New OprM structure highlighting the nature of the N-terminal anchor

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Among the different mechanisms used by bacteria to resist antibiotics, active efflux plays a major role. In Gram-negative bacteria, active efflux is carried out by tripartite efflux pumps that form a macromolecular assembly spanning both membranes of the cellular wall. At the outer membrane level, a well-conserved outer membrane factor (OMF) protein acts as an exit duct, but its sequence varies greatly among different species. The OMFs share a similar tri-dimensional structure that includes a beta-barrel pore domain that stabilizes the channel within the membrane. In addition, OMFs are often subjected to different N-terminal post-translational modifications (PTMs), such as an acylation with a lipid. The role of additional N-terminal anchors is all the more intriguing since it is not always required among the OMFs family. Understanding this optional PTM could open new research lines in the field of antibiotics resistance. In Escherichia coli, it has been shown that CusC is modified with a tri-acylated lipid, whereas TolC does not show any modification. In the case of OprM from Pseudomonas aeruginosa, the N-terminal modification remains a matter of debate, therefore, we used several approaches to investigate this issue. As definitive evidence, we present a new X-ray structure at 3.8 Å resolution that was solved in a new space group, making it possible to model the N-terminal residue as a palmitoylated cysteine.

Keywords: multidrug resistance, efflux pump, membrane protein, post-translational modification, lipoic acid, X-ray structure

Introduction

After several decades of continuous antibiotic therapy success, we are now facing the appearance of multi-drug resistant strains and the near absence of new antibiotic family development for more than 10 years (Fischbach and Walsh, 2009; Hede, 2014). These facts highlight the need for new anti-infection strategies (Walsh, 2003; Olivares et al., 2013), although a promising compound isolated from natural soil bacteria that is able to kill Gram-positive pathogens, was recently reported (Ling et al., 2015). Among the most virulent nosocomial pathogens are Pseudomonas aeruginosa, Escherichia coli, Staphylococcus aureus, Enterococci, and Acinetobacter baumannii (Poole, 2004; Lister et al., 2009; Bereket et al., 2012; Bayram et al., 2013). These strains have developed several resistance strategies including active efflux pumps (Cattoir, 2004; Li and Nikaido, 2009; Nikaido, 2009; Nikaido and Pages, 2012). In Gram-negative bacteria efflux pumps are multimers of three

Abbreviations: βOG, beta-octyl glucopyranoside; DDM, dodecyl maltoside; MexAp, MexA palmitoylated; MexAnp, MexA non-palmitoylated; PTM, posttranslational modification.
different proteins that form a long transmembrane scaffold linking the cytoplasm to the outside of the cell (Nikaido and Pages, 2012). These tripartite assemblies are composed of an inner membrane protein from the RND (Resistance Nodulation cell Division) family, which corresponds to the pumping motor that uses the proton-gradient as an energy source; an outer membrane channel from the OMF (Outer Membrane Factor) family; and a periplasmic protein from the MFP (membrane fusion protein) family, anchored to the inner membrane and connecting the other two proteins. In P. aeruginosa, up to 12 different pumps have been sequenced (Yen et al., 2002), including OprMOMF-MexA MFP- MexB RND which is one of the most studied pumps because of its constitutive expression whereas the others appear under antibiotic pressure (Li and Poole, 2001; Nakajima et al., 2002; Schweizer, 2003). This study will focus on the versatile membrane channel OprM, which has the ability to work with at least four different pumps, including OprM-MexAB, OprM- MexXY (Aires et al., 1999; Morita et al., 2012), OprM-MexJK (Chuan chuen et al., 2002), and OprM-MexMN (Mima et al., 2005).

