# Quarterly Reviews of Biophysics

cambridge.org/qrb

# **Review**

Cite this article: Le Bon C, Michon B, Popot J-L, Zoonens M (2021). Amphipathic environments for determining the structure of membrane proteins by single-particle electron cryomicroscopy. *Quarterly Reviews of Biophysics* **54**, e6, 1–22. https://doi.org/10.1017/ S0033583521000044

Received: 21 August 2020 Revised: 19 February 2021 Accepted: 22 February 2021

#### Key words:

amphipols; detergents; integral membrane protein complexes; lipids; nanodiscs; structural biology

Author for correspondence: Manuela Zoonens, E-mail: manuela.zoonens@ibpc.fr

# Amphipathic environments for determining the structure of membrane proteins by single-particle electron cryo-microscopy

Christel Le Bon<sup>1,2</sup>, Baptiste Michon<sup>1,2</sup>, Jean-Luc Popot<sup>1,2</sup> and Manuela Zoonens<sup>1,2</sup> 💿

<sup>1</sup>Laboratoire de Biologie Physico-Chimique des Protéines Membranaires, Université de Paris, CNRS, UMR 7099, F-75005, Paris, France and <sup>2</sup>Institut de Biologie Physico-Chimique, Fondation Edmond de Rothschild pour le développement de la recherche Scientifique, F-75005, Paris, France

# Abstract

Over the past decade, the structural biology of membrane proteins (MPs) has taken a new turn thanks to epoch-making technical progress in single-particle electron cryo-microscopy (cryo-EM) as well as to improvements in sample preparation. The present analysis provides an overview of the extent and modes of usage of the various types of surfactants for cryo-EM studies. Digitonin, dodecylmaltoside, protein-based nanodiscs, lauryl maltoside-neopentyl glycol, glyco-diosgenin, and amphipols (APols) are the most popular surfactants at the vitrification step. Surfactant exchange is frequently used between MP purification and grid preparation, requiring extensive optimization each time the study of a new MP is undertaken. The variety of both the surfactants and experimental approaches used over the past few years bears witness to the need to continue developing innovative surfactants and optimizing conditions for sample preparation. The possibilities offered by novel APols for EM applications are discussed.

# Table of contents

Introduction	1
The cryo-EM resolution revolution and its impact on MP structural biology	2
Selection of reports	4
An overview of surfactant usage at the vitrification step	5
Surfactant exchange before sample application to the grid	9
Effect of surfactants on image quality	10
Vitrification	13
Can surfactants be used as vitrification helpers?	13
Using functionalized APols for immobilizing MP complexes or labelling	
transmembrane domains	15
Conclusion	15

# Introduction

Membrane proteins (MPs) fulfill essential biological functions in the cell and are important biomedical targets. Obtaining MP high-resolution structures can contribute tremendously to the development of novel therapies. However, structural studies of MPs are notoriously more challenging than those of soluble proteins. Many difficulties affect all steps of sample preparation from protein expression to structural determination. Extraction from the native lipid environment with and stabilization against surfactants are particularly critical steps, whose optimization is time-consuming. Over the years, many strategies and tools have been developed to overcome each type of difficulty.

The first MPs for which atomic models could be built were proteins that are naturally abundant (Deisenhofer *et al.*, 1985; Henderson *et al.*, 1990; Weiss *et al.*, 1990). However, the low natural abundance of most MPs necessitated the development of recombinant expression strategies in either microbial host cells, such as *Escherichia coli* or *Saccharomyces cerevisiae* (Hays *et al.*, 2010; Zoonens and Miroux, 2010; Dilworth *et al.*, 2018), or other expression

© The Author(s), 2021. Published by Cambridge University Press



systems better suited to the production of mammalian MPs, such as baculovirus-infected insect cells or human embryo kidney cells (Bill *et al.*, 2011; Andréll and Tate, 2013; Contreras-Gómez *et al.*, 2014). These host cells have proven very efficient for producing MPs in amounts amenable to structural investigations.

Once they are expressed in suitable quantities, MPs need to be extracted from the membrane and purified in their native state. Unfortunately, their stability in aqueous solutions is often problematic. Perturbation of the protein structure by inadequate surfactants can generate misleading data (Cross et al., 2011; Chipot et al., 2018), which is a major concern in this field. Detergents are historically the first surfactants used for MP solubilization and purification. Detergents disrupt the membrane and adsorb onto the hydrophobic transmembrane surface of MPs, keeping them water-soluble (see e.g. Garavito and Ferguson-Miller, 2001; Otzen, 2015; Sadaf et al., 2015; Orwick-Rydmark et al., 2016; Popot, 2018). However, detergents tend to interfere with molecular interactions in MP transmembrane domains, leading to inactivation (for a recent discussion of causes and remedies, see Chapter 2 in Popot, 2018). The extent of this problem varies both from one detergent and from one MP to another. Quite naturally, the first high-resolution structures of MPs ever obtained were those of abundant and detergent-resistant bacterial MPs. Access to the structures of generally more fragile and less abundant eukaryotic MPs has been made possible by progress in production methods, technical advances in structural approaches, and the development and increasingly frequent usage of novel, less destabilizing surfactants. Detergents and other emerging tools used to prepare MP samples for structural determination by single-particle electron cryo-microscopy (SP cryo-EM) have been described in Mio and Sato (2018) and Autzen et al. (2019).

In the present review, we have carried out a quantitative analysis, for all cryo-EM structures of MPs deposited as of January 1, 2021, of the use of detergents, amphipathic polymers such as amphipols (APols) and styrene-maleic acid (SMA) copolymers, protein-based nanodiscs (NDs), and other emerging compounds. The analysis is based on the list of MP structures available on Stephen White's database (https://blanco.biomol.uci.edu/ mpstruc/), which is designed to be as inclusive as possible. This list contains all unique MP structures, coordinate files, and almost all published reports of MP structures accumulated in the protein data bank (PDB) since 1985. The goal of the present analysis is to give an overview of the evolution of approaches in this rapidly moving field and, more specifically, of current trends in the implementation of conventional and novel surfactants. Recently, a similar analysis has been performed (Choy et al., 2021), which includes structures obtained by X-ray crystallography and NMR but bears on a more restricted corpus of cryo-EM data (see below).

#### The cryo-EM resolution revolution and its impact on MP structural biology

Approaches giving access to the three-dimensional (3D) structure of biological macromolecules include X-ray crystallography, solution and solid-state NMR spectroscopy, electron crystallography and SP cryo-EM. The distribution of the unique MP structures published during each of the past 35 years, sorted out according to the structural approach used, is shown in Fig. 1*a*. Over three decades, X-ray crystallography was the dominant technique in MP structural biology. As a result, more than two-thirds of the total number of currently available unique MP structures have



**Fig. 1.** Distribution of unique MP structures solved by each structural approach between 1985 and January 1, 2021. (*a*) Number of unique MP structures deposited each year using each approach. (*b*) Distribution of the 1195 unique structures extracted from Stephen White's database (https://blanco.biomol.uci.edu/mpstruc/) according to the approach used to solve them. The figures were prepared using the January 8, 2021, update of MP structures from this database. Some MP structures published in 2020 may therefore not be included.

been solved using this approach (Fig. 1b). X-ray crystallography can reveal atomic details of MP structure at a very high resolution. Its major bottleneck remains the production of well-ordered 3D crystals. This is particularly challenging for MPs due to the presence of detergent, instability, and/or intrinsic conformational heterogeneity (Tate, 2010). Significant progress has been achieved by resorting to strategies such as the binding of antibody fragments or nanobodies, MP stabilization through engineered mutations, the use of lipidic cubic phases, and/or novel techniques for collecting diffraction data, such as the use of micrometric synchrotron X-ray beams and X-ray free-electron lasers (for reviews, see e.g. Martin-Garcia et al., 2016; Ishchenko et al., 2017). Over the past decade, however, the most spectacular breakthroughs have been achieved in cryo-EM (Nogales and Scheres, 2015; De Zorzi et al., 2016; Vinothkumar and Henderson, 2016; Cheng, 2018; Nwanochie and Uversky, 2019).

Cryo-EM encompasses a variety of techniques for determining the structures of frozen-hydrated samples, which are based on the analysis of (i) images of single particles of purified biomolecules; (ii) images and diffraction patterns of crystalline arrays; and (iii) images of whole cells or cell sections (tomography). In electron cryo-tomography, although important biological insights can be gained from images of biomolecules within their native environment, the resolution is generally much lower than that achieved with the two other cryo-EM techniques, i.e. electron crystallography and SP analysis, from which structural data at near-atomic resolution can be obtained. Electron crystallography is typically applied to planar or tubular two-dimensional (2D) crystals (Abeyrathne *et al.*, 2012; Kühlbrandt, 2013). A more recent technique, referred to as microcrystal electron diffraction (MicroED), has been developed for the analysis of 3D microcrystals (Nannenga and Gonen, 2019; Nguyen and Gonen, 2020); for the moment, however, microED has not yet yielded any *de novo* MP structure.

Electron crystallography applied to 2D crystals has a long and prestigious history, as it was the approach that enabled building the first medium-resolution structural model ever obtained of a MP, that of the archaebacterial bacteriorhodopsin (at 7 Å resolution in the membrane plane (Henderson and Unwin, 1975)), and, later, yielded the first data (at 3.5 Å in-plane) leading to an atomic model of the protein's transmembrane region (Henderson et al., 1990). Before 2013, deposited cryo-EM coordinates of MPs were all from 2D crystal electron crystallography. In 2013 were published the first near-atomic-resolution MP structures determined by SP cryo-EM (Cao et al., 2013; Liao et al., 2013). Since that date, the number of unique MP structures established by this approach has increased explosively (Fig. 1a). Decisive advances, particularly as regards data collection and correction, as well as software development for model building, have revolutionized the field (Cheng, 2015; Alnabati and Kihara, 2019). A combined approach of SP analysis applied to helical arrays (McCullough et al., 2015; Lopez-Redondo et al., 2018) or 2D crystals (Righetto et al., 2019) has been reported. As a rule, however, SP analysis in cryo-EM refers to images of solubilized, unstained, isolated MPs embedded in a thin layer of vitreous ice, which are classified and averaged for 3D reconstruction.

Because it bypasses crystallization, which is the limiting step in X-ray and electron crystallography, SP cryo-EM has turned into a highly efficient approach to the structural analysis of MPs, to the point that three-quarters of the MP structures determined in 2020 have been obtained in this way (Fig. 1a). However, whereas both crystallography and NMR can solve the structures of small MPs, a size limit (currently ~100 kDa; see Nygaard et al., 2020) exists in SP cryo-EM because, if a MP is too small, aligning particle images becomes difficult (Henderson, 1995). Given that the technique is evolving rapidly, one can expect to reach in a near future a size limit close to ~40 kDa, as for soluble proteins (Zubcevic et al., 2016). Alternatively, the size of small MPs can be artificially increased by binding either a natural (Frauenfeld et al., 2011) or an artificial ligand (Wu et al., 2012), provided that the assembly is well-defined and rigid. This strategy has been widely employed for determining the cryo-EM structures of G protein-coupled receptors (GPCRs) in complex with either their natural G protein or arrestin ligands (Liang et al., 2017; Draper-Joyce et al., 2018; Zhao et al., 2019c; Huang et al., 2020a), an antibody fragment (Fab) and/or a nanobody (Zhao et al., 2019a; Tsutsumi et al., 2020), or a combination of these ligands (Gao et al., 2019; Nguyen et al., 2019; Tsai et al., 2019; Yin et al., 2019a; Ma et al., 2020; Qiao et al., 2020). More recently, another class of chimeric binders, called megabodies, consisting of a nanobody grafted onto a larger scaffold protein via two short peptide linkers, has been engineered to overcome size limitations while improving the particle orientation in ice (Uchański et al., 2021).

Structural studies of eukaryotic MPs have strongly benefited from progresses in SP cryo-EM, to the point that currently  $\sim$ 76% of the unique MP structures solved by this technique are



**Fig. 2.** Distribution of unique MP structures according to biological origin. (*a*) Unique MP structures were classified into five groups (archaea, bacteria, eukaryotes, viruses, and unclassified). The latter group comprises four crystallographic structures of *de novo* designed MPs. (*b*) Repartition of mammalian *versus* other eukaryotic MP structures solved by either SP cryo-EM or X-ray crystallography over the past 6 years.

those of eukaryotic MPs, *versus* ~35% of those solved by X-ray crystallography (Fig. 2*a*). Among eukaryotic cryo-EM MP structures, mammalian proteins represent ~68%, an evolution that also reflects recent progresses in expression strategies (Fig. 2*b*). Challenging protein complexes whose structural determination was not technically possible 10 years ago are now accessible. The distribution of structures obtained by cryo-EM is overwhelmingly in favor of integral MPs with  $\alpha$ -helical (89%) rather than  $\beta$ -barrel transmembrane domains (~9%), whereas monotopic MPs represent ~2%. The main subgroups of  $\alpha$ -helical MPs are channels, ATP-binding cassette (ABC) transporters, electron transport chain supercomplexes, GPCRs, and ATPases (Fig. 3*a*), secretion complexes constituting the main subgroup of  $\beta$ -barrel MPs (Fig. 3*b*).

The resolutions achieved using SP cryo-EM keep improving from year to year. In 2018, the structure with the highest resolution was that of a voltage-gated sodium channel, obtained at 2.8 Å resolution when bound to the gating modifier toxin Dc1a (PDB accession code: 6A90) and at 2.6 Å resolution in the presence of the pore blocker tetrodotoxin (PDB accession code: 6A95) (Shen *et al.*, 2018). In 2019, a resolution of 2.37 Å was achieved for the structure of tetrameric Photosystem I (PDB accession code: 6K61) (Zheng *et al.*, 2019). Finally, in 2020, the threshold of 2 Å resolution was crossed with the structure of the sheep connexin-46 solved at 1.9 Å (PDB accession code: 7JKC) (Flores *et al.*, 2020) and that of a



**Fig. 3.** Family distribution of unique MP structures solved by SP cryo-EM. The groups of (*a*) *a*-helical and (*b*) *β*-barrel MPs gather 281 and 30 unique MP structures, respectively. 'Others', in the *a*-helical MP group, refers to cryo-EM structures of MPs belonging to the membrane-spanning 4-domain (MS4) family, antiporters, cellulose synthases, sterol-sensing domain (SDD) proteins, Type VII secretion systems, intramembrane proteases, adventitious MPs (*a*-helical pore-forming toxins), host-defense proteins, decarboxylases, novel receptors (STRA6 retinol-uptake receptor in complex with calmodulin), transhydrogenases, adenylyl cyclases, chain length determinant proteins, and glycoproteins. 'Other transporters' gather one cryo-EM structure of drug/metabolite transporters belonging to the major facilitator superfamily (MFS), three structures of multi-drug efflux transporters and, finally, fourteen structures of MPs belonging to the solute carrier (SLC) transporter superfamily.

homopentameric  $\beta$ 3  $\gamma$ -aminobutyric acid type-A receptor solved at 1.7 Å (PDB accession code: 7A5 V) (Nakane *et al.*, 2020).

Despite this dramatic progress in improving the resolution of SP cryo-EM data, one should keep in mind that many of the atomic models reported in the PDB have benefited from docking high-resolution structures derived from X-ray crystallography, NMR spectroscopy, or other cryo-EM studies as initial templates into cryo-EM maps. Structures of soluble extramembrane domains or homologous proteins from different species are often used. Rigid or flexible fitting methods are commonly applied for the validation and interpretation of cryo-EM structures (compared to rigid fitting methods, flexible ones allow conformational changes that improve fitting the model into the experimental map). Many atomic models derived from cryo-EM result therefore from a mix between these different approaches rather than being fully ab initio structures. The higher the resolution achieved, of course, the safer is the resulting model, which is a strong incentive for improving the quality of experimental data. Which biochemical approaches are being developed to this end is the main subject of this review.

#### **Selection of reports**

Despite technical advances, substantial barriers still stand in the way of many cryo-EM projects. Obtaining cryo-EM grids of sufficient quality for high-resolution structural analysis is not a trivial task. There is no unique answer to the question, 'Which detergent or detergent substitute is the most appropriate for MP structural determination by SP cryo-EM?' In any study, the final choice of the surfactant(s) in which the structure will be determined depends not only on the target MP, but also on the relative amount of efforts invested optimizing one or another set of experimental conditions. A surfactant that appeared inferior in a first study can thus provide superior results in another. For instance, it was initially observed that the human  $\gamma$ -secretase yielded a higher-resolution cryo-EM structure in APol A8-35 than in digitonin (Lu et al., 2014), whereas, a few years later, using a version of  $\gamma$ -secretase fused with T4 lysozyme, two other structures were obtained in digitonin at a higher resolution than that initially achieved in A8-35 (Yang et al., 2019; Zhou et al., 2019). Beyond some obvious limitations, such as the fact that current membrane scaffold protein (MSP)-based NDs (Denisov and Sligar, 2017; Sligar and Denisov, 2020) cannot accommodate MPs with very large transmembrane domains and thus are not appropriate for studying large supercomplexes (see below), trial-and-error is still the only way to proceed. Yet, examining the set of all MP structures determined to-date by SP cryo-EM reveals some interesting trends that can help selecting the most promising tools.

