Acinetobacter baumannii is a hospital-acquired pathogen that shows an extraordinary capacity to stay in the hospital environment. Adherence of the bacteria to eukaryotic cells or to abiotic surfaces is the first step for establishing an infection. The A. baumannii strain AbH12O-A2 showed an exceptional ability to adhere to A549 epithelial cells. The AbFhaB/FhaC 2-partner secretion (TPS) system involved in adhesion was discovered after the screening of the recently determined A. baumannii AbH12O-A2 strain genome (CP009534.1). The AbFhaB is a large exoprotein which transport to the bacterial surface is mediated by the AbFhaC protein. In the present study, the role of this TPS system in the AbH12O-A2 adherence phenotype was investigated. The functional inactivation of this 2-partner secretion system was addressed by analyzing the outer membrane vesicles (OMV) proteomic profile from the wild-type strain and its derivative mutant AbH12O-A2ΔfhaC demonstrating that AbFhaB is no longer detected in the absence of AbFhaC. Scanning electron microscopy (SEM) and adhesion experiments demonstrated that inactivation of the AbFhaB/FhaC system significantly decreases bacterial attachment to A549 alveolar epithelial cells. Moreover, it has been demonstrated that this 2-partner secretion system is involved in fibronectin-mediated adherence of the A. baumannii AbH12O-A2 isolate. Finally, we report that the AbFhaB/FhaC system is involved in virulence when tested using invertebrate and vertebrate hosts. These data suggest the potential role that this AbFhaB/FhaC secretion system could play in the pathobiology of A. baumannii.

Introduction

Acinetobacter baumannii is an opportunistic human pathogen considered a major cause of nosocomial infections worldwide. A. baumannii causes pneumonia, skin and soft tissue infections, bacteremia, meningitis, endocarditis and urinary tract infections resulting in high rates of morbidity and mortality.1 The emergence of multidrug-resistant (MDR) strains has led to a number of hospital outbreaks that have become a serious health problem worldwide.2-6

Many studies have been focused on the investigation of antimicrobial resistance. Although the clinical importance of A. baumannii infections has increased, the pathogenicity of this microorganism is sparsely understood, since it has been traditionally considered as a low virulent pathogen.2 Several bacterial virulence factors are needed for the pathogenesis of Acinetobacter infections and only a few have been described in this specie. Clinical A. baumannii strains exhibit remarkably variations in virulence-associated phenotypes such as motility, adherence, biofilm formation, invasion, iron uptake, cell capsule development or penicillin binding proteins modifications among others.2,7-14

Some studies have evidenced that Acinetobacter species may reach the human skin and mucosal membranes and then colonize and persist on the host several weeks.15 A. baumannii has an innate capacity to interact with and adhere to diverse types of surfaces. Bacterial adherence constitutes an essential step in the colonization process.
After adhesion, bacteria may form biofilms that are involved in the persistence of this pathogen in the hospital environment. Some components, such as the staphylococcal biofilm-associated protein (Bap), the CsuA/ BABCDE usher-chaperone system or the poly-
\( \beta -1 \rightarrow 6-N \)-acetylglucosamine have been described as involved in the \textit{A. baumannii} biofilm formation and adherence phenotypes.\textsuperscript{16-20} A study made with 52 different clinical strains revealed that the aptitude to form biofilm and the ability to attach to host cells are independent abilities that do not have to be associated.\textsuperscript{21} This could indicate that \textit{A. baumannii} has several independent mechanisms involved in biofilm formation and adherence to different surfaces that also are independently regulated.

In the other hand, the capacity of \textit{A. baumannii} to adhere to and achieve biofilm formation on biotic surfaces is barely known. The outer membrane protein OmpA, besides of being related to biofilm formation in abiotic surfaces, has been shown to promote the adherence to eukaryotic host and invasion.\textsuperscript{22} Also, Bentancor \textit{et al.},\textsuperscript{23} have corroborated that the trimeric autotransporter protein Ata, belonging to the type V secretion system, participates in biofilm formation and intercedes in the attachment of \textit{A. baumannii} cells to immobilized collagen type IV.\textsuperscript{23} The type V secretion systems are classified into either autotransporters or 2-partner secretion systems (TPS). The protein Ata was the first adhesion component that belongs to the type V autotransporters secretion systems described in \textit{A. baumannii}. The filamentous hemagglutinin protein (FHA) from \textit{Bordetella pertussis}, HecA from \textit{Erwinia chrysanthemi} and HMW1/2 from \textit{Haemophilus influenzae} have been described as adhesins belonging to this group of proteins and constitute one of the major virulence factors described in these species.\textsuperscript{24,25} The TPS are widespread secretion pathways of Gram-negative bacteria and involve 2 proteins: TpsA, a large exoprotein, and TpsB, the outer membrane protein. The genes coding for these 2 proteins are usually co-transcribed within the same operon. The TpsB protein forms a \( \beta \)-barrel included into the outer membrane. In the other hand, the N-terminal domain of TpsA connects with the POTRA domain of TpsB proceeding to the secretion. After, TpsA may stay associated at the cell surface or be liberated to the extracellular medium.\textsuperscript{26}

The present work is focused on an \textit{A. baumannii} strain that was responsible for one of the most important nosocomial outbreaks to date.\textsuperscript{5} During more than 30 months more than 350 patients were colonized with MDR \textit{A. baumannii} strains in a hospital in Madrid.\textsuperscript{5} In recent studies, the molecular mechanisms involved in the spread and dissemination of the AbH12O-A2 strain have been characterized.\textsuperscript{3,27,28} This strain was found as the main clone within this outbreak and constituted a target of many studies.\textsuperscript{29,30} Recently, our group determined its complete genome (accession number CP009534.1).\textsuperscript{31} We found that the most remarkable characteristic of the AbH12O-A2 strain is its notable ability to adhere to human cells. An analysis of its genome allowed us to focus on a genome region coding for an FhaB/FhaC-like 2-partner secretion system due to its possible implication in adherence. The main goal of the present work was to analyze the role of the FhaB/FhaC-like 2-partner secretion system in the pathogenicity of \textit{A. baumannii} strain AbH12O-A2.

