Rapid transfer of overexpressed integral membrane protein from the host membrane into soluble lipid nanodiscs without previous purification

Abstract: Structural and functional characterization of integral membrane proteins in a bilayer environment is strongly hampered by the requirement of detergents for solubilization and subsequent purification, as detergents commonly affect their structure and/or activity. Here, we describe a rapid procedure with minimal exposure to detergent to directly assemble an overexpressed integral membrane protein into soluble lipid nanodiscs prior to purification. This is exemplified with recombinant his-tagged rhodopsin, which is rapidly extracted from its host membrane and directly assembled into membrane scaffold protein (MSP) nanodiscs. We further demonstrate that, even when the MSP was his-tagged as well, partial purification of the rhodopsin-nanodiscs could be achieved exploiting immobilized-metal chromatography. Recoveries of rhodopsin up to 80% were achieved in the purified nanodisc fraction. Over 95% of contaminating membrane protein and his-tagged MSP could be removed from the rhodopsin-nanodiscs using a single Ni²⁺-affinity chromatography step. This level of purification is amply sufficient for functional studies. We provide evidence that the obtained rhodopsin-nanodisc preparations are fully functional both photochemically and in their ability to bind the cognate G-protein.

Keywords: baculovirus expression system; his-tagged rhodopsin; membrane protein solubilization; membrane scaffold protein; protein stabilization; recombinant protein.

Introduction

Membrane proteins are involved in numerous biological processes, such as transport, signal transduction, and a variety of metabolic pathways. Up to 30% of the human proteome consists of integral membrane proteins (Wallin and VonHeijne, 1998), and these account for approximately half of all currently marketed therapeutic targets (Wishart et al., 2006). Despite their crucial biological relevance, relatively little is known in detail about their molecular mechanism and interaction patterns. An important bottleneck is the requirement of membrane proteins for a native-like membrane environment to maintain their correct structural folding and flexibility, and thereby, their physiological function.

Unfortunately, for most membrane proteins, studies of structure and function are limited due to either low expression levels and/or the absence of a simple and reproducible procedure for isolating these proteins in a monodisperse form that maintains their native structure. In the first purification step, membrane proteins commonly undergo solubilization with detergents, which nearly always destabilize these proteins and strongly affect their activity (Gohon and Popot, 2003; Zhou and Cross, 2013; Zoonens et al., 2013). The micellar environment, although amphipathic, will by itself destabilize the protein structure. Further, specific lipids that are essential for stability and/or activity may be lost during purification (DeGrip, 1982; DeGrip and Daemen, 1982; Rosenbusch, 2001).

Recently, studies have emerged that use small styrene-malic acid copolymers (SMA) for extracting integral membrane proteins from their natural cellular membrane. This eliminates a detergent solubilization step (Jamshad...
et al., 2011; Orwick-Rydmark et al., 2012; Long et al., 2013). The resulting complexes retain functional properties and they can be readily purified by affinity chromatography. However, this approach has a restricted pH range, the copolymer is highly negatively charged and the membrane protein is surrounded by only a limited number of lipid molecules (Gulati et al., 2014; Swainsbury et al., 2014).

Here, we present an alternative versatile approach exploiting the potential of ‘membrane scaffold protein’ (MSP) nanodiscs. This allows the extraction of integral membrane proteins from cellular membranes under minimal exposure to detergent, and to purify them in a preferentially monodisperse, stable state that maintains an amphipathic environment optimally preserving their structure and activity.

Over the last decade, elegant technology has been developed reconstituting integral membrane proteins into a soluble discoidal phospholipid bilayer environment confined by two copies of a genetically engineered MSP, termed nanodiscs (Bayburt et al., 2002, 2006; Ritchie et al., 2009; Marty et al., 2013). These nanodiscs have diameters ranging from 9.5 nm (nanodiscs) to 30 nm (macrodiscs) (Chromy et al., 2007; Park et al., 2011) and provide a native-like environment surrounding the hydrophobic domains of the membrane proteins. They exhibit high stability in a detergent-free aqueous solution and allow standard chromatographical procedures for protein purification (Bayburt et al., 2007; Whorton et al., 2007; Banerjee et al., 2008; Blanchette et al., 2009; Ritchie et al., 2009). The nanodisc technology has enabled biochemical and biophysical studies of a variety of membrane proteins (Bayburt et al., 2006, 2007; Boldog et al., 2006; Leitz et al., 2006; Whorton et al., 2007; Banerjee et al., 2008; Pandit et al., 2011), and membrane bound enzymes (Civjan et al., 2003; Denisov and Sligar, 2011). However, in almost all studies, the target proteins were purified using detergents before incorporation into nanodiscs. Only in a single early study, N-terminally anchored CYP was assembled into nanodiscs prior to purification (Civjan et al., 2003).