The X-ray structure of OprM was solved in two different space groups showing its trimeric nature [Protein Data Bank (PDB) code: 1WP1 (Akama et al., 2004); 3DSK (Phan et al., 2010)]. The architecture of OprM is composed of a beta-barrel domain that is ~40 Å in height spanning the outer membrane and a periplasmic alpha-helical domain that is ~100 Å in length and bears a central buoy. Most of the structure has been determined with the exception of the eleven C-terminal amino acids, which are not visible in the electronic density maps of both structures. In addition, the structure of the post-translational modification (PTM) that covalently links the Cys-terminal residue to a lipoyl has never been properly characterized at any resolution, despite palmitoylation being suggested some time ago (Nakajima et al., 2000). This lipoyl modification is not commonly shared among OMF family members. For instance, the E. coli homolog TolC (PDB code: 1EK9, Koronakis et al., 2000), has an N-terminal that is 44 residues shorter and does not begin with a cysteine. Other OMFs with known structures, such as VceC from Vibrio cholerae (PDB code: 1YC9, Federici et al., 2005), CusC of the metal effluent pump from E. coli (3PIK, Kulathila et al., 2011; 4K7R, Lei et al., 2014a) and CmeC from E. coli (4MT4, Su et al., 2014) begin with residues Cys-Thr (Figure 1; Supplementary Figure S1) like OprM. The OMF protein with the most recently solved structure, MtrE (4MT0, Lei et al., 2014b) from Neisseria gonorrhoeae, begins with Cys-Thr. The structure of CusC indicates the presence of a di-acylated thiol on the N-terminal cysteine, with or without a supplementary acyl chain on the N-terminal amine (triacylation), depending on the protomer. As the CusC sequence is highly similar to OprM, it has been suggested that the latter may be modified in a similar manner, rather than with only one palmitoyl chain attached by a thio-acyl bond to the N-terminal Cys as previously described (Kulathila et al., 2011). Di-acylation is also found in CmeC, but nothing has been found to be added to the N-terminal cysteine of the MtrE structure. In the VceC structure, the protein sequence was cloned without the first 12 amino acids, therefore, the N-terminal cysteine was not present. In the OprM structure solved in the R32 space group (Akama et al., 2004), the three monomers were found to be identical by the threefold crystallographic symmetry axis, and despite a resolution of 2.6 Å there was no evidence of lipidation to the thiol or amine of the N-terminal cysteine. The same observation applies to the OprM structure solved at 2.4 Å of resolution (Phan et al., 2010) in which the P212121 space group does not stabilize sufficiently the N-terminal modified cysteine.

Consequently, the N-terminal modification of OMFs remains an open question. Post-translational lipidation is particularly essential for the secretion and localization of some membranous proteins, a process involving different biological modifiers such as palmitoyl acyl transferases (Aicart-Ramos et al., 2011), palmitoyl thioesterases, lipoprotein diacylglycerol transferase or lipoprotein N-acyl transferase (Linder and Deschenes, 2007; Kovacs-Simon et al., 2011; Nakayama et al., 2012). These modifying enzymes could potentially lead to new therapeutic targets. In addition, it is not known why some OMF proteins need to undergo PTM in addition to their trans-membrane beta-barrel insertion, or what is the nature of this modification. We asked this question using OprM from P. aeruginosa with specific N-terminal chemical probes to investigate whether it is actually palmitoylated via a thio-acyl or subjected to other PTMs, and resolved a new OprM structure in a new crystallographic space group.

Materials and Methods

Expression and Purification of OprM, MexAp, and MexAnp

The three proteins were produced following the protocol described by Phan et al. (2010) and Ferrandez et al. (2012) with some modifications. The protein genes were inserted into the pBAD33-GFPuv plasmid with a C-terminal 6-histidines tag and the N-terminus extremity being dedicated to the signal peptide. For the non-palmitoylated form of MexA (MexAnp), the signal peptide was deleted from the construct, resulting in a non-membranous protein bearing a free N-terminal cysteine. Wt MexA has the same amino acid sequence as MexAnp, but its starting cysteine is palmitoylated via a thio-acyl or subjected to other PTMs.

The plasmids were transformed into the C43-DE3 E. coli strain (Miroux and Walker, 1996). For each protein 6 L of cultures were grown; they were begun at OD600 = 0.05 from dilution of an overnight pre-culture at 37°C in LB medium containing 25 μg/ml chloramphenicol, and were then grown at 30°C. Cells were induced at OD600 = 0.6–0.8 by the addition of 0.02% L-arabinose and grown for 2 h before centrifugation for 20 min at 9,000 g. The cell pellet was resuspended in 45 ml of buffer containing 20 mM Tris-HCl (pH 8.0), 5 mM MgCl2, 1 μl/ml cocktail of anti-protease inhibitors set III (Calbiochem) and 50 units of benzonase (Promega).

Cells were broken using a French pressure cell at 69 MPa and then centrifuged twice for 30 min at 8,500 g to remove inclusion bodies and unbroken cells. As for OprM, the supernatant was applied to a sucrose step gradient (0.5 and 1.5 M) and then centrifuged for 3 h at 200,000 g at 4°C for membrane separation. The pellet, corresponding to the outer membrane fraction, was...
FIGURE 1 | Sequence alignment of the six outer membrane factor (OMF) proteins with known structures. Only the N- and C-terminal portions of the alignment (extract from Supplementary Figure S1) corresponding to the most divergent 3D structure regions of these OMF proteins are shown. The numbering corresponds to the OprM sequence after cleavage of the targeting signal presented in grey letters. The secondary structure of OprM is indicated at the top of the aligned sequences. The red brackets indicate the beginning and end of each protein resolved PDB structure.

