

Detergent free purification of ABC transporters

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Abstract:

ABC (ATP Binding Cassette) transporters carry out many vital functions and are involved in numerous diseases, but study of the structure and function of these proteins is often hampered by their large size and membrane location. Membrane protein purification usually utilises detergents to solubilise the protein from the membrane, effectively removing it from its native lipid environment. Subsequently lipids have to be added back and detergent removed to reconstitute the protein into a lipid bilayer. We present here the application of a new methodology for the extraction and purification of ABC transporters without the use of detergent, instead using a styrene maleic acid co-polymer (SMA). SMA inserts in a bilayer and assembles into discrete particles, essentially solubilising the membrane into small discs of bilayer encircled by polymer, termed SMA lipid particles (SMALPs). We show that this polymer can extract several eukaryotic ABC transporters; P-glycoprotein (ABCB1), MRP1 (ABCC1), MRP4 (ABCC4), ABCG2 and CFTR (ABCC7), from a range of different expression systems. The SMALP encapsulated ABC transporters can be purified by affinity chromatography, and are able to bind ligands comparably to those in native membranes or detergent micelles. A greater degree of purity and enhanced stability is seen compared to detergent solubilisation. This study demonstrates that eukaryotic ABC transporters can be extracted and purified without ever being removed from their lipid bilayer environment, opening up a wide range of possibilities for the future study of their structure and function.

Summary statement:

A styrene maleic acid copolymer can be effectively used to extract and purify large eukaryotic transmembrane proteins in the absence of detergents, forming small bilayer discs encapsulating the protein, which have great potential for future structure & function studies.

Short title: Detergent free purification of ABC transporters

Keywords: membrane protein, solubilisation, purification, polymer, nanodisc, detergent

Introduction:

The ABC (ATP Binding Cassette) transporter superfamily is found in all organisms from bacteria to man, with humans possessing 48 different members [1]. The human transporters are involved in a wide range of functions within the body including protection from toxins, metabolism, controlling drug distribution in the body, mediating inflammatory responses and transporting lipids [2-5]. Several members are involved in causing drug resistance during cancer treatment [6], while others are responsible for genetic diseases such as cystic fibrosis and adrenoleukodystrophy [7, 8].

The study of the function and structure of these vital proteins is often hampered by difficulties in expressing, extracting and purifying them, as they are large transmembrane proteins requiring a lipid bilayer environment. Traditionally purification of membrane proteins such as these has used detergent to solubilise the protein from the membrane, thus removing it from its native lipid environment. Following purification lipids need to be added back in and detergent removed to reconstitute the protein into a lipid bilayer [9]. These reconstituted systems include proteoliposomes and planar membranes, and more recently several forms of lipid nano-particles, such as bicelles or nanodiscs. The last of these, nanodiscs use membrane associated amphipathic peptides to produce small discs of membrane bilayer into which membrane proteins can be inserted [10]. Several ABC transporters have been successfully incorporated into nanodiscs, to enable many downstream applications [11-14]. However formation of these particles still requires detergent solubilisation of the protein before reconstitution into the nanodiscs. Detergent solubilisation requires an expensive, trial and error process, and detergents that work well for solubilisation may not be compatible with downstream purification or analytical methods [15, 16].

Here we report the application of a recently developed system for detergent-free extraction of ABC transporters from their native membranes using a readily available, biocompatible and chemically stable polymer, polystyrene-co-maleic acid (SMA) [17-19]. The polymer inserts into the membrane isolating small discs of lipid bilayer that are encircled by the polymer, termed SMA lipid particles (SMALPs) [17]. We demonstrate that the polymer is able to effectively solubilise several different eukaryotic ABC transporters from a range of different cell types, including mammalian cells, insect cells and yeast. ABC transporters within the SMALPs can then be purified using standard affinity chromatography methods. We demonstrate that the proteins within the SMALPs can bind substrates comparably to those in native membranes or detergent micelles, and we characterise the biophysical properties of a SMALP-purified transporter.

Overall this study demonstrates that the SMALP method can be successfully applied to eukaryotic ABC transporters to extract and purify them without ever removing them from their lipid environment.

Experimental:

SMA preparation

SMA polymer was prepared as described previously [17]. Briefly, a 10% (w/v) solution of SMA copolymer in 1M NaOH was refluxed for 2 hours, allowed to cool to room temperature, then dialysed against 20mM Tris buffer pH8 before use.

Cell culture, expression & membrane preparation

High Five (*Trichoplusia ni*) insect cells were cultured and infected with recombinant baculovirus to express either human P-glycoprotein (Pgp/ABCB1) with a C-terminal 12-His tag or human ABCG2 with an N-terminal 6-His-tag, and membrane preparations carried out as described previously [15, 20]. Murine CFTR (cystic fibrosis transmembrane regulator/ABCC7) with two hexa-histidine tags and a GFP tag was expressed in *Sacchomyces cerevisiae* and microsomes prepared as described elsewhere [21]. MRP1 (ABCC1) overexpressing drug selected H69AR small cell lung cancer cell line was cultured as described previously [22]. MRP1 was also expressed in HEK cells by transfection of the pcDNA3.1-MRP1 plasmid [23] using JetPRIME transfection reagent (Polyplus-transfection). Plates (10cm) were seeded with 1.3×10^6 cells transfected after 24 hours with 10 μ g DNA/plate and 20 μ l JetPRIME/plate, and harvested 48 hours later. ABCG2 was expressed in either MCF7-FLV cells cultured as described [24], or following transfection of HEK293T cells with pcDNA3.1_His6_ABCG2 [25]. MRP4 (ABCC4) with a C-terminal hexa-histidine tag was expressed in HEK cells and membranes prepared as described before [26]. MRP4 with a C-terminal hexa-histidine tag was also expressed in *Spodoptera frugiperda* (Sf9) insect cells by infection with a recombinant baculovirus. The cDNA for MRP4 was amplified from the pcDNA3.1-MRP4 plasmid [27], and the stop codon mutated to an AgeI site by PCR (using the following primers 5' ctacgtcgacgacctagatgctgccc and 3' ctaagatttagtcgcgccgctcaatggtgatg). The PCR product was digested with EcoRI and AgeI and subcloned into a pFastBac-1 vector, resulting in the generation of a C-terminally hexahistidine tagged construct. Recombinant baculovirus was formed using the Bac-to-Bac kit (Invitrogen). Sf9 cells were maintained at $0.5\text{--}2.5 \times 10^6$ cells/mL in shaker cultures at 28 °C in Insect Xpress media (Lonza) supplemented with 10% (v/v) fetal bovine serum, 100 U/mL penicillin, and 100 mg/mL streptomycin. MRP4 expressing baculovirus was amplified by infection of Sf9 cells (1×10^6 cells/mL) at a multiplicity of infection (MOI) of 0.2 for 5 days. MRP4 expression in Sf9 cells was achieved by infection with the baculovirus at an MOI of 5 for 3 days. Cells were harvested by centrifugation (6000 g, 20 min, 4°C). Whole cell pellets of MCF7-FLV, HEK293T-ABCG2, H69AR, HEK-MRP1 or Sf9-MRP4 were resuspended in buffer 1 (50 mM Tris pH7.4, 250 mM sucrose) supplemented with 0.25 mM CaCl₂ and protease inhibitors (Roche or Pierce), then lysed by nitrogen cavitation (500 psi, 15 min, 4°C). Cell debris was removed by low speed centrifugation (750 g, 20 min, 4°C), then membranes were harvested by ultracentrifugation (100000 g, 20 min, 4°C). Membranes were resuspended in buffer 1, aliquoted and stored at -70°C for up to 6 months.