In this report, we describe the first study in which an integral membrane protein is extracted from its host membrane and directly assembled into MSP nanodiscs. The nanodiscs with the target protein can subsequently be purified, for example, by affinity chromatography (Figure 1B). This strongly minimizes exposure to detergents, and the resulting preparation retains full functional activity. The particular protein we selected is rhodopsin, the visual pigment of the vertebrate rod photoreceptor cell, an established paradigm for the G protein-coupled receptor (GPCR) family. Rhodopsin folds into the membrane-spanning seven-helical bundle typical of GPCRs. Its ligand and chromophore, 11-cis retinal, is covalently bound to Lys-296 in helix VII via a protonated Schiff-base linkage (Wald, 1968; Palczewski et al., 2000).

The presented approach is amenable to any membrane protein, and in particular useful for those proteins that are susceptible to perturbation by detergents. We further demonstrate that, even when his-tagged MSPs are employed for nanodisc formation, a single immobilized metal affinity chromatography (IMAC) purification step can already isolate the his-tagged target-protein containing nanodiscs from most of the contaminating nanodisc population.

### Results

#### Nanodisc formation: strategy

Whereas the lipid bilayer in a nanodisc presents an appropriate environment for membrane proteins, the target-protein is routinely solubilized and it is purified using a suitable detergent prior to incorporation into nanodiscs (Figure 1A). However, detergent solubilization and lipid removal seriously compromise structural and functional integrity of membrane proteins (Gohon and Popot, 2003; Zhou and Cross, 2013; Zoonens et al., 2013). These negative effects often cannot be fully repaired upon reconstitution of the purified protein within a bilayer environment. Here we propose an approach to minimize this problem by incorporating membrane proteins into nanodiscs immediately after brief solubilization of the cellular membranes with detergent. This has the major advantage that the target protein is only briefly exposed to detergents and that stabilizing lipids remain present. The nanodiscs are assembled with the available lipid pool in the cellular membranes used for heterologous expression, creating a system that closely mimics a natural membrane environment. We expect that this should provide sufficient lipids, since the lipid to protein ratio in natural membranes is close to 1:1 (w/w) and large proteins and complexes will not easily incorporate into nanodiscs. In fact, it has been recently demonstrated that nanodisc solubilization does represent the entire membrane proteome (Marty et al., 2013). Our approach is schematically explained in Figure 1B.

In order to model our approach, we used rhodopsin, as it can be easily monitored by its spectral properties and was shown before to retain full functionality in a nanodisc environment (Leitz et al., 2006; Bayburt...
et al., 2007, 2011; Banerjee et al., 2008; Whorton et al., 2008; Tsukamoto et al., 2011). Native rhodopsin can be extracted from purified rod outer segment membranes (DeGrip et al., 1980). His-tagged opsin can be heterologously expressed in Sf9 insect cells (Janssen et al., 1995; Klaassen et al., 1999) and incubation with 11-cis retinal generates his-tagged rhodopsin. For the generation of nanodiscs we opted for the membrane scaffold protein zebrafish Apo A-1 (ZAP1), shown before to form functional nanodiscs with purified rhodopsin (Banerjee et al., 2008). To estimate the minimal quantity of ZAP1 required for proper nanodisc formation of the membrane protein population of Sf9 cells, we used a membrane protein content of 225±25 μg per 10^6 cells (DeCaluwe et al., 1993) and estimated the average mass of the membrane protein population at 55±5 kDa (Prioret et al., 2011; Marty et al., 2013). This results in approximately 4 nmol of membrane protein per 10^6 cells. For nanodisc assembly with purified membrane proteins, a minimal molar ratio of two MSP per membrane protein (MP) is required, but in practice, ratios of 2–25 have been employed (Bayburt et al., 2007; Banerjee et al., 2008; Ritchie et al., 2009; Pandit et al., 2011; Inagaki et al., 2013; Mors et al., 2013). A higher ratio may result in a more homogeneous population of MP-nanodiscs, but also generates larger contamination with empty nanodiscs and single MSPs. As the entire membrane protein population will not insert into nanodiscs, and to be cost-effective with respect to MSP, we considered a 2–3-fold molar excess of MSP, that is, 8–12 nmol of MSP per 10^6 cells. For solubilization of rhodopsin, we selected n-dodecyl-β-D-maltoside (DDM), a detergent commonly used for

Figure 1: Schematic representation of our approach. Compared are the classic approach (panel A) (target-protein incorporation into nanodiscs after purification) and our approach (panel B). In the latter, first, the entire membrane population is incorporated into nanodiscs and out of this total mixture, the target-protein nanodiscs are purified. Note that the liposomes that can be generated with the purified target-protein (panel A), in reality are much larger (diameter varying between 100 and 5000 nm) than the nanodiscs (12–30 nm).
membrane protein solubilization (Rigaud et al., 1998) and shown to be effective as well as mild towards rhodopsin (DeGrip, 1982). Detergent extraction to generate nanodiscs is usually accomplished by incubation with a lipophilic matrix (Rigaud et al., 1998; Civjan et al., 2003; Leitz et al., 2006; Bayburt et al., 2007; Banerjee et al., 2008). However, this is a relatively slow process and, in our hands, led to considerable loss of protein. To effectuate rapid detergent extraction with minimal loss of protein, we selected the cyclodextrin-inclusion procedure, which also affords quantitative removal of detergent, but allows better control over kinetics and time investment (DeGrip et al., 1998).