re-suspended in a solution containing 20 mM Tris-HCl (pH 8.0), 10% glycerol (v/v) and 2% βOG (w/v) (Anatrace) and then stirred overnight at 23°C. The solubilized membrane proteins were recovered by centrifugation for 30 min at 50,000 g. For MexAp, the lysis supernatant was directly centrifuged at 100,000 g for 1 h at 4°C, and the pellet was resuspended in a solution containing 20 mM Tris-HCl (pH 8.0), 10% glycerol (v/v), 2% βOG (w/v) (Anatrace) 0.2% N-lauryl sarkosyl, and 15 mM imidazole, then stirred overnight at 23°C. The solubilized membrane proteins were recovered by centrifugation at 4°C for 1 h at 100,000 g. For MexAnp, because it is produced directly in the cytoplasm, there is no need for a detergent solubilization step. After lysis and centrifugation, 15 mM imidazole was added to the supernatant before loading onto the column. For the three proteins, the same protocol was then used. The proteins were loaded onto a Ni-NTA resin column pre-equilibrated with 20 mM Tris-HCl (pH 8.0), 200 mM NaCl, 10% glycerol (v/v), 15 mM imidazole for MexAnp, the same buffer plus 0.9% βOG (w/v) for OprM, and the addition of 0.2% N-lauryl sarkosyl for MexAp. After washing of the column, the proteins were eluted in the same respective buffers containing 300 mM imidazole, and then desalted using a PD-10 desalting column (GE) to remove the imidazole. MexAp and MexAnp were concentrated up to 2.5 mg/ml. OprM was concentrated up to 8 mg/ml using the 30-kDa cutoff Amicon system (Millipore).

Labeling of the N-Terminal Cysteine Sulfur

The fluorescent compound MTS-EMCA [N-(2-Methanethiosulfonyl)ethyl]-7-methoxycoumarin-4-acetamide, Toronto Research, Chemicals Inc.] is a sulfhydryl active reagent that covalently attaches to the reduced cysteine via a disulfide bond. 25 mM proteins (MexAp, MexAnp, and OprM) were incubated for 2 h with 2 mM MTS-EMCA in 10 mM HEPES pH 7.5, 150 mM NaCl, βOG 1% (w/v), at room temperature away from light. Reactions were stopped by adding an equivalent volume of SDS-PAGE loading buffer. After electrophoresis, gels were visualized with a UV transilluminator at 312 nm before coomassie blue staining. MTS-EMCA fluorescence appeared for the labeled proteins.

Crystallization and Data Collection

OprM crystals were grown by vapor diffusion using the hanging drop method. Two different crystal forms were obtained, both leading to the recording of a diffraction dataset after 100s of crystal tests on the different beam lines of both SOLEIL and ESRF synchrotrons. One crystal was obtained in a 1 M sodium citrate (pH 5-6) precipitation solution and was found to belong to the P2₁2₁2₁ space group. The structure in this space group has been previously published (Phan et al., 2010). The second
Crystal was obtained in 100 mM sodium acetate (pH 4.5), 6% PEG 20 000 (w/v), 300 mM ammonium citrate, 25-30% glycerol (v/v), and 0.9% BOC (w/v). These rhombohedral crystals (100 μm × 100 μm × 30 μm) belong to the C2 space group and diffracted to 3.8 Å resolution. A complete dataset was collected on beamline ID29 (ESRF, Grenoble) with an exposure time of 10 s per degree of oscillation. Owing to its low resolution, this data set has been kept for a long time without solving the structure, but the recent question of the N-terminal modification nature prompted us to ultimately solve the OprM structure in the C2 space group.

### Data processing, Model Building, and Refinement

Reflections were integrated, scaled and reduced using the programs XDS (Kabsch, 1993) and TRUNCATE from the CCP4 suite (1994). Data collection statistics are summarized in Table 1.

We have solved the C2 structure of OprM by molecular replacement with PHASER (McCoy et al., 2007) in automatic mode using our previously solved structure (PDB code 3D5K, Phan et al., 2010) as a model. Refinements were conducted using Phenix (Adams et al., 2002) and the protein was rebuilt with COOT (Emsley and Cowtan, 2004). The Bfactors were refined by groups. These groups were determined after a refinement step using the two Bfactors per residues option, but the resulting structure was not kept for the continuation. The TLS option was not used.

The last 19 C-terminal residues could not be assigned, probably due to the large flexibility of this region. The validity of our model was checked using MolProbity (Davis et al., 2007) and the polygon tool (Urzhumtseva et al., 2009) from Phenix (see Supplementary Figure S3). The OprM structure model of the C2 crystal was deposited in the PDB (4Y1K).

Figures were created with PyMol (DeLano, 2002).

### Results

### Chemical Analysis of the Lipoyl Position

Among the different PTMs that can occur on an N-terminal cysteine (Chalker et al., 2009) N- or S-palmitoylation or acetylation are readily observed (Resh, 1999; Tooley and Schaner Tooley, 2014). As these different modifications are regulated by specific transferases, it is important to characterize the exact nature of OprM PTM. Thus, two questions needed to be clarified: what is the chemical nature of the modification, and which group of the amino acid is modified? To address these two questions, two different types of chemical labeling were performed on purified OprM. To analyze the occupancy of the N-terminal amine, this protein was labeled with the fluorogenic molecule 4-chloro-7-nitrobenzofurazan (NBD-Cl) at neutral pH. This molecule has been proven to be a specific probe for the N-terminal amine only (Bernal-Perez et al., 2012). After incubation with NBD-Cl, the protein was analyzed on an SDS gel, revealing a bright fluorescent band when exposed at 475 nm (Figure 2A), thus showing that the N-terminal amine of OprM is actually accessible.