### Results

**The \textit{A. baumannii} AbH12O-A2 strain harbor a genomic island involved in adherence**

The \textit{A. baumannii} strain AbH12O-A2 was the main clone isolated from a large nosocomial outbreak occurring in a hospital in Madrid (Spain) from 2006 to 2008. This \textit{A. baumannii} isolate has the capacity to survive and persist for long periods in the hospital settings. Our group recently demonstrated the molecular mechanisms implicated in the reply to desiccation and persistence of this pathogen.\textsuperscript{29} Several phenotype analyses were performed to find out the molecular strategies that may be involved in persistence and disease potential of \textit{A. baumannii} strain AbH12O-A2.\textsuperscript{3,27-31} Attachment assays showed that the AbH12O-A2 strain had a higher ability to adhere to A549 alveolar epithelial cells compare with the ATCC 17978 strain, being 37-fold more adherent with a \( P \)-value < 0.001 (Fig. 1A). This adherence phenotype may explain its propensity to remain in colonized patients, this being one of the keys of the prolonged nosocomial outbreak. Protein coding genes involved in the adherence phenotype shown by AbH12O-A2 strain were analyzed based on the fully sequenced and annotated genome of this clinical isolate (GenBank accession code CP009534.1).\textsuperscript{31} The whole-genome analysis of the AbH12O-A2 strain revealed the presence of a \textit{ca. 18-kb} chromosomal region containing genes involved in adherence, which also appears in the genomes of the \textit{A. baumannii} 3207 and AB030 strains that showed 99% and 100% identity to the AbH12O-A2 chromosomal region, respectively. This \textit{ca. 18-kb} region was also found in the genomes of the \textit{A. baumannii} IOMTU 433, 6200 and SDF strains and, as assessed by the Blast tool of the NCBI, it was found coverage of 87, 87 and 70% with an identity of 96, 94 and 96%, respectively. It was observed that the \textit{ca. 18-Kb} region was partially conserved in the genomes of the \textit{A. baumannii} strains R2091, CIP70.10, AB031 and ZW85–1 where the genes flanking the island (LX00_12065 and LX00_12105: \textit{AbfhaC}) are present and
the genes LX00_12075, LX00_12080, LX00_12085, LX00_12090 and LX00_12095 are absent and only the ending region of the LX00_12100 gene (AbfhaB) appears. Fig. 2A shows that this ca. 18-kb region described in the AbH12O-A2 genome is flanked by repeated DNA sequences. The LX00_12060 locus coding for a LuxR family transcriptional regulator and the LX00_12110 locus coding for a tRNA-Trp were found upstream and downstream of this region, respectively. Eight coding sequences were identified within this region: LX00_12065 coding for a TetR family transcriptional regulator (AIS07096.1), LX00_12075, LX00_12080, LX00_12085, LX00_12090, LX00_12095 coding for hypothetical proteins (AIS07097.1, AIS07098.1, AIS07099.1, AIS07100.1 and AIS07101.1, respectively), LX00_12100 coding for an adhesion protein (AKB90480), and LX00_12105 coding for a membrane protein (AIS07102.1), as shown in Fig. 2A. Further bioinformatic analysis revealed a TPS, named here as AbFhaB/AbFhaC, encoded by the LX00_12100 and LX00_12105 genes, respectively (Fig. 2B). The LX00_12100 gene has been predicted to code for a 392.72-kDa exoprotein (TpsA, named here as AbFhaB) with a 25-amino acid residue signal peptide, typical of Type V secretion systems. A deeper examination of this sequence showed the appearance of an N-terminus haemagglutination activity domain (amino acids 101 to 221), highly conserved in the TPS domain of the TpsA exoproteins (IPR008638 code from InterPro database). In addition, multiple copies of a 20-amino acid residue repeat (IPR010069 code from InterPro database), also found in filamentous haemagglutinins, were detected in the AbFhaB protein, as shown in Fig. 2A. An Arg-Gly-Asp triplet (RGD) that has been related to adhesion functions occurred once in the predicted sequence (amino acids 2029 to 2031) in the TpsA ortholog. The LX00_12105 gene was expected to code for a 65.54-kDa outer membrane protein (TpsB, named here as AbFhaC), that belongs to the Omp85/TpsB transporter family. This protein contains a polypeptide transport-associated domain named POTRA_2 (IPR005565 code from InterPro database) localized between amino acids 87 and 166. Structure prediction analysis using Phyre2 suggested that the LX00_12105 gene may have a similar structure (100% confidence and 86% coverage) to the FhaC.