Nanodisc formation: experimental implementation

Opsin expression, rhodopsin regeneration and solubilization

Bovine opsin extended with a C-terminal deca-histidine tag was produced in Sf9 insect cells using recombinant baculovirus, as described (Klaassen et al., 1999; Shirzad-Wasei et al., 2013). Full-length opsin expression was confirmed by immunoblotting (Figure 2, lane 1). The triad pattern represents diglycosylated (upper band), mono glycosylated (middle band) and unglycosylated (bottom band) opsin (Janssen et al., 1991; DeCaluwé and DeGrip, 1996). To convert the apo-protein opsin into the holo-pigment rhodopsin, which allows for a straightforward spectroscopic assay (Wald, 1968; DeGrip, 1982), membranes from infected Sf9 cells were incubated with 11-cis retinal. This reaction is completed at room temperature (RT) within 60 min. Subsequently, hydroxylamine was added to 50 mM final concentration to convert excess retinal into the chemically more stable retinaloxime (Wald, 1968; Groenendijk et al., 1980). The expression level of rhodopsin can be assessed after solubilization from the absorbance and its molar absorbance coefficient at 500 nm [40 600 M⁻¹ cm⁻¹; (Wald, 1968; Daemen et al., 1970)]. To minimize solubilization time and exposure of the target protein to detergent prior to nanodisc formation, a relatively low membrane density was employed (equivalent to 20×10⁴ cells/ml) in combination with a moderate detergent concentration (20 mM DDM). Near-complete solubilization of rhodopsin was achieved within 30 min at RT or within 60 min at 4°C. Immunoblot analysis demonstrates that predominantly the diglycosylated upper band is solubilized (Figure 2, lane 2). This is in line with earlier conclusions that the majority of the mono- and the unglycosylated protein is not correctly folded and aggregates in the presence of a mild detergent (Janssen et al., 1991; DeCaluwé and DeGrip, 1996).

Nanodisc assembly

For assembly of a homogeneous population of nanodiscs, a defined ratio of phospholipids to MSP is recommended (Bayburt et al., 2002). We utilized the available lipid pool of cellular membranes and first tested this on bovine rod outer segment membranes, which mainly contain rhodopsin (DeGrip et al., 1980). After brief solubilization, addition of ZAP1 to a 3-fold molar excess over rhodopsin, formation of nanodiscs and centrifugation to precipitate larger structures, the supernatant was analyzed for the presence of rhodopsin by spectroscopy. As shown in Figure 3, over 90% of the rhodopsin was recovered in the soluble nanodisc fraction, indicating that the natural membrane lipid pool can be sufficient to accommodate the nanodiscs.

Figure 2: Immobilized metal affinity chromatography purification profile of rhodopsin-nanodiscs analyzed by immunoblotting. The blot was screened with anti-his-tag antibody. Lane M represents marker proteins with molecular weights indicated. Lane 1 shows the Sf9 cell membrane fraction (see the text for explanation of the triad pattern for opsin in the 30–40 kDa range). Lane 2 contains the n-dodecyl-β-D-maltoside extract of Sf9 membranes after regeneration of rhodopsin. The sample in lane 3 is taken after addition of the membrane scaffold protein zebrafish Apo A-1 (ZAP1) to the extract. Note the big blurred band below rhodopsin that represents the large excess of ZAP1. Lane 4 shows the supernatant after nanodisc formation and spinning down the insoluble components, confirming that soluble rhodopsin-nanodiscs have been generated. The majority of the protein contamination including ZAP1 elutes in the non-bound fraction and the additional washes with buffer and low concentrations of histidine (cf. Figure 3). Most of the remaining ZAP1 is eluted in the 50 mM histidine fraction (lane 5) where rhodopsin slowly starts to elute. The remaining rhodopsin co-elutes with the remaining ZAP1 in the 200 mM histidine fraction (lane 6) and the 270 mM histidine fraction (lane 7). Lanes 8 and 9 present purified ZAP1 and his-tagged rhodopsin, respectively. ZAP1 is expressed and purified in three major forms that are all competent in nanodisc formation (Pandit et al., 2011).
Subsequently, this was tested on his-tagged rhodopsin heterologously produced in Sf9 insect cells, where rhodopsin constitutes <0.5% of the membrane protein population (Klaassen et al., 1999; Bosman et al., 2003). In this case, lipid content and composition of the cellular membranes may vary somewhat over different cultures. We observed that under those conditions the recovery of rhodopsin in nanodiscs varied considerably. Instead, addition of extra lipid [lipid:MP ratio of 1:4 (w/w)] produced quite reproducible results. For the extra lipid, we selected asolectin, a natural lipid source that has a good variety in lipid species and fully maintains the functionality of a variety of membrane proteins, including rhodopsin (Sone et al., 1977; Klaassen and DeGrip, 2000; Ratnala et al., 2004). Asolectin could be added before or after solubilization of the membrane fraction.