SMA extraction

Membranes were pelleted (100000 g, 20 min, 4°C) then resuspended in buffer 2 (20 mM Tris pH8, 150 mM NaCl, 20% (v/v) glycerol) at a final concentration of 30 mg/ml membrane (wet membrane weight) and SMA was then added to a final concentration of 2.5% (w/v). Samples were incubated at room temperature with gentle shaking for 30 - 60 minutes. Insoluble material was removed by centrifugation (100000 g, 20 min, 4°C). For ABCG2 from High Five insect cells only, this method was modified slightly in that 1% (w/v) DMPC was also included alongside the SMA, the sample was briefly probe sonicated and incubated at 37°C.

The insoluble pellet was resuspended in buffer containing 2.5% SDS (w/v) to the same volume as the soluble fraction. Samples of both the soluble and insoluble fractions were analysed by Western blotting. For most of the proteins studied an anti-His primary antibody was used (R & D Systems, typically at 1:500 dilution). For the H69AR samples the primary antibody used was QCRL-1 (1:2000). For the MCF7-FLV and HEK-ABCG2 samples the primary antibody was Bxp-21 (Merck 1:2000). In all cases either an anti mouse-Alkaline Phosphatase (Sigma) or anti mouse-HRP (Sigma) secondary antibody was used, and visualised using BCIP/NBT colorimetric detection (Sigma) or chemiluminescence (Pierce) respectively. The efficiency of solubilisation was quantified from the Western blots using densitometry (Image J).

Ni²⁺-NTA affinity purification

SMA-solubilised membranes were incubated overnight at 4°C with Ni²⁺-NTA agarose (Qiagen) using 100 µl resin/ml solubilised membrane and slowly rotated. The sample was transferred to a gravity flow column (Biorad), washed 5 times with 10 bed volumes (bv) of buffer 2 supplemented with 20 mM imidazole, then twice with 10 bv of buffer 2 supplemented with 40 mM imidazole, then once with 2 bv of buffer 2 supplemented with 60 mM imidazole. Proteins were eluted using buffer 2 supplemented with 120 mM imidazole and 1 bv fractions were collected. Samples of various stages of the purification were analysed by SDS-PAGE and visualised by silver staining (Pierce). The concentration of purified proteins were determined using Coomassie stained SDS-PAGE gels as described previously (Rothnie et al 2004 JBC).

Following purification, samples were concentrated and/or buffer exchanged using centrifugal concentrators (Amicon Ultra, Millipore, 30000 MWCO) or subjected to gel filtration spin columns to remove imidazole (Biorad, Micro Bio-Spin columns).

Protein A and antibody affinity purification

Immunoaffinity purification of MRP1 from H69AR membranes using the monoclonal antibody QCRL-1 was based on the procedure of Mao *et al* [28]. Briefly, QCRL-1 ascites fluid (50 µl) was diluted to 1ml in buffer 2, then slowly injected on to a protein A AcroSep pre-packed 1ml column (PALL). After incubation at room temperature for 30 min the column was washed 10 x 1 ml with buffer 2. SMALP-encapsulated H69AR membranes (1 ml) were then slowly injected on to the column. After 30 mins the column was washed 10 x 1 ml with buffer 2. MRP1 was eluted from the column using buffer 2 supplemented with the peptide SSYSGDI (ThermoFisher Custom Biopolymers) at 3.3 mg/ml, and 1 ml fractions were collected.

Drug binding to Pgp: Fluorescence quenching assay

Formation of the Pgp:drug complex was studied using the fluorescence quenching method of Liu and Sharom [29]. In this method the fluorophore, MIANS, was first used to label cysteine residues within the two nucleotide binding domains of ABC transporters. Purified Pgp (100 µg/mL) encapsulated within SMALPs was incubated with 20 µM MIANS (Invitrogen) at room temperature for 1 h in the dark. DTT (1 mM) was added to quench unreacted fluorophore. Separation of MIANS-labelled Pgp-SMALPs from DTT-MIANS was performed by gel filtration chromatography on Micro Bio-Spin columns (Bio-rad). Fluorescence quenching of MIANS-labelled Pgp-SMALP (50 µg/mL) by successive additions of doxorubicin (2.5-25 µM) or verapamil (0.5-12 µM) was measured using a Perkin Elmer LS55 fluorimeter at an excitation wavelength of 322±10 nm and emission wavelength of 420 nm (λ_{max}). Fluorescence intensities were corrected for dilution, scattering and inner filter effect as described previously [29]. Results were analysed by non-linear regression using Graphpad Prism (Graphpad Software Inc.) to fit a one-site binding hyperbolic equation 1:

$$B = B_{\max} \cdot [L] / (K_d + [L])$$

Drug binding to MRP1: Radioligand binding assay

The binding of 40 nM ³H-estrone sulphate (Perkin Elmer) to SMALP encapsulated membranes from H69AR cells, in the presence or absence of 3 mM GSH was performed as described previously [30]. However instead of using rapid filtration to separate bound and unbound radioligand, samples were instead loaded onto pre-equilibrated Micro Bio-Spin columns (Biorad) and the void volumes collected by centrifugation (3000 rpm, 4 min). Scintillation counting was performed using a Packard Tri-Carb Liquid Scintillation counter.

Statistical analysis used a student t-test (GraphPad Prism), and a P<0.05 was considered significant.

Drug binding to ABCG2: Pheophorbide A fluorescence

Binding of SMALP encapsulated ABCG2 to the fluorescent substrate pheophorbide A was detected by an enhancement of the fluorescence intensity. Pheophorbide A (10 µM, 1 ml) fluorescence emission spectra (640 – 720 nm) were measured at an excitation wavelength of 390±10 nm using a Perkin Elmer LS55 fluorimeter. Successive additions (5 – 100 µl) of SMALP encapsulated membranes from cells overexpressing ABCG2 were added to the pheophorbide A and incubated for 10 minutes before reading the fluorescence spectra again. Specific binding to ABCG2 was determined by subtracting the fluorescence intensity obtained when membranes lacking ABCG2 were added to pheophorbide A, or the ABCG2 membranes were pre-incubated with chrysin (30 µM) for 10 minutes before addition to pheophorbide A. Results were analysed by non-linear regression using Graphpad Prism (Graphpad Software Inc.) to fit equation 1.

Nucleotide binding to Pgp; tryptophan fluorescence quenching

Binding of nucleotides to purified Pgp within SMALPs was assayed by measuring the quenching of intrinsic tryptophan fluorescence as described previously [31]. Briefly, the fluorescence of purified Pgp (50 µg/ml) encapsulated within SMALPs was measured using a Perkin Elmer LS55 fluorimeter at an excitation wavelength of 290±10 nm and emission wavelength of 330 nm (λ_{max}), and the quenching by successive additions of ATP (0.05-2 mM) or the non-hydrolysable fluorescent ATP analog, TNP-ATP (9.8-160 µM) was measured. Fluorescence intensities were corrected for dilution, scattering and inner filter effect as described previously [29]. Results were analysed by non-linear regression using Graphpad Prism (Graphpad Software Inc.) to fit a one-site binding hyperbolic (equation 1).

Circular Dichroism Spectroscopy

Purified Pgp within SMALPs was buffer exchanged into 10 mM Tris pH 8 and concentrated to 0.65 mg/ml. The circular dichroism spectra was measured using a JASCO J-715 CD spectrophotometer with a path length of 0.1 cm at room temperature. The sample was scanned 8 times at 200 nm/min with a bandwidth of 2 nm. A spectrum of the buffer used in the experiment was used as a background.

