Tripartite multidrug efflux systems of Gram-negative bacteria are composed of an inner membrane transporter, an outer membrane channel and a periplasmic adaptor protein. They are assumed to form ducts inside the periplasm facilitating drug exit across the outer membrane. Here we present the reconstitution of native 
\textit{Pseudomonas aeruginosa} MexAB–OprM and 
\textit{Escherichia coli} AcrAB–TolC tripartite Resistance Nodulation and cell Division (RND) efflux systems in a lipid nanodisc system. Single-particle analysis by electron microscopy reveals the inner and outer membrane protein components linked together via the periplasmic adaptor protein. This intrinsic ability of the native components to self-assemble also leads to the formation of a stable interspecies AcrA–MexB–TolC complex suggesting a common mechanism of tripartite assembly. Projection structures of all three complexes emphasize the role of the periplasmic adaptor protein as part of the exit duct with no physical interaction between the inner and outer membrane components.
Gram-negative efflux pumps of the Resistance Nodulation cell Division (RND) superfamily are exporters of biological metabolites and antimicrobial compounds, thus playing a prominent role in the bacterial resistance, which has nowadays become a major health concern.22,25–28. The inner membrane located RND pumps are driven by the proton motive force and as a part of a tripartite system, working in conjunction with an outer membrane factor (OMF), and a periplasmic membrane fusion protein (MFP), the latter assumed to link the RND component to the OMF.23. Well-studied examples of tripartite RND systems are MexAB–OprM of Pseudomonas aeruginosa and AcrAB–ToLC of Escherichia coli.24. For these systems, high-resolution X-ray structures are available for substrates and are in one case also transported by the efflux function as competitive inhibitors or impair the proper binding of in vitro cross-linking studies.34,35

**Results**

**Protocol of tripartite assembly using NDs.** The rationale for the reconstitution of tripartite complexes was based on the insertion of the integral membrane proteins (that is, OMF or RND) into NDs. On detergent removal, the membrane proteins (MexB, AcrB, OprM and TolC) were inserted into a 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC)-containing NDs whose size is limited by the membrane scaffold protein (MSP) wrapped around the hydrophobic core of the lipids (Fig. 1a). The control of the assembly process relied on the insertion of a single molecule per ND, which necessitated the use of two MSP differing in size (MSP1D1 or MSP1E3D1) because of the respective diameters of the transmembrane domains of the RND and OMF proteins (RND ≈ 80 Å, OMF ≈ 40–55 Å (refs 7,11)). Subsequently, the separately ND-reconstituted efflux components were mixed with native lipidated MFP (AcrA or MexA, Fig. 1b).

This two-step reconstitution protocol was successfully applied to MexAB–OprM and AcrAB–ToLC. In the following sections, the reconstitution is detailed by first characterizing OprM or MexB into ND, followed by the process of whole-tripartite assembly reconstitution. Moreover, reconstitution of cognate AcrAB–ToLC and non-cognate AcrA–MexB–ToLC complexes are presented highlighting the generic approach of our protocol. The ND-reconstituted native tripartite complexes were visualized by

Figure 1 | Tripartite assembly based on RND and OMFI inserted into nanodiscs. (a) After detergent removal, the integral membrane proteins are reconstituted into a small lipid bilayer wrapped by two MSPs (purple) forming the nanodisc. Lipids are red/yellow and detergent is grey. (b) Self-assembly of RND (blue) and OMF (orange) in nanodiscs in the presence of native lipid-modified MFP (green) leading to the tripartite complex in lipid membrane.
EM and single-particle analysis, resulting in elongated structures of 33 nm along their main axis.

**OprM and MexB molecules inserted into NDs.** The ND reconstitution of OprM was achieved with construct MSP1D1 and POPC lipids reported to form 10-nm-diameter sized NDs. Analysis of these OprM–NDs by EM was done on negatively stained samples. At a MSP:lipid:OprM molar ratio of 1:36:0.4, OprM–ND mainly contained one OprM molecule per ND, with their long axis preferentially oriented parallel to the carbon support (Fig. 2a). EM observations were consistent with the trimeric assembly of OprM. An average image (from 446 particles) of the OprM–ND revealed a 11 nm in diameter ND spanned by a duct formed by the OprM β-barrel domain (visible due to the presence of uranyl acetate within the β-barrel), followed by the 10-nm-long OprM periplasmic domain including the equatorial domain (Fig. 2a inset). Note that MSP1E3D1 construct produced larger ND leading to the insertion of two OprM molecules (Supplementary Fig. 1).