Analyzing unique MP structures provides an overview of the types of structures solved and their biological origin. Based on this dataset, an analysis of the surfactants used for solubilization and structure determination of MPs (whatever the structural approach used) has been recently published (Choy et al., 2021). However, the structure of a given MP has often been published more than once, generating several PDB files for the same protein. In the present study, we chose to include all of these data, hereafter called 'published reports'. This raises the number of experimental situations from 1195 unique structures to 2212 published reports (Fig. 4). As regards cryo-EM alone (including both SP and electron crystallography), this number increases from 328 unique structures (187 in Choy et al. (2021)) to 616 reports, thus providing a more exhaustive view of the extent to which each type of surfactant is used. Most of the structures have been obtained by SP cryo-EM analysis (316 unique MP structures, 587 reports). The number of  $\alpha$ -helical MPs almost doubles, with 536 reports *versus* 281 unique structures (six reports missing from S. White's database have been included in the list (Wang and Sigworth, 2009; Glavier et al., 2020; Hua et al., 2020; McDowell et al., 2020; Yao et al., 2020; Zhang et al., 2020)). Here again, the most representative subgroups include channels, ABC transporters, electron transport chain supercomplexes, ATPases, and GPCRs. The latter comprise 67 structures overall for only 18 unique ones. The resolution of the vast majority of structures reported lies between 2 and 5 Å (554 over 593 files, i.e. >90%), with a median around 3.4-3.5 Å. Knowing that the resolution of cryo-EM maps is usually not uniform due to protein flexibility and dynamics, which tend to degrade the resolution at the periphery of the protein as compared to its core, the resolution value is usually given on an indicative basis. Somewhat arbitrarily, a threshold of 5 Å resolution (sufficient to fit  $\alpha$ -helices into EM maps) was chosen for a structure to be included in our analysis, the aim being to obtain as broad an overview as possible



Fig. 4. Number of unique cryo-EM MP structures and published reports analyzed in this study. Among the cryo-EM structures, SP analysis is the predominant technique, only nine unique MP structures having been solved using electron crystallography. Most cryo-EM MP structures have been solved by SP analysis applied to images of isolated MPs embedded in vitreous ice, one and two structures having been obtained by SP analysis applied to 2D crystals and helical assemblies, respectively. The list of reports analyzed comes from Stephen White's database, manually completed with 60 SP cryo-EM structures for which a PDB and/or EMDB accession number was available (see below). Two crvo-EM structures of MPs reconstituted into liposomes have been included. In addition, examination of each of the 587 studies revealed that some cryo-EM structures were missing. They have been manually integrated into the list of reports as follows: (a) in four studies, an additional cryo-EM structure of a distinct MP or MP complex was identified and added to the list of reports, (b) in 50 studies, a 3D reconstruction of a MP has been obtained in two different surfactants, yielding two distinct refined structures, generating two sets of experimental conditions that have been included in our analysis as 'duplicates', and (c) in two studies, the cryo-EM structure of a MP has been obtained in three different surfactants, yielding 'triplicates'. These additional experimental conditions were gathered in three groups: 'Duplicates 1': distinct cryo-EM structures were obtained in two (or three) different surfactants, whereas the detergent used for the purification was unchanged; 'Duplicates 2': both the purification and vitrification surfactants differed; 'Duplicates 3': the vitrification surfactant was the same but the purification was performed in two distinct detergents.

of the diversity of surfactants that can be used to achieve near-atomic resolution, while leaving aside those that have not yet proven up to this task. In certain studies, SP cryo-EM analysis of a given MP was performed in two or three distinct environments, leading to the refinement of several models deposited in the PDB. This is a very interesting situation, whose lessons will be discussed below. To obtain the widest possible view of all usable environments, such 'duplicate' structures have been included when appropriate (i.e. whenever several distinct structures have been deposited in the PDB and/or the electron microscopy data bank (EMDB)), increasing the number of experimental situations that have yielded 2-5 Å resolution structures from 554 to 604 (Fig. 4).

#### An overview of surfactant usage at the vitrification step

Among SP cryo-EM structures obtained at a resolution of 5 Å or better (hereafter noted  $\leq 5$  Å), the six surfactants most frequently used for vitrification are, in order of decreasing frequency of use, (1) digitonin, (2) detergents with a single maltose-based polar moiety and a single alkyl chain (mostly *n*-dodecyl- $\alpha$ - (or  $\beta$ -) D-maltoside (DDM), occasionally undecyl- or decyl-maltoside (respectively UDM and DM)), (3) MSP-based NDs, (4) detergents belonging to the maltoside-neopentyl glycol (MNG) family, which feature two maltose-based polar heads and two alkyl chains, (5) glyco-diosgenin (GDN), and (6) classical APols (mainly A8-35 and PMAL-C8) (Figs 5–7).

Digitonin, the most popular surfactant, was used for ~18% of MP structures determined at resolutions  $\leq 5$  Å (Fig. 7*a*). This natural detergent is a pentasaccharide derivative of steroidal aglycon digitogenin, produced by the purple foxglove (Digitalis purpurea) (Fig. 5). Commercial digitonin is in fact a mixture consisting of about five glycosides, the main components of this mixture being digitonin and digalonin, a saponin of similar structure, in which the 2-hydroxy digitogenin is missing (Fukunaga et al., 1988). Commercial preparations of digitonin cannot be used directly, because even the best preparations contain 30-50% w/w of material that cannot be dispersed evenly in water, and the effects of unpurified digitonin are unpredictable. Therefore, purification of commercial digitonin by recrystallization is recommended (Kun et al., 1979). Whereas digitonin presents quite a few drawbacks, such as batch-to-batch variations, formation of large micelles, crystallization at low temperature and ill-defined critical micellar concentration (CMC <0.5 mM), it is widely used in cryo-EM due to its mildness. GDN is a recent synthetic detergent (Chae et al., 2012), whose chemical structure mimics that of digitonin (Fig. 5). Due to its homogeneity, the CMC of GDN, which is very low  $(\sim 18 \,\mu\text{M})$ , is better defined than that of digitonin (Chae et al., 2012). SP cryo-EM structures of GDN-solubilized MPs appeared for the first time in 2017. Three years later, its use is reported more often than that of digitonin (in 41 structures versus 33 for digitonin) (Fig. 7b).

The second most widely used surfactants in cryo-EM are maltoside detergents such as DDM, UDM and DM, with a clear preference for DDM (86 structures out of 103). DDM (Fig. 5) was developed in the mid '70s to early '80s (Rosevear *et al.*, 1980) and has been widely used in X-ray crystallography (Dilworth *et al.*, 2018; Choy *et al.*, 2021). The polar head of DDM, UDM and DM is provided by a disaccharide (maltose), whereas the hydrophobic parts are 12-, 11-, and 10-carbon *n*-alkyl chains, respectively. The CMC value of DDM is low (~0.17 mM for  $\beta$ -DDM) and it forms well-defined micelles; UDM and DM have higher CMC values (0.59 and 1.8 mM, respectively) (Bhairi and Mohan, 2007; Stetsenko and Guskov, 2017; and see Table 2 in Abeyrathne *et al.* (2012)).

MNG detergents were synthesized and initially used mainly for GPCR crystallization (Chae *et al.*, 2010). These molecules carry two alkyl chains and two hydrophilic groups derived from maltose, linked *via* a central quaternary carbon atom (Fig. 5). Because of the presence of two long alkyl chains, MNGs have very low CMCs (e.g.  $\sim 10 \,\mu$ M for LMNG, which is by far the most widely used MNG, with 73 structures out of 75). While MNGs appear only at the fourth position of the most frequently used surfactants, with an overall frequency of  $\sim 12\%$  (Fig. 7*a*), they represent one of the surfactant classes most frequently used in 2020 (Fig. 7*b*), partly due to their use for solving GPCR structures.

The third and sixth most often used surfactant classes are, respectively, protein-based NDs and amphipathic polymers called APols, both developed in the '90s. These two detergent substitutes are structurally very different, but both aim at solving the problem of detergent-induced MP instability by eliminating the detergent altogether.



Fig. 5. Some of the amphipathic environments used for MP stabilization. Chemical structures of detergents: digitonin, glyco-diosgenin (GDN), n-dodecyl- $\beta$ -p-maltoside (DDM) and Lauryl Maltoside-Neopentyl Glycol (LMNG); APols: A8-35, PMAL-C8, SMA, and CyclAPols C<sub>8</sub>-C<sub>0</sub>-50 and C<sub>6</sub>-C<sub>2</sub>-50; hydrosoluble cholesterol analog: cholesteryl hemisuccinate (CHS).

APols constitute a family of short and flexible amphipathic polymers specially developed for MP biochemistry (Tribet et al., 1996; Popot et al., 2011; Zoonens and Popot, 2014; Popot, 2018) (Fig. 5). For a long time, when X-ray crystallography was the predominant technique in MP structural biology, APols were considered relatively unattractive. This was due to the mistaken view that the use of polydisperse molecules would hinder crystallization. This point is not necessarily relevant, given that most of the molecules comprising the surfactant belt are disordered anyway (for discussions, see Charvolin et al. (2014), and Chapter 11 in Popot, 2018). Indeed, a recent study has shown that a mix of DDM and A8-35 (the latter used as a stabilizing agent) allows one to solve the X-ray structure of herpes simplex virus glycoprotein B at a higher resolution than can be achieved with DDM alone (Cooper et al., 2018; Cooper and Heldwein, 2020). More to the point is the fact that early APols carried charged polar moieties, which tend to hamper 3D crystallization, whereas non-ionic APols (NAPols) (Bazzacco et al., 2012) have only recently become commercially available. The use of APols has been reported in >100 EM studies (those published from 1998 until late 2017 are reviewed in Chapter 12 of Popot, 2018). In the present analysis, only the cryo-EM structures of APol-complexed MPs included in Stephen White's database with resolutions ≤5 Å have been taken into consideration. Out of 60 cryo-EM MP structures determined using APols, 34 were obtained in A8-35 (Tribet et al., 1996) (Fig. 5) and 19 in PMAL-C8 (Nagy et al., 2001) (Fig. 5). A8-35 first left its mark in SP cryo-EM with the low-resolution structure (at  $\sim$ 19 Å) of a mammalian MP supercomplex, that of the bovine mitochondrial respirasome (Althoff et al., 2011). It also led to the first near-atomic resolution structures (at 3.8 and 3.27 Å) of a MP ever solved by SP cryo-EM, those of the transient receptor potential cation channel V1 (TRPV1) (Cao et al., 2013; Liao et al., 2013). To date, the use of NAPols has been reported in only one cryo-EM structure, that of the outer mitochondrial membrane translocase (TOM) core complex (PDB accession code: 5O8O); however, due to indistinguishable 3D maps of the complex in DDM and NAPols, the two datasets have been merged into a single map refined to 6.8 Å (Bausewein et al., 2017) and therefore have not been included among the reports analyzed in this review.

The co-polymerization of styrene and maleic acid yields APol variants known as SMA co-polymers (Fig. 5). SMAs form with lipids and MPs the so-called SMALPs. Their interest derives



Fig. 6. Membrane protein structures established in protein-based nanodiscs. (a) Molecular model of an MSP-based ND (side view). The two MSPs are shown in red and blue, the 160 associated lipids in orange and grey. From Shih et al. (2005). (b) Molecular model of a saposin-based ND (top view). The saposin A dimer is in purple, the 26 associated lipid molecules in reddish brown. From Li et al. (2016). (c) Family distribution of MPs whose structures have been solved after reconstitution in protein-based NDs. 'Other transporters' comprise cryo-EM structures of three multi-drug efflux transporters, three amino acid secondary transporters, three major facilitator superfamily (MFS) transporters, and three transporters distributed among the three superfamilies of drug/metabolite transporters (DMT), CorA ion transporters and solute carrier (SLC) transporters. (d) Distribution of MSPs and lipids used to reconstitute MPs in protein-based NDs. The MSP called zap1 is derived from Zebra fish apo-lipoprotein A-1; MSP2X was constructed by linking two MSP1E3D1 chains with a two-amino acid linker; csMSP1E3D1 is a circularized version of MSP1E3D1. In three studies, the MSP used has not been clearly mentioned ('unspecified'). When the lipids were a mix of natural and synthetic lipids, they are noted as 'natural and synthetic lipids'; when their nature has not been clearly mentioned, as 'natural or synthetic lipids'.

mainly from their ability to extract MPs from membranes without resorting to detergents (Knowles *et al.*, 2009; Stroud *et al.*, 2018). As of January 1, 2021, only four MP structures had been solved at high resolution using SMAs (Qiu *et al.*, 2018; Sun *et al.*, 2018; Tascón *et al.*, 2020; Yoder and Gouaux, 2020), probably due to a set of limitations that have already been discussed elsewhere (Autzen *et al.*, 2019). For instance, the isolation conditions with SMAs (typically, high ionic strength buffers such as 500 mM



**Fig. 7.** Surfactants used at the vitrification step. (*a*) Distribution of all surfactants in which MP structures have been solved by SP cryo-EM to a resolution  $\leq 5 Å$  (604 reports overall, taking all duplicates into account). (*b*) Number of near-atomic resolution MP structures reported each year since 2013 according to the type of surfactant used at the vitrification step. 'Others' gather two cryo-EM structures obtained in Brij-35 and ten other structures obtained in one of the following surfactants: Tween 20, Igepal CA-630 (NP40), lauryldimethylamine oxide, MNA-C12, PCC-a-M, sodium cholate, Facade-EM (FA-3), octyl glucoside, sodium deoxycholate, and liposomes. Sixteen of the cryo-EM structures gathered in Stephen White's database are either MP fragments, toxins, or peripheral MPs vitrified in the absence of detergent.

NaCl or L-arginine) and/or the purification conditions (12–24 h of incubation with affinity resins) tend to be too harsh for fragile MPs of mammalian origin.

CyclAPols, a novel generation of APols that carry cyclic, saturated hydrocarbon chains instead of alkyl ones (Fig. 5), have also proven able to directly extract MPs without resorting to detergents while stabilizing target MPs to the same extent as classical APols (Marconnet *et al.*, 2020). With these molecules, the usual protocol of detergent/APol exchange is therefore no longer mandatory. CyclAPols have been recently validated for high-resolution structure determination by SP cryo-EM (Higgins *et al.*, submitted for publication).

Finally, protein-based NDs are nanometric-sized objects of discoidal shape formed by lipid bilayer patches stabilized by two encircling amphipathic helical proteins derived from apolipoprotein A1, called MSPs (Fig. 6a; for recent reviews, see Denisov and Sligar, 2017; Denisov *et al.*, 2019; Sligar and Denisov, 2020). The total number of MP structures established in MSP-based NDs at resolutions  $\leq 5$  Å is 100. Seven additional structures have been reported using a related system, called Salipro, where another protein, saposin A, replaces MSPs (Fig. 6b) (Frauenfeld et al., 2016). Furthermore, three other structures have been obtained with a system called peptidiscs, based on reconstituting the detergentsolubilized MP into short amphipathic bi-helical peptides derived from the apolipoprotein A1 in the absence of additional lipids (Carlson et al., 2018). The main groups of MPs whose structures have been solved after reconstitution in protein-based NDs are channels, ABC transporters, ATPases, Cys-loop receptors and members of the TMEM16 family (Fig. 6c). At variance with other amphipathic environments, which depend on securing surfactants from commercial sources, protein-based NDs can be produced in the laboratory. MSPs, which can be expressed in E. coli, have been engineered to form NDs of various diameters compatible with the incorporation of MPs of different transmembrane domain sizes. MSP2N2, MSP1E3D1 and MSP1D1 form NDs with diameters comprised between ~10 and ~17 nm (Denisov et al., 2004; Grinkova et al., 2010) and are the most frequently used MSPs, with 37, 24 and 22 cryo-EM structures, respectively (Fig. 6d). Up to now, the largest transmembrane domains embedded in MSP-based NDs are those of the yeast vacuolar ATPase  $V_{0}$ proton channel (Roh et al., 2018, 2020) and of the human calcium homeostasis modulator 5 (CALHM5) (Liu et al., 2020), which comprise 48 and 44 transmembrane  $\alpha$ -helices, respectively. Despite a lower number of transmembrane  $\alpha$ -helices, the diameter of CALHM5 is wider than that of ATPase  $V_0$  due to the presence of a large 60 Å diameter pore at the center. In this study, circularized NDs in which the N and C termini of MSPs are covalently linked were used, whereas conventional MSP1E3D1 yielded NDs of sufficient size to accommodate the transmembrane domain ATPase  $V_{0}$ . Circularized NDs were engineered to be more homogeneous and stable than standard NDs (Nasr et al., 2017; Nasr and Wagner, 2018). Circularized MSPs can form NDs of larger diameter (50 nm, and up to 80 nm) as compared to standard MSPs, but these wider NDs have yet to yield highresolution cryo-EM structures.