Figure 1. (A) Determination of the attachment ability of the AbH12O-A2 clinical strain, the AbH12O-A2ΔfhaC isogenic mutant derivative strain and the complemented strain to A549 epithelial cells. T-student tests were performed and 6 independent replicates were done. Bars indicate the standard deviation and asterisks indicate P values under 0.001. (B) SEM visualization of bacterial attachment to A549 human alveolar epithelial cells infected with A. baumannii. a) A. baumannii AbH12O-A2 clinical strain. b) A. baumannii mutant derivative AbH12O-A2ΔfhaC. c) Healthy uninfected A549 cells covered by surfactant as a negative control. Micrographs were taken at 10.000X magnifications. Bars indicated the scale (1 μm).
protein, member of the Omp85/Tpsb2 transporter family.

The AbH12O-A2ΔfhaC mutant derivative strain does not produce the AbFhaB and AbFhaC proteins

In order to investigate the role of the TPS system in the pathogenicity of the AbH12O-A2 strain, this system was inactivated by deleting the POTRA_2 domain of the fhaC gene, which is responsible for the transport and linkage of FhaB in the cell surface, obtaining the AbH12O-A2ΔfhaC mutant derivative. Briefly, the POTRA_2 domain-coding region of the LX00_12105 gene (fhaC) was deleted by double crossover recombination employing the pMo130TelR. The construction of the AbH12O-A2ΔfhaC mutant strain was verified by PCR and qRT-PCR results demonstrated the lack of LX00_12105 gene expression in this mutant strain (data not shown). The AbH12O-A2ΔfhaC isogenic deletion derivative mutant of AbH12O-A2 was constructed by deleting a region encompassing the fhaC gene without affecting the upstream and downstream surrounding genes, as assessed by RNA expression analysis, since no polar effects were observed (data not shown).

The inactivation of the TPS system was also confirmed through proteomic approaches. OMVs protein profile from AbH12O-A2 and its isogenic derivative mutant AbH12O-A2ΔfhaC were analyzed using an Information Dependent Analysis (IDA) Enhanced MS–Enhanced Resolution (EM ER) method. With this approach a total of 43 proteins from the AbH12O-A2 strain and 55 proteins from the AbH12O-A2ΔfhaC strain were identified in the OMV fraction. The FhaB protein was identified in the wild type sample using the protein database of the AbH12O-A2 strain available in GenBank (protein accession codes from A1S04698.1 to A1S8323.1), whereas the FhaC protein was not identified using this database. Besides, none of these 2 proteins were detected in the mutant sample when the same database was used. However, for the wild type strain, the 2 proteins matched several peptides when a specific database (FhaBFhaC_M.) was used and, as expected, none of these 2 proteins was identified in the mutant samples. The best proteotypic peptides were chosen for the 2 target proteins to create a MRM method where 3 peptides per protein and at least 3 transitions per peptide were considered.

In the AbH12O-A2 samples the 2 proteins were present, the FhaB protein was readily identified by MS/MS spectra but the FhaC protein, which is a minor component, was not identified with IDA under the confidence parameters of the study (Confidence > 99%). However, their peptides were identified and its spectrum of fragmentation was present with less confidence (Confidence > 95%), enough to set the conditions for the targeted analysis. In the MRM study the co-elution of at least 3 transitions for each of the 3 peptides belonging to each protein clearly demonstrates that both FhaB and FhaC are present in the AbH12O-A2 sample, as it is showed in Fig. 3. In the AbH12O-A2ΔfhaC samples, none of these 2 proteins
were observed. In addition, no fragmentation spectrum data are available from these proteins in the AbH12O-A2ΔfhaC sample. As a conclusion, these 2 proteins were not present in the AbH12O-A2ΔfhaC samples, being identified in the AbH12O-A2 samples using targeted proteomics methodology.

Figure 3. (A) Identification of tryptic peptides using SRM/MRM. Extracted ion chromatogram (XIC) of 3 peptides from the protein AbFhaC (FhaC) in the AbH12O-A2ΔfhaC mutant strain (a) and in the AbH12O-A2 wild type strain (b) samples. The name of the co-eluted transitions for each peptide is showed in the legend. (B) Identification of tryptic peptides using SRM/MRM. Extracted ion chromatogram (XIC) of 3 peptides from the protein AbFhaB (FhaB) in the AbH12O-A2ΔfhaC mutant strain (a) and the AbH12O-A2 wild type strain (b) samples. The name of the co-eluted transitions for each peptide is showed in the legend.
The two-partner secretion system AbFhaB/AbFhaC is related to attachment to human alveolar epithelial

Fig. 4 showed that the AbH12O-A2 strain is a poor biofilm former compared with the ATCC 17978 strain (micrographs A and B), being unable to form multilayered biofilm structures on abiotic surfaces. When the fhaC-deficient strain was analyzed (micrograph C) the adherence pattern showed well-separated cells. In this case, no evidence of tridimensional structures was visualized. For the complemented strain, this phenotype was somehow restored (micrograph D) showing some tridimensional cell aggregation, also observed in the wild type AbH12O-A2 strain. However, the low ability of biofilm formation of this strain does not allow a clear interpretation of the role of the AbFhaB/AbFhaC system on biofilm formation on abiotic surfaces.