The formation of MexB–ND was achieved using POPC and MSP1E3D1 as scaffold resulting in 12- to 14-nm-diameter sized NDs (that is, a diameter larger than the transmembrane domain of MexB). At a MSP:lipid:MexB molar ratio of 1:27:1, EM revealed side views of MexB–ND containing one molecule per ND (Fig. 2b) in accordance with the trimeric organization of MexB (Protein Data Bank entry: 2V50). Clearly visible is also the exposed periplasmic domain that protrudes 7 nm away from the lipid-containing ND. Averaging 341 single particles revealed a continuous layer of electron density of the ND, including the 36-transmembrane helix domain of trimeric MexB. The periplasmic part of MexB exhibited furthermore two clearly distinguishable layers of density, assigned to the porter domain and to the more distal funnel domain (Fig. 2b inset).

**Formation of a tripartite complex.** Tripartite complex formation was achieved by mixing OprM–ND, MexB–ND and lipidated MexA in a 1:1:10 molar ratio (Fig. 1b). Formation of tripartite complexes was visualized using native PAGE resulting in an electrophoretic mobility shift on complex formation (Fig. 3a). ND-reconstituted MexB and OprM migrated as a single band strongly stained by silver (Fig. 3a, lanes 1 and 2). MexB–ND migration was less than OprM–ND due to its larger size and hydrodynamic radius. Mixing of MexB–ND with OprM–ND in a 1:1 molar ratio yielded two separate stained bands (Fig. 3a, lane 3) with corresponding electrophoretic mobilities of the two single ND-reconstituted components (Fig. 3a, lanes 1 and 2). However, in a mixture containing MexB–ND, OprM–ND and lipidated MexA (39 kDa monomer; 1:1:10 molar ratio), a significant upshifted band was observed after staining with silver (Fig. 3a, lane 6). This extra band was only observed when all three components were mixed. When lipidated MexA was mixed with either MexB–ND (Fig. 3a, lane 4) or with OprM–ND (Fig. 3a, lane 5), no upshift of bands could be observed. Our
interpretation is that MexB–ND and OprM–ND do not form bipartite complexes and only when the three components of the tripartite complex (that is, MexB–ND, OprM–ND and lipidated MexA) are present in the sample, a larger complex is formed.

To get structural details on the assembly of the efflux pump, the (1:1:10) mixture of OprM–ND, MexB–ND and lipidated MexA was analysed by EM (Supplementary Fig. 2). Strikingly, within the population of single particles, ca. 10% were elongated structures of 33 nm at their longest expansion. Clearly different from isolated MexB–ND and OprM–ND (Fig. 2a,b), these new structures likely correspond to complex formation evidenced on the silver-stained native polyacrylamide gel (Fig. 3a, lane 6). To improve the yield of tripartite complex, the assembly formed with OprM–ND, MexB–ND and lipidated MexA in a 1:1:20 molar ratio was purified by size-exclusion chromatography (SEC) and analysis of the fractions by SDS–PAGE (Fig. 3c) revealed the presence of the three partners, in particular, in the first peak (Fig. 3b, fraction A12). Note that OprM–ND and MexB–ND were eluted in B4 and B3 fractions, respectively when applied alone on the same column (Supplementary Fig. 3). EM analysis of A12 fraction exhibited a vast majority of elongated structures viewed from their sides (Fig. 4a). The majority of class averages (125 over 200 classes) from single-particle average image analysis revealed an edifice of protein densities at both ends resembling ND densities ~23 nm apart (Fig. 4b). The upper part of the complex resembled OprM–ND with its central duct, including the ND-surrounded β-barrel domain. Adjacent to the ND density, a bulky knot is visible, which we interpret as the equatorial domain (Fig. 4b,c). At the other end of the elongated particle, we observe the recognizable features of MexB, that is, the ND-embedded transmembrane domain, and the protruding MexB porter and funnel domains (Fig. 4b,c). In between the OprM–ND and MexB–ND densities, additional densities are present, which we assign to connecting MexA molecules. Isocontours of OprM–ND and MexB–ND (derived from the average images of Fig. 2), overlaid on the isocontours of the putative tripartite complex average image, showed a good match of densities of ND-embedded OprM and MexB molecules (Fig. 4c). The 6-nm long, non-overlapping densities most likely corresponded to MexA molecules that are interacting with both OprM and MexB (blue contours in Fig. 4c). Hence, in accordance with this analysis, it can be concluded that the tripartite MexB–MexA–OprM complex was successfully formed using NDs reconstitution. Comparison of the intermembrane distance (23 nm, Fig. 4c) with the known dimensions of the periplasmic domains of MexB (7 nm (ref. 15)) and OprM (10 nm (refs 11,12)) and the observed non-overlapping density of ~6 nm suggest that the RND (MexB) and OMF (OprM) components are not in direct contact. This observation is in accordance with the results recently obtained with genetically fused RND–MFP constructs36,38, supporting the role of MexA as a part of the periplasmic duct formed by the MexB–MexA–OprM complex, bridging the gap between MexB and OprM (Supplementary Fig. 4).