Regarding the lipid components, lipid extracts from natural sources (soybean, brain, *E. coli*, yeast and thylakoids) are found in almost half of the MP structures established in protein-based NDs (46 cases), whereas pure or mixed synthetic phospholipids are found in 44 cases (Fig. 6d). The most frequently used NDs are those resulting from a combination of MSP2N2 and natural lipids.

In 2019 and 2020, MSP-based NDs have been the most widely used system in SP cryo-EM (Fig. 7b) because the important role of lipid/MP interactions is being paid increasing attention to. This justifies the choice of NDs, which, at the cost of a lengthy optimization process required to obtain high-quality samples, allows one to solve MP structures in an artificial lipid bilayer more similar to the native membrane than any of the other surfactants used in SP cryoEM, barring liposomes. In the past, analyses of MPs embedded into liposomes were essentially performed by subtomography cryo-EM, yielding medium- to low-resolution structural data. More recently, imaging liposome-embedded MPs using SP cryo-EM has been developed based on the method of random spherically constrained SP reconstruction. This approach relies on fitting and subtracting a model of the membrane contribution to each image and reconstituting the MP particles alone (Wang and Sigworth, 2010; Tonggu and Wang, 2020). It has yielded cryo-EM structures of the large conductance voltage- and calcium-activated potassium (BK) channels first at 17 Å (Wang



**Fig. 8.** Addition of cholesteryl hemisuccinate (CHS) to MP samples. (*a*) Purification surfactants used in combination with CHS. (B) Distribution of MP structures solved in the presence of CHS among protein classes. 'Others' are as described in the legend to Figure 7.

and Sigworth, 2009), and then at 3.5 Å (Tonggu and Wang, 2018). Recently, another method of cryo-EM data processing was developed to determine high-resolution structures of MPs reconstituted into liposomes, as exemplified by the 3.9 Å structure of the multidrug-resistant transporter AcrB (Yao *et al.*, 2020). These approaches have the unique potential of making it possible to examine the structure of MPs under conditions where they experience a gradient of solutes or a transmembrane electric field. One should nevertheless bear in mind that none of these systems, be they liposomes, MSP-based NDs, SMALPs or MP/ APol/lipid complexes, provides MPs with an environment identical to that of the native membrane, if only because, in the latter, the two monolayers have different compositions (for a discussion, see Chapter 3 in Popot, 2018).

A practice worth mentioning is the addition, to surfactantsolubilized MPs, of cholesteryl hemisuccinate (CHS; Fig. 5), a water-soluble cholesterol analog that improves the stability of many purified MPs (Weiss and Grisshammer, 2002; Magnani *et al.*, 2008; Thompson *et al.*, 2011). According to published reports, 207 MPs out of 604 have been solubilized in the presence of CHS, and in 90 cases, CHS was kept along with the purification surfactants, which are preferentially maltoside-derived detergents and MNGs (Fig. 8*a*). In total, 71% of the MP structures solved in the presence of CHS, many of which are GPCRs and channels (Fig. 8*b*), have been obtained with these two detergents. With respect to the total number of MP structures solved in these two surfactant classes, the use of CHS represents 37% of the cases.



(b) Surfactants used for solubilization



**Fig. 9.** Surfactant exchange between MP purification and sample deposit onto cryo-EM grids. (*a*) Detergents used for MP solubilization and the initial steps of purification. In three cases (labeled 'Maltoside or MNGs'), the detergent used for solubilization has not been clearly specified. (*b*) Surfactants used for vitrification, depending on which detergent was used for solubilization, namely either maltoside detergents (mainly DDM), MNGs (mainly LMNG), or digitonin. In Panel A, the analysis includes duplicates provided that the surfactants used for solubilization were different (i.e. including duplicates 2 and 3), whereas in panel (*b*) all duplicates leading to  $\leq 5$  Å structures have been included. References to studies using the combinations of surfactants shown in panel (*b*) are given in the Supplementary Information in the legend to Table S1.

For the sake of exhaustiveness, the present analysis has considered all SP cryo-EM structures solved to resolutions  $\leq 5$  Å. One may argue that a threshold of 4 Å is more appropriate for building high-resolution models (Cheng, 2015). However, this would have reduced the number of experimental conditions to be analyzed from 604 to 487. We have checked that lowering the threshold from 5 to 4 Å has no impact on the general conclusions, i.e. the six most frequently used surfactants remain the same, their ranking changing as follows: MSP-based NDs now come first, with 18% of the structures, closely followed by maltoside detergents and digitonin, with almost 18 and 17%, respectively, then MNGs and GDN, each with 12%, and APols with ~9%. One clear lesson is that, as of now, no surfactant has emerged as the universal best environment for high-resolution SP cryo-EM.



**Fig. 10.** Surfactants used for solubilization and the initial steps of purification, depending on which surfactant was used for vitrification (either digitonin, GDN, MSP-based NDs, APols, maltoside detergents, or MNGs). All duplicates are included. In one case, the detergent used for vitrification has not been clearly specified and could be either maltosides or MNGs.

### Surfactant exchange before sample application to the grid

The surfactant used for protein purification is very often exchanged before the sample is applied onto the grids (Fig. 9; for references, see Table S1 in the Supplementary Information). Most often, extraction from the membrane and the first steps of purification are performed using one surfactant (hereafter referred to as the 'purification surfactant'), whereas the last steps are used to exchange it for that in which the sample will be vitrified ('vitrification surfactant'). Maltoside detergents are most frequently used for purification (~44%), followed by MNGs (~23%) and digitonin (~11%) (Fig. 9a). However, in ~68% of cases, MPs purified in maltoside detergents (essentially DDM) were subsequently transferred to a different amphipathic environment, most often MSP-based NDs, APols, digitonin, or GDN, before applying the sample to the grids (Fig. 9b). In nine cases, the maltoside detergent used for purification was exchanged for another maltoside detergent. Exchange of MNGs is also very common (~59% of cases), most often for digitonin, GDN or APols. Digitonin is less often exchanged. Out of 67 structures of MPs purified in digitonin (taking into account duplicates), 52 were obtained without resorting to surfactant exchange (Fig. 9b). Figure 10 summarizes, for each of the six types of surfactants most commonly used at the vitrification step, which medium the proteins were initially purified in. Slightly more than half of MPs whose structure was solved in digitonin were purified in other detergents, mainly maltoside-based detergents and MNGs. The frequent transfer to digitonin of MPs extracted with another detergent is not unexpected, given that digitonin is both less solubilizing and less destabilizing than most detergents. A large fraction of those MPs whose structure was solved in GDN were initially purified in maltoside detergents or in MNGs (Fig. 10).

When MSP-based NDs or APols are used for vitrification, surfactant exchange is most often obligatory, given that neither of these surfactants is effective for direct solubilization. In the case of APols, one can expect that the number of structures solved without surfactant exchange will increase should the use of SMAs and other APols suitable for direct MP extraction, like CyclAPols, develop in the future. Currently, near two-thirds and half of the MPs whose structures were established in MSP-based NDs or in APols, respectively, were purified in maltoside detergents (Fig. 10).

MNGs are also often chosen as final surfactants. Almost threequarters of the MP structures established in MNGs were obtained without detergent exchange. In 16% of the cases, the MP was first extracted using a maltoside-based detergent. In four cases, the MNG used for purification was exchanged before vitrification for another MNG.

DDM is, by far, the detergent that is the most often used for MP extraction and purification (Fig. 9*a*). As such, it is, most likely, the first detergent tested in initial imaging attempts. It is therefore noteworthy that DDM is, nevertheless, replaced with another surfactant in more than two-thirds (~68%) of the studies we examined (Fig. 9*b*). Conversely, DDM and the other maltoside detergents are seldom used to replace the detergent used for purification (Fig. 10). There are likely two main reasons for moving or keeping away from DDM for imaging: (i) the desire to transfer the protein to less perturbing and/or more native-like an environment, and (ii) the higher quality of images obtained with other surfactants. These two issues, of course, are not independent of each other.

Overall, the analysis summarized in Figs 9 and 10 indicates that it is more the exception than the rule that the surfactant used for extraction and purification be the one kept for preparing cryo-EM specimens, as it is exchanged in 60% of the cases. Table S1 of the Supplementary Information provides references to those studies in which the protein was initially obtained in one of the three types of surfactants most commonly used for purification, maltosides, MNGs and digitonin, sorted out according to what the medium used for vitrification was.

In any original study of a new MP, optimization of the purification and vitrification conditions is likely to stop once the quality of images appears sufficient for obtaining near-atomic resolution. In subsequent studies, however, when a better structural preservation and/or resolution is desired, or a different conformational state is explored, the search for improved conditions is broadened and may well lead to favoring another surfactant than that used in the first study. Because surfactants that have given good results in the past are tested first, there is initially a bias in their favor, which does not necessarily reflect an actual superiority, and the use of novel molecules tends to lag behind. A notable exception to this pattern is GDN, which, certainly by virtue of its similarity to digitonin and the advantages it presents over it, seems to have been adopted with remarkable rapidity.

#### Effect of surfactants on image quality

Many parameters affect the resolution of cryo-EM structures, including the number of particles used for the 3D reconstruction, grid preparation, imaging conditions, etc. A straightforward relationship between the surfactants used and image quality is therefore difficult to establish. However, a few cryo-EM studies have reported duplicate (or, in two cases, triplicate) structures of the same MP in two (or three) distinct amphipathic environments. One can expect that, in each of these studies, where the starting material is the same, all efforts have been made to optimize image quality and that, as a consequence, differences of resolution from one surfactant to the next can be more likely related to the surfactant itself than to any other factor.

Whereas high-CMC detergents are known to degrade the quality of images due to (i) their tendency to destabilize MPs (cf. Chapter 2 in Popot, 2018) and (ii) the presence of abundant micelles in the background, which lowers the contrast, resorting to low-CMC detergents tends to improve the resolution. These detergents, such as LMNG, exhibit an extremely slow off-rate, which has been used to develop a gradient-based detergent removal (GraDeR) approach enabling an extensive elimination of free detergent micelles and detergent monomers to further improve image quality (Hauer *et al.*, 2015). Empty NDs and APol excess can also be removed from the samples, before vitrification, by size exclusion chromatography.

Because it is not known beforehand which amphipathic environment will yield the best data, using in parallel two or more vitrification surfactants, which, a few years ago, was an anecdotal practice, is becoming increasingly frequent. Table 1 shows a comparison of the resolutions achieved for a given MP or MP complex, in a given study, in the presence of various amphipathic environments (the best resolution achieved in a given study is noted in *red*, the surfactant used indicated on the top line, whereas other data obtained in the same study using an alternative surfactant - reported in the left column - are noted in *black*). It reveals a clear tendency in favor of MSP-based NDs (12 of the best structures), followed by GDN (6), maltoside-based detergents, MNGs and APols (5 each). An additional benefit of such comparative approaches is that two structures solved at similar resolution in distinct environments may reveal different conformations, which could provide a glimpse into the conformational landscape of the protein. A case in point is mouse TMEM16A, whose structures in LMNG and NDs show what seems to be two distinct closed conformations, featuring either one or two bound Ca<sup>2+</sup> ions per monomer (Dang et al., 2017). There are other such examples in the literature, such as, for instance, the cryo-EM structures of the full-length rabbit TRPV2 channel obtained in APols versus MSP-based NDs (Zubcevic et al., 2019). This suggests that different amphipathic environments exert different constraints on the transmembrane domain of a given MP, leading to the stabilization of different conformational states. Molecular dynamics data show that APols may slow down MP dynamics (Perlmutter et al., 2014). On the one hand, this probably contributes to stabilizing MPs as compared to conventional detergents (Picard et al., 2006; Pocanschi et al., 2013); on the other, it may favor low-energy conformational states. This can be seen as an advantage to reduce the conformational heterogeneity of samples, like nanobodies and/or Fabs do. Compared to APols, NDs may restrict transmembrane domain conformational states to a lesser extent. Nevertheless, a tight wrapping of MPs by MSPs has been reported (for examples, see Arkhipova et al., 2020; Roh et al., 2020), which may potentially give rise to conformational constraints. The presence of lipids in MP/ND or MP/polymer complexes may increase the dynamics of the system, allowing MPs to adopt a range of conformations, which can be relevant to the protein's function.

In other studies, on the contrary, no significant differences were observed, as is the case of the 3.8 Å resolution structures of the innexin-6 gap junction proteins in an undocked hemichannel observed in either NDs or MNG (Burendei *et al.*, 2020), or of the 2.9 Å resolution structures of the human potassium-chloride cotransporter (KCC1) in NDs *versus* GDN (Liu *et al.*, 2019b).

Table 1. A comparison of resolutions achieved for a given MP or MP complex vitrified in various amphipathic environments

	Surfactants giving the highest resolution (Å)								
Surfactants used for vitrification	Digitonin	Maltosides	MSP-based NDs	MNGs	Amphipathic polymers	GDN	Others <sup>k</sup>		
Digitonin	3.2; 3.8 <sup>a</sup> (Tao and MacKinnon, 2019)	4 (Sui <i>et al.</i> , 2018)	3.2; 3.5 (Feng <i>et al.</i> , 2019) 3; 3.9 (Bloch <i>et al.</i> , 2020)		4.2; 4.3 (Yin <i>et al.</i> , 2019 <i>a</i> , 2019 <i>b</i> )	4.24; 4.3 (Singh et al., 2018) 3.3; 3.5 (Yan et al., 2019) 2.6; 2.9 (Zhao et al., 2019a, 2019b, 2019c, 2019d)			
Maltosides	4 (Sui <i>et al.</i> , 2018)	<ul> <li>3.3; 3.9<sup>b</sup> (Gopalasingam et al., 2019)</li> <li>3.3; 3.9<sup>c</sup> (Niu et al., 2020)</li> </ul>	3.1; 3.4 (Reddy <i>et al.</i> , 2019)	3.2; 4 (Parey <i>et al.</i> , 2019)	6.8 (Bausewein <i>et al.</i> , 2017) 3.1 (Johnson <i>et al.</i> , 2020)		3.1 (Johnson et al., 2020)		
MSP-based NDs	3.2; 3.5 (Alvadia <i>et al.</i> , 2019)	<mark>3.7</mark> (3.6); 3.8 (3.6) <sup>g</sup> (Kalienkova <i>et al.</i> , 2019)	<ul> <li>3.6; 3.8<sup>d</sup> (Srivastava et al., 2018)</li> <li>3.5; 3.93 (Kishikawa et al., 2020)</li> <li>3.5; 8.25 (Guan et al., 2020)</li> </ul>	<b>3.67</b> ; 4.92 (Wang <i>et al.</i> , 2020 <i>b</i> ) 3.8; 3.8 <sup>h,i</sup> (Dang <i>et al.</i> , 2017; Burendei <i>et al.</i> , 2020) <b>4.1</b> ; 6.6 (Jin <i>et al.</i> , 2020)	3.7 (Yoo et al., 2018) 2.9; 3.8 (Zubcevic et al., 2019) 3.4; 3.4 (Maeda et al., 2020)	3.96; 7.77 (Walter et al., 2019) 2.9; 2.9 (Liu et al., 2019b) 3; 3.39 (Deng et al., 2020)			
Saposin-based NDs		3.8; 5 (Nguyen <i>et al.</i> , 2018) 3.34; 4.7 (Gharpure <i>et al.</i> , 2019)							
MNGs			<ul> <li>3.1; 3.5 (Jojoa-Cruz et al., 2018)</li> <li>3.8; 3.8<sup>h;i</sup> (Zhou et al., 2017; Burendei et al., 2020)</li> <li>3.4; 4 (Maeda et al., 2020)</li> </ul>		4.06; 4.65 (Zhou et al., 2017) 4; 4.1 (Kuhlen et al., 2020) 3.6; 3.6 (Vinayagam et al., 2020) 3.4; 4 (Maeda et al., 2020)		4.1; 4.2 (Benton et al., 2018)		
Amphipathic polymers		6.8 (Bausewein <i>et al.</i> , 2017) 3.4; 3.5 (Zubcevic <i>et al.</i> , 2018) 3.1 (Johnson <i>et al.</i> , 2020)	3; 4 (Shen <i>et al.</i> , 2016) 3.7 (Yoo <i>et al.</i> , 2018) 3.6; 4.2 (McGoldrick <i>et al.</i> , 2018) 3.15; 3.16(Sauer <i>et al.</i> , 2020) 3.4; 3.4 (Maeda <i>et al.</i> , 2020)	3.58; 6.25 (Alam <i>et al.</i> , 2018) 2.6; 3.3 (Zhao <i>et al.</i> , 2020) 3.6; 3.6 (Vinayagam <i>et al.</i> , 2020)	<b>3.63;</b> 4.2 <sup>e</sup> (Wang et al., 2019) <b>2.17;</b> 3 <sup>f</sup> (Owji et al., 2020)	2.83; 6.04 (Ruan et al., 2020a)	3.1 (Johnson et al., 2020)		
GDN		3.06; 4.1 (Tucker and Park, 2019)	4; 4 (7.9) <sup>j</sup> (Koehl <i>et al.</i> , 2019) 3.45; 3.53 (McGoldrick <i>et al.</i> , 2019) 2.9; 2.9 (Liu <i>et al.</i> , 2019 <i>b</i> ) 3.1; 3.7 (Gao <i>et al.</i> ,						

	Surfactants giving the highest resolution (Å)							
Surfactants used for vitrification	Digitonin	Maltosides	MSP-based NDs	MNGs	Amphipathic polymers	GDN	Others <sup>k</sup>	
			2020 <i>a</i> ) 3; 3.6 (Qian <i>et al.</i> , 2020) 3.4; 3.6 (Diederichs <i>et al.</i> , 2020)					
CHAPS	3.6; 4.2 (Gong <i>et al.</i> , 2019)							
Triton-X100		3.1 (Johnson <i>et al.</i> , 2020)			3.1 (Johnson <i>et al.</i> , 2020)			
Total	2	5	12	5	5	6	1	

Only the resolutions (in Å) of MP cryo-EM structures published in the same study and for which PDB or EMDB accession codes are available are reported. These reports are referred to in the text and figures as 'duplicates'. The columns list the surfactants whose use led to the best resolution for the MP considered (in a given study), printed in *red*, the lines the alternative surfactants used in the same study, the resolution achieved being indicated in *black*. In four studies, the analysis resulted in indistinguishable 3D reconstructions regardless of the environment and the data were pooled to yield a single structure, in which cases the resolution is printed in *green*. These four studies and those which resorted to two or more surfactants belonging to the same surfactant family have not been included in the total number of best-resolution structures achieved with each surfactant family (last line).