Upon removal of detergent, nanodiscs self-assemble from a mixture of phospholipid/detergent mixed micelles and MSP (Bayburt et al., 2002). Many GPCRs show a time-dependent loss of activity in detergents (Navratilova et al., 2005); hence, long-term exposure to detergents needs to be avoided. The cyclodextrin-inclusion procedure described earlier (DeGrip et al., 1998) allows rapid assembly of membrane proteins (MP), lipid and MSP into nanodiscs with good recovery of MP (Pandit et al., 2011). In the example presented here (Figure 2), we used ZAP1 as MSP (Banerjee et al., 2008) and an estimated 1:2 molar ratio of membrane protein to ZAP1. Similar results were obtained with the MSPs MSPD1 and MSPE3D1. Molar ratios of membrane protein to MSP of 1:3 yielded comparable recoveries. Generation of nanodiscs was initiated by addition of a slight excess of heptakis-2,6-di-O-methyl-β-cyclodextrin (β-cyclodextrin), which will extract detergent from mixed micelles under formation of inclusion complexes (DeGrip et al., 1998). Addition in three steps allows gradual self-assembly of the nanodiscs. Even more rapid formation of nanodiscs, that is, within 15 min, can be accomplished by single-step addition of β-cyclodextrin, but this may result in a lower recovery or a more heterogeneous population of nanodiscs (Pandit et al., 2011).

Partial purification of rhodopsin-nanodiscs

Following nanodisc assembly, insoluble contamination, such as protein aggregates and accidental proteoliposomes, was removed by centrifugation. The resulting supernatant contained empty and MP-nanodiscs, possibly excess MSP, cyclodextrin-detergent complexes, and excess cyclodextrin (Figure 2, lane 4). The most straightforward procedure to purify the target-protein nanodiscs from this mixture would be exploiting dedicated affinity chromatography, for example, using an immobilized antibody against the target protein or an immobilized ligand. This is well documented in the literature (Bayburt et al., 2007; Banerjee et al., 2008; Whorton et al., 2008; D’Antona et al., 2014) and does not need to be further demonstrated here. However, such options are not often available. A his-tag is the most popular tag for recombinant proteins, allowing easy, affordable and high-capacity purification over immobilized-Ni²⁺ or Co²⁺-matrices (Janssen et al., 1995; Klaassen et al., 1999). In our case, we anticipated that MSPs and other nanodiscs would be co-purified owing to the his-tag carried by the MSP, but we observed that empty nanodiscs assembled with his-tagged MSP (HT-ND) have a relatively low affinity for the Ni²⁺-matrix. This allows for a differential elution from the IMAC matrix of HT-ND containing his-tagged rhodopsin, empty HT-ND and HT-ND containing non-his-tagged membrane protein.

A typical example is presented in Figures 2 and 4. While most of the rhodopsin-nanodiscs are bound to the nickel matrix (Table 1), most of the protein contamination including the ZAP1 proteins appears in the flow-through and the wash (Figure 4, lane 2). Most of the remaining ZAP1 elutes in the 50 mM histidine fraction, where rhodopsin only slowly starts to elute (Figure 2, lane 5; Figure 4, lanes 5 and 6). Subsequent treatment with 200 and 270 mM histidine elutes the remaining rhodopsin, co-purifying with
Figure 4: Immobilized metal affinity chromatography (IMAC) purification profile of rhodopsin-nanodiscs analyzed using silver staining. Fractions were separated by SDS-PAGE, and analyzed for protein by silver staining. Membrane fractions, extracts, supernatant after nanodisc formation and IMAC flow through are not shown, as they are overabundant with Sf9 membrane proteins. Lane M represents marker proteins with molecular weight indicated. Lanes 1 and 10 represent purified his-tagged rhodopsin and purified zebrafish Apo A-1 (ZAP1), respectively. Lanes 2–4 present wash fractions showing that most of the protein contamination and part of the ZAP1 proteins are removed at that stage. Lanes 5 and 6 contain the 50 mm histidine elution fractions, demonstrating elution of most remaining contaminating proteins and ZAP1, and rhodopsin beginning to elute. Lanes 7 and 8 (200 mm histidine fractions) and lane 9 (270 mm histidine fraction) represent elution of rhodopsin-nanodiscs, as rhodopsin and ZAP1 co-elute.

Table 1: Recovery of rhodopsin-nanodiscs.