Interestingly, few class averages (about 400 particles) showed that the contact between MexA and OprM are apparently different. OprM appears to contact the MexAB complex over MexA via small links (black arrows Fig. 4b). These extremities arising from the β-barrel of OprM and engaged the interaction with MexA resemble those observed in images of isolated OprM, which in an isolated state is present in a closed β-barrel conformation11 (Fig. 2). The complexes showing the closed OprM engaged in a presumably looser manner with the MexAB complex may correspond to intermediate steps in the formation of the tripartite system, even though we cannot exclude that these particles represent dissociated complexes due to their interaction on the EM grid.

**Figure 4 | TEM analysis of tripartite MexAB–OprM assembly.** (a) Field of view revealing elongated complexes when OprM–ND and MexB–ND were mixed in the presence of MexA. Scale bar, 30 nm. (b) Gallery of five class average side views of a 33-nm-long tripartite MexAB–OprM complex delineated by two nanodiscs (157, 192, 185, 99, 151 images, respectively) and one top view class average (56 images). Lower row, two average classes of atypical complexes showing faint contacts between OprM and MexAB (black arrows). Scale bar, 10 nm. (c) Isocontours of MexB–ND and OprM–ND (red) overlaid on isocontours of tripartite complex (blue). Characteristic features are displayed: OprM β-barrel and ND (1), equatorial domain (2), tip of β barrel (3'), MexB funnel (4') and porter (5') domains anchored to ND (6). The remaining blue densities correspond to MexA that linked OprM to MexB and interacts with the domains marked with ('). The tripartite assembly ND (6) has a smaller size compared with ND of MexB probably because of the detergent carried with MexA that may extract some lipids.
an overall appearance similar to the cognate tripartite complexes by EM. Surprisingly, for the MexB–AcrA–TolC mixture, we also analysed mixtures of non-cognate TolC–ND, MexB–ND and AcrB (Fig. 5d). Clearly visible from this comparison, the distance was similar to that of MexAB–OprM tripartite complex, to AcrA molecules were observed in the negative-stained AcrB–ND (lower part) and connecting those, densities assigned showed that a complex was composed of TolC–ND (upper part), side chain substitutions in the helical hairpin of AcrA, which resistance of the non-cognate efflux systems could be extended by MexA or AcrA are known for their high flexibility most likely explain the partial resistance observed in an observed only few hybrid AcrA–MexB–TolC complexes may also observed that AcrA–MexB–TolC hybrid system could confer transport by the cognate efflux systems. The fact that we observed that AcrA–MexB–TolC hybrid system could confer resistance to *E. coli* for only a limited subset of drugs normally transported by the cognate efflux systems. The fact that we observed only few hybrid AcrA–MexB–TolC complexes may also explain the partial resistance observed in an *E. coli ΔacrB* background complemented with heterologous expressed *mexB*. MexA or AcrA are known for their high flexibility most likely facilitating tripartite formation. The spectrum of antibiotic resistance of the non-cognate efflux systems could be extended by side chain substitutions in the helical hairpin of AcrA, which interacts with TolC or by substitutions in the AcrA β-barrel domain interacting with MexB. The use of variants of AcrA, TolC or MexB may provide a better fit with the two other