Following the same approach, a different probe was used to verify the occupancy of the sulfhydryl group of the cysteine. The fluorescent compound MTS-EMCA, which specifically links to this group, was used to label OprM. As a control we also analyzed MexA (wt palmitylated MexA), MexAnp (a mutated form of MexA lacking the signal peptide and bearing a free N-terminal cysteine) and MexAnp in the presence of DTT to reverse thiol acylation. After migration on an SDS gel, UV exposure of the gel at 312 nm (Figure 2B, lower panel) revealed a bright band for MexAnp and a faint band for MexAnp with DTT but no signal for OprM, demonstrating that the N-terminal cysteine of OprM is occupied on its thiol by lipoyl modification. Therefore, it can be concluded that the lipoyl modification is anchored exclusively to the sulfhydryl group of OprM.

### Sequence Alignment

Sequence alignment of the different OMF proteins whose structures were deposited in the PDB was performed using the program MUSCLE (MULTiple Sequence Comparison by Log-Expectation)\(^1\). The alignment was submitted to ESPript 3.0 (Robert and Gouet, 2014) for customization.

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\(^{1}\)http://www.ebi.ac.uk/Tools/msa/muscle/
As a third approach, measuring the precise protein mass has also been considered. Indeed, in our case, one needs to distinguish between a palmitoyl (chemical composition $\text{C}_{16}\text{H}_{32}\text{O}_2$ resulting in a mass of 256 Da) and a tri- or di-acyl that can adopt a variable length. As an example, in the CusC structure can be found a tri-acyl composed of $\text{C}_{19}\text{O}_5\text{H}_{33}$ (PDB code: 3PIK) resulting in a mass of 341 Da, or a di-acyl (PDB code: 4K7R) of chemical composition $\text{C}_{14}\text{O}_4\text{H}_{25}$, resulting in a mass of 257 Da, which is close to the mass of a palmitoyl. This result illustrates how difficult it is to obtain an answer using this technique as several combinations result in the same mass and it is necessary to measure mass with a precision as high as one Dalton, which is far to be routinely achieved to date with membrane proteins of that size. Attempts to address this question in the proteolized protein using the electrospray and MALDI techniques have been unsuccessful because the N-terminus peptide was not detected despite the use of different protease enzymes, and even thought 90% of the OprM sequence was covered by the analysis (data not shown).

**OprM Crystal Structure in the C2 Space Group and Comparison with the Structure Solved in P2$_1$$2_1$$2_1$**

As the question about the nature of the modification remained unanswered, it has been envisaged to refine the OprM crystallographic structure in a different space group, as different crystal packing could stabilize the N-terminus and eventually reveal the complete structure of the added lipid. We previously solved the structure of OprM at 2.4 Å resolution in the P2$_1$$2_1$$2_1$ space group (Phan et al., 2010) but this structure showed only the beginning of the N-terminal cysteine (lane 3), and boiled OprM (lane 4). A coomassie blue coloration shows all of the purified proteins. Fluorescence visualization of NBD-Cl ($\lambda = 475$ nm) that shows that the N-terminal amine of OprM is available (A) whereas visualization of MTS-EMCA ($\lambda = 312$ nm) shows that the N-terminal thiol of OprM is not available for labeling (B) and is thus occupied by the palmitoyl moiety. Only MexAnp with a free N-terminal thiol is available for MTS-EMCA labeling (B).

As the question about the nature of the modification remained unanswered, it has been envisaged to refine the OprM crystallographic structure in a different space group, as different crystal packing could stabilize the N-terminus and eventually reveal the complete structure of the added lipid. We previously generated several OprM datasets in the C2 space group, but they were set aside without solving their structures owing to their lower resolution. To investigate the lipoyl structure, we decided to solve the structure of the best diffracting dataset limited to 3.8 Å resolution.