Note that for duplicate structures obtained using the same surfactant family, the difference of resolution could be attributed to the following reasons: <sup>a</sup>detergent exchange in one case and not in the other, the best resolution being obtained when the sample was purified and kept in digitonin rather than purified in MNG and transferred to digitonin; <sup>b</sup>two versions of the target MP (wild-type *versus* mutant); <sup>c</sup>the vitrification detergent was either DDM or DM, the latter giving the best resolution; <sup>d</sup>the lipid compositions of NDs differed; <sup>e</sup>two pH values (6.2 and 7.5) were used, the best resolution being obtained at the highest pH; <sup>f</sup>the vitrification surfactant was either A8-35 or PMAL-C8, the latter giving the best resolution.

In some cases, the same resolution was achieved in two distinct amphipathic environments: <sup>8</sup>the structures of the apo and halo forms of TMEM16 were obtained at different resolutions, namely 3.7 and 3.8 Å for the calcium-free structure in DDM and MSP-based NDs, respectively, and 3.6 Å for the calcium-bound one regardless to the environment; <sup>1</sup>the detergent sample was prepared using the GraDeR approach (Hauer *et al.*, 2015); <sup>1</sup>the same resolution (3.8 Å) was reported for the innexin-6 hemichannel in LMNG and NDs; however, the resolution in LMNG can be improved to 3.4 Å by reducing the anisotropy of the reconstruction thanks to the addition in the refinement process of particles from a dataset in which a Fab fragment was bound to the target MP; <sup>1</sup>the structures of two conformational states of the metabotropic glutamate receptor (a GPCR) were solved to the same resolution (4 Å), namely the active conformation in GDN and the inactive (apo) form in MSP-based NDs; that of the apo form in GDN could be solved to only 7.9 Å resolution; note that a large population of the receptor in its apo form was observed to be dissociated from its dimeric state into monomeric forms, suggesting detergent-induced destabilization; <sup>k-</sup>Others' in the last column correspond to MNA-C12 (influenza virus hemagglutinin) and Triton-X100 (MS-ring of bacterial flagellum).

In one case, three cryo-EM structures of the same MP, the autophagy-related 9 protein, were resolved in three different environments, namely MSP-based NDs, APols, and LMNG. Whereas no significant differences were observed between NDs and APols, both surfactants leading to 3.4 Å structures with an r.m.s.d. over all  $C\alpha$  atoms of 0.68 Å, the use of LMNG resulted in a lesser resolution (Maeda *et al.*, 2020). In four cases, the absence of differences between distinct amphipathic environments led to merging different datasets to build a single cryo-EM structure (whose resolution is noted in *green* in Table 1).

Over the years, groups that initially used a particular surfactant to study a given MP or MP complex may move to another one. A case in point is that of mitochondrial respiratory supercomplexes, historically solubilized and purified in digitonin. Different types of supercomplexes have been identified, the biggest one being the respirasome, which results from the association of complexes I<sub>1</sub>III<sub>2</sub>IV<sub>1</sub>, comprising altogether 80 subunits and featuring 132 transmembrane helices. The digitonin/protein ratio used during solubilization considerably affects the types and relative amounts of supercomplexes (Pérez-Pérez et al., 2016), pointing to a dissociating effect. The first 3D reconstruction ever achieved of a whole mammalian respirasome, which predated the 'resolution revolution' in SP cryo-EM, was obtained by stabilizing the supercomplex in APol A8-35 after extracting it with digitonin. Whereas the resolution of the map was limited to  $\sim 19$  Å, it made it possible to position the three respiratory complexes with respect to each other within the supramolecular assembly (Althoff et al., 2011). Five years later, thanks to progress in cryo-EM technology, four structures of respirasomes from three different organisms were obtained at subnanometer resolution and published nearly simultaneously (Gu et al., 2016; Letts et al., 2016; Sousa et al., 2016; Wu et al., 2016). In one study, it was reported that 42% of the particles had lost complex IV when purified in digitonin, whereas the particles extracted with a DDM analog, PCC-a-M (Hovers et al., 2011), and transferred to A8-35 were homogeneous and stable (Sousa et al., 2016). In another study, where digitonin was used both for purification and vitrification, two 3D reconstructions of the ovine respirasome in 'tight' and 'loose' forms were obtained at resolutions of 5.8 and 6.7 Å, respectively (Letts et al., 2016). Over storage, the ratio of the tight to loose respirasomes changed in favor of the loose form, whose population doubled after overnight incubation at 4 °C before grid preparations. In the same study, the authors also presented the cryo-EM structure of supercomplex  $I_1III_2$  in digitonin at 7.8 Å. More recently, the same authors reported that transferring the digitonin-solubilized supercomplex I<sub>1</sub>III<sub>2</sub> to APol A8-35 preserved all expected enzymatic activities and significantly improved the supercomplex stability over time, leading to cryo-EM structures solved, in four distinct states, at resolutions ranging from 4.6 to 3.8 Å (Letts et al., 2019). Nevertheless, replacing digitonin with APol is most likely not the only reason explaining the improvement of resolution, as other parameters such as the number of particles, grid preparation, and imaging conditions may have affected the final resolution of the 3D reconstruction. In addition, one should keep in mind that, despite their lesser dissociating character as compared to detergents, too high an excess of APols can also result in breaking up large MP complexes into subfragments (Popot et al., 2011; Sverzhinsky et al., 2014). This process can be limited by (i) finetuning the APol concentration following a rather simple method (for a detailed protocol, see Le Bon et al., 2018), (ii) using less dissociating APols such as NAPols, and (iii) adding lipids.

#### Vitrification

The usual approach to vitrification is as follows: the carbon film of a crvo-EM grid is rendered hydrophilic by plasma treatment, the sample is deposited on the grid and incubated in a chamber under controlled humidity and temperature, excess liquid is removed (blotted) by touching the surface of the grid with a filter paper, and the resulting thin supported liquid film is flash-frozen by immersion into a cryogen (typically liquid ethane) (Dobro et al., 2010; Sgro and Costa, 2018). Blotting is considered one of the critical steps for achieving reproducible ice quality and thickness (Armstrong et al., 2020). Processes that avoid the hard-to-standardize contact with a filter paper are thus being developed. These include self-blotting grids (nano-wire support grids absorbing the excess liquid due to an increased adsorption surface (Razinkov et al., 2016)), removal of excess liquid by applying a pressure gradient ('Preassis'; Zhao et al., 2019b), microfluidic isolation and controlled deposition of the protein of interest (Schmidli et al., 2019), or automated deposition of minimal sample volumes, either directly ('Spotiton' inkjet dispenser (Arnold et al., 2017); 'Vitrojet' pin-printing (Ravelli et al., 2020)) or through microfluidic-based (Feng et al., 2017) or ultrasoundbased (Ashtiani et al., 2018) spraying. To our knowledge, no new structure has so far been solved using any of the latter prototypes, despite promising preliminary results. Commercial vitrification devices based on blotting (sold by FEI, Leica, and Gatan) remain overwhelmingly dominant (~95% of the studies). Including home-made devices, more than 97% of the sample preparation is performed with a paper-based blotting step (the missing 3% reflecting a lack of information rather than blotting alternatives). This might be attributable to a lack of visibility/ availability of newly developed prototypes, or to still ongoing proof-of-concept studies. A point to keep in mind is that, at the blotting stage, most of the surfactant adsorbed at the air/water interface will be removed. The rate of reformation of a surfactant monolayer will depend on the nature and concentration of the surfactant, which may affect such factors as thinning of the film and MP adsorption at the interface.

#### Can surfactants be used as vitrification helpers?

The preparation of high-quality cryo-EM grids depends on being able to preserve the target macromolecule in a vitreous thin film, with the particles exhibiting an even distribution and, ideally, adopting random orientations within the ice layer. The key to achieving high resolution is a careful optimization of cryospecimen preparation (Sgro and Costa, 2018). For MP samples, the presence of surfactant represents a critical factor. Several types of difficulties have been identified, including uneven dispersion of MPs throughout the ice film (aggregation at the edge of the holes in carbon films, particularly if the ice film is too thin, and/or preferential orientations) and lowering of the contrast due to the presence of detergent micelles in the background. The latter problem has encouraged the use of APols, NDs or low-CMC detergents to minimize the presence of free surfactant. MP dispersion within the ice film constitutes the most difficult parameter to control (Drulyte et al., 2018). Commercial cryo-EM grids are metal mesh grids, typically copper, with a support film of amorphous carbon layered over the top, which can be either continuous or perforated. Amorphous carbon support films remain the preferred cryo-EM support material because it is relatively inexpensive. However, it is prone to bending and

Additive: Main surfactant	Fos-choline-8 (0.5–3 mM)	Octyl-maltoside (0.01–0.5%)	Fos-choline 8 (1.5 mM) or octyl maltoside (0.7 mM)	n-Octyl-β-⊳-glucopyranoside (0.05 or 0.1%)	CHAPS/ CHAPSO (0.05-0.2%)	Brij-35 (0.02%)	Digitonin (0.1%)	DDM (0.06 mM)	Fos-choline-8	A8-35 (0.0005%)	PMAL-C8 (0.1– 0.2%)	Total
MSP-based NDs	(Falzone <i>et al.</i> , 2019; Khelashvili <i>et al.</i> , 2019; Zubcevic <i>et al.</i> , 2019; Fan <i>et al.</i> , 2020; Wang and Boudker, 2020; Yin <i>et al.</i> , 2020; Huang <i>et al.</i> , 2020b)	(Gatsogiannis <i>et al.</i> , 2016; Willegems and Efremov, 2018; Kater <i>et al.</i> , 2019; Reddy <i>et al.</i> , 2019; Qian <i>et al.</i> , 2020; Ruan <i>et al.</i> , 2020b)			(Pleiner <i>et al.</i> , 2020)				(Sun and MacKinnon, 2020)	(Koehl et al., 2019)		16
MNGs	(Wasilko <i>et al.</i> , 2020)	(Deme <i>et al.</i> , 2020)	(Lauber <i>et al.</i> , 2018)	(Yu <i>et al.</i> , 2019)	(Kampjut and Sazanov, 2019; Kampjut and Sazanov, 2020; Maeda <i>et al.</i> , 2020; Pinke <i>et al.</i> , 2020)							8
Digitonin	(Guo and MacKinnon, 2017; Liu et al., 2017; Zhang et al., 2017; Johnson Chen, 2018; Zhang et al., 2018b; Liu et al., 2019a; Wang et al., 2020a)											7
Amphipols	(Wang et al., 2019)					(Letts <i>et al.,</i> 2019)	(Maldonado et al., 2020)	(Owji <i>et al.</i> , 2020)			(Owji et al., 2020)	5
GDN	(Itskanov and Park, 2019; Tucker and Park, 2019)			(Koehl <i>et al.</i> , 2019; Kim <i>et al.</i> , 2020 <i>b</i> )								4
Maltosides	(Tucker and Park, 2019; Niu <i>et al</i> ., 2020)	(Coleman <i>et al.</i> , 2020)										3
Saposin-based NDs	(Gharpure <i>et al.</i> , 2019; Rahman <i>et al.</i> , 2020; Kim <i>et al.</i> , 2020a)											3
POE-glycols	(Basak <i>et al.</i> , 2019)											1
<i>n</i> -Octyl-β-□-glucopyranoside				(Benton et al., 2020)								1
No surfactant <sup>a</sup>		(Linsky <i>et al.</i> , 2020; Piton <i>et al.</i> , 2020)		(Gao <i>et al.</i> , 2020 <i>b</i> )								3
Total	24	10	1	5	5	1	1	1	1	1	1	51

Hydrogenated detergents

 Table 2. Studies in which the addition of secondary surfactants to solubilized MP samples has been carried out before vitrification

 Fluorinated surfactants

<sup>a</sup>No surfactant has been used for the purification of SARS-CoV-2S ectodomain (residues16–1206) (Linsky *et al.*, 2020), the mycobacterial EspB heptamer, which appears as an external membrane protein (Piton *et al.*, 2020), and the visual G protein-effector complex, a peripherally attached complex to the rod photoreceptor outer segment membranes (Gao *et al.*, 2020*b*).

Amphipathic polymers

deformation as a result of exposure to the electron beam (Brilot et al., 2012), entailing afterwards a correction of beam-induced movements. All-gold cryo-EM grids, which have been shown to significantly reduce specimen motion during illumination with the electron beam (Russo and Passmore, 2014), are now more frequently tested. Chemical modification of gold-coated grids with a thiol bearing a PEG group has been developed as a mean to reduce the aggregation of soluble proteins on the support and at the edges of the holes (Meyerson et al., 2014), an approach that has also been shown to improve the distribution and orientations of MPs (Blaza et al., 2018). Another approach is the chemical modification of the MP itself by reacting its surface-exposed lysines with an appropriately functionalized low-molecular-mass PEG (Wu et al., 2019; Wu et al., 2020). Obtaining thicker films can alleviate the orientation problem and it seems that APols can help (Flötenmeyer et al., 2007). An additional risk, which affects both MPs and soluble proteins, is protein adsorption at the air/water interface, which generally results in partial or complete denaturation (Noble et al., 2018). Immobilizing proteins on a graphene-coated surface is a way to prevent this adsorption, as shown with soluble proteins (D'Imprima et al., 2019). Binding of megabodies has also been reported to randomize the particle distribution in ice for MPs that normally exhibit preferential orientations (Uchański et al., 2021).

Surfactant molecules adsorb at the air/water interface, which has two interesting consequences: on the one hand, this reduces the surface tension of the buffer and likely influences the thinning kinetics; on the other, it creates a barrier that limits protein adsorption and denaturation at the interface (Glaeser *et al.*, 2016; Glaeser and Han, 2017). The first effect adds to the complexity of grid preparation because the rate at which the film drains, its thickness at the time of vitrification and, as a result, the even or uneven distribution and orientation of MPs can be difficult to control. It is, however, a factor that can be harnessed to improve particle distribution (cf. Flötenmeyer *et al.*, 2007).

It has been reported that the addition of small amounts of surfactants to a sample of soluble proteins can improve particle distribution and/or orientation within the ice film, which facilitates collecting the various views required for 3D reconstruction. This improvement has been observed for instance with detergents such as CHAPSO (Chen *et al.*, 2019) as well as with APol A8-35 (Chowdhury *et al.*, 2015; Zhang *et al.*, 2018a). Table 2 lists the cases where a secondary surfactant has been added to solubilized MP samples in order to improve the quality of the samples. NDs and MNGs are the two systems in which the addition of extra surfactants has been most often resorted to (Table 2).

Fluorinated surfactants have proven particularly useful for this task because (i) they mix poorly with hydrogenated alkyl chains of detergent molecules and (ii) they are strongly attracted to the air/water interface, where they oppose the adsorption of hydrogenated molecules (cf. Chapter 3 in ref. Popot (2018)). Fluorinated Fos-choline 8, with a CMC of 2.5 mM, is most frequently used, followed by fluorinated octylmaltoside (CMC = 0.7 mM). MP cryo-EM structures have also been obtained using as additives hydrogenated detergents, such as Fos-choline 8, DDM, digitonin, CHAPS/CHAPSO, Brij-35, or octylglucoside (Table 2). An interesting observation is that A8-35 can also facilitate sample vitrification for ND-trapped MPs (Koehl *et al.*, 2019). As with small surfactants, the addition of A8-35 seems to oppose the adsorption of MP/ND complexes at the air/water interface, most likely due to its forming an interfacial film (cf. Giusti *et al.*, 2012).