**Discussion**

We have devised a generic approach allowing the formation of tripartite RND efflux pumps as characterized by native gel and EM analysis. By means of a ND toolkit, integral membrane components of two tripartite RND efflux systems were incorporated into a lipid membrane ND and were further assembled after adding purified lipidated MFPs. Similar tripartite assemblies were obtained for the two different pumps from *E. coli* and from *P. aeruginosa*, that is, AcrB–AcrA–TolC and MexB–MexA–OprM, respectively. With an overall height of 33 nm, the tripartite complexes were composed of an OMF molecule and an RND molecule facing each other with their periplasmic domains without being in direct contact, linked by MFP molecules. These self-assembled tripartite complexes strongly resembled the AcrAB–TolC edifice obtained using protein fusion and suggests a 3:6:3 stoichiometric assembly of the components. In this report, the tripartite assemblies comprised native proteins and self-assembly of the three components occurred without any additional protein(s) or chemical cross-linking. Interestingly, both tripartite setups based on native (this work) or fused components indicated a common organization of the MFP connecting OMF and RND components at this level of detail (Supplementary Fig. 4). The three-dimensional reconstruction of MexAB–OprM (Supplementary Fig. 4a) at an estimated resolution of 25 Å (Supplementary Fig. 4b) displays the distinct features of the tripartite system (that is, OMF, MFP-hairpin, -lipoyl, -β-barrel and -membrane proximal domains, in addition to the RND transporter), which is in good agreement with the cryo-EM structure by Du et al. (Supplementary Fig. 4c). This organizational resemblance also suggests that our tripartite pump systems were assembled with similar stoichiometry, that is, 3:6:3 as has been previously proposed for other tripartite pumps as well. It is worth noting that by mixing MexB–ND and OprM–ND, complex formation was not observed on native gel or on EM grid suggesting that OprM does not strongly bind to MexB in solution as was previously observed by isothermal titration calorimetry for the homologous partners TolC and AcrB.

Our results also provided evidence that RND tripartite systems were able to self-assemble in a mixture of non-cognate components. When MexB was interchanged with AcrB in the self-assembling process, tripartite AcrA–MexB–TolC complexes were observed but with a lower occurrence compared with the formation of cognate complexes. The formation of hybrid tripartite complexes has a significant meaning since it has been observed that AcrA–MexB–TolC hybrid system could confer resistance to *E. coli* for only a limited subset of drugs normally transported by the cognate efflux systems. The fact that we observed only few hybrid AcrA–MexB–TolC complexes may also explain the partial resistance observed in an *E. coli ΔacrB* background complemented with heterologous expressed *mexB*. MexA or AcrA are known for their high flexibility most likely facilitating tripartite formation. The spectrum of antibiotic resistance of the non-cognate efflux systems could be extended by side chain substitutions in the helical hairpin of AcrA, which interacts with TolC or by substitutions in the AcrA β-barrel domain interacting with MexB. The use of variants of AcrA, TolC or MexB may provide a better fit with the two other
components. Since similar tripartite assemblies were formed with cognate and non-cognate components, as schematically shown in Fig. 5e, we suggest that there might be a general mechanism triggering tripartite assembly, the formation efficiency being dependent on the binding affinities of the MFPs, therefore modulating drug efflux.

As discussed above, the MFP component possesses a high adaptability in the maintenance of interactions between the OMF and RND components. It has been proposed that during drug export, conformational changes of the MFP component, triggered by substrate and/or H⁺ binding and/or release to/from the RND component could catalyse the assembly/disassembly of the tripartite complex47. The role of the MFP as a central element might indicate that the tripartite assembly could be a multi-stage process. The observed assemblies in this study and in Du and Luisi’s work56 might represent the final stage formation of a fully functional tripartite setup. However, we cannot exclude that the observed setup might be an intermediate formation that, under in vivo conditions, is subsequently subjected to a yet unknown conformational step leading towards fully functional tripartite complexes, where the OMF and RND components are in direct contact. This might explain the previously reported direct cysteine disulﬁde cross-linking between AcrB and TolC34 and would possibly indicate an energy requirement (for example, substrate-binding energy and/or proton motive force) for functional tripartite complex formation.