The AbH12O-A2ΔfhaC isogenic mutant was used to investigate the role of the AbFhaB/AbFhaC system in adherence to biotic surfaces. Thus, A549 human alveolar epithelial cells were infected with A. baumannii AbH12O-A2, the fhaC mutant derivative (AbH12O-A2ΔfhaC) and the complemented mutant (AbH12O-A2ΔfhaC complemented) strains for 3 h (Fig. 1A). Fig. 1A shows that adherence of AbH12O-A2 to A549 was significantly reduced (2.3-fold less, P value < 0.0001) when the fhaC gene was deleted. The adherence phenotype was restored when the fhaC gene was overexpressed in the mutant strain (Fig. 1A). At this point, it is important to remark that the wild type strain and the mutant derivatives grow at the same rate in rich medium without selective pressure (Figure S1).

In agreement with the attachment assay, the AbH12O-A2 wild type strain showed a high attachment phenotype and appeared to form biofilm structures over the A549 cells (micrograph a, Fig. 1B). In addition, the fhaC-deficient AbH12O-A2 strain had reduced adherence to A549 cells and was unable to form such biofilm structures (micrograph b, Fig. 1B) as compared with the parental strain (micrograph a, Fig. 1B). Interestingly, the wild type strain (micrograph a, Fig. 1B) caused more damage to the cell layer than the fhaC-deficient strain (micrograph b, Fig. 1B), demonstrated by the higher loss of the protective surfactant layer, which remained intact in the uninfected cells (micrograph c, Fig. 1B). In agreement with these results, the data obtained through the LIVE/DEAD assay also revealed that the fhaC-deficient strain caused less damage to the A549 cells while the complemented mutant restored the wild type phenotype (Fig. 5).

The two-partner secretion system AbFhaB/AbFhaC interacts with the host cell protein fibronectin

Adherence to extracellular matrix proteins, such as fibronectin, may have a role in bacterial adhesion and internalization.33-36 An analysis of the interaction of the

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Figure 4. SEM visualization of biofilm on abiotic surfaces of the A. baumannii strains (a) ATCC 17978, (b) AbH12O-A2, (c) AbH12O-A2ΔfhaC and d) AbH12O-A2ΔfhaC complemented. Micrographs were taken at 10,000x magnification. Bars indicate the scale marks (2 μm).
The two-partner secretion system AbFhaB/AbFhaC plays a role in virulence

The function of the AbFhaB/AbFhaC system in the virulence of the A. baumannii AbH12O-A2 strain was assessed by a fertility assay using a C. elegans model and a survival assay using a systemic mouse infection model. The C. elegans model showed that worms infected with the A. baumannii AbH12O-A2 strain had a drop of 50% in fertility compared with those worms infected with the knockout strain AbH12O-A2ΔfhaC (Fig. 7). The progeny production was twice higher when C. elegans were infected the fhaC-deficient A. baumannii derivative strain compared with the parental strain AbH12O-A2, being the difference statistically significant. Complementation of the mutant strain with the parental allele restored the wild-type phenotype.

Moreover, the ability of the knockout strain to establish a systemic infection in an experimental murine model was evaluated. To assess the role of this 2-partner secretion system AbFhaB/AbFhaC on virulence, BALB/c mice were intraperitoneally infected with the AbH12O-A2 strain or...
its isogenic knockout strain AbH12O-A2ΔfhaC. The survival rates were monitored for several days as shown in Fig. 8. The results indicated a higher mortality (17 out of 18 mice) associated with wild-type strain after about 20 h of infection (Fig. 8). In contrast, the mortality rate was significantly reduced (12 out of 18 mice) when mice were infected with the mutant strain within the same period of time.

Discussion

The role of *Acinetobacter baumannii* as a nosocomial pathogen is well known, but the molecular aspects of its pathogenicity remain poorly understood. The *A. baumannii* clinical significance is influenced by its extraordinary ability to acquire resistance mechanisms. Besides this emerging resistance profile, *A. baumannii* has an amazing capacity to survive for long periods in a hospital environment, thus giving advantage to its efficiency for dissemination. Adherence is the crucial first step for colonization and therefore for the development of complex communities and it is an important virulence factor for extracellular bacteria that facilitates their persistence in the host. Previous works have analyzed the capacity of several clinical isolates of *A. baumannii* to adhere to surfaces and to form biofilm.\(^1\)\(^5\),\(^2\)\(^1\)\(^)\ Here, we have investigated, using genotypic and phenotypic approaches, the characteristics that may be involved in the persistence and disease potential of the MDR *A. baumannii* AbH12O-A2 isolate, which caused a large outbreak for more than 30 months producing 65 bacteremia episodes.\(^3\) Phenotypic analysis of the AbH12O-A2 isolate revealed its remarkable ability to adhere to human alveolar epithelial cells. However, this *A. baumannii* clinical strain is a non-motile isolate that does not form multilayered biofilm structures on abiotic surfaces, as assessed by SEM.

It has been previously reported that the filamentous hemagglutinin (FHA) participates in biofilm maturing by encouraging the formation of microcolonies in *Bordetella pertussis*.\(^3\)\(^7\) Also, the TPS system described by Darvish *et al.*\(^3\)\(^8\) was shown to be involved in biofilm formation in the strain *A. baumannii* 19606\(^T\). However, the role of the TPS system from the AbH12O-A2 strain in biofilm formation cannot be clearly elucidated in the present work. Although this strain showed a remarkably attachment ability, it is indeed a poor biofilm former not being able to form a mature biofilm, which distress the detection of any biofilm formation change in its isogenic derivative mutants. In contrast, this strain showed a high attachment ability which suggests that the mechanisms of adherence of *A. baumannii* AbH12O-A2 strain to either abiotic or biotic surfaces could be different. This supposition is in agreement with data previously found by Eijkelkamp *et al.*\(^2\)\(^1\) who reported that there is not a mandatory relationship between the capacity to form biofilms and eukaryotic cell adherence.