Preferential orientation has been reported in the case of the ryanodine receptor RyR1 trapped in A8-35 (Baker *et al.*, 2015). Random orientation of the particles was achieved following addition to the sample of a low concentration of octylglucoside (one-tenth of the CMC). The mechanism underlying this effect is not clear (for a discussion, see Chapter 12 in Popot, 2018). Combining two or more surfactants can thus have favorable effects, although it complicates optimizing grid preparation.

#### Using functionalized APols for immobilizing MP complexes or labelling transmembrane domains

One of the advantages of APols derives from the fact that grafting a small percentage of a tag or label onto the polymer does not affect its solution properties, while functionalizing it for specific purposes. Thereby, transferring a MP from detergents to a functionalized APol results in functionalizing the MP of interest without the need for any chemical nor genetic modification (Della Pia et al., 2014a). This can be achieved either by exchanging detergents for labelled APols (Le Bon et al., 2018) or, given that APols are miscible (Zoonens et al., 2007; Della Pia et al., 2014b), by adding a functionalized APol to a MP directly extracted using the newly developed CyclAPols (Marconnet et al., 2020). Many applications of such a mild labeling can be contemplated. On the one hand, chemically functionalized carbon films carrying biotin, nickel-charged nitrilotriacetic acid (Ni<sup>2+</sup>-NTA), DNA, and so on have been described (Llaguno et al., 2014). Functionalized APols carrying biotin moieties (Charvolin et al., 2009), hexahistidine (His<sub>6</sub>)-tags (Giusti et al., 2015), or a single strand of DNA (Le Bon et al., 2014) could be used to immobilize MPs on the corresponding films, so as to provide an even distribution and prevent protein contact with the air/water interface. In addition, functionalized APols can be used to label the transmembrane domain of some challenging MP complexes for which only medium- to low-resolution structures are currently available. The principle of this approach has been validated using negative-stain EM (Perry et al., 2019). It relies on trapping the target MP with an APol carrying biotin moieties (BAPol) (Charvolin et al., 2009) and localizing the BAPol-coated transmembrane regions using monovalent streptavidin. This labelling strategy could presumably help in the initial steps of a cryo-EM study, even though it should be kept in mind that streptavidin monomers, while bound to the surface of the APol layer, will not occupy identical positions from one particle to the next. This approach could be extended to other functionalized APols. For instance, APols carrying His<sub>6</sub>-tags (Giusti et al., 2015) can bind Ni<sup>2+</sup>-functionalized gold nanoparticles or quantum dots, whereas those carrying an oligonucleotide (Le Bon et al., 2014) could be used to create or bind to origami motifs (Martin et al., 2016).

#### Conclusion

The present analysis provides an overview of how surfactants are being used in the field of SP cryo-EM MP studies. One remarkable observation is that, even when a mild detergent such as DDM, LMNG, digitonin, or GDN is used for MP extraction and purification, microscopists often find it advantageous to exchange it for another surfactant such as a milder detergent, NDs or APols in order to improve the quality of the reconstructions and/or to catch distinct conformational states. One can expect that, in the future, developing still better surfactants will be one of the major avenues to improving cryo-EM data. Two major directions will likely be explored in parallel: (i) improving surfactants that can extract MPs directly while also providing excellent images, so that no surfactant exchange would be necessary, and (ii) improving the final surfactant used to maintain MPs in solution during grid preparation, irrespective of whether it is also used for the initial solubilization and purification steps. Further progress will likely also depend on a better control of the factors that govern the thickness of the ice film and the distribution and orientation of MP/surfactant complexes within it. Here as well, a better understanding of the behavior of surfactants and liquid films at the blotting step will probably prove critical. Finally, the use of functionalized surfactants offers intriguing perspectives.

**Supplementary material.** The supplementary material for this article can be found at https://doi.org/10.1017/S0033583521000044.

Acknowledgements. We would like to thank Daniel Picot for interesting discussions on the 3D reconstruction of structural models solved either by X-ray crystallography or by SP cryo-EM and Mark E. Dumont, Richard Henderson, Pierre D. McCrea, and the referees of *QRB* for very useful comments on earlier versions of the review. Financial support came from the Centre National de la Recherche Scientifique (CNRS), including specific '80PRIME' grant, and the Université de Paris (Université Paris-7). Additional funding came from the 'Initiative d'Excellence' program from the French State (Grant 'DYNAMO', ANR-11-LABX-0011-01).

Conflict of interest. None.

#### References

- Abeyrathne PD, Arheit M, Kebbel F, Castano-Diez D, Goldie KN, Chami M, Stahlberg H, Renault L and Kühlbrandt W (2012) Analysis of 2-D crystals of membrane proteins by electron microscopy. *Comprehensive Biophysics* 1, 277–310.
- Alam A, Küng R, Kowal J, McLeod RA, Tremp N, Broude EV, Roninson IB, Stahlberg H and Locher KP (2018) Structure of a zosuquidar and UIC2-bound human-mouse chimeric ABCB1. Proceedings of the National Academy of Sciences 115, e1973–e1982.
- Alnabati E and Kihara D (2019) Advances in structure modeling methods for cryo-electron microscopy maps. *Molecules* 25, 82.
- Althoff T, Mills DJ, Popot J-L and Kühlbrandt W (2011) Arrangement of electron transport chain components in bovine mitochondrial supercomplex I<sub>1</sub>III<sub>2</sub>IV<sub>1</sub>. The EMBO journal **30**, 4652–4664.
- Alvadia C, Lim NK, Clerico Mosina V, Oostergetel GT, Dutzler R and Paulino C (2019) Cryo-EM structures and functional characterization of the murine lipid scramblase TMEM16F. *eLife* 8, e44365.
- Andréll J and Tate CG (2013) Overexpression of membrane proteins in mammalian cells for structural studies. *Molecular Membrane Biology* 30, 52–63.
- Arkhipova V, Guskov A and Slotboom DJ (2020) Structural ensemble of a glutamate transporter homologue in lipid nanodisc environment. *Nature Communications* 11, 1–9.
- Armstrong M, Han B-G, Gomez S, Turner J, Fletcher DA and Glaeser RM (2020) Microscale fluid behavior during cryo-EM sample blotting. *Biophysical Journal* 118, 708–719.
- Arnold SA, Albiez S, Bieri A, Syntychaki A, Adaixo R, McLeod RA, Goldie KN, Stahlberg H and Braun T (2017) Blotting-free and lossless cryo-electron microscopy grid preparation from nanoliter-sized protein samples and single-cell extracts. *Journal of Structural Biology* 197, 220–226.
- Ashtiani D, Venugopal H, Belousoff M, Spicer B, Mak J, Neild A and de Marco A (2018) Delivery of femtolitre droplets using surface acoustic wave based atomisation for cryo-EM grid preparation. *Journal of Structural Biology* 203, 94–101.
- Autzen HE, Julius D and Cheng Y (2019) Membrane mimetic systems in CryoEM: keeping membrane proteins in their native environment. *Current Opinion in Structural Biology* 58, 259–268.

- Baker MR, Fan G and Serysheva II (2015) Single-particle cryo-EM of the ryanodine receptor channel in an aqueous environment. *European Journal of Translational Myology* **25**, 4803.
- Basak S, Gicheru Y, Kapoor A, Mayer ML, Filizola M and Chakrapani S (2019) Molecular mechanism of setron-mediated inhibition of full-length 5-HT3A receptor. *Nature Communications* **10**, 3225.
- Bausewein T, Mills DJ, Langer JD, Nitschke B, Nussberger S and Kühlbrandt W (2017) Cryo-EM structure of the TOM core complex from *Neurospora crassa. Cell* 170, 693–700.e7.
- Bazzacco P, Billon-Denis E, Sharma KS, Catoire LJ, Mary S, Le Bon C, Point E, Banères J-L, Durand G, Zito F, Pucci B and Popot J-L (2012) Nonionic homopolymeric amphipols: application to membrane protein folding, cell-free synthesis, and solution nuclear magnetic resonance. *Biochemistry* 51, 1416–1430.
- Benton DJ, Nans A, Calder LJ, Turner J, Neu U, Lin YP, Ketelaars E, Kallewaard NL, Corti D, Lanzavecchia A, Gamblin SJ, Rosenthal PB and Skehel JJ (2018) Influenza hemagglutinin membrane anchor. Proceedings of the National Academy of Sciences of the USA 115, 10112– 10117.
- Benton DJ, Gamblin SJ, Rosenthal PB and Skehel JJ (2020) Structural transitions in influenza haemagglutinin at membrane fusion pH. *Nature* 583, 150–153.
- Bhairi SM and Mohan C (2007) Detergents. A guide to the properties and uses of detergents in biology and biochemistry (Calbiochem booklet), Darmstadt: EMD Biosciences, an affiliate of Merck KGaA.
- Bill RM, Henderson PJF, Iwata S, Kunji ERS, Michel H, Neutze R, Newstead S, Poolman B, Tate CG and Vogel H (2011) Overcoming barriers to membrane protein structure determination. *Nature Biotechnology* 29, 335–340.
- Blaza JN, Vinothkumar KR and Hirst J (2018) Structure of the deactive state of mammalian respiratory complex I. *Structure* 26, 312–319.e3.
- Bloch JS, Pesciullesi G, Boilevin J, Nosol K, Irobalieva RN, Darbre T, Aebi M, Kossiakoff AA, Reymond J-L and Locher KP (2020) Structure and mechanism of the ER-based glucosyltransferase ALG6. *Nature* 579, 443–447.
- Brilot AF, Chen JZ, Cheng A, Pan J, Harrison SC, Potter CS, Carragher B, Henderson R and Grigorieff N (2012) Beam-induced motion of vitrified specimen on holey carbon film. *Journal of Structural Biology* 177, 630–637.
- Burendei B, Shinozaki R, Watanabe M, Terada T, Tani K, Fujiyoshi Y and Oshima A (2020) Cryo-EM structures of undocked innexin-6 hemichannels in phospholipids. *Science Advances* **6**, eaax3157.
- Cao E, Liao M, Cheng Y and Julius D (2013) TRPV1 structures in distinct conformations reveal activation mechanisms. *Nature* 504, 113–118.
- Carlson ML, Young JW, Zhao Z, Fabre L, Jun D, Li J, Li J, Dhupar HS, Wason I, Mills AT, Beatty JT, Klassen JS, Rouiller I and Duong F (2018) The Peptidisc, a simple method for stabilizing membrane proteins in detergent-free solution V. Dötsch & R. Aldrich, eds. *eLife* 7, e34085.
- Chae PS, Rasmussen SGF, Rana RR, Gotfryd K, Chandra R, Goren MA, Kruse AC, Nurva S, Loland CJ, Pierre Y, Drew D, Popot J-L, Picot D, Fox BG, Guan L, Gether U, Byrne B, Kobilka B and Gellman SH (2010) Maltose-neopentyl glycol (MNG) amphiphiles for solubilization, stabilization and crystallization of membrane proteins. *Nature Methods* 7, 1003–1008.
- Chae PS, Rasmussen SGF, Rana RR, Gotfryd K, Kruse AC, Manglik A, Cho KH, Nurva S, Gether U, Guan L, Loland CJ, Byrne B, Kobilka BK and Gellman SH (2012) A new class of amphiphiles bearing rigid hydrophobic groups for solubilization and stabilization of membrane proteins. Chemistry (Weinheim an Der Bergstrasse, Germany) 18, 9485–9490.
- Charvolin D, Perez J-B, Rouvière F, Giusti F, Bazzacco P, Abdine A, Rappaport F, Martinez KL and Popot J-L (2009) The use of amphipols as universal molecular adapters to immobilize membrane proteins onto solid supports. *Proceedings of the National Academy of Sciences of the* USA 106, 405–410.
- **Charvolin D, Picard M, Huang L-S, Berry EA and Popot J-L** (2014) Solution behavior and crystallization of cytochrome bc<sub>1</sub> in the presence of amphipols. *The Journal of Membrane Biology* **247**, 981–996.
- Chen J, Noble AJ, Kang JY and Darst SA (2019) Eliminating effects of particle adsorption to the air/water interface in single-particle cryo-electron

microscopy: bacterial RNA polymerase and CHAPSO. *Journal of Structural Biology X* **1**, 100005.

- Cheng Y (2015) Single-particle cryo-EM at crystallographic resolution. *Cell* **161**, 450–457.
- Cheng Y (2018) Single-particle cryo-EM How did it get here and where will it go. *Science* **361**, 876–880.
- Chipot C, Dehez F, Schnell JR, Zitzmann N, Pebay-Peyroula E, Catoire LJ, Miroux B, Kunji ERS, Veglia G, Cross TA and Schanda P (2018) Perturbations of native membrane protein structure in alkyl phosphocholine detergents: a critical assessment of NMR and biophysical studies. *Chemical Reviews* 118, 3559–3607.
- Chowdhury S, Ketcham SA, Schroer TA and Lander GC (2015) Structural organization of the dynein-dynactin complex bound to microtubules. *Nature Structural & Molecular Biology* **22**, 345–347.
- Choy BC, Cater RJ, Mancia F and Pryor EE (2021) A 10-year meta-analysis of membrane protein structural biology: detergents, membrane mimetics, and structure determination techniques. *Biochimica et Biophysica Acta* (*BBA*) *Biomembranes* 1863, 183533.
- Coleman JA, Navratna V, Antermite D, Yang D, Bull JA and Gouaux E (2020) Chemical and structural investigation of the paroxetine-human serotonin transporter complex. *eLife* 9, e56427.
- Contreras-Gómez A, Sánchez-Mirón A, García-Camacho F, Molina-Grima E and Chisti Y (2014) Protein production using the baculovirus-insect cell expression system. *Biotechnology Progress* **30**, 1–18.
- Cooper RS and Heldwein EE (2020) Expression, purification, and crystallization of full-length HSV-1 gB for structure determination. *Methods in Molecular Biology (Clifton, N.J.)* 2060, 395–407.
- Cooper RS, Georgieva ER, Borbat PP, Freed JH and Heldwein EE (2018) Structural basis for membrane anchoring and fusion regulation of the herpes simplex virus fusogen gB. *Nature Structural & Molecular Biology* 25, 416–424.
- Cross TA, Sharma M, Yi M and Zhou H-X (2011) Influence of solubilizing environments on membrane protein structures. *Trends in Biochemical Sciences* 36, 117–125.
- Dang S, Feng S, Tien J, Peters CJ, Bulkley D, Lolicato M, Zhao J, Zuberbühler K, Ye W, Qi L, Chen T, Craik CS, Jan YN, Minor DL, Cheng Y and Jan LY (2017) Cryo-EM structures of the TMEM16A calcium-activated chloride channel. *Nature* 552, 426–429.
- Deisenhofer J, Epp O, Miki K, Huber R and Michel H (1985) Structure of the protein subunits in the photosynthetic reaction centre of *Rhodopseudomonas viridis* at 3 Å resolution. *Nature* **318**, 618–624.
- Della Pia EA, Hansen RW, Zoonens M and Martinez KL (2014*a*) Functionalized amphipols: a versatile toolbox suitable for applications of membrane proteins in synthetic biology. *The Journal of Membrane Biology* 247, 815–826.
- Della Pia EA, Holm JV, Lloret N, Le Bon C, Popot J-L, Zoonens M, Nygård J and Martinez KL (2014b) A step closer to membrane protein multiplexed nanoarrays using biotin-doped polypyrrole. ACS Nano 8, 1844–1853.
- Deme JC, Johnson S, Vickery O, Aron A, Monkhouse H, Griffiths T, James RH, Berks BC, Coulton JW, Stansfeld PJ and Lea SM (2020) Structures of the stator complex that drives rotation of the bacterial flagellum. *Nature Microbiology* 5, 1553–1564.
- Deng Z, Maksaev G, Schlegel AM, Zhang J, Rau M, Fitzpatrick JAJ, Haswell ES and Yuan P (2020) Structural mechanism for gating of a eukaryotic mechanosensitive channel of small conductance. *Nature Communications* 11, 3690.
- Denisov IG and Sligar SG (2017) Nanodiscs in membrane biochemistry and biophysics. *Chemical Reviews* 117, 4669–4713.
- **Denisov IG, Grinkova YV, Lazarides AA and Sligar SG** (2004) Directed selfassembly of monodisperse phospholipid bilayer nanodiscs with controlled size. *Journal of the American Chemical Society* **126**, 3477–3487.
- Denisov IG, Schuler MA and Sligar SG (2019) Nanodiscs as a new tool to examine lipid-protein interactions. *Methods in Molecular Biology* **2003**, 645–671.
- De Zorzi R, Mi W, Liao M and Walz T (2016) Single-particle electron microscopy in the study of membrane protein structure. *Microscopy* 65, 81–96.