The reconstitution of the tripartite assembly opens new perspectives for the development of efflux pump inhibitors (EPIs), precluding antibiotic efflux and resulting in an increase of intracellular antibiotic concentration. Efforts have been concentrated up till now on the search for compounds that could compete for antibiotic-binding sites and/or block drug translocation48,49. Nevertheless, such inhibitors are particularly difﬁcult to identify owing to the broad variety of substrates that the multidrug pumps can accommodate. Another strategy would be to target the assembly of the tripartite system. The generation of a leak in the channel by preventing the protein/protein recognitions or by hampering their structural adaptability could also affect the pump efficiency. Compounds having such effect would represent a class of EPIs acting on the assembly process of efflux pumps, instead of acting on drug translocation. Interestingly, unlike RND proteins, OMFs have often been observed to be shared among different tripartite systems, independent of the nature of the inner membrane protein or periplasmic adaptor component50. In the case of P. aeruginosa, OprM can operate with MexAB, MexXY51, and MexMN52. An inhibitor compound precluding OprM–MexA interaction could therefore also inhibit the activity of the other pumps.

Our protocol opens the way towards a better understanding of the tripartite assembly of AcrAB–ToIC and MexAB–OprM. RND and OMF components inserted into NDS are able to self-assemble in a tripartite complex in the presence of a lipidated MFP component. We provide evidence for a common mechanism of tripartite assembly with cognate and non-cognate components, whereby MFP molecules link their cognate OMF partners to cognate and non-cognate RND components. From the single-particle analysis, there is clear evidence that the RND and OMF components are not in direct contact within the tripartite assembly. This approach can be extended to a vast number of RND pump systems and open the field for further structural analysis at atomic level. In addition, the development of an EPI class targeting the tripartite assembly process can be approached more systematically. Although our protocol is not suitable for a large screening of compounds, it is a proof of concept for inhibitor compound analysis.

**Methods**

**Materials and reagents.** POPC was purchased from Avanti Polar Lipids (USA), sodium cholate hydrate, octyl-poly-oxyethylene, n-octyl-β-D-glucopyranoside (OG) and n-Dodecyl β-D-maltoside (DDM) were purchased from Sigma. SM2 Bio-beads and 4–15% Mini-PROTEAN TGX gel were obtained from Bio-Rad. Supersore 6.3/2/300 column and PlusOne Silver Staining Kit were purchased from GE Healthcare. Cu 300 mesh grids were obtained from Agar Scientific.

**Lipid preparation.** POPC lipids were dissolved in methanol/chloroform (v/v), dried on a glass tube under steady flow of nitrogen and followed by exposure to nitrogen for 1 h. The lipid film was suspended in the lysozyme buffer containing 100 mM NaCl, 10 mM Tris/Cl, pH 7.4 and subjected to five rounds of sonication for 30 s each. Lipid concentration was quantiﬁed by phosphate analysis.

**Protein preparation.** Two MSPs, MSP1D1 and MSP1E3D1 (genetic constructs available from AddGene) expressed and puriﬁed from bacteria46, were used to make OprM/ToIC and MexB/AcrB NDs, respectively. The aacrA and tolC genes were individually cloned into pET24a after ampliﬁcation with primers aacrA_forward 5'-GATTGCAGGCACCACAAACACAGAGGTTTACG-3', aacrA_reverse 5'-ATAATAGAGACTTTAAGACCTGACTTGTCAGGCTG-3', tolC_forward 5'-AAAAAACATAAAGAAATGATGCCCTCC-3' and tolC_reverse 5'-AAAAATCCTGGTGTTGTTGCTACGGAATGTACGCTT-3' containing Ndel and Xhol restriction sites, analogous to the cloning of aacrC as described previously34. AcrA, AcrB and TolC were separately expressed in E. coli C43(DE3)/AcrAB and subsequently puriﬁed as previously described for AcrB53. In brief, cells were grown in TB medium at 37°C at an r.p.m of 180,000 for 4°C. The supernatant was subsequently centrifuged for 1 h at 180,000g at 4°C to collect the membranes. Membranes were solubilised, until a 10 mg per g wet weight membranes in lysis buffer containing 10 mM imidazole, pH 7.5, and 2% DDM for 1 h at 4°C while stirring. Samples were centrifuged for 1 h at 180,000g and the supernatant was subjected to Ni-NTA afﬁnity chromatography. The proteins were eluted in a 10 ml buffer (Tris/Cl 20 mM, pH 7.5, NaCl 150 mM, imidazole 200 mM pH 8.0, and 0.03% DDM). AcrA and AcrB were then subjected to SEC using a Superpose 6 HR column (GE Healthcare) with Tris/Cl 20 mM pH 7.5, NaCl 150 mM and 0.05% DDM as running buffer. The ToIC sample was passed through a desalting column (NAP-10, GE Healthcare) using Tris/Cl 20 mM pH 7.5, NaCl 150 mM and 1.5% OG as buffer.

