteins were phosphorylated on serine, threonine or tyrosine, it can be envisaged that phosphorylation can act as a switch to modulate protein function or protein secretion by either the addition of negative charge or via steric hindrance.

CONCLUSION

Due to the crucial role of phosphorylation in bacterial virulence, resistance or adaptation, phosphorylation mapping is an important prerequisite for the discovery of novel regulatory mechanisms. Investigation of protein phosphorylation is essential to understand its role in bacterial physiology and to identify new control strategies for combating the spread of MDR bacteria. In this study, we used the reference strain ATCC 17978 and the clinical isolate AB0057 to investigate for the first time the phospho-secretome of A. baumannii when in the biofilm mode of growth, which promotes bacterial environmental persistence and antibiotic tolerance. The comparative study of planktonic with biofilm modes of growth showed an enhanced number of phosphorylated proteins in cells grown within a biofilm. In addition, a higher number of phosphorylation events was observed in proteins derived from the virulent and antibiotic resistant strain AB0057, compared to those identified in the reference strain ATCC 17978.

Our data show a pivotal role of phosphorylation in many biological processes, including antibiotic resistance, biofilm formation and virulence. Indeed, phosphorylation of residues located within the catalytic site of several β-lactamases indicates
this activity modulates their enzymatic activity. Phosphorylation of pili proteins could modulate polymerization, contribute to the stabilization of their soluble form, and could play a role in pili-facilitated adhesion to eukaryotic cells. Many phosphosites were detected on the T6SS tube protein Hcp and we showed that phosphorylation at amino acid S18 is required for Hcp secretion and therefore T6SS activity. Phosphorylation at three other sites on Hcp also contributes significantly to T6SS activity. This work is a promising starting point for further investigations into the biological role of phosphorylation on secreted proteins during growth in biofilms.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/online repositories.

AUTHOR CONTRIBUTIONS

ED, JB, and JH contributed to conception and design of the study. SM, BR, MM, AW, MH, BH, and IB performed the experiments. SM, MM, MH, IB, JB, ED, and JH wrote the manuscript. All authors contributed to manuscript revision, read, and approved the submitted version.

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