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Percentage rhodopsin</th>
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<tr>
<td></td>
<td>Immunoblot</td>
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<tr>
<td>Membrane extract</td>
<td>=100</td>
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<tr>
<td>Non-bound fraction</td>
<td>11±7</td>
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<tr>
<td>Wash fraction</td>
<td>17±15</td>
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<tr>
<td>Purified nanodiscs</td>
<td>72±8</td>
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The amount of rhodopsin solubilized in detergent is taken as 100%. Immobilized metal affinity chromatography (IMAC) fractions are quantified with quantitative immunoblot analysis using primary anti-His and anti-rhodopsin antibodies in combination with the Odyssey imaging system (left column). Calibrated amounts of purified recombinant bovine his-tagged rhodopsin were applied as an internal standard (Klaassen et al., 1999). Rhodopsin-nanodiscs present in the purified IMAC fractions were also quantified by spectroscopy (right column). As immunoblotting is significantly more sensitive than spectroscopy, the latter probably underestimates the rhodopsin levels in the IMAC fractions. Values are given with SD, based upon three experiments; n.d., not detectable.

Figure 5: Spectral analysis of immobilized metal affinity chromatography purification fractions.

Typical UV/Vis absorbance spectra of a diluted wash fraction (dashed curve) mainly presenting a protein band at 280 nm and a purified rhodopsin-nanodisc fraction (continuous curve) displaying the additional typical rhodopsin band at 500 nm and excess ligand-derived retinaloxime at 365 nm.

the remaining ZAP1 (Figure 2, lanes 6 and 7; Figure 4, lanes 7, 8, and 9). Thus, a single IMAC procedure can purify his-tagged rhodopsin-nanodiscs from the large majority of contaminating proteins and nanodiscs, even if the MSP is his-tagged as well. Preliminary evidence suggests that this purification can be fine-tuned by using a slightly concave continuous gradient of histidine (10–270 mm). We surmise that in the nanodiscs the his-tag is not sufficiently accessible or is structurally too restrained to bind with high affinity to the matrix-bound metal ion. Quantitative immunoblot analysis using primary anti-his and anti-rhodopsin antibodies revealed excellent recoveries of up to 80% of the rhodopsin solubilized from the cellular membranes (Table 1, left column). This is at least comparable to earlier results obtained via path A in Figure 1, where following reconstitution into proteoliposomes or nanodiscs after purification 40–70% of the originally solubilized rhodopsin is recovered (Klaassen et al., 1999; Klaassen and DeGrip, 2000; Mors et al., 2013).

Spectra of the purified rhodopsin-nanodisc fractions are characterized by three absorbance bands, peaking near 280, 365, and 500 nm (Figure 5). The absorbance at 280 nm represents the total protein population, including rhodopsin and ZAP1. The 365 nm band is due to retinaloxime, generated from hydroxylamine and excess retinal, which became inserted in the lipid phase of nanodiscs. Retinaloxime is not a ligand for opsin and does not affect the functionality of rhodopsin (Wald, 1968; Hubbard et al., 1971). The absorbance band around 500 nm is typical for rhodopsin. Upon illumination in the presence of hydroxylamine it is converted to a band at 365 nm (Figure 6), caused...
by conversion of all photoproducts to retinaloxime (Wald, 1968). We estimated the rhodopsin recovery in the IMAC fractions by spectroscopy. However, this assay has much lower sensitivity and did not detect rhodopsin in the flow-through and wash fractions. Thus, the estimated recovery of rhodopsin in the purified nanodiscs was somewhat lower than that estimated by immunoblot (Table 1, right column).

The ratio A280/A500 in the spectra of the purified rhodopsin-nanodiscs allows us to estimate the efficacy of the purification procedure. For a highly purified rhodopsin-ZAP1-nanodisc preparation, this ratio was reported to be approximately 5 (Banerjee et al., 2008). In our partially purified rhodopsin-ZAP1-nanodisc samples, this ratio varies between 15 and 20, indicating that, as anticipated, we did not yet achieve high-level purification. Nevertheless, considering that in the host membranes recombinant rhodopsin accounts for maximally 0.5% (w/w) of the total MP population, and that MSP has been added in a 2-fold molar ratio to total MP, we estimate that over 95% of contaminating protein has been removed in a single IMAC step. This level of purification is more than sufficient for spectral, as well as functional studies, as we will demonstrate in the next section.

**Rhodopsin-nanodiscs: characterization**

**Dynamic light scattering: size distribution**

For empty ZAP1-nanodiscs, diameters of 10–12 nm have been measured, while protein-containing ZAP1-nanodiscs vary in size between 12 and 17 nm, probably depending on the type of assay and the lipid content (Banerjee et al., 2008). Dynamic light scattering analysis of our purified rhodopsin-nanodisc preparations yielded a bimodal distribution of a major component (>98% in particle number) with a fairly narrow size distribution (17±3 nm) and a minor larger component (82±7 nm; ≤2% in particle number). The size distribution of the smaller component agrees with a single nanodisc, while the larger component probably consists of a more composite build-up (Banerjee et al., 2008; Ritchie et al., 2009; Inagaki et al., 2013).