Analytical Ultracentrifugation

Purified Pgp encapsulated within SMALPs (0.65 mg/ml) in 10 mM Tris pH 8 was subjected to analysis using sedimentation velocity analytical ultracentrifugation. The sample was centrifuged at 40000 rpm/129000g at 4°C in a Ti50 rotor within a Beckman XL-1 analytical ultracentrifuge (Beckman 693 Coulter, Palo Alto, CA, USA). The movement of the protein within the cell was monitored using absorbance at 280 nm with 1780 datasets (890 each for absorbance and velocity)

being collected over 23 hr 22 min. These data were analysed using SEDFIT [32] and the $c(s)$ and $c(M)$ method. Values for V_{bar} and buffer viscosity and density were calculated using SEDNTERP [33].

Gel filtration

Purified Pgp encapsulated within SMALPs was run on a Superdex 200 gel filtration column (diameter 1.6 cm, column height 55 cm) pre-equilibrated in 20 mM Tris pH8 using an AKTA Purifier system. A total of 50 μ g in a volume of 0.5 ml 20 mM Tris pH 8 was loaded and the column was run at a flow rate of 0.5 ml/min. The column was calibrated using the following standards: Blue Dextran, Bovine Serum Albumin, Carbonic Anhydrase, Cytochrome c and Aprotinin (Sigma).

Electron microscopy

Holey electron microscopy grids (2micron holes, Quantifoil) were prepared of purified Pgp encapsulated within SMALPs (1 mg/ml) using an FEI Vitrobot freezing device with plunge freezing into liquid ethane. Grids were transferred into a Polara FEG transmission electron microscope and maintained at <100K throughout. Areas with sufficiently thin ice were selected at low magnification and low dose, and then micrographs were recorded on a 4k x 4k CCD device with 1 sec exposure and low dose mode imaging conditions. Micrographs were analysed using the EMAN2 software package [34], with >8000 particles selected from 20 images. Strong defocus was employed because of the relatively small mass of the particles (defocus range 3.5 to 5.9 micron). Correction for the contrast transfer function was carried out and then reference-free classification of the particles was performed with unique 32 classes identified. Preliminary 3D structures were generated from these classes using a common-lines algorithm and with no symmetry applied. The ten low resolution structures were similar and one was selected for refinement of the 3D structure using all the single particles. The resolution of the structure after 5 rounds of refinement was estimated by Fourier shell correlation between two structures generated from even- and odd-numbered particles from the data set. The features observed in the final structure corresponded to the resolution estimate and with the strong suppression of higher resolution frequencies in the images as a result of the need for strong underfocus.

Thermostability

Pgp was extracted from High Five membranes either using 2.5% (w/v) SMA as described above or 2% (w/v) octyl-glucoside plus 0.4% (w/v) lipids (4:1 *E.coli* lipid extract:cholesterol) as detailed in [20], then purified by Ni-NTA affinity chromatography as described above, except supplementing the buffers with 1.25% (w/v) octyl-glucoside and 0.1% (w/v) lipids. Relative thermostability of the purified proteins was monitored using the thiol-reactive flourophore 7-diethylamino-3-(4-maleimidophenyl)-4-methylcoumarin (CPM, Sigma) as described by Alexandrov *et al* [35]. CPM is essentially non-fluorescent until it reacts with a sulphhydryl, thus it can report on the relative accessibility of cysteine residues within a protein. As a protein denatures, buried cysteine residues become accessible to CPM and the fluorescence signal increases. Purified Pgp (5 μ g) was mixed with CPM (8 μ g/ml) in a total volume of 200 μ l. The reaction mixture was heated continuously from 10°C - 90°C in increments of 5°C for 2 minutes each. After 2 minutes at each temperature the fluorescence spectra was measured (λ_{ex} 463 \pm 10 nm, λ_{em} 410-560 \pm 20 nm) in a Perkin Elmer LS55 spectrophotometer. The fluorescence intensity (at 470 nm) at each time point was calculated as a % of the final fluorescence intensity. The assays were repeated twice for each sample.

Results & Discussion:

Extraction of ABC transporters using SMA

The initial aim was to determine if the SMALP method for extraction of membrane proteins could be successfully applied to eukaryotic ABC transporters. For each protein tested membrane fractions were prepared and mixed with 2.5% (w/v) SMA for 30-60 mins in the absence of any detergent, and insoluble material was removed by centrifugation. It can be seen in Figure 1 that for each of the proteins tested, Pgp, MRP1, MRP4, CFTR and ABCG2, successful extraction was observed using SMA. These samples included all of the commonly used cell types for eukaryotic ABC transporter expression, namely insect cells, mammalian cells and yeast. It also encompassed a range of sizes, with MRP1 at 190 kDa and comprising 17 transmembrane helices, being the largest, and both the conventional and reversed topology ABC transporters (i.e. ABCG2 having an NBD-TMD topology). The degree of extraction varied but there were no discernible patterns between expression systems or protein size.

For Pgp from High Five insect cells the extraction efficiency with SMA (60%) was comparable to previous reports using octyl-glucoside (50%) [20], and the extraction of MRP1 from H69AR cell membranes with SMA was comparable to previous reports using the detergent CHAPS (90%) [28]. MRP4 expressed in HEK cells is 63% solubilised by SMA whereas when expressed in Sf9 insect cells 86% is solubilised. The efficiency of MRP4 solubilisation with detergents has not previously been reported, therefore we also examined the ability of two commonly used detergents, octyl-glucoside and dodecylmaltoide, to solubilise MRP4 from Sf9 cells (Supplementary Figure 1). Octyl glucoside was unable to solubilise MRP4 at all, whereas with DDM 78% of the MRP4 was successfully solubilised, which is comparable with the SMA. SMA solubilisation of ABCG2 from High Five insect cells was the least efficient at just 36%, however it was previously found that this protein is completely insoluble with most detergents and only the long chain foscholine detergents were capable of solubilising this protein at all [15]. One possible reason for the difficulty in solubilising ABCG2 from High Five cells may be the formation of large oligomers in this system [15], which are too large for the lipid discs formed by SMA. However the successful solubilisation of ABCG2 from both HEK cells and MCF7-FLV cells show that it is not a feature of ABCG2 *per se*. Interestingly the two most efficient extractions were from drug selected cancer cell lines (H69AR and MCF7-FLV) which overexpress transporters due to development of drug resistance. It is well known that such drug resistant cell lines display altered membrane constituents and properties [36-38], and it seems that this improves the ability of SMA to solubilise these transporters.

Purification of ABC transporters encapsulated within SMALPs

The SMA extracted membranes were then subjected to affinity purification. The SMALPs once formed are stable and, unlike detergent solubilised proteins, did not require addition of further polymer during the purification procedure. Figure 2 shows silver stained gels for the successful purification of each protein. The Pgp, MRP4, CFTR and ABCG2 proteins were purified using single step Ni²⁺-NTA affinity chromatography, whereas MRP1 (which did not have a His-tag) was purified using a Protein A column and QCRL-1 antibody. An unexpected outcome of using SMA to solubilise the ABC transporters was a much higher degree of purity was obtained following just a single step affinity purification than has previously been observed with detergent solubilised proteins, which tend to require a second purification step of gel filtration [15, 21, 39] or ion exchange [40].