MexB, MexA and OprM membrane proteins were separately produced in E. coli C43/AcrB and puriﬁed as described for MexB57, MexA13 and OprM12,55. In brief, cells were grown in TY medium at 37 °C at 200 r.p.m. till OD600 of 0.6 before induction with 1 mg ml⁻¹ IPTG. After induction, the cells were grown overnight at 20°C at 200 r.p.m. After harvest, cells were suspended in lysis buffer containing 10 ml per g wet weight membranes in lysis buffer containing 10 ml per g wet weight membranes (10 ml per g wet weight membranes). The supernatant was subsequently subjected to an ultracentrifugation at 100,000g for 1 h at 4°C. For OprM, the supernatant is solubilized with 2% n-octyl-poly-oxyethylene before ultracentrifugation to get rid of the inner membranes. Membranes were subsequently subjected to an overnight detergent solubilization in Bio-Tris 10 mM pH 7.4, glycerol 20%, imidazole 10 mM and NaCl 500 mM and at a 2:1 and 40:1 (w/v) detergent-to-protein ratio for MexB and MexA/OprM, respectively (protein concentration is determined using the Bicinchoninic acid test from Sigma). Samples were subjected to an ultracentrifugation at 100,000g for 1 h at 4°C. Protein purifications were performed by afﬁnity chromatography on a HiTrap HP column followed by a gel ﬁltration (supersorze 6 HR, 10/300 GE Healthcare). After puriﬁcation, protein buffers contained 0.03% DDM for MexA and MexB and 0.9% OG for OprM.

**Preparation of MexB/AcrB and OprM/ToIC in NDS.** MexB and OprM were inserted into NDS according to the standard protocol46,49. Brieﬂy, to obtain MexB–ND, MexB solution was mixed with POPC solution and MSP solution at a ﬁnal 36:1:0.4 lipids/MSP/protein molar ratio in a 10 mM Tris/Cl, pH 7.4, 100 mM NaCl, 0.009% DDM and 15 mM Na-cholate solution. Detergent was removed by the addition of SM2 Bio-beads into the mixture shaken overnight at 4°C. For OprM–ToIC, OprM solution was mixed with POPC solution and MSP solution at a ﬁnal 36:1:0.4 lipids/MSP/protein molar ratio in a 10 mM Tris/Cl, pH 7.4, 100 mM NaCl, 0.009% DDM and 15 mM Na-cholate solution. For MexB–ND, detergent was removed with SM2 Bio-beads overnight at 4°C. The same protocol was used for AcrB and ToIC proteins.

**Puriﬁcation of the tripartite complex.** To enrich the tripartite complexes from the initial mixed samples, we subjected the MexB–OprM (or AcrB–ToIC) containing samples to SEC using a Supersorze 6 column pre-equilibrated with reconstruction buffer, which was also used as running buffer. Peak fractions were
collected and subjected to SDS–PAGE analysis. The resulting gel was stained with PlusOne Silver Staining Kit.

**Native gel analysis.** For native gel electrophoresis, samples (0.5 to 2 μl) were separated on 4–15% continuous gradient polyacrylamide Mini-PROTEAN TGX gel. The gel was run in 1x TBE buffer at 1300 V (18 cm), 90 min, and 188 V for 90 min in Tris-Glycine running buffer. The gel was stained with PlusOne Silver Staining Kit.