The here-solved structure comprises two trimers in the asymmetric unit with the second trimer being poorly defined. The crystal packing of our C2 OprM structure is slightly different from the P2$_1$$2_1$$2_1$ structure (see Supplementary Figure S2) but they share common type I crystal packing in which the homo-trimer channels interact in a head-to-head manner through the hydrophobic beta-barrel domains mimicking a lipid bilayer plane. Despite its higher resolution (2.4 Å), the crystal structure of OprM in the P2$_1$$2_1$$2_1$ space group does not reveal the entire palmitoyl moiety because all the three N-termini are oriented toward the solvent and this results in high thermal motions. For two monomers, the closest amino acid is located more than 10 Å away from the expected palmitate main chain, and for the third, only the cysteine portion is clearly constrained, but this is not sufficient to build in the palmitoyl tail. In contrast, when stabilized by a more packed environment around the N-terminus, the entire fatty acid chain appears in the density map of the C2 form where two different orientations are observed (Figure 3). This is the sole, but crucial, advantage of this structure, because the crystal packing of OprM is equivalent to that previously published in the R32 space group (Akama et al., 2004) with the exception that the three monomers are not linked by crystallographic symmetries. Attempts were made to highlight some eventual local differences between the monomers even if at low resolution. Superposition of the six different
FIGURE 3 | Palmitoyl modified N-terminal cysteine of OprM. Packing of the C2 crystal structure of OprM showing the interactions around the palmitate from monomers A (A) and B and C (B). The red lines indicate an artificial membrane. Enlargements show the respective electron densities of the palmitoyl modified cysteines (contour at 1 and 0.8 sigma, respectively).

monomers from the C2 asymmetric unit onto monomer A demonstrates a mean Cα-atom RMSD of 0.2 Å for monomers B and C, and 0.58 Å for monomers D, E, and F. These monomer superposition values have to be compared to those obtained for the two other structures of OprM in different space groups (a mean RMSD of 0.50 Å for 440 Cα atoms in both cases) and that of the other OMF solved structures (a mean RMSD of 0.50 Å for 440 Cα atoms in both cases) and that of the other OMF solved structures (a mean RMSD of 0.50 Å for 440 Cα atoms in both cases) and that of the other OMF solved structures (a mean RMSD of 0.50 Å for 440 Cα atoms in both cases). Although no striking differences among the alternative crystal structures of OprM monomers were revealed, this analysis highlights the highly conserved folding within the OMF family with the most divergent being TolC in accordance with the sequence alignment (Supplementary Figure S1). Thus it appears that the only reason OprM would crystallize in either the R32 or C2 space groups is that the N-terminus lipoyl adopts a different conformation within the three monomers. To understand the different orientations of the palmitoyl, we compared their respective environments (Figure 4). For the three N-termini, the main chain is stabilized by hydrogen bonds with R133 and the carboxyl group of L128. In each case, the palmitoyl tail then turns around two hydrophobic residues, L128 and F129. The palmitoyl from the B monomer (subsequently called Palm-B) is located near the Palm-C of a symmetrical molecule but not close enough to form van der Waals contact because the distance between them is greater than 7 Å. Interestingly, Palm-A instead makes short van der Waals contacts with its own symmetric structure, justifying the quality of the electron density map for this region (see Figure 3) even at low resolution. The quality of the electron density for this particular monomer makes us confident about the exact nature of the fatty acid modification of OprM. As a final control the N-terminal PTMs that were generated in other OMF protein structures, namely CusC and CmeC (Figure 4E) were superimposed on our structure and these demonstrate a longer acyl chain for OprM, supporting its palmitoyl nature.

Discussion

Post-translational modifications (PTMs) play an important role in cell life as they govern most signaling events. Among the different PTMs, lipidation ranks as the second most common modification after phosphorylation2 (dbPTM – database of protein PTMs; Beltrao et al., 2013), anchoring proteins to the membrane and stabilizing their interactions with the lipid bilayer.

It is not well understood why proteins that are embedded within cellular membranes via a large hydrophobic structural domain need supplementary PTMs such as the N-terminal lipidation of OMF proteins. Akama et al. (2004) suggested that these proteins first have to be anchored to the membrane by an N-terminal lipid so that the insertion of their large hydrophobic domain may be triggered. This later step is critical for the correct folding of OMFs. This hypothesis has been reinforced by structural determination of two CusC mutants for which the signal peptide has been conserved but the first cysteine residue after processing is replaced with a serine (C1S-CusC) or is deleted (ΔC1-CusC), (Lei et al., 2014a). The structures result in a random folding of the beta-barrel domain of both mutants together with inappropriate opening of the periplasmic helices. These structures support the essential role of OMF N-terminal lipidation in the membrane insertion mechanism; moreover, they also highlight the possible involvement of membrane-interacting components in the OMF opening process and consequently the efflux-pump function.

2http://dbptm.mbc.nctu.edu.tw
In addition to this function, other functions have been attributed to lipid PTMs. It has been shown that the size of the hydrophobic regions of membrane proteins does not necessarily match the thickness of the cellular membrane (Mitra et al., 2004). Additional hydrophobic elements could then help membrane proteins to fit into membranes of variable thickness, which would explain why some proteins require the addition of 16 carbons, whereas others require only fourteen, depending on the transmembrane domain shape and size.