To ascertain whether this higher purity using SMA altered the final purified protein yield we quantified this and present the data in Supplementary Table 1. For purified Pgp obtained using SMA the yield was 631 ± 46 µg per litre of baculovirus infected High Five cell culture which is very

similar to that previously reported using octyl glucoside with these same cell membranes ($792 \pm 197 \mu\text{g}$ [20]), and the SMA purified Pgp is significantly more pure than the octyl glucoside preparation. Other reports which have used DDM to solubilise Pgp from these membranes have used a more stringent purification procedure including gel filtration as well as Ni^{2+} -NTA [41] and thus the Pgp produced was highly pure, however this resulted in a significantly lower yield ($249 \pm 15 \mu\text{g/L}$ culture) than that reported here with SMA. The yield of purified ABCG2 using SMA was $116 \pm 32 \mu\text{g/L}$ culture, which is almost 3-fold higher than that previously obtained using foscholine (Supplementary Table S1, [15]). For MRP4 the yield of purified protein is relatively low at $35 \pm 5 \mu\text{g/L}$ culture. This reflects a lower expression level in these cells, but it should be noted that the SMA allows an efficient extraction from these low expressing cells and an effective purification, whereas when using the detergent DDM with these same membranes the eluted fractions from the Ni-NTA column contain a large number of contaminating proteins in levels comparable or higher than the MRP4 (Supplementary Figure 3), showing that the SMA method is effective even for poorly expressed membrane proteins.

Ligand binding by ABC transporters within SMALPs

Next we determined whether the structure and function of the ABC transporters was retained following SMA extraction by investigating their ligand binding efficiency. A fluorescence quenching method devised by Liu & Sharom [29] was used to measure the binding of both doxorubicin (a transported substrate) and verapamil (a competitive inhibitor) to purified Pgp within SMALPs (Figure 3A). The data were best fitted by a single site binding curve with a K_d for doxorubicin of $9.5 \pm 1.5 \mu\text{M}$ (mean \pm sem, $n=3$) and for verapamil $K_d = 2.0 \pm 0.4 \mu\text{M}$ (mean \pm sem, $n = 4$), which compare favourably to the previous results for detergent purified Pgp of $4.4 \mu\text{M}$ for doxorubicin and $2.4 \mu\text{M}$ for verapamil [29]. Furthermore the amplitude of the fluorescence change induced by drug binding shows that for both the SMA solubilised and previous detergent solubilised preparations the change in intensity for doxorubicin binding is close to twice that observed for verapamil [29], showing that the characteristics of doxorubicin and verapamil binding to SMALP-encapsulated Pgp are consistent with detergent solubilised Pgp.

The results of an estrone sulphate radioligand binding assay for SMA solubilised H69AR membranes in the absence or presence of GSH are shown in Figure 2. It has previously been shown that GSH binding causes a conformational change in MRP1 which increases the affinity for estrone sulphate binding [30]. SMALP-extracted MRP1 showed the same effect with the presence of 3 mM GSH causing a significant increase (2.3-fold, $P<0.05$) in the binding of estrone sulphate to SMALP encapsulated MRP1 which is comparable to that seen in native membranes.

A fluorescent chlorophyll metabolite, Pheophorbide A, is a known substrate of ABCG2 [42]. Upon binding to SMALP encapsulated ABCG2 a large increase was detected in the emission fluorescence of pheophorbide A (Figure 3C & Supplementary figure S2A). This increase was dose-dependent (Figure 3C), specific to ABCG2 (Supplementary figure S2B) and could be displaced by the ABCG2 inhibitor chrysin (Figure 3C inset) with an IC_{50} of $3.7 \mu\text{M}$, which is comparable with previous reports [43].

Nucleotide binding to purified Pgp encapsulated within SMALPs

ABC transporters undergo nucleotide driven conformational changes to reorient substrate binding sites from high to low affinity conformations. For at least two of the transporters purified here it is ATP binding that provides this “power-stroke” [44, 45] and so we investigated whether the SMALP purified Pgp retained ATP binding. Figure 4 shows the quenching of intrinsic Pgp tryptophan fluorescence upon addition of either ATP or the fluorescent analogue TNP-ATP. The degree of tryptophan quenching and the affinity for TNP-ATP was much higher than for ATP itself, as has been reported previously [31,

46]. The data were fit with a single site binding curve with a K_d for ATP of 0.49 ± 0.08 mM and for TNP-ATP of 84 ± 9 μ M, which are comparable to previously reported values of 0.28 mM for ATP and 50.6 μ M for TNP-ATP [31].

Biophysical characterisation of Pgp encapsulated within SMALPs

Next we wanted to characterise the biophysical properties of the purified ABC-SMALPs. For these studies we used Pgp as an example, because it is one of the best characterized ABC transporters [41]. The CD spectra shown in Figure 5A shows that the purified Pgp is folded, and the characteristic minima at 208 nm and 222 nm show that the structure is predominantly α -helical, as expected from recent structural studies [47, 48]. Analytical ultracentrifugation (Figure 5B) gives a single major peak showing that the Pgp within the SMALPs is monomeric. This is also shown with the gel filtration in Figure 5C, where the main peak corresponds to a molecular weight of approx 164 kDa, in agreement with the predicted molecular weight (140 kDa unglycosylated). The gel filtration trace shows that the SMALP encapsulated Pgp does not show the levels of aggregation previously observed with dodecylmaltoside solubilised Pgp [39, 41].

Cryo-electron microscopy (Figure 6) shows discrete particles of 5nm width and 15 nm length, consistent with monomers of Pgp. Classification of >8000 selected particles showed that the majority of particles were probably monomeric with a few classes indicating two Pgp particles associating to form hinge-shaped particles (Figure 6B). An initial low resolution 3D structure (Figure 6C) produced from the single particles showed an envelope which was consistent with monomeric Pgp. Refinement of the structure (Figure 6D) was possible, generating a final (low) resolution of about 3.5nm (Figure 6E). The final structure was consistent with the published Pgp structure (3G5U) [49], although the resolution of the electron microscopy-derived structure did not allow unambiguous discrimination between the inward- and outward facing states (for example the outward-facing Sav1866 structure [50] could equally well be fitted within the low resolution envelope). Notably, the cryo-EM structure of Pgp within SMALPs lacks a large disc-shaped belt, which might have been expected. However we have observed previously that the polymer itself shows almost no signal in unstained cryo-EM samples. Secondly it suggests the disc does not contain a large belt of lipid molecules, just a small annulus, which agrees with a previous study showing approximately 11 lipid molecules per protein [17].

Thermostability of Pgp encapsulated within SMALPs

We surmised that the extraction of an ABC transporter with SMA, maintaining the lipid bilayer surrounding it, would not only preserve its structural integrity, but could also provide some advantages in terms of protein stability compared to detergent extraction. To investigate this we examined the thermostability of purified Pgp encapsulated within SMALPs and compared it to Pgp solubilised using octyl glucoside as previously published [20]. Thermostability was monitored using the method of Alexandrov *et al.* [35] where reactivity with the fluorophore CPM was monitored over time as the temperature was systematically increased (Figure 7). This demonstrates that the Pgp-SMALP sample was significantly more stable than the detergent solubilised sample, with the apparent melting temperature increasing by more than 10 °C.

Conclusion

Until now the solubilisation and purification of ABC transporters has relied on the use of detergents to disrupt the membrane environment surrounding membrane proteins. This study has demonstrated that the polymer SMA can instead be used to extract eukaryotic ABC transporters from a wide range of different cell types and expression systems without the need for detergents.