- Diederichs KA, Ni X, Rollauer SE, Botos I, Tan X, King MS, Kunji ERS, Jiang J and Buchanan SK (2020) Structural insight into mitochondrial  $\beta$ -barrel outer membrane protein biogenesis. *Nature Communications* 11, 3290.
- Dilworth MV, Piel MS, Bettaney KE, Ma P, Luo J, Sharples D, Poyner DR, Gross SR, Moncoq K, Henderson PJF, Miroux B and Bill RM (2018) Microbial expression systems for membrane proteins. *Methods* 147, 3–39.
- D'Imprima E, Floris D, Joppe M, Sánchez R, Grininger M and Kühlbrandt W (2019) Protein denaturation at the air-water interface and how to prevent it. *eLife* **8**, e42747.
- Dobro MJ, Melanson LA, Jensen GJ and McDowall AW (2010) Plunge freezing for electron cryomicroscopy. *Methods in Enzymology* **481**, 63–82.
- Draper-Joyce CJ, Khoshouei M, Thal DM, Liang Y-L, Nguyen ATN, Furness SGB, Venugopal H, Baltos J-A, Plitzko JM, Danev R, Baumeister W, May LT, Wootten D, Sexton PM, Glukhova A and Christopoulos A (2018) Structure of the adenosine-bound human adenosine A 1 receptor-G i complex. *Nature* 558, 559–563.
- Drulyte I, Johnson RM, Hesketh EL, Hurdiss DL, Scarff CA, Porav SA, Ranson NA, Muench SP and Thompson RF (2018) Approaches to altering particle distributions in cryo-electron microscopy sample preparation. Acta Crystallographica. Section D, *Structural Biology* 74, 560–571.
- Falzone ME, Rheinberger J, Lee B-C, Peyear T, Sasset L, Raczkowski AM, Eng ET, Di Lorenzo A, Andersen OS, Nimigean CM and Accardi A (2019) Structural basis of Ca<sup>2+</sup>-dependent activation and lipid transport by a TMEM16 scramblase. *eLife* **8**, e43229.
- Fan C, Sukomon N, Flood E, Rheinberger J, Allen TW and Nimigean CM (2020) Ball-and-chain inactivation in a calcium-gated potassium channel. *Nature* 580, 288–293.
- Feng X, Fu Z, Kaledhonkar S, Jia Y, Shah B, Jin A, Liu Z, Sun M, Chen B, Grassucci RA, Ren Y, Jiang H, Frank J and Lin Q (2017) A fast and effective microfluidic spraying-plunging method for high-resolution singleparticle cryo-EM. Structure 25, 663–670.e3.
- Feng S, Dang S, Han TW, Ye W, Jin P, Cheng T, Li J, Jan YN, Jan LY and Cheng Y (2019) Cryo-EM studies of TMEM16F calcium-activated Ion channel suggest features important for lipid scrambling. *Cell Reports* 28, 567–579.e4.
- Flores JA, Haddad BG, Dolan KA, Myers JB, Yoshioka CC, Copperman J, Zuckerman DM and Reichow SL (2020) Connexin-46/50 in a dynamic lipid environment resolved by CryoEM at 1.9Å. Nature Communications 11, 4331.
- Flötenmeyer M, Weiss H, Tribet C, Popot J-L and Leonard K (2007) The use of amphipathic polymers for cryo-electron microscopy of NADH:ubiquinone oxidoreductase (Complex I). *Journal of Microscopy* 227, 229–235.
- Frauenfeld J, Gumbart J, van der Sluis EO, Funes S, Gartmann M, Beatrix B, Mielke T, Berninghausen O, Becker T, Schulten K and Beckmann R (2011) Cryo-EM structure of the ribosome-SecYE complex in the membrane environment. *Nature Structural & Molecular Biology* 18, 614–621.
- Frauenfeld J, Löving R, Armache J-P, Sonnen AF-P, Guettou F, Moberg P, Zhu L, Jegerschöld C, Flayhan A, Briggs JAG, Garoff H, Löw C, Cheng Y and Nordlund P (2016) A saposin-lipoprotein nanoparticle system for membrane proteins. *Nature Methods* 13, 345–351.
- Fukunaga K, Szabo L, Fales HM and Pitha J (1988) 2-Hydroxypropyldigitonin: synthesis and properties of preparations differing in degree of substitution. *Journal of Pharmaceutical Sciences* 77, 640–642.
- Gao Y, Hu H, Ramachandran S, Erickson JW, Cerione RA and Skiniotis G (2019) Structures of the rhodopsin-transducin complex: insights into G-protein activation. *Molecular Cell* **75**, 781–790.e3.
- Gao S, Valinsky WC, On NC, Houlihan PR, Qu Q, Liu L, Pan X, Clapham DE and Yan N (2020a) Employing NaChBac for cryo-EM analysis of toxin action on voltage-gated Na<sup>+</sup> channels in nanodisc. Proceedings of the National Academy of Sciences of the USA 117, 14187–14193.
- Gao Y, Eskici G, Ramachandran S, Poitevin F, Seven AB, Panova O, Skiniotis G and Cerione RA (2020b) Structure of the visual signaling complex between transducin and phosphodiesterase 6. *Molecular Cell* 80, 237–245.e4.
- Garavito RM and Ferguson-Miller S (2001) Detergents as tools in membrane biochemistry. The Journal of Biological Chemistry 276, 32403–32406.

- Gatsogiannis C, Merino F, Prumbaum D, Roderer D, Leidreiter F, Meusch D and Raunser S (2016) Membrane insertion of a Tc toxin in near-atomic detail. *Nature Structural & Molecular Biology* 23, 884–890.
- Gharpure A, Teng J, Zhuang Y, Noviello CM, Walsh RM, Cabuco R, Howard RJ, Zaveri NT, Lindahl E and Hibbs RE (2019) Agonist selectivity and ion permeation in the  $\alpha 3\beta 4$  ganglionic nicotinic receptor. *Neuron* **104**, 501–511.e6.
- Giusti F, Popot J-L and Tribet C (2012) Well-defined critical association concentration and rapid adsorption at the air/water interface of a short amphiphilic polymer, amphipol A8-35: a study by Förster resonance energy transfer and dynamic surface tension measurements. *Langmuir* 28, 10372–10380.
- Giusti F, Kessler P, Hansen RW, Della Pia EA, Le Bon C, Mourier G, Popot J-L, Martinez KL and Zoonens M (2015) Synthesis of a polyhistidinebearing amphipol and its use for immobilizing membrane proteins. *Biomacromolecules* 16, 3751–3761.
- Glaeser RM and Han B-G (2017) Opinion: hazards faced by macromolecules when confined to thin aqueous films. *Biophysics Reports* 3, 1–7.
- Glaeser RM, Han B-G, Csencsits R, Killilea A, Pulk A and Cate JHD (2016) Factors that influence the formation and stability of thin, cryo-EM specimens. *Biophysical Journal* 110, 749–755.
- Glavier M, Puvanendran D, Salvador D, Decossas M, Phan G, Garnier C, Frezza E, Cece Q, Schoehn G, Picard M, Taveau J-C, Daury L, Broutin I and Lambert O (2020) Antibiotic export by MexB multidrug efflux transporter is allosterically controlled by a MexA-OprM chaperone-like complex. *Nature Communications* 11, 4948.
- Gong D, Chi X, Wei J, Zhou G, Huang G, Zhang L, Wang R, Lei J, Chen SRW and Yan N (2019) Modulation of cardiac ryanodine receptor 2 by calmodulin. *Nature* 572, 347–351.
- Gopalasingam CC, Johnson RM, Chiduza GN, Tosha T, Yamamoto M, Shiro Y, Antonyuk SV, Muench SP and Hasnain SS (2019) Dimeric structures of quinol-dependent nitric oxide reductases (qNORs) revealed by cryo-electron microscopy. *Science Advances* 5, eaax1803.
- Grinkova YV, Denisov IG and Sligar SG (2010) Engineering extended membrane scaffold proteins for self-assembly of soluble nanoscale lipid bilayers. *Protein Engineering, Design & Selection* 23, 843–848.
- Gu J, Wu M, Guo R, Yan K, Lei J, Gao N and Yang M (2016) The architecture of the mammalian respirasome. *Nature* 537, 639–643.
- Guan C, Niu Y, Chen S-C, Kang Y, Wu J-X, Nishi K, Chang CCY, Chang T-Y, Luo T and Chen L (2020) Structural insights into the inhibition mechanism of human sterol O-acyltransferase 1 by a competitive inhibitor. *Nature Communications* 11, 2478.
- Guo YR and MacKinnon R (2017) Structure-based membrane dome mechanism for piezo mechanosensitivity. *eLife* 6, e33660.
- Hauer F, Gerle C, Fischer N, Oshima A, Shinzawa-Itoh K, Shimada S, Yokoyama K, Fujiyoshi Y and Stark H (2015) GraDeR: membrane protein complex preparation for single-particle cryo-EM. *Structure* 23, 1769–1775.
- Hays FA, Roe-Zurz Z and Stroud RM (2010) Overexpression and purification of integral membrane proteins in yeast. *Methods in Enzymology* 470, 695–707.
- Henderson R (1995) The potential and limitations of neutrons, electrons and X-rays for atomic resolution microscopy of unstained biological molecules. *Quarterly Reviews of Biophysics* 28, 171–193.
- Henderson R and Unwin PNT (1975) Three-dimensional model of purple membrane obtained by electron microscopy. *Nature* 257, 28–32.
- Henderson R, Baldwin JM, Ceska TA, Zemlin F, Beckmann E and Downing KH (1990) Model for the structure of bacteriorhodopsin based on high-resolution electron cryo-microscopy. *Journal of Molecular Biology* **213**, 899–929.
- Higgins AJ, Flynn AJ, Marconnet A, Musgrove LJ, Postis VLG, Lippiat JD, Chung C, Ceska T, Zoonens M, Sobott F and Muench SP Cycloalkane-modified amphiphilic polymers provide direct extraction of membrane proteins for CryoEM analysis. Submitted for publication. doi:10.21203/rs.3.rs-131488/v1
- Hovers J, Potschies M, Polidori A, Pucci B, Raynal S, Bonneté F, Serrano-Vega M, Tate C, Picot D, Pierre Y, Popot J-L, Nehmé R, Bidet M, Mus-Veteau I, Bußkamp H, Jung K-H, Marx A, Timmins PA and Welte W (2011) A class of mild surfactants that keep integral membrane proteins water-soluble for functional studies and crystallization. *Molecular Membrane Biology* 28, 171–181. https://blanco.biomol.uci.edu/mpstruc/.

- Hua T, Li X, Wu L, Iliopoulos-Tsoutsouvas C, Wang Y, Wu M, Shen L, Johnston CA, Nikas SP, Song F, Song X, Yuan S, Sun Q, Wu Y, Jiang S, Grim TW, Benchama O, Stahl EL, Zvonok N, Zhao S, Bohn LM, Makriyannis A and Liu Z-J (2020) Activation and signaling mechanism revealed by cannabinoid receptor-Gi complex structures. *Cell* 180, 655–665.e18.
- Huang W, Masureel M, Qu Q, Janetzko J, Inoue A, Kato HE, Robertson MJ, Nguyen KC, Glenn JS, Skiniotis G and Kobilka BK (2020*a*) Structure of the neurotensin receptor 1 in complex with  $\beta$ -arrestin 1. *Nature* 579, 303–308.
- Huang Y, Wang X, Lv G, Razavi AM, Huysmans GHM, Weinstein H, Bracken C, Eliezer D and Boudker O (2020b) Use of paramagnetic 19F NMR to monitor domain movement in a glutamate transporter homolog. *Nature Chemical Biology* 16, 1006–1012.
- Ishchenko A, Abola EE and Cherezov V (2017) Crystallization of membrane proteins: an overview. *Methods in Molecular Biology* **1607**, 117–141.
- Itskanov S and Park E (2019) Structure of the posttranslational Sec proteintranslocation channel complex from yeast. *Science* **363**, 84–87.
- Jin Q, Zhang B, Zheng X, Li N, Xu L, Xie Y, Song F, Bhat EA, Chen Y, Gao N, Guo J, Zhang X and Ye S (2020) Cryo-EM structures of human pannexin 1 channel. *Cell Research* 30, 449–451.
- Johnson ZL and Chen J (2018) ATP binding enables substrate release from multidrug resistance protein 1. *Cell* 172, 81–89.e10.
- Johnson S, Fong YH, Deme JC, Furlong EJ, Kuhlen L and Lea SM (2020) Symmetry mismatch in the MS-ring of the bacterial flagellar rotor explains the structural coordination of secretion and rotation. *Nature Microbiology* 5, 966–975.
- Jojoa-Cruz S, Saotome K, Murthy SE, Tsui CCA, Sansom MS, Patapoutian A and Ward AB (2018) Cryo-EM structure of the mechanically activated ion channel OSCA1.2. *eLife* 7, e41845.
- Kalienkova V, Clerico Mosina V, Bryner L, Oostergetel GT, Dutzler R and Paulino C (2019) Stepwise activation mechanism of the scramblase nhTMEM16 revealed by cryo-EM. *eLife* 8, e44364.
- Kampjut D and Sazanov LA (2019) Structure and mechanism of mitochondrial proton-translocating transhydrogenase. *Nature* 573, 291–295.
- Kampjut D and Sazanov LA (2020) The coupling mechanism of mammalian respiratory complex I. *Science* **370**, eabc4209.
- Kater L, Frieg B, Berninghausen O, Gohlke H, Beckmann R and Kedrov A (2019) Partially inserted nascent chain unzips the lateral gate of the Sec translocon. *EMBO Reports* 20, e48191.
- Khelashvili G, Falzone ME, Cheng X, Lee B-C, Accardi A and Weinstein H (2019) Dynamic modulation of the lipid translocation groove generates a conductive ion channel in Ca<sup>2+</sup>-bound nhTMEM16. *Nature Communications* **10**, 1–15.
- Kim JJ, Gharpure A, Teng J, Zhuang Y, Howard RJ, Zhu S, Noviello CM, Walsh RM, Lindahl E and Hibbs RE (2020a) Shared structural mechanisms of general anaesthetics and benzodiazepines. *Nature* 585, 303–308.
- Kim Y, Jeong E, Jeong J-H, Kim Y and Cho Y (2020b) Structural basis for activation of the heterodimeric GABAB receptor. *Journal of Molecular Biology* 432, 5966–5984.
- Kishikawa J-I, Nakanishi A, Furuta A, Kato T, Namba K, Tamakoshi M, Mitsuoka K and Yokoyama K (2020) Mechanical inhibition of isolated V<sub>o</sub> from V/A-ATPase for proton conductance. *eLife* **9**, e56862.
- Knowles TJ, Finka R, Smith C, Lin Y-P, Dafforn T and Overduin M (2009) Membrane proteins solubilized intact in lipid containing nanoparticles bounded by styrene maleic acid copolymer. *Journal of the American Chemical Society* 131, 7484–7485.
- Koehl A, Hu H, Feng D, Sun B, Zhang Y, Robertson MJ, Chu M, Kobilka TS, Laeremans T, Steyaert J, Tarrasch J, Dutta S, Fonseca R, Weis WI, Mathiesen JM, Skiniotis G and Kobilka BK (2019) Structural insights into the activation of metabotropic glutamate receptors. *Nature* 566, 79–84.
- Kühlbrandt W (2013) Introduction to electron crystallography. Methods in Molecular Biology 955, 1–16.
- Kuhlen L, Johnson S, Zeitler A, Bäurle S, Deme JC, Caesar JJE, Debo R, Fisher J, Wagner S and Lea SM (2020) The substrate specificity switch FlhB assembles onto the export gate to regulate type three secretion. *Nature Communications* 11, 1296.