Regarding OprM, several constructs were previously designed for N-terminal labeling, membrane targeting and antibiotic response experiments (Nakajima et al., 2000) and demonstrated that only when residue 18 is a cysteine is the protein labeled by radioactive palmitate and targeted to the outer membrane. Nevertheless, the three tested OprM mutants (C18G, C18F, C18W) were functional, although none were properly targeted to the outer membrane.

Concerning MexA, the MFP component from the OprM-MexAB efflux pump that is also palmitoylated at its N-terminus and attached to the inner membrane, when this PPM is missing MexA becomes unable to interact with OprM as highlighted by in vitro blue native gel experiments (Ferrandez et al., 2012).

All of these data highlight interest in the study of these lipoproteins and the protein partners involved in their modification. To gain insight into these lipoproteins, it seemed important to identify which modification occurs on OprM. To that end, we analyzed the chemical accessibility of the N-terminal cysteine thiol and amine using different fluorescent probes, which revealed that only the thiol was occupied. In addition, the nature of the attached lipoyl was proven to be a palmitoyl by our determination of the structure of OprM in a new space group that trapped the lipoyl chain at the interface of one monomer with its own symmetric structure.

We now know the nature of OprM N-terminal PTM, and that other OMFs such as CusC and CmeC from different bacterial strains, have similar but different N-terminal modifications. Nevertheless, if anchorage was important for the function of this class of proteins, why would the analog TolC not undergo PTMs? The sequences of OprM and TolC were submitted to the prediction of palmitoylation site3, which confirmed that C18 was a palmitoylation target in OprM and that there was no such site in the TolC sequence. When comparing the structure of TolC with that of OprM, CusC, and CmeC (Figure 5), it appears

3http://csspalm.biocuckoo.org/
that only TolC is different at the N-terminus. In particular, TolC is 44 residues shorter in sequence than OprM (Figure 1; Supplementary Figure S1), meaning that the TolC structure starts at the buoy level. Nevertheless, it should be noted that although it possesses a shortened N-terminus, TolC has a longer C-terminal tail (Figure 1; Supplementary Figure S1), which is not present in its solved structure. Even without knowing anything about the role of this long C-terminal region, it can be hypothesized that it could lead to an additional interaction with the membrane. This hypothesis is supported by the fact that the C-terminal structure of CmeC (an OMF protein with the longest C-terminal sequence of those known) is oriented toward the outer membrane, and the C-terminal alpha-helix of TolC is structurally equivalent to the N-terminal alpha-helix of the other OMFs according to structure superposition (Figure 5). In addition, the distance between the last visible C-terminal amino acid of the TolC structure and the membrane proximal N-terminus of the superposed OprM monomer is approximately 60 Å, a distance that could be covered by a 42 residue-long helix, which corresponds to the number of residues missing at the C-terminus of the TolC construct.

The fact that TolC can be properly inserted in the membrane despite its particularity, the absence of lipid anchor, can be due to the differences in the peptidoglycan (PG) structure between Gram-negative bacteria. The NMR structure of a 2 kDa synthetic fragment composed of NAG–NAM (pentapeptides; Meroueh et al., 2006) allowed to visualize TolC as completely embedded in the PG, its periplasmic alpha-helical domain just flushing the limit of the PG. Measurements of the cell wall dimensions of both E. coli and P. aeruginosa were performed by cryo-transmission electron microscopy (Matias et al., 2003), revealing large size differences between the different cell wall constituents, particularly the PG. The empty space between the PG and the outer-membrane d(PG-OM), and the total thickness, PG + d(PG-OM), were estimated to be 53 ± 9 and 61 ± 9 Å for d(PG-OM), and 117 ± 14 and 85 ± 14 Å for PG + d(PG-OM) in E. coli and P. aeruginosa, respectively.
(see Figure 5). Consequently, ToIC is more likely to be tightly embedded in the membrane barrier than OprM. Nevertheless, no published experiments can support these hypotheses to date.

Conclusion

In this study, we have shown that OprM is palmitoylated at its N-terminal cysteine by thio-palmitoylation. It is now necessary to search for the different proteins involved in this acylation, lipid-transferases, and signal peptidases. These lipidoprotein modifiers could represent new interesting targets for the fight against antibiotic resistance.

Author Contributions

Conceived and designed the experiments: GP, HB, and IB. Performed the experiments: LM, GP, HB, M-BL, YN, and IB. Analyzed the data: LM, GP, HB, M-BL, YN, MP, and IB. Wrote the paper: GP, MP, and IB.