**Photochemical properties: pH-dependent photolysis**

In this and the following section, we show that our rhodopsin-nanodisc preparation not only is photosensitive, but also fully active in signal transmission. The rationale of this assay is described in the Supplementary Information. The photochemical integrity of rhodopsin in the nanodisc preparation was evaluated by photolysis at two pH-values (pH 6.0 and pH 8.0). The spectral transitions observed in illuminated rhodopsin-nanodiscs at 10°C and pH 6.0 are shown as two difference spectra (Figure 7A, curves 1 and 2). Curve 1 is obtained by subtracting the ‘dark’ spectrum from the spectrum recorded immediately after illumination. Curve 2 is obtained by subtracting the latter spectrum from that recorded 30 min after illumination. Curve 1 shows the immediate disappearance of rhodopsin upon illumination (negative band around 500 nm), together with a slight positive inflection near 480 nm (Meta I) and a strong new band at 380 nm (Meta II). Subsequently, Meta II decays under release of retinal into several products (Figure 7A, curve 2), one of which is represented by an absorbance band peaking around 450 nm (VanBreugel et al., 1979). Thus, the rhodopsin-nanodisc preparation presents the established late-phase photocascade (Matthews et al., 1963; Emelis et al., 1982; Sato et al., 2010; Tsukamoto et al., 2011). The same conclusion can be drawn from a similar experiment at pH 8.0 (Figure 7B), where we expect Meta I and Meta III as the major late photoproducts, (Wald, 1968; Hofmann, 1986; Vogel et al., 2004). Indeed, curve 1 in Figure 7B presents a more red-shifted negative band near 525 nm (photolyzed rhodopsin), because now the photoproducts Meta I (480 nm) and Meta III (470 nm) dominate and there is no evidence for formation of Meta II (380 nm). Subsequently, slow decay of Meta I into Meta III and free retinal (380 nm) is observed (Figure 7B, curve 2).
We next examined whether the Meta II photoproduct in our nanodisc preparation is able to bind its cognate heterotrimeric GTP binding protein, transducin (Gt). The principle again is explained in the Supplementary Information. For convenience, we employed an analog of the C-terminal peptide of the transducin α-subunit (NH₂-VLEDLKSCGLF-COOH) that has high affinity for Meta II (Aris et al., 2001; Sato et al., 2010). Photolysis of the rhodopsin-nanodisc preparation at pH 6.0 in the presence of this peptide clearly enhances the formation of Meta II, and there is no longer evidence for the presence of Meta I (Figure 7C, curve 1). The relative amount of Meta II generated in the absence of peptide varies between 60% and 70% of that in the presence of peptide, which is in good agreement with a pKa of 6.5±0.2 (see Supplementary Information). The peptide does not influence the subsequent decay pattern of Meta II; the same patterns are observed with and without the peptide (curve 2 in Figure 7A and C).

**G-protein interaction: effect of a transducin peptide on rhodopsin photolysis**

We next examined whether the Meta II photoproduct in our nanodisc preparation is able to bind its cognate heterotrimeric GTP binding protein, transducin (Gt). The principle again is explained in the Supplementary Information. For convenience, we employed an analog of the C-terminal peptide of the transducin α-subunit (NH₂-VLEDLKSCGLF-COOH) that has high affinity for Meta II (Aris et al., 2001; Sato et al., 2010). Photolysis of the rhodopsin-nanodisc preparation at pH 6.0 in the presence of this peptide clearly enhances the formation of Meta II, and there is no longer evidence for the presence of Meta I (Figure 7C, curve 1). The relative amount of Meta II generated in the absence of peptide varies between 60% and 70% of that in the presence of peptide, which is in good agreement with a pKa of 6.5±0.2 (see Supplementary Information). The peptide does not influence the subsequent decay pattern of Meta II; the same patterns are observed with and without the peptide (curve 2 in Figure 7A and C).
Even more importantly, photolysis at pH 8.0 in the presence of peptide completely matches the data at pH 6.0 with peptide, that is, single formation of Meta II (Figure 7D, curve 1). Hence, even at pH 8.0 binding of the peptide to Meta II completely shifts the equilibrium to the Meta II side, and the subsequent decay follows the same pattern as that at pH 6.0 (Figure 7D, curve 2).

Discussion

Various procedures have been developed to study membrane proteins in vitro using detergents and/or lipid-based systems. It has gradually become apparent that the natural phospholipid bilayer presents the optimal micro-environment to investigate the physiological function and structure of membrane proteins (Silvius, 1992; Popot, 2010). For such studies, membrane proteins have to be properly purified. However, for unrestricted extraction of membrane proteins from their membrane environment, detergents have proven to be indispensable. Here, we describe an approach to directly transfer the target-protein from the cellular membrane into MSP nanodiscs, employing the natural lipid pool and requiring only short exposure to detergent. The target-protein nanodiscs can then be selectively purified from the resulting total MP-nanodisc population.

We show that our model protein, recombinant histagged rhodopsin, can be transferred directly and with good recovery from host membranes into MSP nanodiscs through brief exposure to detergent in the presence of MSP and some extra lipid, followed by rapid extraction of detergent via β-cyclodextrin inclusion. Exposure to detergent can be limited to as brief as 60 min. This approach, which strongly reduces the risk of detergent-induced irreversible loss of structural and/or functional integrity, is in principle applicable to any recombinant membrane protein.