The genome analysis of the AbH12O-A2 strain\(^3\)\(^1\) revealed that the AbFhaB/FhaC 2-partner secretion system was only present, with the 100% of coverage, in the case of the MDR *A. baumannii* strains 3207 and AB030, being also highly conserved in the case of the strains IOMTU 433, 6200 and SDF. Our genomic analysis suggests a low prevalence of this genomic island in *A. baumannii* spp and showed that the AbFhaB/FhaC operon was located into an 18 kb-region flanked by repetitive sequences. This fact along together with the lower GC content (ca. 38%) suggested that this genome 18 kb-sequence may have been transmitted by lateral transfer as a mobile genetic element to form what is known as a pathogenicity island.\(^3\)\(^9\) AbFhaB/FhaC is a type V 2-partner secretion system with a TPS domain containing 2 proteins; AbFhaB (TpsA protein) and AbFhaC (TpsB protein), where AbFhaB includes the secretion domain and the cleavage site for its transporter AbFhaC. The amino acid sequence analysis of AbFhaB showed similarity to other bacterial TpsA proteins reported, such as *Bordetella pertussis* FHA\(^3\)\(^7\) or *Erwinia chrysanthemi* HecA.\(^4\)\(^0\) Moreover, this protein has a RGD motif, which is found in many adhesive proteins associated with an integrin-mediated attachment to mammalian cells, although its role in pathogenesis of *A. baumannii* has not been determined yet. The predictive structure of AbFhaC protein showed that it is a transmembrane β-barrel channel protein hypothesized to serve as the AbFhaB-conducting pore through the outer membrane. It contains a conserved polypeptide transport-associated domain (POTRA_2), which is expected to participate in
the TpsA recognition mediating the translocation of its TpsA partner across the membrane, or their integration into the outer membranes. Overall, the data suggest that AbFhaC, which belongs to the Omp85-TpsB transporter superfamily, could mediate the secretion of AbFhaB, which ultimately would stay in the outer membrane. Our findings showed that bacteria producing the AbFhaB/FhaC system adhered to A549 cells significantly higher than the fhaC-deficient cells. The functional inactivation of this 2-partner secretion system was addressed by analyzing the OMV proteomic profile from both the wild-type and its derivative mutant AbH12O-A2ΔfhaC strains. Considering the OMVs as a representative sample of the bacterial membrane, we may conclude that AbFhaB is no longer present in the membrane in the absence of AbFhaC.

In previous works, the role of different 2-partner secretion systems from different genera was reported finding that they play an important role in attachment to biotic surfaces. In the present work we demonstrated that when the AbFhaB/FhaC system of the AbH12O-A2 strain was functionally inactivated, the adherence phenotype was significantly reduced. Our results are in agreement with those found by Serra et al. and Darvish et al. wherein they demonstrate that the filamentous hemagglutinin protein has an important function in adherence to epithelial cells in both Bordetella pertussis and A. baumannii ATCC 19606, respectively.

Structural and functional analyses of bacteria-host cell interactions provided insights into cellular processes implicated in the pathogenicity of A. baumannii AbH12O-A2. This strain resulted to show not only a remarkable ability to attach to but also a high capacity to cause serious damage in the A549 human alveolar epithelial cell monolayer, which was assessed both by SEM analysis and cell death assay, probably involving the surface exposure of AbFhaB.

Further analysis on the interplay of A. baumannii with host cells revealed that the AbFhaB/FhaC system could be involved in fibronectin-binding. Adhesion to extracellular matrix proteins, such as fibronectin, gives an advantage to pathogens to adhere to host cells, which is related with microorganism’s virulence.

It has been reported that fibronectin is an important receptor for several bacterial species. Smani et al. confirmed that A. baumannii has specific linkages for fibronectin revealing that fibronectin could act as a host receptor for A. baumannii cells. Here, we reported that fibronectin could act as a host receptor for the MDR A. baumannii AbH12O-A2 strain since the binding to immobilized fibronectin was shown to be mediated by the AbFhaB/AbFhaC system. Indeed, its inactivation reduced significantly the attachment of the AbH12O-A2 strain to immobilized fibronectin. Thus, we demonstrated that this AbFhaB/AbFhaC TPS system plays a clear role in fibronectin-mediated adherence of the A. baumannii AbH12O-A2 isolate.

Several A. baumannii clinical isolates caused pneumonia, sepsis and soft tissue infections in murine experimental infections. Since the AbFhaB/AbFhaC 2-partner system had a key role in adherence, we wanted to evaluate its involvement on A. baumannii virulence. To address this issue, 2 experimental infection models were performed demonstrating the importance of this 2-partner system in virulence. The fertility assay performed in C. elegans has been previously validated as a sensitive and reproducible model to determine subtle differences between isogenic derivative strains of A. baumannii ATCC 17978. Data revealed that the fhaC gene deletion caused a significant increase in the fertility of the worm, showing the implication of the AbFhaB/AbFhaC 2-partner system in A. baumannii pathogenesis. The results obtained using an acute systemic infection model also supported the involvement of the AbFhaB/AbFhaC 2-partner system of the A. baumannii AbH12O-A2 strain in virulence. These results are in agreement with those found by Melvin et al. who reported that FhaB is involved in Bordetella virulence.