The extracted proteins retain the ability to bind substrates and nucleotide and can be purified by affinity chromatography. This is important for two key reasons: firstly detergents are expensive and the requirement to supplement all purification buffers with detergent to maintain solubility can become financially limiting when scaling up procedures. In contrast SMA is mass produced and can be used in a single step to form stable lipid disc structures that do not require addition of further polymer to any buffers. Secondly, and perhaps more importantly, the SMA method doesn't disrupt the lipid environment surrounding a membrane protein in the way that detergents do. Instead a segment of membrane is extracted, and the native lipid bilayer surrounding a membrane protein is maintained.

Furthermore the resultant purified transporters encapsulated within SMALPs are more pure, more stable and less prone to aggregation than detergent solubilised. It was also possible to concentrate the purified Pgp-SMALPs to > 8 mg/ml without significant loss of protein or aggregation.

In the current study we have not attempted to demonstrate ATPase of SMALP purified ABC transporters, (although we have showed that the proteins retain their interaction with nucleotide), because the SMALP is destabilised by divalent metal ions. Reconstitution from the lipid disc structure into proteoliposomes will be required to fully characterise the ATP hydrolysis kinetics and this will also allow the measurement of substrate transport.

However, as demonstrated by the work of Sligar and co-workers with nanodiscs, the small lipid-disc structure is amenable to many different techniques [51]. The amphipathic polymer-based SMALPs do not suffer from signal interference from the apolipoprotein surrounding the lipid disc. Finally the EM structural data shows that there is not a large belt of lipids, and thus it may be compatible with structural studies such as cryo-electron microscopy and in principle could be used in 3D crystallisation trials. Thus this work opens up several new opportunities for the structural and pharmacological study of ABC transporters, and their annular lipid environment.

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Figure legends:

Figure 1. Extraction of ABC transporters using SMA polymer. Membrane preparations (30 mg/ml wet weight) were mixed with 2.5 % (w/v) SMA in 20 mM Tris pH8, 150 mM NaCl, 10 % (v/v) glycerol for 1 hour at room temperature, then centrifuged at 100,000 g for 20 min. Sample F also had 1 % (w/v) DMPC added alongside the SMA, and was incubated at 37°C rather than room temperature. Samples of the soluble and insoluble material were assayed by Western blotting, and the percentage of the target protein in each was measured using densitometry.

Figure 2. Purification of ABC transporters within SMA-lipid particles. A; SMA solubilised Pgp membranes (Sol) were incubated with Ni-NTA resin at 4°C overnight. Unbound material (FT) was removed and the resin was washed with buffer supplemented with 20-60mM imidazole before being eluted with buffer containing 120mM imidazole. Samples from each fraction were run on SDS-PAGE and silver stained. B-D; elution fractions from purifications carried out as in A but for ABCG2 (B), MRP4 (C) and CFTR (D). E; QCRL-1 antibody was bound to a protein A column, then SMA solubilised H69AR membranes were incubated with the column for 30 min at 20°C. Unbound proteins were washed out and MRP1 was eluted upon addition of the peptide SSYSGDI.

Figure 3. Ligand binding assays. A; Purified Pgp-SMALPs were labelled with the fluorophore MIANS, then fluorescence quenching upon binding of the inhibitor verapamil (open circles) and transport substrate doxorubicin (closed circles) were measured. B; Binding of ³H-estrone sulphate to SMA solubilised H69AR membranes (MRP1) was measured in the presence and absence of 3mM GSH (*P<0.05). E; Specific binding of pheophorbide A to SMA-solubilised HEK-ABCG2 membranes detected by an enhanced fluorescence of pheophorbide A at 681 nm. All data are mean±sem, n≥3. Inset; dose-dependent displacement of pheophorbide A binding by chrysin.

Figure 4. Nucleotide binding to purified Pgp-SMALPs. Quenching of the intrinsic tryptophan fluorescence of purified Pgp (50µg/ml) within SMALPs upon binding of nucleotides (A) ATP and (B) TNP-ATP. Data are mean±sem and n≥3.

Figure 5. Biophysical characterisation of purified Pgp-SMALPs. A; Circular dichroism spectra at 0.65 mg/ml with a 0.1mm pathlength. B; Analytical ultracentrifugation trace. C; Gel filtration on a superdex 200 column (diameter 1.6cm, height 55cm), solid line is absorbance at 280nm, dashed line is conductivity.

Figure 6. Structural characterisation of purified Pgp-SMALPs. A; Pgp particles solubilised in SMALPs and visualised (unstained) using cryo-electron microscopy. The edge of the hole in the carbon support film of the electron microscopy grid is visible top right. The particles are 10-15nm diameter and are relatively monodisperse. B; Classification of the particles reveals elongated particles of about 5nm width and 15nm length. Some classes appear to be formed by two particles associating. Box size is 21.6nm. C; An initial low resolution 3D envelope generated from single particle classes, and D; further refinement of the structure against all the raw particles. Orthogonal views are shown, with the ABCB1 crystal structure (3G5U) docked within it (blue ribbon). The refined structure was estimated to be at about 3.5nm resolution (E).

Figure 7. Thermostability of purified Pgp. Purified Pgp (5 µg) was mixed with 8 µg/ml coumarin maleimide in a total volume of 200 µl. The reaction mixture was heated continuously from 10°C - 90°C in increments of 5°C for 2 minutes each and the fluorescence spectra measured.

Fluorescence intensity (at 470nm) at each time point was calculated as a % of the final fluorescence intensity. Pgp was either solubilised with 2.5% SMA (open circles) or 2% octyl glucoside (closed circles).