- Kun E, Kirsten E and Piper WN (1979) Stabilization of mitochondrial functions with digitonin. *Methods in Enzymology* 55, 115–118.
- Lauber F, Deme JC, Lea SM and Berks BC (2018) Type 9 secretion system structures reveal a new protein transport mechanism. *Nature* 564, 77–82.
- Le Bon C, Della Pia EA, Giusti F, Lloret N, Zoonens M, Martinez KL and Popot J-L (2014) Synthesis of an oligonucleotide-derivatized amphipol and its use to trap and immobilize membrane proteins. *Nucleic Acids Research* **42**, e83.
- Le Bon C, Marconnet A, Masscheleyn S, Popot J-L and Zoonens M (2018) Folding and stabilizing membrane proteins in amphipol A8-35. *Methods* **147**, 95–105.
- Letts JA, Fiedorczuk K and Sazanov LA (2016) The architecture of respiratory supercomplexes. *Nature* 537, 644–648.
- Letts JA, Fiedorczuk K, Degliesposti G, Skehel M and Sazanov LA (2019) Structures of respiratory supercomplex I + III<sub>2</sub> reveal functional and conformational crosstalk. *Molecular Cell* 75, 1131–1146.e6.
- Li J, Richards MR, Bagal D, Campuzano IDG, Kitova EN, Xiong ZJ, Privé GG and Klassen JS (2016) Characterizing the size and composition of saposin A lipoprotein picodiscs. *Analytical Chemistry* 88, 9524–9531.
- Liang Y-L, Khoshouei M, Radjainia M, Zhang Y, Glukhova A, Tarrasch J, Thal DM, Furness SGB, Christopoulos G, Coudrat T, Danev R, Baumeister W, Miller LJ, Christopoulos A, Kobilka BK, Wootten D, Skiniotis G and Sexton PM (2017) Phase-plate cryo-EM structure of a class B GPCR-G-protein complex. *Nature* 546, 118–123.
- Liao M, Cao E, Julius D and Cheng Y (2013) Structure of the TRPV1 ion channel determined by electron cryo-microscopy. *Nature* 504, 107–112.
- Linsky TW, Vergara R, Codina N, Nelson JW, Walker MJ, Su W, Barnes CO, Hsiang T-Y, Esser-Nobis K, Yu K, Reneer ZB, Hou YJ, Priya T, Mitsumoto M, Pong A, Lau UY, Mason ML, Chen J, Chen A, Berrocal T, Peng H, Clairmont NS, Castellanos J, Lin Y-R, Josephson-Day A, Baric RS, Fuller DH, Walkey CD, Ross TM, Swanson R, Bjorkman PJ, Gale M, Blancas-Mejia LM, Yen H-L and Silva D-A (2020) De novo design of potent and resilient hACE2 decoys to neutralize SARS-CoV-2. Science 370, 1208-1214.
- Liu F, Zhang Z, Csanády L, Gadsby DC and Chen J (2017) Molecular structure of the human CFTR ion channel. *Cell* 169, 85–95.e8.
- Liu F, Zhang Z, Levit A, Levring J, Touhara KK, Shoichet BK and Chen J (2019*a*) Structural identification of a hotspot on CFTR for potentiation. *Science* **364**, 1184–1188.
- Liu S, Chang S, Han B, Xu L, Zhang M, Zhao C, Yang W, Wang F, Li J, Delpire E, Ye S, Bai X-C and Guo J (2019b) Cryo-EM structures of the human cation-chloride cotransporter KCC1. *Science* 366, 505–508.
- Liu J, Wan F, Jin Q, Li X, Bhat EA, Guo J, Lei M, Guan F, Wu J and Ye S (2020) Cryo-EM structures of human calcium homeostasis modulator 5. *Cell Discovery* **6**, 1–4.
- Llaguno MC, Xu H, Shi L, Huang N, Zhang H, Liu Q and Jiang Q-X (2014) Chemically functionalized carbon films for single molecule imaging. *Journal of Structural Biology* 185, 405–417.
- Lopez-Redondo ML, Coudray N, Zhang Z, Alexopoulos J and Stokes DL (2018) Structural basis for the alternating access mechanism of the cation diffusion facilitator YiiP. Proceedings of the National Academy of Sciences of the USA 115, 3042–3047.
- Lu P, Bai X-C, Ma D, Xie T, Yan C, Sun L, Yang G, Zhao Y, Zhou R, Scheres SHW and Shi Y (2014) Three-dimensional structure of human γ-secretase. *Nature* **512**, 166–170.
- Ma S, Shen Q, Zhao L-H, Mao C, Zhou XE, Shen D-D, de Waal PW, Bi P, Li C, Jiang Y, Wang M-W, Sexton PM, Wootten D, Melcher K, Zhang Y and Xu HE (2020) Molecular basis for hormone recognition and activation of corticotropin-releasing factor receptors. *Molecular Cell* 77, 669–680.e4.
- Maeda S, Yamamoto H, Kinch LN, Garza CM, Takahashi S, Otomo C, Grishin NV, Forli S, Mizushima N and Otomo T (2020) Structure, lipid scrambling activity and role in autophagosome formation of ATG9A. *Nature Structural & Molecular Biology* 27, 1194–1201.
- Magnani F, Shibata Y, Serrano-Vega MJ and Tate CG (2008) Co-evolving stability and conformational homogeneity of the human adenosine A2a receptor. *Proceedings of the National Academy of Sciences of the USA* **105**, 10744–10749.

- Maldonado M, Padavannil A, Zhou L, Guo F and Letts JA (2020) Atomic structure of a mitochondrial complex I intermediate from vascular plants. *eLife* **9**, e56664.
- Marconnet A, Michon B, Le Bon C, Giusti F, Tribet C and Zoonens M (2020) Solubilization and stabilization of membrane proteins by cycloalkane-modified amphiphilic polymers. *Biomacromolecules* 21, 3459– 3467.
- Martin-Garcia JM, Conrad CE, Coe J, Roy-Chowdhury S and Fromme P (2016) Serial femtosecond crystallography: a revolution in structural biology. Archives of Biochemistry and Biophysics 602, 32–47.
- Martin TG, Bharat TAM, Joerger AC, Bai X-C, Praetorius F, Fersht AR, Dietz H and Scheres SHW (2016) Design of a molecular support for cryo-EM structure determination. *Proceedings of the National Academy of Sciences of the USA* 113, e7456–e7463.
- McCullough J, Clippinger AK, Talledge N, Skowyra ML, Saunders MG, Naismith TV, Colf LA, Afonine P, Arthur C, Sundquist WI, Hanson PI and Frost A (2015) Structure and membrane remodeling activity of ESCRT-III helical polymers. *Science* 350, 1548–1551.
- McDowell MA, Heimes M, Fiorentino F, Mehmood S, Farkas Á, Coy-Vergara J, Wu D, Bolla JR, Schmid V, Heinze R, Wild K, Flemming D, Pfeffer S, Schwappach B, Robinson CV and Sinning I (2020) Structural basis of tail-anchored membrane protein biogenesis by the GET insertase complex. *Molecular Cell* 80, 72–86.e7.
- McGoldrick LL, Singh AK, Saotome K, Yelshanskaya MV, Twomey EC, Grassucci RA and Sobolevsky AI (2018) Opening of the human epithelial calcium channel TRPV6. *Nature* 553, 233–237.
- McGoldrick LL, Singh AK, Demirkhanyan L, Lin T-Y, Casner RG, Zakharian E and Sobolevsky AI (2019) Structure of the thermo-sensitive TRP channel TRP1 from the alga *Chlamydomonas reinhardtii*. Nature Communications 10, 4180.
- Meyerson JR, Rao P, Kumar J, Chittori S, Banerjee S, Pierson J, Mayer ML and Subramaniam S (2014) Self-assembled monolayers improve protein distribution on holey carbon cryo-EM supports. *Scientific Reports* 4, 7084.
- Mio K and Sato C (2018) Lipid environment of membrane proteins in cryo-EM based structural analysis. *Biophysical Reviews* 10, 307–316.
- Nagy JK, Kuhn Hoffmann A, Keyes MH, Gray DN, Oxenoid K and Sanders CR (2001) Use of amphipathic polymers to deliver a membrane protein to lipid bilayers. *FEBS Letters* 501, 115–120.
- Nakane T, Kotecha A, Sente A, McMullan G, Masiulis S, Brown PMGE, Grigoras IT, Malinauskaite L, Malinauskas T, Miehling J, Uchański T, Yu L, Karia D, Pechnikova EV, de Jong E, Keizer J, Bischoff M, McCormack J, Tiemeijer P, Hardwick SW, Chirgadze DY, Murshudov G, Aricescu AR and Scheres SHW (2020) Single-particle cryo-EM at atomic resolution. *Nature* 587, 152–156.
- Nannenga BL and Gonen T (2019) The cryo-EM method microcrystal electron diffraction (MicroED). Nature Methods 16, 369–379.
- Nasr ML and Wagner G (2018) Covalently circularized nanodiscs; challenges and applications. *Current Opinion in Structural Biology* **51**, 129–134.
- Nasr ML, Baptista D, Strauss M, Sun ZJ, Grigoriu S, Huser S, Plückthun A, Hagn F, Walz T, Hogle JM and Wagner G (2017) Covalently circularized nanodiscs for studying membrane proteins and viral entry. *Nature Methods* 14, 49–52.
- Nguyen C and Gonen T (2020) Beyond protein structure determination with MicroED. *Current Opinion in Structural Biology* 64, 51–58.
- Nguyen NX, Armache J-P, Lee C, Yang Y, Zeng W, Mootha VK, Cheng Y, Bai X-C and Jiang Y (2018) Cryo-EM structure of a fungal mitochondrial calcium uniporter. *Nature* 559, 570–574.
- Nguyen AH, Thomsen ARB, Cahill TJ, Huang R, Huang L-Y, Marcink T, Clarke OB, Heissel S, Masoudi A, Ben-Hail D, Samaan F, Dandey VP, Tan YZ, Hong C, Mahoney JP, Triest S, Little J, Chen X, Sunahara R, Steyaert J, Molina H, Yu Z, des Georges A and Lefkowitz RJ (2019) Structure of an endosomal signaling GPCR-G protein-β-arrestin megacomplex. Nature Structural & Molecular Biology 26, 1123–1131.
- Niu Y, Tao X, Touhara KK and MacKinnon R (2020) Cryo-EM analysis of PIP<sub>2</sub> regulation in mammalian GIRK channels. *eLife* **9**, e60552.
- Noble AJ, Dandey VP, Wei H, Brasch J, Chase J, Acharya P, Tan YZ, Zhang Z, Kim LY, Scapin G, Rapp M, Eng ET, Rice WJ, Cheng A, Negro CJ,

Shapiro L, Kwong PD, Jeruzalmi D, des Georges A, Potter CS and Carragher B (2018) Routine single particle CryoEM sample and grid characterization by tomography. *eLife* 7, e34257.

- Nogales E and Scheres SHW (2015) Cryo-EM: a unique tool for the visualization of macromolecular complexity. *Molecular Cell* 58, 677–689.
- Nwanochie E and Uversky VN (2019) Structure determination by singleparticle cryo-electron microscopy: only the sky (and intrinsic disorder) is the limit. *International Journal of Molecular Sciences* **20**, 4186.
- Nygaard R, Kim J and Mancia F (2020) Cryo-electron microscopy analysis of small membrane proteins. *Current Opinion in Structural Biology* 64, 26–33.
- Orwick-Rydmark M, Arnold T and Linke D (2016) The use of detergents to purify membrane proteins. *Current Protocols in Protein Science* 84, 4.8.1– 4.8.35.
- Otzen DE (2015) Proteins in a brave new surfactant world. *Current Opinion in Colloid & Interface Science* **20**, 161–169.
- Owji AP, Zhao Q, Ji C, Kittredge A, Hopiavuori A, Fu Z, Ward N, Clarke OB, Shen Y, Zhang Y, Hendrickson WA and Yang T (2020) Structural and functional characterization of the bestrophin-2 anion channel. *Nature Structural & Molecular Biology* **27**, 382–391.
- Parey K, Haapanen O, Sharma V, Köfeler H, Züllig T, Prinz S, Siegmund K, Wittig I, Mills DJ, Vonck J, Kühlbrandt W and Zickermann V (2019) High-resolution cryo-EM structures of respiratory complex I: mechanism, assembly, and disease. *Science Advances* 5, eaax9484.
- Pérez-Pérez R, Lobo-Jarne T, Milenkovic D, Mourier A, Bratic A, García-Bartolomé A, Fernández-Vizarra E, Cadenas S, Delmiro A, García-Consuegra I, Arenas J, Martín MA, Larsson N-G and Ugalde C (2016) COX7A2L is a mitochondrial complex III binding protein that stabilizes the III<sub>2</sub> + IV supercomplex without affecting respirasome formation. *Cell Reports* 16, 2387–2398.
- Perlmutter JD, Popot J-L and Sachs JN (2014) Molecular dynamics simulations of a membrane protein/amphipol complex. *The Journal of Membrane Biology* 247, 883–895.
- Perry TN, Souabni H, Rapisarda C, Fronzes R, Giusti F, Popot J-L, Zoonens M and Gubellini F (2019) BAmSA: visualising transmembrane regions in protein complexes using biotinylated amphipols and electron microscopy. *Biochimica Et Biophysica Acta. Biomembranes* 1861, 466–477.
- Picard M, Dahmane T, Garrigos M, Gauron C, Giusti F, le Maire M, Popot J-L and Champeil P (2006) Protective and inhibitory effects of various types of amphipols on the Ca<sup>2+</sup>-ATPase from sarcoplasmic reticulum: a comparative study. *Biochemistry* 45, 1861–1869.
- Pinke G, Zhou L and Sazanov LA (2020) Cryo-EM structure of the entire mammalian F-type ATP synthase. *Nature Structural & Molecular Biology* 27, 1077–1085.
- Piton J, Pojer F, Wakatsuki S, Gati C and Cole ST (2020) High resolution CryoEM structure of the ring-shaped virulence factor EspB from Mycobacterium tuberculosis. Journal of Structural Biology X 4, 100029.
- Pleiner T, Tomaleri GP, Januszyk K, Inglis AJ, Hazu M and Voorhees RM (2020) Structural basis for membrane insertion by the human ER membrane protein complex. *Science* 369, 433–436.
- Pocanschi CL, Popot J-L and Kleinschmidt JH (2013) Folding and stability of outer membrane protein A (OmpA) from *Escherichia coli* in an amphipathic polymer, amphipol A8-35. *European biophysics journal* 42, 103–118.
- **Popot J-L** (2018) Membrane Proteins in Aqueous Solutions: From Detergents to Amphipols. New York: Springer.
- Popot J-L, Althoff T, Bagnard D, Banères J-L, Bazzacco P, Billon-Denis E, Catoire LJ, Champeil P, Charvolin D, Cocco MJ, Crémel G, Dahmane T, de la Maza LM, Ebel C, Gabel F, Giusti F, Gohon Y, Goormaghtigh E, Guittet E, Kleinschmidt JH, Kühlbrandt W, Le Bon C, Martinez KL, Picard M, Pucci B, Sachs JN, Tribet C, van Heijenoort C, Wien F, Zito F and Zoonens M (2011) Amphipols from A to Z. Annual Review of Biophysics 40, 379-408.
- Qian H, Wu X, Du X, Yao X, Zhao X, Lee J, Yang H and Yan N (2020) Structural basis of low-pH-dependent lysosomal cholesterol egress by NPC1 and NPC2. *Cell* **182**, 98–111.e18.
- Qiao A, Han S, Li X, Li Z, Zhao P, Dai A, Chang R, Tai L, Tan Q, Chu X, Ma L, Thorsen TS, Reedtz-Runge S, Yang D, Wang M-W, Sexton PM, Wootten D, Sun F, Zhao Q and Wu B (2020) Structural basis of Gs and Gi recognition by the human glucagon receptor. *Science* **367**, 1346–1352.

- Qiu W, Fu Z, Xu GG, Grassucci RA, Zhang Y, Frank J, Hendrickson WA and Guo Y (2018) Structure and activity of lipid bilayer within a membrane-protein transporter. *Proceedings of the National Academy of Sciences of the USA* 115, 12985–12990.
- Rahman MM, Teng J, Worrell BT, Noviello CM, Lee M, Karlin A, Stowell MHB and Hibbs RE (2020) Structure of the native muscle-type nicotinic receptor and inhibition by snake venom toxins. *Neuron* 106, 952–962.e5.
- Ravelli RBG, Nijpels FJT, Henderikx RJM, Weissenberger G, Thewessem S, Gijsbers A, Beulen BWAMM, López-Iglesias C and Peters PJ (2020) Cryo-EM structures from sub-nl volumes using pin-printing and jet vitrification. Nature Communications 11, 2563.
- Razinkov I, Dandey V, Wei H, Zhang Z, Melnekoff D, Rice WJ, Wigge C, Potter CS and Carragher B (2016) A new method for vitrifying samples for cryoEM. Journal of Structural Biology 195, 190–198.
- Reddy B, Bavi N, Lu A, Park Y and Perozo E (2019) Molecular basis of force-from-lipids gating in the mechanosensitive channel MscS. *eLife* 8, e50486.
- Righetto RD, Biyani N, Kowal J, Chami M and Stahlberg H (2019) Retrieving high-resolution information from disordered 2D crystals by single-particle cryo-EM. *Nature Communications* **10**, 1722.
- Roh S-H, Stam NJ, Hryc CF, Couoh-Cardel S, Pintilie G, Chiu W and Wilkens S (2018) The 3.5-Å CryoEM structure of nanodisc-reconstituted yeast vacuolar ATPase Vo proton channel. *Molecular Cell* 69, 993–1004.e3.
- Roh S-H, Shekhar M, Pintilie G, Chipot C, Wilkens S, Singharoy A and Chiu W (2020) Cryo-EM and MD infer water-mediated proton transport and autoinhibition mechanisms of V<sub>o</sub> complex. *Science Advances* 6, eabb9605.
- **Rosevear P, VanAken T, Baxter J and Ferguson-Miller S** (1980) Alkyl glycoside detergents: a simpler synthesis and their effects on kinetic and physical properties of cytochrome c oxidase. *Biochemistry* **19**, 4108–4115.
- Ruan Z, Orozco IJ, Du J and Lü W (2020*a*) Structures of human pannexin 1 reveal ion pathways and mechanism of gating. *Nature* 584, 646–651.
- Ruan Z, Osei-Owusu J, Du J, Qiu Z and Lü W (2020b) Structures and pH-sensing mechanism of the proton-activated chloride channel. *Nature* 588, 350–354.
- Russo CJ and Passmore LA (2014) Ultrastable gold substrates for electron cryomicroscopy. Science 346, 1377–1380.
- Sadaf A, Cho KH, Byrne B and Chae PS (2015) Amphipathic agents for membrane protein study. *Methods in Enzymology* 557, 57–94.
- Sauer DB, Trebesch N, Marden JJ, Cocco N, Song J, Koide A, Koide S, Tajkhorshid E and Wang D-N (2020) Structural basis for the reaction cycle of DASS dicarboxylate transporters. *eLife* 9, e61350.
- Schmidli C, Albiez S, Rima L, Righetto R, Mohammed I, Oliva P, Kovacik L, Stahlberg H and Braun T (2019) Microfluidic protein isolation and sample preparation for high-resolution cryo-EM. *Proceedings of the National Academy of Sciences of the USA* 116, 15007–15012.
- Sgro GG and Costa TRD (2018) Cryo-EM grid preparation of membrane protein samples for single particle analysis. *Frontiers in Molecular Biosciences* 5, 74.
- Shen PS, Yang X, DeCaen PG, Liu X, Bulkley D, Clapham DE and Cao E (2016) The structure of the polycystic kidney disease channel PKD2 in lipid nanodiscs. *Cell* 167, 763–773.e11.
- Shen H, Li Z, Jiang Y, Pan X, Wu J, Cristofori-Armstrong B, Smith JJ, Chin YKY, Lei J, Zhou Q, King GF and Yan N (2018) Structural basis for the modulation of voltage-gated sodium channels by animal toxins. *Science* 362, eaau2596.
- Shih AY, Denisov IG, Phillips JC, Sligar SG and Schulten K (2005) Molecular dynamics simulations of discoidal bilayers assembled from truncated human lipoproteins. *Biophysical Journal* 88, 548–556.
- Singh AK, McGoldrick LL and Sobolevsky AI (2018) Structure and gating mechanism of the transient receptor potential channel TRPV3. *Nature Structural & Molecular Biology* 25, 805–813.
- Sligar SG and Denisov IG (2020) Nanodiscs: a toolkit for membrane protein science. Protein Science 30, 297–315.
- Sousa JS, Mills DJ, Vonck J and Kühlbrandt W (2016) Functional asymmetry and electron flow in the bovine respirasome. *eLIFE* 5, e21290.
- Srivastava AP, Luo M, Zhou W, Symersky J, Bai D, Chambers MG, Faraldo-Gómez JD, Liao M and Mueller DM (2018) High-resolution

cryo-EM analysis of the yeast ATP synthase in a lipid membrane. *Science* **360**, eaas9699.