The AbFhaB/AbFhaC 2-partner system of the A. baumannii AbH12O-A2 strain plays a role in virulence as confirmed by the infection models mediating the adhesion of bacteria to human epithelial cells. These findings could help to encourage the development of new therapeutic options.

**Materials and methods**

**Bacterial strains, culture conditions, plasmids and DNA extraction**

A. baumannii AbH12O-A2 was isolated at the Hospital 12 de Octubre, Madrid (Spain). A. baumannii and E. coli strains were grown in LB broth, adding 2% of agar when necessary. Strains were grown at 37°C and 180 rpm. The storage was done at −80°C in LB broth with 10% glycerc. Sodium tellurite (Sigma-Aldrich, St. Louis, MO) was used for selection of transformant strains at a concentration of 10 μg/mL for E. coli and 30 μg/mL for A. baumannii. Genomic DNA was obtained using the Wizard Genomic DNA purification Kit (Promega Corporation, Madison, WI, USA). Plasmids were extracted using the High Pure Isolation Kit (Roche Diagnostics GmbH, Mannheim, Germany). E. coli TG1 was employed for cloning approaches. A. baumannii ATCC 17978 was used as a control strain for adhesion and SEM analysis.
The plasmid pMo130-Tel\(^R\) kindly gave up by Chua Kim Lee (University of Singapore) harbouring a tellurite resistance gene was used to generate the double crossover knockout derivatives as previously described.\(^{48}\) The pWH1266-Tel\(^R\) plasmid was used to genetically complement the mutant strain was obtained by cloning the tellurite resistance gene with its own promoter sequence into the pWH1266 plasmid.\(^{51}\) The pWH1266-Tel\(^R\) plasmid used to genetically complement the mutant derivative strain lacking the \(fhaC\) gene. The \(fhaC\) gene was PCR-amplified including its own promoter sequence (locus \_tag ACIAD2922), amplified by PCR from A. \(\text{calcoaceticus}\) ADP1 using primers listed in Table 1 into the \(Hind\text{III}\) site of the pWH1266 plasmid.\(^{49}\)

### Construction of knockout strain

A mutant derivative strain lacking the \(fhaC\) gene (AbH12O-A2\(\Delta fhaC\)) was constructed using the suicide plasmid pM0130-Tel\(^R\), which contains the genes \(xylE, sacB\), and the kanamycin and tellurite resistance cassettes, as detailed previously.\(^{48}\) Briefly, fragments of 608 bp and 889 bp located upstream and downstream of the POTRA\_2 domain of the LX00\_12105 locus (\(fhaC\) gene), respectively, were inserted into the pM0130-Tel\(^R\) vector using the oligonucleotides detailed in Table 1. The construction obtained was introduced in A. \(\text{baumannii}\) AbH12O-A2 by electroporation. Recombinant colonies were selected using tellurite and cathecol, following the method previously described by Hamad et al.\(^{50}\) Colonies representing the second crossover events were obtained as described previously\(^{48,50}\) and checked by PCR using the oligonucleotides detailed in Table 1.

### Complementation of the mutant derivative strain AbH12O-A2\(\Delta fhaC\)

The \(fhaC\) gene was PCR-amplified including its own promoter from AbH12O-A2 strain genomic DNA and then cloned into the \(SalI\) restriction site of the pWH1266-Tel\(^R\) plasmid for complementation using the primers listed in Table 1. This plasmid is a derivative of the pWH1266\(^{51}\) that harbors tellurite resistance marker which was PCR amplified as described above. The new derivative pWH1266-Tel\(^R\)-\(fhaC\) was introduced in the AbH12O-A2\(\Delta fhaC\) strain by electroporation. Transformants were selected on 30 \(\mu\g/mL\) tellurite plates and PCR was used for checking the constructions using oligonucleotides detailed on Table 1.

### RNA extraction and real-time RT-PCR

RNA isolation and expression level determination of the target genes were performed following the procedures and using the materials previously described by Álvarez-Fraga et al.\(^{48}\) The oligonucleotides and probes used are detailed in Table 1.

### Isolation and purification of OMVs

OMVs were isolated following the protocol described by Méndez et al.\(^{30}\) Briefly, cells were centrifuged and the supernatant was filtered to remove residual bacteria. The OMV pellet was obtained by ultracentrifugation and then resuspended in PBS. The OMV suspension was