Figure 1

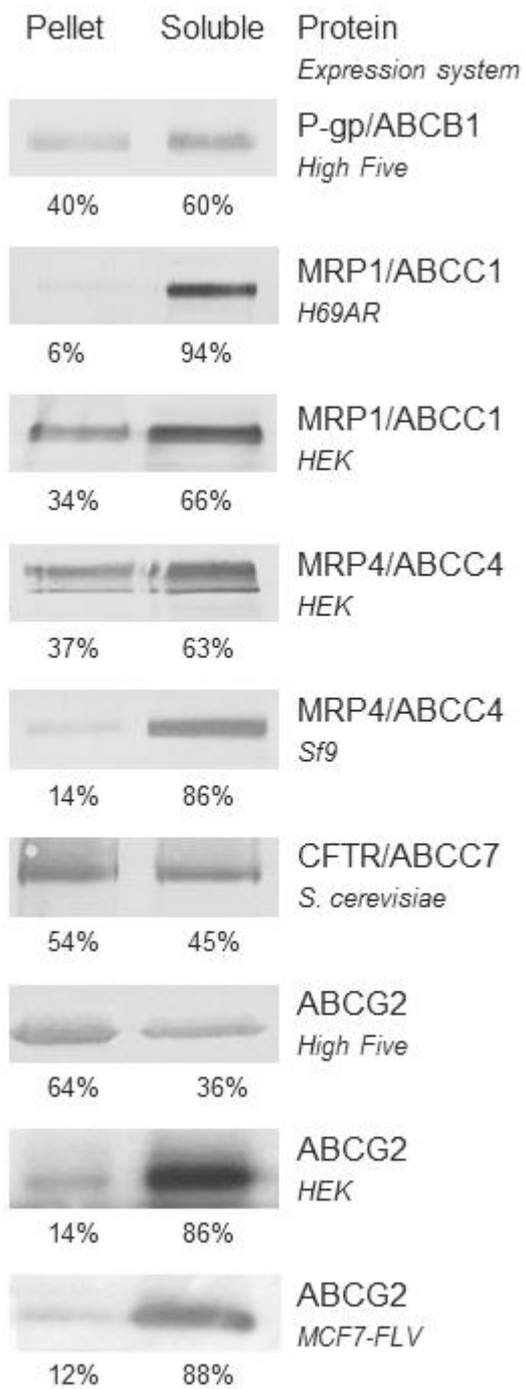


Figure 2

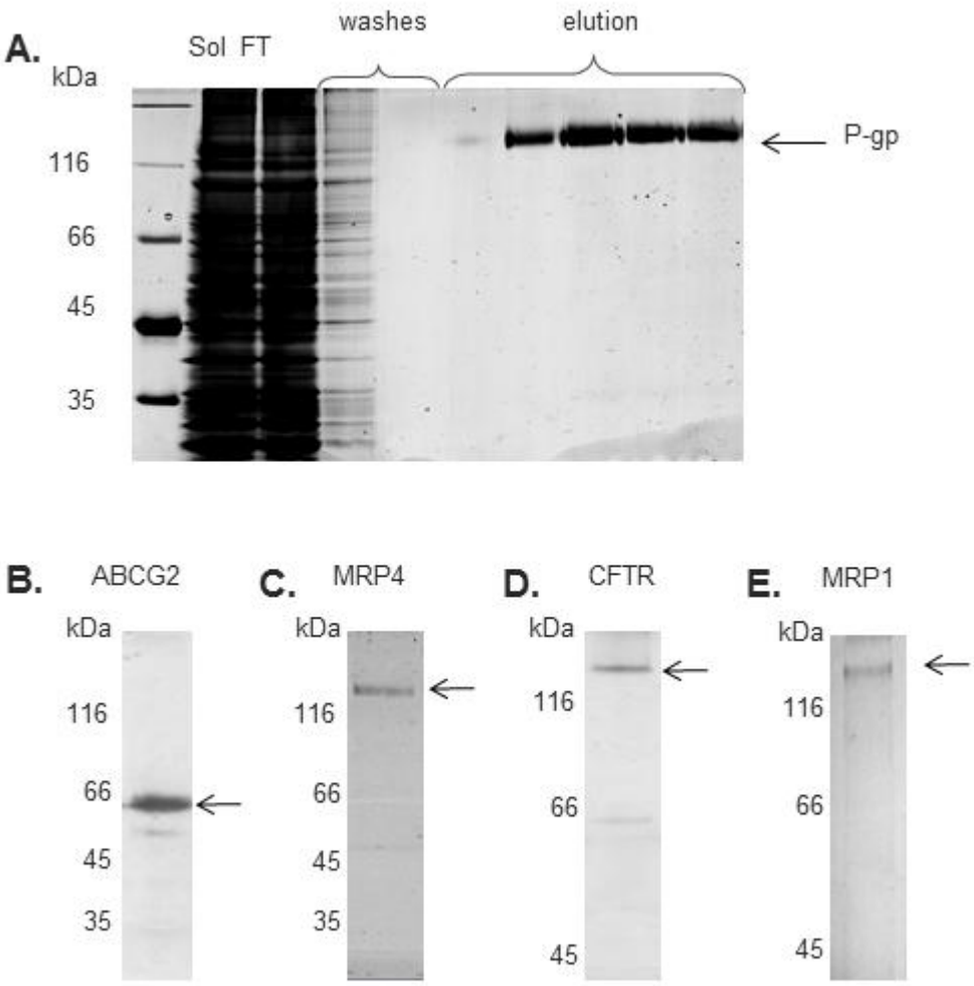


Figure 3

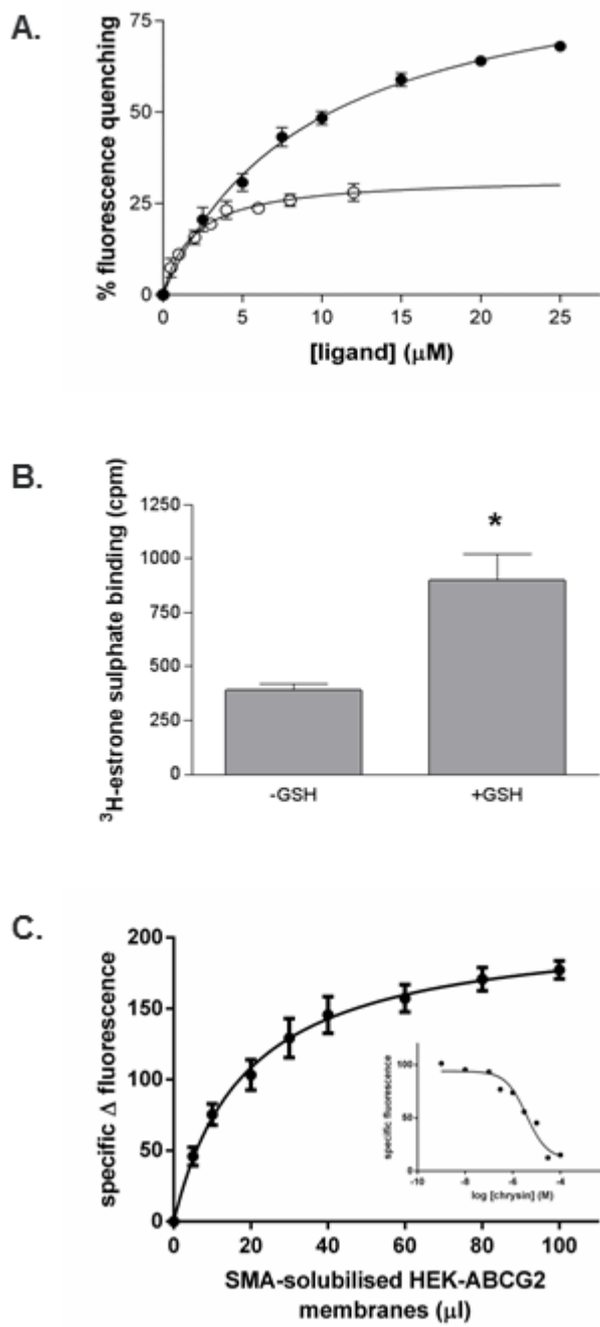


Figure 4