**Electron microscopy acquisition and analysis.** For EM grid preparations, a diluted mixture of the samples suspension was applied to a glow-discharged carbon-coated copper 300 mesh grids and stained with 2% uranyl acetate (w/v) solution. Images were recorded under low-dose conditions on electron microscope (Tecnai 12 and F20 FEG, FEI) using a FEG Eagle 4k × 4k and a Gatan USG100–2k × 2k cameras. Each image was acquired and two-dimensional averages were performed with SPIDER using a reference-free alignment procedure. For MexAB–OprM and for AcrAB–ToLC, a total of 22,979 and 28,863 particles, respectively, were automatically selected and processed for class averaging.

**For assessing the occurrence of complex formation, OprM–ND, MexB–ND and lipidated MexB-ND were mixed at a ratio of 1:1:1, and sampled after 1, 6, 12, 24, 48, 96 and 168 h (1 week) and from there on every 168 h up to 6 weeks.** For each sample, two negatively stained grids were prepared and subjected to transmission electron microscopy. Twenty-five micrographs were randomly collected for each gel (total 50 micrographs per sample). The occurrence of the complexes was expressed as a ratio of complexes present on the grid corresponding to the number of complexes determined by a manual picking divided by the number of objects (ND alone; MexB–ND, OprM–ND and complexes) calculated by automatic picking based on the finding of maximum values in a Difference of Gaussian filtered micrograph.

**References**

1. May, M. Drug development: time for teamwork. Nature 509, 54–55 (2014).
2. Li, X.-Z., Plesiat, P. & Nikaido, H. The challenge of efflux-mediated antibiotic resistance in Gram-negative bacteria. Clin. Microbiol. Rev. 28, 337–418 (2015).
3. Nikaido, H. Multidrug resistance in bacteria. Annu. Rev. Biochem. 78, 119–146 (2009).
4. Koronakis, V., Sharff, A., Koronakis, E., Luisi, B. & Hughes, C. Crystal structure of the bacterial membrane protein ToLC central to multidrug efflux and protein export. Nature 405, 914–919 (2000).
5. Mikolosko, J., Bobyk, K., Zgurskaya, H. I. & Ghosh, P. Conformational flexibility in the multidrug efflux system protein AcrA. Structure 14, 577–587 (2006).
6. Eicher, T. et al. Transport of drugs by the multidrug transporter AcrB involves an access and a deep binding pocket that are separated by a switch-loop. Proc. Natl Acad. Sci. USA 106, 16416–16421 (2009).
7. Nakashima, E. et al. Structural basis for the inhibition of bacterial multidrug resistance efflux pumps in Pseudomonas aeruginosa: novel agents for combination therapy. Antimicrob. Agents Chemother. 45, 105–116 (2001).
8. Pos, K. M. Drug transport mechanism of the AcrB efflux pump. Nature 405, 578–593 (2000).
9. Tamura, N., Murakami, S., Oyama, Y., Ishiguro, M. & Yamaguchi, A. Direct interaction of multidrug efflux transporter AcrB and outer membrane channel ToLC detected via site-directed disulfide cross-linking. Biochemistry 44, 11113–11121 (2005).
10. Sennhauser, G., Amstutz, P., Brändli, A., Grütter, M. G. Drug export pathway of multidrug exporter AcrB revealed by DARPin inhibitors. PLoS Biol. 5, e7 (2007).
11. Akama, H. et al. Crystal structure of the drug discharge outer membrane protein, OprM, of Pseudomonas aeruginosa: dual modes of membrane anchoring and occluded cavity end. J. Biol. Chem. 279, 52816–52819 (2004).
12. Hancock, R. E. W. Multidrug efflux pumps. Annu. Rev. Pharmacol. Toxicol. 41, 295–331 (2001).
13. Husain, F., Humbard, M. & Misra, R. Interaction between the ToLC and AcrA proteins of a multidrug efflux system of Escherichia coli. J. Bacteriol. 186, 8523–8536 (2004).
14. Husain, F., Humbard, M. A. & Misra, R. Evolution of a bacterial efflux pump: adaptation of the E. coli ToLC exit duct to the Pseudomonas MexAB translocase. FEBS Lett. 580, 5339–5343 (2006).
15. Husain, F., Humbard, M. A. & Misra, R. Evolution of a bacterial efflux pump: adaptation of the E. coli ToLC exit duct to the Pseudomonas MexAB translocase. FEBS Lett. 580, 5339–5343 (2006).
46. Tikhonova, E. B., Wang, Q. & Zgurskaya, H. I. Chimeric analysis of the multicomponent multidrug efflux transporters from gram-negative bacteria. J. Bacteriol. 184, 6499–6507 (2002).