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Supplementary Material

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2015.00667

References

Adams, P. D., Grosse-Kunstleve, R. W., Hung, L. W., Ioerger, T. R., McCoy, A. J., Moriarty, N. W., et al. (2002). PHENIX: building new software for automated crystallographic structure determination. Acta Crystallogr. D Biol. Crystallogr. 58, 1948–1954. doi: 10.1107/S0907444902106657

Ahmed, A., Al-Khalili, S., Al-Tamimi, A., Al-Khalili, R., Al-Rafee, A., et al. (2001). A novel thio-palmitoylation of the glycosylated membrane protein OprM of Pseudomonas aeruginosa: dual modes of membrane anchoring and occluded cavity end. J. Biol. Chem. 279, 52816–52819. doi: 10.1074/jbc.C400445200

Bayram, Y., Parlak, M., Aypek, C., and Bayram, I. (2013). Three-year review of bacteriological profile and antibiogram of burn wound isolates in Van, Turkey. Int. J. Med. Sci. 10, 19–23. doi: 10.7150/ijms.4723

Beltrao, P., Bork, P., Krogn, N. J., and van Noort, V. (2013). Evolution and functional cross-talk of protein post-translational modifications. Mol. Syst. Biol. 9, 714. doi: 10.1002/msb.201304521

Bereket, W., Hemalathra, K., Getenet, B., Wondwossen, T., Solomon, A., Zeynudin, A., et al. (2012). Update on bacterial nosocomial infections. Eur. Rev. Med. Pharmacol. Sci. 16, 1039–1044.

Bernal-Perez, L. F., Prokai, L., and Ryu, Y. (2012). Selective N-terminal fluorescent labeling of proteins using 4-chloro-7-nitrobenzofurazan: a method to distinguish protein N-terminal acetylation. Anal. Biochem. 428, 13–15. doi: 10.1016/j.ab.2012.05.026

Cattoir, V. (2004). [Efflux-mediated antibiotics resistance in bacteria]. Pathol. Biol. 52, 607–616. doi: 10.1016/j.pathbio.2004.09.001

CCP4 suite (1994). The CCP4 suite: programs for protein crystallography. Acta Crystallogr. D Biol. Crystallogr. 50, 760–763. doi: 10.1107/S0907444994003112

Chalker, J. M., Bernardes, G. J., Lin, Y. A., and Davis, B. G. (2009). Chemical modification of proteins at cysteine: opportunities in chemistry and biology. Chem. Asian J. 4, 630–640. doi: 10.1002/asia.200800427

Chuanchuen, R., Narasaka, C. T., and Schweizer, H. P. (2002). The MexJk efflux pump of Pseudomonas aeruginosa requires OprM for antibiotic efflux but not for efflux of triclosan. J. Bacteriol. 184, 5036–5044. doi: 10.1128/JB.184.18.5036-5044.2002

Davis, I. W., Lever-Fay, A., Chen, V. B., Block, J. N., Kapral, G. J., Wang, X., et al. (2007). MolProbity: all-atom contacts and structure validation for proteins and nucleic acids. Nucleic Acids Res. 35, W375–W383. doi: 10.1093/nar/gkm216

DeLano, W. L. (2002). The PyMOL Molecular Graphics System. Available at: http://pymol.sourceforge.net

Emsley, P., and Cowtan, K. (2004). Coot: model-building tools for molecular graphics. Acta Crystallogr. D Biol. Crystallogr. 60, 2126–2132. doi: 10.1107/S0907444904019158

Fedreci, L., Du, D., Walas, F., Matsumura, H., Fernandez-Recio, J., McKeegan, K. S., et al. (2005). The crystal structure of the outer membrane protein VceC from the bacterial pathogen Vibrio cholerae at 1.8 Å resolution. J. Biol. Chem. 280, 15307–15314. doi: 10.1074/jbc.M500401200

Ferrandez, Y., Monlezun, L., Phan, G., Benabdellah, H., Benas, P., Ulryck, N., et al. (2012). Stoichiometry of the MexA-OprM binding, as investigated by blue native gel electrophoresis. Electrophoresis 33, 1282–1287. doi: 10.1002/elps.201100541

Fischbach, M. A., and Walsh, C. T. (2009). Antibiotics for emerging pathogens. Science 325, 1089–1093. doi: 10.1126/science.1176667

Ghosh, P. B., and Whitehouse, M. W. (1968). 7-chloro-4-nitrobenzo-2-oxa-1,3-diazole: a new fluorogenic reagent for amino acids and other amines. Biochem. J. 108, 155–156.

Hede, K. (2014). Antibiotic resistance: an infectious arms race. Nature 509, 52–53. doi: 10.1038/509S2a

Kabsch, W. (1993). Automatic processing of rotation diffraction data from crystals of initially unknown symmetry and cell constants. J. Appl. Crystallogr. 26, 795–800. doi: 10.1107/S0021889893005588

Koroknai, V., Sharif, A., Koronakis, E., Luisi, B., and Hughes, C. (2000). Crystal structure of the bacterial membrane protein ToIC central to multidrug efflux and protein export. Nature 405, 914–919. doi: 10.1038/35016007

Kovacs-Simon, A., Tithall, R. W., and Michell, S. L. (2011). Lipoproteins of bacterial pathogens. Infect. Immun. 79, 548–561. doi: 10.1128/IAI.00682-10