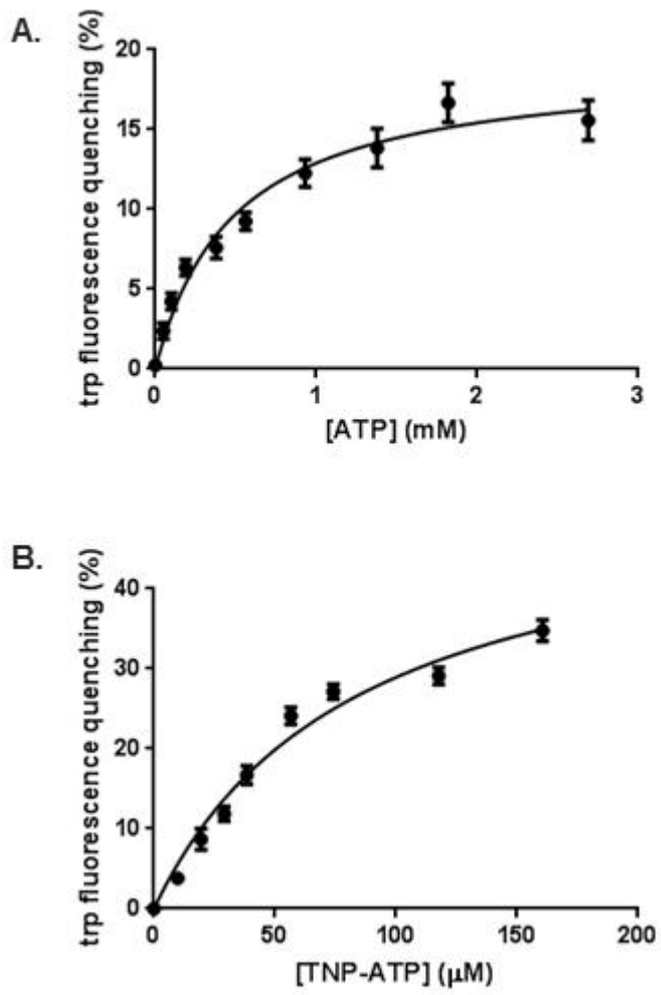


Figure 5

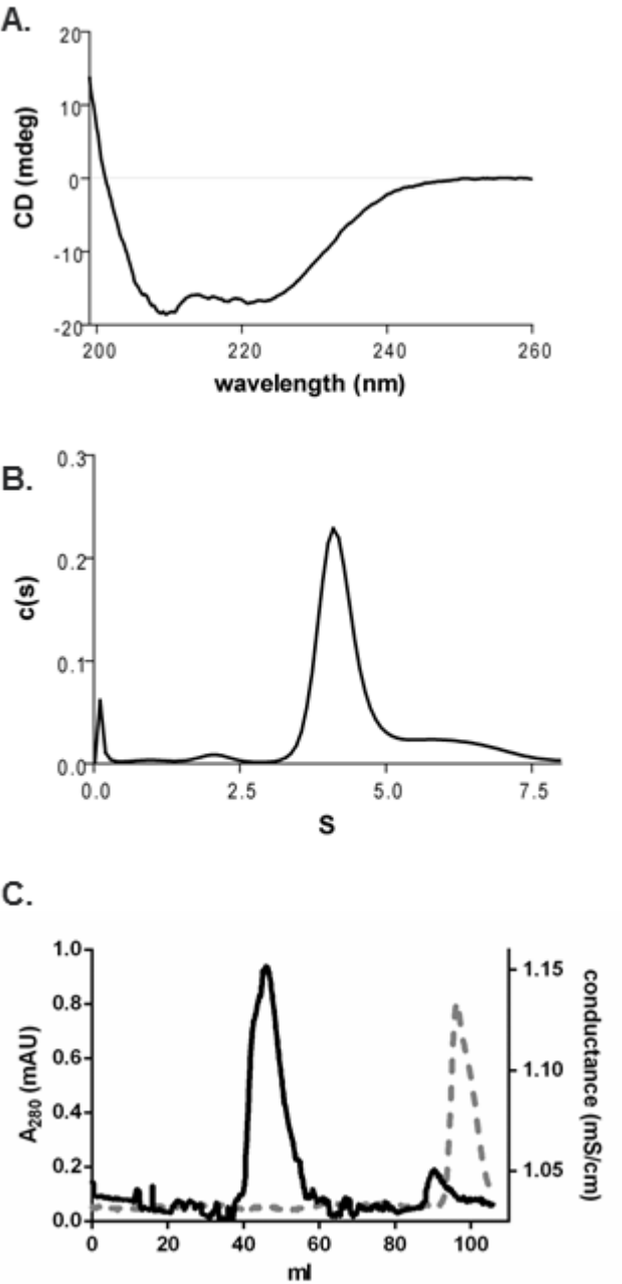


Figure 6

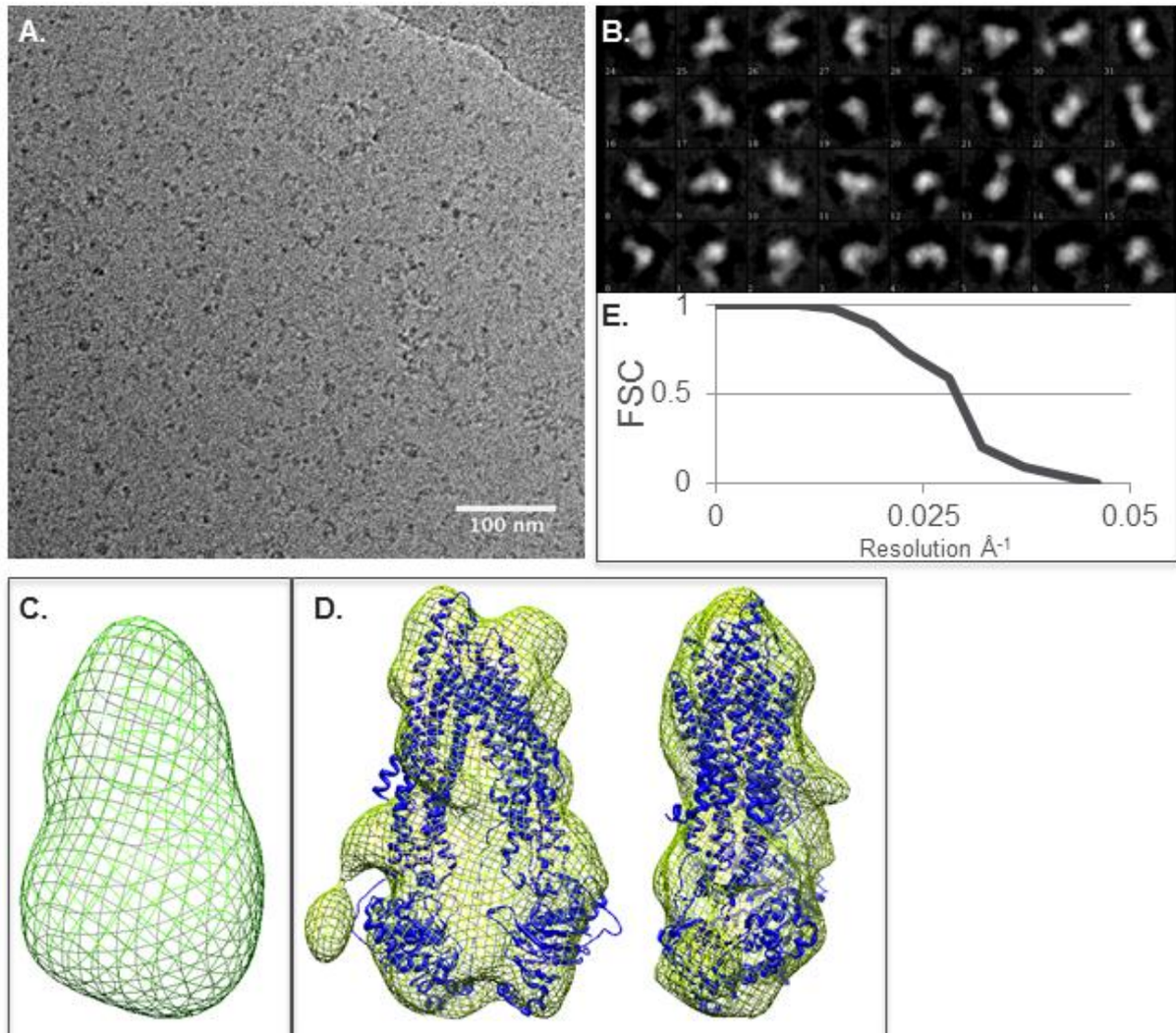
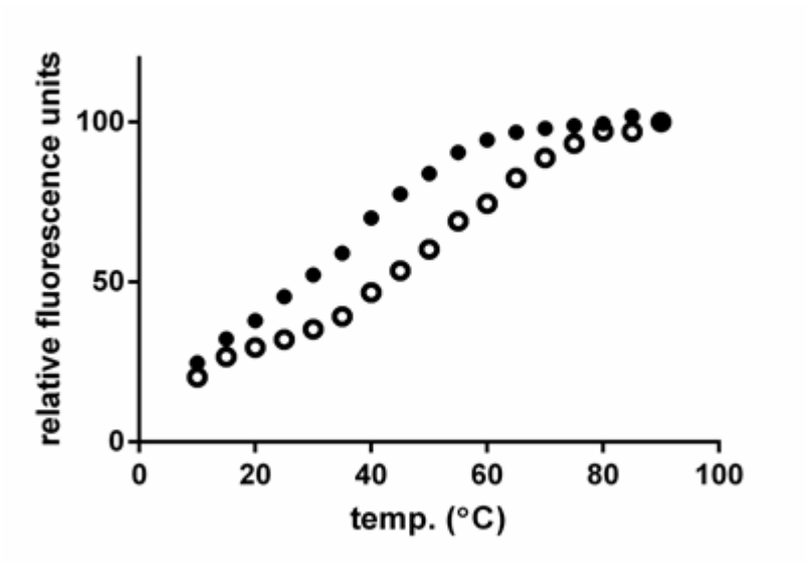