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**Table 1. Oligonucleotides and probes used in the present work.**

| Primer/Probe name | Sequence | Use in the present study |
|-------------------|----------|--------------------------|
| LX00\_12105-Up-NotI-Fw | ggcgcgccgccataagaaagaa | Construction of AbH12O-A2\(\Delta fhaC\) knockout strain |
| LX00\_12105-Up-BglI-Rv | gcagacgtttctagctagtaag | Construction of AbH12O-A2\(\Delta fhaC\) knockout strain |
| LX00\_12105-Down-BglI-Fw | gcagacgtttctagctagtaag | Construction of AbH12O-A2\(\Delta fhaC\) knockout strain |
| LX00\_12105-Down-SphiI-Rv | gcagacgtttctagctagtaag | Construction of AbH12O-A2\(\Delta fhaC\) knockout strain |
| LX00\_12105-ext-Fw | gttgaaagttcagtaagc | Construction of the AbH12O-A2\(\Delta fhaC\) knockout strain |
| LX00\_12105-ext-Rv | cggtttctagctagtaag | Construction of the AbH12O-A2\(\Delta fhaC\) knockout strain |
| LX00\_12105-int-Fw | ccaacacgctgcaacagc | Construction of the AbH12O-A2\(\Delta fhaC\) knockout strain |
| LX00\_12105-int-Rv | gttgacgtgcaaaatgac | Construction of the AbH12O-A2\(\Delta fhaC\) knockout strain |
| pMo130 site2 Fw | attcatgaccgtgctgac | Construction of the AbH12O-A2\(\Delta fhaC\) knockout strain |
| pMo130 site2 Rv | cttgtctgttagcag | Construction of the AbH12O-A2\(\Delta fhaC\) knockout strain |
| LX00\_12105-\(Sal\)Fw | gcgcgcgccgcccatagaaagaa | Cloning of the \(\text{fa}\)C gene into the pWH1266 plasmid for complementation |
| LX00\_12105-\(Sal\)Rv | gcgcgcgccgcccatagaaagaa | Cloning of the \(\text{fa}\)C gene into the pWH1266 plasmid for complementation |
| Tel\(^R\)-\(Pprom\)-\(Hind\text{III}\)Fw | gcgccctgatacgttagctagtaag | Cloning of the \(\text{fa}\)C gene into the pWH1266 plasmid for complementation |
| Tel\(^R\)-\(Pprom\)-\(Hind\text{III}\)Rv | gcgcgcgccgcccatagaaagaa | Cloning of the \(\text{fa}\)C gene into the pWH1266 plasmid for complementation |
| RT73\_LX00\_12105-Fw | caaaaaggttgtgtaacacgga | qRT-PCR |
| RT73\_LX00\_12105-Rv | cacgcgcgccgcccatagaaagaa | qRT-PCR |
| RT3\_LX00\_12100-Fw | ctcgtcaagcttgatacgttagctagtaag | qRT-PCR |
| RT3\_LX00\_12100-Rv | ctcgtcaagcttgatacgttagctagtaag | qRT-PCR |
| RT76\_gyrB-Fw | gttgcttgatgcaagatatggaaggtc | qRT-PCR |
| RT76\_gyrB-Rv | gggtatactgcttcgacgttc | qRT-PCR |
filtered and processed for protein extraction. Trichloroacetic acid was added to precipitate the proteins from the OMV suspension. Then, for proteomic approaches, proteins were solubilized and their concentration was measured using the Bio-Rad protein assay (Bio-Rad, Munich, Germany).

**In-solution protein digestion**

In-solution digestions were performed as described by Méndez et al. Two biological samples of the AbH12O-A2 and AbH12O-A2ΔfhaC were analyzed. In all cases the resulting tryptic peptide mixture was acidified with 1 µL of 10% of trifluoroacetic acid and desalted using home-made stage-tips as previously described. Samples were aliquoted, subjected to a speed-vacuum and kept at −80°C until used.

**Nanoscale liquid chromatography and mass spectrometry**

Both enhanced MS (EM) and enhanced resolution (ER) methods were performed injecting 2 µg of AbH12O-A2 and AbH12O-A2ΔfhaC samples in the LC-MS/MS, using a nanoLC system (Tempo, Eksigent, Dublin, (CA), USA coupled to a 5500QTRAP instrument (ABSciex). After precolumn desalting using a C18 column, (5 µm, 300A, 100 µm, 2 cm, Acclaim PepMap, Thermo Scientific, USA) at a flow of 3 µL/min during 10 min, tryptic digests (2 µg) were separated on C18 nanocolumns (75 µm id, 15 cm, 3 µm particle size) (Acclaim PepMap 100, Thermo Scientific, USA) at a flow rate of 300 nL/min. A standard 120 min gradient from 5 to 40% of buffer B (0.1% formic acid in 95% acetonitrile) was used.

**Peptide selection for multiple reaction monitoring (MRM)**

The proteotypic peptides detected with the highest spectral counts and that follow the selection criteria, (fully tryptic, with no missed cleavages, unique to a particular protein, length between 8 and 30 amino acids) were chosen for MRM assay development using Skyline target proteomics Environment. We selected the top transitions for MRM assay development using Skyline target version 2.0.4 (http://www.psort.org/psortb2/index.html) was used for prediction of sub-cellular locations of the proteins.

**Adhesion to A549 human alveolar cells**

A549 epithelial cells were grown in 5% CO₂ at 37°C in DMEM medium (Sigma-Aldrich) with 10% heat-inactivated fetal bovine serum and 1% of penicillin-streptomycin (Gibco). Confluent monolayers were washed following the protocol previously described. The multiplicity of infection (MOI) of 10 was used; in each
well 10⁴ A549 cells were infected with 10⁵ bacterial cells. Infected cells were then incubated during 3 h at 37°C in mHBSS medium. Bacterial adhesion was determined as previously described by Álvarez-Fraga et al.⁴⁸ The number of colony forming units (CFUs) was established to assess the percentage of bacteria attached to eukaryotic cells in comparison with a growth control (bacteria grown in HBSS and 37°C for 3 h without eukaryotic cells). T-student tests were done to determine the statistical significance of the differences. Six biological replicates were performed.