Kulathila, R., Indic, M., and van den Berg, B. (2011). Crystal structure of Escherichia coli CusC, the outer membrane component of a heavy metal efflux pump. PLoS ONE 6:e15610. doi: 10.1371/journal.pone.0015610

Lei, H. T., Bolla, J. R., Bishop, N. R., Su, C. C., and Yu, E. W. (2014a). Crystal structures of CusC review conformational changes accompanying...
Li, X. Z., and Nikaido, H. (2009). Efflux-mediated drug resistance in bacteria: an update. Drug 69, 1555–1623. doi: 10.1007/s12518-012-00087-6

Nakajima, A., Kurokawa, K., and Lee, B. L. (2012). Lipoproteins in bacteria: structures and biosynthetic pathways. FEBS J. 279, 4247–4268. doi: 10.1111/j.1742-4658.2012.08616.x

Nikaido, H. (2009). Multidrug resistance in bacteria. Annu. Rev. Biochem. 78, 119–146. doi: 10.1146/annurev.biochem.78.082907.145923

Nikaido, H., and Pages, J. M. (2012). Broad-specificity efflux pumps and their role in multidrug resistance of Gram-negative bacteria. FEMS Microbiol. Rev. 36, 340–363. doi: 10.1111/j.1574-6976.2011.00290.x

Olivesa, J., Bernardini, A., Garcia-Leon, G., Corona, F., Sanchez, B. M., and Martinez, J. L. (2013). The intrinsic resistome of bacterial pathogens. Front. Microbiol. 4:103. doi: 10.3389/fmicb.2013.00103

Phan, G., Benabdellah, H., Lascombe, M. B., Benas, P., Rety, S., Picard, M., et al. (2010). Structural and dynamical insights into the opening mechanism of P. aeruginosa OprM channel. Structure 18, 507–517. doi: 10.1016/j.str.2010.01.018

Poole, K. (2004). Efflux-mediated multidrug resistance in Gram-negative bacteria. Clin. Microbiol. Infect. 10, 12–26. doi: 10.1111/j.1348-0421.2004.00763.x

Resh, M. D. (1999). Fatty acylation of proteins: new insights into membrane targeting of myristoylated and palmitoylated proteins. Biochim. Biophys. Acta 1451, 1–16. doi: 10.1016/S0167-4889(99)00075-0

Robert, X., and Gouet, P. (2014). Deciphering key features in protein structures with the new ENDScriptr server. Nucleic Acids Res. 42, W320–W324. doi: 10.1093/nar/gku316

Schweizer, H. P. (2003). Efflux as a mechanism of resistance to antimicrobials in Pseudomonas aeruginosa and related bacteria: unanswered questions. Genet. Mol. Res. 2, 48–62.

Su, C. C., Radhakrishnan, A., Adams, P. D., Winn, M. D., Storoni, L. C., and Read, R. J. (2007). Phaser crystallographic software. J. Appl. Crystallogr. 40, 658–674. doi: 10.1107/S0021889807021206

Meronch, S. O., Benze, K. Z., Hesen, D., Lee, M., Fisher, J. F., Stemmer, T. L., et al. (2006). Three-dimensional structure of the bacterial cell wall peptidoglycan. Proc. Natl. Acad. Sci. U.S.A. 103, 4404–4409. doi: 10.1073/pnas.0510182103

Mima, T., Sekiya, H., Mizushima, T., Kuroda, T., and Tsuchiya, T. (2005). Gene cloning and properties of the RND-type multidrug efflux pump MexPQ-OpmE from Pseudomonas aeruginosa. J. Bacteriol. 187, 6112–6118. doi: 10.1128/JB.187.21.6112-6118.2003

Mitra, K., Ubarretxena-Belandia, I., Taguchi, T., Warren, G., and Engelman, D. M. (2003). CmeC outer membrane channel. J. Mol. Biol. 329, 1040–1053. doi: 10.1016/j.jmb.1999.11.022

Morita, Y., Tomida, J., and Kawamura, Y. (2012). Mefx multidrug efflux system of Pseudomonas aeruginosa. Front. Microbiol. 3:408. doi: 10.3389/fmicb.2012.00408

Nakajima, A., Sugimoto, Y., Yoneyama, H., and Naka, T. (2000). Localization of the outer membrane subunit OprM of resistance-nodulation-cell division family multidrug efflux pump in Pseudomonas aeruginosa. J. Biol. Chem. 275, 30604–30608. doi: 10.1074/jbc.M005742200

Nakajima, A., Sugimoto, Y., Yoneyama, H., and Naka, T. (2002). High-level fluoroquinolone resistance in Pseudomonas aeruginosa due to interplay of the MexAB-OprM efflux pump and the DNA gyrase mutation. Microbiol. Immunol. 46, 391–395. doi: 10.1111/j.1348-0421.2002.tb02711.x