Furthermore, we demonstrate that, even if both the target-protein and the MSP are his-tagged, extensive purification of the target-protein nanodiscs can be accomplished using a single IMAC step, removing over 95% of contaminating protein, with high recovery of target-protein nanodiscs. This affords a level of purification that is sufficient for spectral and other functional studies. We demonstrate in this way, purified rhodopsin-nanodisc preparation, which upon illumination generates the active photointermediate Meta II, and can transduce the activation signal to its downstream signaling partner.

If required, high-level purification of the target-protein nanodiscs can be realized by a variety of established techniques, such as size-exclusion or ion-exchange chromatography, velocity or density centrifugation, or more dedicated methods like ligand- or immuno-affinity chromatography [e.g., Inagaki et al. (2013)].

Our approach provides a mild and generic platform for rapidly assembling membrane proteins into a soluble unit with maintenance of structural stability and functionality. This will be particularly useful for GPCRs as well as other integral membrane proteins that are relative sensitive to perturbation by detergent action. Here we demonstrate that this approach works successfully for rhodopsin. It remains to be shown that it is equally well applicable to potentially less stable membrane proteins. This, however, is better left to research groups with the required expertise with such proteins and outside the scope of this paper. With this paper, we trust to have laid a foundation for further research into this promising direction.

Materials and methods

Materials

Penicillin and streptomycin were from Gibco-BRL (Breda, The Netherlands). Leupeptin was obtained from MP Biomedicals (Santa Ana, CA, USA), heptakis (2,6-di-O-methyl)-β-cyclodextrin (β-cyclodextrin) and asolectin from Sigma-Aldrich Co. (St. Louis, MO, USA). Grace’s and Insect-Xpress insect media were from Lonza (Breda, The Netherlands); 11-cis retinal was a generous gift from Dr. Rosalie Crouch (Medical University of South Carolina, Charleston, SC, USA) through financial support from NEI; DDM and n-nonyl-β-glucoside (NG) were obtained from Serva Electrophoresis GmbH (Heidelberg, Germany). The plasmid for ZAP1 (Banerjee et al., 2008) was provided by Dr. Thomas Sakmar (The Rockefeller University, New York, NY, USA). The plasmids for MSP1D1 and MSP1E3D1 (Bayburt et al., 2002) were acquired through Addgene (Cambridge, MA, USA). Ni-NTA Superflow beads were purchased from Qiagen Benelux B.V. (Venlo, The Netherlands) and the high affinity analog of the Gtα C-terminal peptide (NH2–VLEDLSCGLF–COOH) from United Peptide Corporation (Chantilly, VA, USA).

Buffer A: 40 mm Tris, 0.3 m NaCl, and 1 mm NaN3 (pH 8.0). Buffer B: 20 mm Bis/Tris propane, 5 mm MgCl2, 1 mm dithioerythritol (DTE), 1 mm benzamidine, and 5 μm leupeptin (pH 7.6). Buffer C: 20 mm Bis/Tris propane, 1 m NaCl, 1 m histidine, and 5 μm leupeptin (pH 7.2). Buffer D: buffer B+100 mm NG.

Construction of recombinant baculovirus

Bovine opsin cDNA was ligated into the transfer vector pAcGP67 (BD Biosciences Clontech, Palo Alto, CA, USA). With the resulting vector recombinant baculovirus was generated, as described (Shirzad-Wasei et al., 2013). For relevant details, see the Supplementary Information.
**Sf9 cell culture and rhodopsin expression**

Amplification of the rhodopsin-baculovirus and large-scale production of rhodopsin were performed as described (Klaassen and DeGrip, 2000; Shirzad-Wasei et al., 2013). Relevant details are given in the Supplementary Information. The culture was infected with rhodopsin-baculovirus at a multiplicity of infection = 0.1. The cells were routinely harvested at 4 days post infection (dpi) and directly processed for regeneration with 11-cis retinal.

**Overexpression and purification of membrane scaffold proteins (MSP)**

ZAPI, MSPID1, and MSP1E3D1 were overexpressed in the BL21(DE3)-Rosetta2 Escherichia coli strain and purified over a Ni-NTA matrix using their N-terminal his-tag as described (Pandit et al., 2011). The yield of purified MSP varied between 60 and 200 mg/l culture. According to immunoblot analysis, the purified ZAPI consisted of two major bands and a minor lower band between 25 and 30 kDa (Figure 2), that all are competent in nanodisc formation (Pandit et al., 2011). The purified MSPID1 and MSP1E3D1 consisted of a single band at ca 25 and ca 30 kDa, respectively. Purified MSPs were concentrated on a 10 K filter (Amicon® Ultra-15; EMD-Millipore, Billerica, MA, USA) and mixed with 1 volume of buffer A containing 5 mM DDM to a final concentration of 10–20 mg/ml.