**Scanning electron microscopy (SEM)**

SEM analysis of biofilms formed on polystyrene coverslips was performed following the methods described by Gaddy et al.²² Briefly, A. baumannii cultures were incubated with sterile polystyrene coverslips for 48 h at 37°C without shaking and prepared for SEM.²² A Zeiss Supra Gemini Series 35V scanning electron microscope was used to view each sample.

**SEM analysis of bacterial attachment to A549 human alveolar epithelial cells infected with A. baumannii**

A549 human alveolar cells were maintained as previously described.²² Human epithelial cells were grown at a liquid-air interface following the methods described by Álvarez-Fraga et al.⁴⁸ Briefly, each membrane was seeded with 10⁵ eukaryotic cells. Cells were grown and processed during 3 weeks. Then, each membrane containing the A549 cells was infected with a 1 µL suspension of 10⁶ bacteria in HBSS, as previously described.⁴⁸ After 72 h of infection, the membranes were washed, fixed with 4% formaldehyde-HBSS, and prepared for SEM using the previously described methods.¹⁹⁴⁸

**Cell death assay on infected A549 monolayers**

A549 human alveolar epithelial cells were grown with 10% CO₂ at 37°C in DMEM supplemented with 100 mg/L of penicillin, 10% fetal bovine serum, and 100 mg/L streptomycin. For LIVE/DEAD assays, A549 cells were grown to a density of 1 × 10⁵ cells per well in 24-well plates. Then, the cells were infected with 3.6 × 10⁸ CFU/well of A. baumannii in HBSS without antibiotics and grown for 24 h at 37°C.⁵⁴ The cell death assay was performed as previously described by Alvarez-Fraga et al.⁴⁸ using the LIVE/DEAD Cell Double Staining Kit (Sigma-Aldrich) according to the manufacturer’s instructions. Cell viability was measured after infection. Cell viability was determined as the number of red cells related to the total cells visualized. The results were presented as a percentage. Each assay was done by triplicate and Student’s t test was used to determine the statistical significance.

**Fibronectin-binding assay**

The fibronectin-binding assay was performed as described previously.³⁶⁵⁵ Briefly, wells were coated with fibronectin. After washing and blocking the wells bacteria were added. Controls were done with BSA and PBS as previously described.³⁶ Overnight cultures of A. baumannii AbH120-A2 and derivative strains grown in LB at 180 rpm and 37°C were used. Grown bacteria were washed 3 times in PBS and resuspended in a double-volume of the same buffer. One-hundred µL of bacterial suspension were added to each coated or non-coated wells (controls) and incubated 3 h at room temperature. Then, non-adhered bacteria were removed by washing the plates 6 times with PBS. Adherent bacteria were then collected with PBS and Triton X-100 as previously described.³⁶ Finally, bacterial suspensions were plated onto LB agar and incubated at 37°C for 24 h in order to determine the number of CFU and the percentage of bacteria that had attached to immobilized fibronectin compared with the growth control (bacteria grown in LB at 37°C for 3 h in non-coated wells). Four independent replicates were performed. Student’s t test was done and bars indicate the standard deviation.

**Caenorhabditis elegans virulence assay**

Fertility assays were performed as previously described.⁴⁷ Briefly, both the AbH120-A2 and the mutant derivative strains were grown overnight in LB and then cultured at 37°C for 24 h in nematode growth medium (NG). The eggs of C. elegans N2 Bristol were hatched in M9 medium and L1 worms were arrested overnight at 20°C. Then, the L1 worms were incubated in NG medium plates in presence of the bacterial strains. One L4 worm was placed on a peptone-glucose-sorbitol medium (PGS) plate individually seeded with each A. baumannii strain and incubated at 25°C for 24 h. The worms were transferred to new plates seeded with the same bacterial strain and the worm progeny was counted daily for 3 d to determine their viability. Six independent replicates were done with each strain. The differences were statistically evaluated by using Student’s t test. Means of the differences between strains are reported.
**Virulence assays in a mouse systemic-infection model**

*A. baumannii* strains were evaluated for virulence in terms of mortality and survival time of infected mice as previously described. Groups of 18 BALB/c female mice, weighting 20–25 g, 6–9 weeks old, were inoculated by intra-peritoneal injection with 250 μL of bacterial suspension with $33 \times 10^{7}$ CFU/mouse of the wild type strain or $46 \times 10^{7}$ CFU/mouse of the knockout strain. Mice were monitored for signs of disease during a week. Animals that survived after this period were euthanized. Results were analyzed using the log rank test. A $P$ value of 0.05 was considered statistically significant. Mice were maintained in the specific pathogen-free facility at the Technology Training Center of the Hospital of A Coruña (CHUAC, Spain). All experiments were done with the approval of and in accordance with regulatory guidelines and standards set by the Animal Ethics Committee (CHUAC).

**Bioinformatic analysis**

The MAUVE 2.3.1 software was used for multiple genome analysis. Functional analysis of proteins were done using the The Interpro program. The Phyre2 server was used for predicting the 3-dimensional structure of the protein. VectorNT11 (Thermo Fisher Scientific) was used to analyze the DNA sequences and the genetic constructions. The blast tool of the National Center of Biotechnology Information was used for genetic analysis procedures.

**Disclosure of potential conflicts of interest**

No potential conflicts of interest were disclosed.

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