- Stetsenko A and Guskov A (2017) An overview of the top ten detergents used for membrane protein crystallization. *Crystals* 7, 197.
- Stroud Z, Hall SCL and Dafforn TR (2018) Purification of membrane proteins free from conventional detergents: SMA, new polymers, new opportunities and new insights. *Methods* 147, 106–117.
- Sui X, Arlt H, Brock KP, Lai ZW, DiMaio F, Marks DS, Liao M, Farese RV and Walther TC (2018) Cryo-electron microscopy structure of the lipid droplet-formation protein seipin. *The Journal of Cell Biology* 217, 4080– 4091.
- Sun J and MacKinnon R (2020) Structural basis of human KCNQ1 modulation and gating. *Cell* 180, 340–347.e9.
- Sun C, Benlekbir S, Venkatakrishnan P, Wang Y, Hong S, Hosler J, Tajkhorshid E, Rubinstein JL and Gennis RB (2018) Structure of the alternative complex III in a supercomplex with cytochrome oxidase. *Nature* 557, 123–126.
- Sverzhinsky A, Qian S, Yang L, Allaire M, Moraes I, Ma D, Chung JW, Zoonens M, Popot J-L and Coulton JW (2014) Amphipol-trapped ExbB-ExbD membrane protein complex from *Escherichia coli*: a biochemical and structural case study. *The Journal of Membrane Biology* 247, 1005– 1018.
- **Tao X and MacKinnon R** (2019) Molecular structures of the human Slo1 K<sup>+</sup> channel in complex with  $\beta 4$ . *eLife* **8**, e51409.
- Tascón I, Sousa JS, Corey RA, Mills DJ, Griwatz D, Aumüller N, Mikusevic V, Stansfeld PJ, Vonck J and Hänelt I (2020) Structural basis of proton-coupled potassium transport in the KUP family. *Nature Communications* 11, 626.
- Tate CG (2010) Practical considerations of membrane protein instability during purification and crystallisation. Heterologous expression of membrane proteins: methods and protocols. In Mus-Veteau I (ed.), *Methods in Molecular Biology*. Totowa, NJ: Humana Press, pp. 187–203.
- Thompson AA, Liu JJ, Chun E, Wacker D, Wu H, Cherezov V and Stevens RC (2011) GPCR stabilization using the bicelle-like architecture of mixed sterol-detergent micelles. *Methods* 55, 310–317.
- Tonggu L and Wang L (2018) Broken Symmetry in the Human BK Channel, *bioRxiv*.
- Tonggu L and Wang L (2020) Cryo-EM sample preparation method for extremely low concentration liposomes. Ultramicroscopy 208, 112849.
- Tribet C, Audebert R and Popot J-L (1996) Amphipols: polymers that keep membrane proteins soluble in aqueous solutions. *Proceedings of the National Academy of Sciences* 93, 15047–15050.
- Tsai C-J, Marino J, Adaixo R, Pamula F, Muehle J, Maeda S, Flock T, Taylor NM, Mohammed I, Matile H, Dawson RJ, Deupi X, Stahlberg H and Schertler G (2019) Cryo-EM structure of the rhodopsin- $G\alpha i$ - $\beta\gamma$  complex reveals binding of the rhodopsin C-terminal tail to the  $g\beta$  subunit. *eLife* **8**, e46041.
- Tsutsumi N, Mukherjee S, Waghray D, Janda CY, Jude KM, Miao Y, Burg JS, Aduri NG, Kossiakoff AA, Gati C and Garcia KC (2020) Structure of human Frizzled5 by fiducial-assisted cryo-EM supports a heterodimeric mechanism of canonical Wnt signaling. *eLife* 9, e58464.
- Tucker K and Park E (2019) Cryo-EM structure of the mitochondrial protein-import channel TOM complex at near-atomic resolution. *Nature Structural & Molecular Biology* 26, 1158–1166.
- Uchański T, Masiulis S, Fischer B, Kalichuk V, López-Sánchez U, Zarkadas E, Weckener M, Sente A, Ward P, Wohlkönig A, Zögg T, Remaut H, Naismith JH, Nury H, Vranken W, Aricescu AR, Pardon E and Steyaert J (2021) Megabodies expand the nanobody toolkit for protein structure determination by single-particle cryo-EM. Nature Methods 18, 60–68.
- Vinayagam D, Quentin D, Yu-Strzelczyk J, Sitsel O, Merino F, Stabrin M, Hofnagel O, Yu M, Ledeboer MW, Nagel G, Malojcic G and Raunser S (2020) Structural basis of TRPC4 regulation by calmodulin and pharmacological agents. *eLife* 9, e60603.
- Vinothkumar KR and Henderson R (2016) Single particle electron cryomicroscopy: trends, issues and future perspective. *Quarterly Reviews of Biophysics* **49**, e13.

- Walter JD, Sawicka M and Dutzler R (2019) Cryo-EM structures and functional characterization of murine Slc26a9 reveal mechanism of uncoupled chloride transport. *eLife* **8**, e46986.
- Wang X and Boudker O (2020) Large domain movements through the lipid bilayer mediate substrate release and inhibition of glutamate transporters. *eLife* 9, e58417.
- Wang L and Sigworth FJ (2009) Structure of the BK potassium channel in a lipid membrane from electron cryomicroscopy. *Nature* **461**, 292–295.
- Wang L and Sigworth FJ (2010) Liposomes on a streptavidin crystal: a system to study membrane proteins by cryo-EM. *Methods in Enzymology* 481, 147–164.
- Wang K, Preisler SS, Zhang L, Cui Y, Missel JW, Grønberg C, Gotfryd K, Lindahl E, Andersson M, Calloe K, Egea PF, Klaerke DA, Pusch M, Pedersen PA, Zhou ZH and Gourdon P (2019) Structure of the human ClC-1 chloride channel. *PLoS Biology* 17, e3000218.
- Wang L, Johnson ZL, Wasserman MR, Levring J, Chen J and Liu S (2020a) Characterization of the kinetic cycle of an ABC transporter by singlemolecule and cryo-EM analyses. *eLife* 9, e56451.
- Wang Z, Hu W and Zheng H (2020b) Pathogenic siderophore ABC importer YbtPQ adopts a surprising fold of exporter. Science Advances 6, eaay7997.
- Wasilko DJ, Johnson ZL, Ammirati M, Che Y, Griffor MC, Han S and Wu H (2020) Structural basis for chemokine receptor CCR6 activation by the endogenous protein ligand CCL20. *Nature Communications* 11, 3031.
- Weiss HM and Grisshammer R (2002) Purification and characterization of the human adenosine A(2a) receptor functionally expressed in *Escherichia coli. European Journal of Biochemistry* **269**, 82–92.
- Weiss MS, Wacker T, Weckesser J, Weite W and Schulz GE (1990) The three-dimensional structure of porin from *Rhodobacter capsulatus* at 3 Å resolution. *FEBS Letters* **267**, 268–272.
- Willegems K and Efremov RG (2018) Influence of lipid mimetics on gating of ryanodine receptor. *Structure* 26, 1303–1313.e4.
- Wu S, Avila-Sakar A, Kim J, Booth DS, Greenberg CH, Rossi A, Liao M, Li X, Alian A, Griner SL, Juge N, Yu Y, Mergel CM, Chaparro-Riggers J, Strop P, Tampé R, Edwards RH, Stroud RM, Craik CS and Cheng Y (2012) Fabs enable single particle cryoEM studies of small proteins. Structure 20, 582–592.
- Wu M, Gu J, Guo R, Huang Y and Yang M (2016) Structure of mammalian respiratory supercomplex I<sub>1</sub>III<sub>2</sub>IV<sub>1</sub>. *Cell* **167**, 1598–1609.e10.
- Wu X, Cabanos C and Rapoport TA (2019) Structure of the posttranslational protein translocation machinery of the ER membrane. *Nature* 566, 136–139.
- Wu X, Siggel M, Ovchinnikov S, Mi W, Svetlov V, Nudler E, Liao M, Hummer G and Rapoport TA (2020) Structural basis of ER-associated protein degradation mediated by the Hrd1 ubiquitin ligase complex. *Science* 368, eaaz2449.
- Yan R, Zhao X, Lei J and Zhou Q (2019) Structure of the human LAT1-4F2hc heteromeric amino acid transporter complex. *Nature* **568**, 127–130.
- Yang G, Zhou R, Zhou Q, Guo X, Yan C, Ke M, Lei J and Shi Y (2019) Structural basis of notch recognition by human γ-secretase. *Nature* **565**, 192–197.
- Yao X, Fan X and Yan N (2020) Cryo-EM analysis of a membrane protein embedded in the liposome. Proceedings of the National Academy of Sciences 117, 18497–18503.
- Yin W, Li Z, Jin M, Yin Y-L, de Waal PW, Pal K, Yin Y, Gao X, He Y, Gao J, Wang X, Zhang Y, Zhou H, Melcher K, Jiang Y, Cong Y, Edward Zhou X, Yu X and Eric Xu H (2019*a*) A complex structure of arrestin-2 bound to a G protein-coupled receptor. *Cell Research* 29, 971–983.
- Yin Y, Wu M, Hsu AL, Borschel WF, Borgnia MJ, Lander GC and Lee S-Y (2019b) Visualizing structural transitions of ligand-dependent gating of the TRPM2 channel. *Nature Communications* 10, 1–14.
- Yin J, Chen K-YM, Clark MJ, Hijazi M, Kumari P, Bai X-C, Sunahara RK, Barth P and Rosenbaum DM (2020) Structure of a D2 dopamine receptor-G-protein complex in a lipid membrane. *Nature* 584, 125–129.
- Yoder N and Gouaux E (2020) The His-Gly motif of acid-sensing ion channels resides in a reentrant 'loop' implicated in gating and ion selectivity. *eLife* 9, e56527.
- Yoo J, Wu M, Yin Y, Herzik MA, Lander GC and Lee S-Y (2018) Cryo-EM structure of a mitochondrial calcium uniporter. *Science* **361**, 506–511.

- Yu X, Plotnikova O, Bonin PD, Subashi TA, McLellan TJ, Dumlao D, Che Y, Dong YY, Carpenter EP, West GM, Qiu X, Culp JS and Han S (2019) Cryo-EM structures of the human glutamine transporter SLC1A5 (ASCT2) in the outward-facing conformation. *eLife* **8**, e48120.
- Zhang Z, Liu F and Chen J (2017) Conformational changes of CFTR upon phosphorylation and ATP binding. *Cell* **170**, 483–491.e8.
- Zhang C, Konermann S, Brideau NJ, Lotfy P, Wu X, Novick SJ, Strutzenberg T, Griffin PR, Hsu PD and Lyumkis D (2018a) Structural basis for the RNA-guided ribonuclease activity of CRISPR-Cas13d. Cell 175, 212–223.e17.
- Zhang Z, Liu F and Chen J (2018b) Molecular structure of the ATP-bound, phosphorylated human CFTR. Proceedings of the National Academy of Sciences of the USA 115, 12757–12762.
- Zhang L, Zhao Y, Gao Y, Wu L, Gao R, Zhang Q, Wang Y, Wu C, Wu F, Gurcha SS, Veerapen N, Batt SM, Zhao W, Qin L, Yang X, Wang M, Zhu Y, Zhang B, Bi L, Zhang X, Yang H, Guddat LW, Xu W, Wang Q, Li J, Besra GS and Rao Z (2020) Structures of cell wall arabinosyltransferases with the anti-tuberculosis drug ethambutol. *Science* 368, 1211–1219.
- Zhao DY, Pöge M, Morizumi T, Gulati S, Van Eps N, Zhang J, Miszta P, Filipek S, Mahamid J, Plitzko JM, Baumeister W, Ernst OP and Palczewski K (2019a) Cryo-EM structure of the native rhodopsin dimer in nanodiscs. *The Journal of Biological Chemistry* 294, 14215–14230.
- Zhao J, Xu H, Carroni M, Lebrette H, Wallden K, Moe A, Matsuoka R, Högbom M and Zou X (2019b) A simple pressure-assisted method for cryo-EM specimen preparation. *bioRxiv*, 665448.
- Zhao L-H, Ma S, Sutkeviciute I, Shen D-D, Zhou XE, de Waal PW, Li C-Y, Kang Y, Clark LJ, Jean-Alphonse FG, White AD, Yang D, Dai A, Cai X, Chen J, Li C, Jiang Y, Watanabe T, Gardella TJ, Melcher K, Wang M-W, Vilardaga J-P, Xu HE and Zhang Y (2019c) Structure and dynamics of the active human parathyroid hormone receptor-1. Science 364, 148–153.

- Zhao Y, Huang G, Wu J, Wu Q, Gao S, Yan Z, Lei J and Yan N (2019d) Molecular basis for ligand modulation of a mammalian voltage-gated Ca<sup>2</sup> <sup>+</sup> channel. *Cell* **177**, 1495–1506.e12.
- Zhao J, Lin King JV, Paulsen CE, Cheng Y and Julius D (2020) Irritant-evoked activation and calcium modulation of the TRPA1 receptor. *Nature* 585, 141–145.
- Zheng L, Li Y, Li X, Zhong Q, Li N, Zhang K, Zhang Y, Chu H, Ma C, Li G, Zhao J and Gao N (2019) Structural and functional insights into the tetrameric photosystem I from heterocyst-forming cyanobacteria. *Nature Plants* 5, 1087–1097.
- Zhou X, Li M, Su D, Jia Q, Li H, Li X and Yang J (2017) Cryo-EM structures of the human endolysosomal TRPML3 channel in three distinct states. *Nature Structural and Molecular Biology* 24, 1146–1154.
- Zhou R, Yang G, Guo X, Zhou Q, Lei J and Shi Y (2019) Recognition of the amyloid precursor protein by human γ-secretase. *Science* **363**, eaaw0930.
- Zoonens M and Miroux B (2010) Expression of membrane proteins at the *Escherichia coli* membrane for structural studies. *Methods in Molecular Biology* **601**, 49–66.
- Zoonens M and Popot J-L (2014) Amphipols for each season. The Journal of Membrane Biology 247, 759–796.
- Zoonens M, Giusti F, Zito F and Popot J-L (2007) Dynamics of membrane protein/amphipol association studied by Förster resonance energy transfer: implications for in vitro studies of amphipol-stabilized membrane proteins. *Biochemistry* 46, 10392–10404.
- Zubcevic L, Herzik MA, Chung BC, Liu Z, Lander GC and Lee S-Y (2016) Cryo-electron microscopy structure of the TRPV2 ion channel. *Nature Structural & Molecular Biology* 23, 180–186.
- Zubcevic L, Herzik MA, Wu M, Borschel WF, Hirschi M, Song AS, Lander GC and Lee S-Y (2018) Conformational ensemble of the human TRPV3 ion channel. *Nature Communications* **9**, 1–12.
- Zubcevic L, Hsu AL, Borgnia MJ and Lee S-Y (2019) Symmetry transitions during gating of the TRPV2 ion channel in lipid membranes. *eLife* 8, e45779.