Figure 7



Supplementary Information

ABC protein (expression cell type)	Yield of purified protein (μg) per 1L culture		
	SMA	detergents	
Pgp/ABCB1 (<i>High Five</i>)	631 \pm 46	OG ^a 792 \pm 197	DDM ^b 249 \pm 15
ABCG2 (<i>High Five</i>)	116 \pm 32	FC-16 ^c 39 \pm 14	-
MRP4/ABCC1 (<i>Sf9</i>)	35 \pm 3	-	-
CFTR/ABCC7 (<i>S.cerevisiae</i>)	33	DDM ^d 67	LPG ^d 133
		Yield of purified protein (μg) per 10 ⁸ cells	
MRP1/ABCC1 (<i>H69AR</i>)	126	CHAPS ^e 250	-

Supplementary Table S1. Comparison of yield of purified protein using SMA or detergents

The final yield of purified protein produced using SMA was calculated for each of the proteins studied per litre of cultured cells (Pgp, ABCG2, MRP4 and CFTR) or per 10⁸ cells (MRP1). Where errors are shown data are mean \pm sem, n \geq 3. For comparison yields previously obtained with these membranes using various different detergents are also shown. OG (octyl β -D-glucoside), DDM (dodecyl- β -D- maltoside), FC-16 (foscholine-16), LPG (lysophosphatidylglycerol), CHAPS (3-(3-cholamidopropyl)dimethylammonio-1-propanesulfonate).

^aTaylor *et al* (2001) Br J Pharmacol 134;1609-1618, R. Callaghan (personal communication)

^bPollock *et al* (2014) Biochim Biophys Acta 1838;134-47, R. Callaghan (personal communication)

^cMcDeviit *et al* (2006) Structure 14;1623-32, R. Callaghan (personal communication)

^dO’Ryan *et al* (2012) J Vis Exp 61, e3860, doi:10.3791-3860

^eMao *et al* (1999) Biochim Biophys Acta 1461;69-82.

Supplementary Figure 1

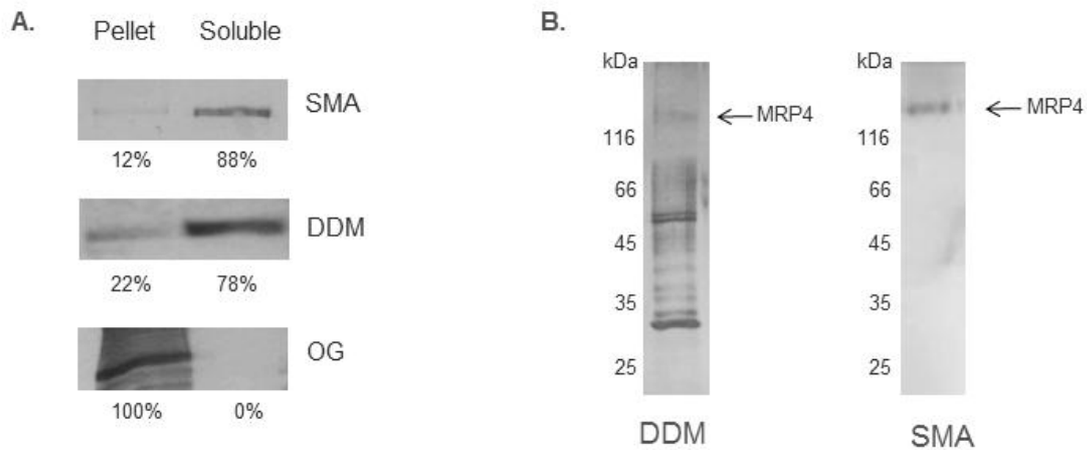


Figure S1. Comparison of MRP4 solubilisation and purification from Sf9 cells using SMA or detergent. A. Sf9-MRP4 membranes were solubilised in either 2.5 % (w/v) SMA, 2 % (w/v) DDM or 2 % (w/v) OG for 1 hour and then centrifuged at 100,000g for 20min at 4°C. Samples of the soluble and insoluble material were assayed by Western blotting, and the percentage of the target protein in each was measured using densitometry. B. Soluble material was incubated with Ni-NTA resin overnight at 4°C, transferred to a mini column and washed extensively with buffer supplemented with 20mM imidazole. Samples shown were eluted using 120mM imidazole. SDS-PAGE gels are silver stained.

Supplementary Figure 2

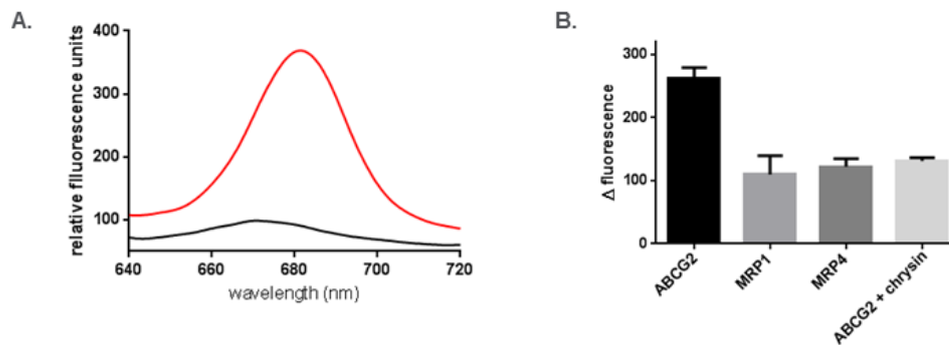


Figure S2. Enhanced fluorescence of pheophorbide A upon binding to ABCG2. A; Pheophorbide A (10 μ M, 1 ml) fluorescence with an excitation wavelength of 390 ± 10 nm, before (black) and after (red) a 10 minute incubation with ABCG2 containing SMA-solubilised membranes (40 μ l). B; The increase in fluorescence emission at 681 nm of pheophorbide A (as in A) when SMA-solubilised membranes with overexpressed ABCG2, MRP1 or MRP4 are added, or when ABCG2 membranes are preincubated for 10 minutes with 30 μ M chrysin. Data are mean \pm sem, $n \geq 3$.