47. Janganan, T. K., Bavro, V. N., Zhang, L., Borges-Walmsley, M. I. & Walmsley, A. R. Tripartite efflux pumps: energy is required for dissociation, but not assembly or opening of the outer membrane channel of the pump. Mol. Microbiol. 88, 590–602 (2013).

48. Lomovskaya, O., Zgurskaya, H. I., Totrov, M. & Watkins, W. J. Waltzing transporters and 'the dance macabre' between humans and bacteria. Nat. Rev. Drug Discov. 6, 56–65 (2007).

49. Nikaido, H. & Pagès, J.-M. Broad-specificity efflux pumps and their role in multidrug resistance of Gram-negative bacteria. FEMS Microbiol. Rev. 36, 340–363 (2012).

50. Yum, S. et al. Crystal structure of the periplasmic component of a tripartite macrolide-specific efflux pump. J. Mol. Biol. 387, 1286–1297 (2009).

51. Morita, Y., Tomida, J. & Kawamura, Y. MexXY multidrug efflux system of Pseudomonas aeruginosa. Front. Microbiol. 3, 408 (2012).

52. Mima, T., Sekiya, H., Mizushima, T., Kuroda, T. & Tsuchiya, T. Gene cloning and properties of the RND-type multidrug efflux pumps MexPQ-OpmE and MexMN-OprM from Pseudomonas aeruginosa. Microbiol. Immunol. 49, 999–1002 (2005).

53. Pos, K. M. & Diederichs, K. Purification, crystallization and preliminary diffraction studies of AcrB, an inner-membrane multi-drug efflux protein. Acta Crystallogr. D Biol. Crystallogr 58, 1865–1867 (2002).

54. Mokhonov, V. et al. Multidrug transporter MexB of Pseudomonas aeruginosa: overexpression, purification, and initial structural characterization. Protein Expr. Purif. 40, 91–100 (2005).

55. Ferrandez, Y. et al. Amphipol-mediated screening of molecular orthoses specific for membrane protein targets. J. Membr. Biol. 247, 925–940 (2014).

Acknowledgements

This work was supported by French national agency ANR (Assembly ANR-12BSV8-0010-03) and Conseil Régional d’Aquitaine (20121301007) grants. We thank A. Luciani, T. Days, C. Garnier and M. Desi for technical assistance. We thank Cordouan Technologies for the use of Elmo. K.M.P. and R.K.R.M. are supported by the German Research Foundation (SFB 807, Transport and Communication across Biological Membranes), the DFG-EXC115 (Cluster of Excellence Frankfurt—Macromolecular Complexes) and a research grant from the Human Frontier Science Program (HFSP-RGP004/2013). L.M. was supported by a grant from Vaincre la Mucoviscidose. A.V. was supported by a grant from the Région Ile-de-France (DIM-Malinfa 110058). The research of R.K.R.M and K.M.P was conducted as part of the Translocation Consortium (http://www.translocation.eu) and has received support from the Innovative Medicines Joint Undertaking under Grant Agreement no.115525, resources which are composed of financial contribution from the European Union’s Seventh Framework Program (FP7/2007–2013) and EFPIA companies in kind contribution.

Author contributions

L.D., I.C.T. and O.L. designed and interpreted the experiments with I.B., M.P and K.M.P. A.V., L.M., M.P., I.B. and R.K.R.M produced and purified membrane proteins. L.D. and F.O. produced proteins in nanodisc and performed EM characterizations. C.G. and L.D. purified and characterized the complexes. L.D. carried out native gel analysis. I.C.T. performed EM analysis. O.L. and K.M.P. wrote the manuscript with input from all authors.

Additional information

Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

Competing financial interests: The authors declare no competing financial interests.

Reprints and permission information is available online at http://npg.nature.com/reprintsandpermissions/

How to cite this article: Daury, L. et al. Tripartite assembly of RND multidrug efflux pumps. Nat. Commun. 7:10731 doi: 10.1038/ncomms10731 (2016).

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/