**Regeneration of opsins with ligand**

At 4 dpi, the infected Sf9 cells were centrifuged by sedimentation (10 min, 3000 g, 4°C) and a membrane suspension was prepared as described (Shirzad-Wasei et al., 2013) in buffer C at a density equivalent to 10⁶ cells/ml (see the Supplementary Information for full details). All subsequent manipulations were performed under dim red light (λ>620 nm, RG620 cut-off filter; Schott, Menden, Germany). The ligand 11-cis retinal was added as a concentrated solution in dimethylformamide (30 μM final concentration) and the suspension was incubated for 1-2 h with constant rotation at RT. Subsequently, hydroxylamine was added from a 1 M solution (pH 7) to 50 mM final concentration. The membrane suspension was then directly used for solubilization of the components and nanodisc assembly.

**Formation of nanodisks**

A membrane suspension, equivalent to 10⁶ cells/ml, containing regenerated rhodopsin, was diluted with buffer C containing 20 mM DDM to a density equivalent to 20×10⁶ cells/ml and mixed properly. After incubation under rotation at 4°C for maximally 1 h, the insoluble material was removed by centrifugation (30 min, 100 000 g, 4°C). Asolectin was dissolved in buffer D and added to the solubilized membrane constituents at a 1:4 (w/w) ratio to total membrane protein, that is, 60 μg per 10⁶ cells. This mixture was incubated for 15 min under constant rotation at 4°C. ZAPI was added in a molar ratio of 2:1 to total membrane protein, that is, 8 nmol of ZAPI per 10⁶ cells harvested, followed by incubation for maximally 30 min at 4°C under constant rotation. Nanodisks were then formed through extraction of the detergent by addition of heptakis-(2,6-di-O-methyl)-β-cyclodextrin (DeGrip et al., 1998). Solid β-cyclodextrin was added in three steps to a final molar ratio of 1:2:1 β-cyclodextrin to detergent. Every addition of β-cyclodextrin was followed by incubation for 15 min at 4°C under constant rotation. Finally, insoluble material was pelleted (100 000 g, 30 min, 4°C). The collected supernatant was directly used for purification or stored at -80°C until further use.

**Purification of rhodopsin-nanodisks**

His-tagged rhodopsin-nanodisks were partially purified by batch-wise chromatography over Ni-NTA agarose beads pre-equilibrated with buffer C containing 0.5 mM histidine (details are given in the Supplementary Information). All eluted fractions were assayed by SDS-PAGE and silver-staining or by quantification via immunoblotting, as described (Shirzad-Wasei et al., 2013). Fractions containing rhodopsin-nanodisks were stored at -80°C.

**UV/Vis spectroscopy and photolysis**

UV/Vis spectroscopy was performed on a Lambda 18 spectrophotometer (Perkin Elmer, Norwalk, CT, USA). To quantify rhodopsin, spectra were recorded before and after illumination (150 W halogen light with fiber optics, 30 s, Schott OG350 cut-off filter) in the presence of 50 mM hydroxylamine to convert all rhodopsin photoproducts into retinaloxime (Hubbard et al., 1971). If functional rhodopsin is present, difference spectra (‘dark’ spectrum subtracted from the spectrum after illumination) show a negative band around 500 nm, representing the photolyzed rhodopsin, and a positive band around 365 nm representing the retinaloxime formed upon illumination.

The late photointermediates Meta I and Meta II (Wald, 1968) were generated by brief illumination (10 s, Schott OG350 cut-off filter) at 10°C using buffer B at pH 8.0 and pH 6.0, respectively. To mimic interaction with the cognate G protein transducin (Gt) that only binds to Meta II, the high affinity analog of the C-terminal peptide of the Gtα-subunit was used (NH₂-VLEDKSCGFL–COOH). This was added to the rhodopsin-nanodisk samples at 20 μM final concentration just before the assay. A ‘dark’ spectrum was recorded and then photolysis was triggered by illumination for 10 s followed by immediately recording the first spectrum after illumination. Subsequently, spectra were recorded every 3 min. After 30 min, hydroxylamine was added to a final concentration of 50 mM and the presence of remaining rhodopsin was checked via 5 min of illumination and a final recording. This protocol was executed under four conditions, namely at pH 6.0 and at pH 8.0, and with and without peptide. In order to clearly display the various transitions, difference spectra were generated by subtracting the ‘dark’ spectrum from the first spectrum after illumination, and the latter from the spectrum recorded 30 min after illumination.

**Dynamic light scattering**

The size distribution in the purified nanodisc samples was analyzed by dynamic light scattering (DLS) using the ZetasizerNanoZS.
(Malvern Instruments Ltd, Malvern, UK), equipped with a 633 nm laser. If required, the rhodopsin-nanodisc samples were concentrated on a 10 k filter (EMD-Millipore). The DLS signals were analyzed by the internal software of the equipment and converted into a size distribution with scatter intensity.

Acknowledgments: We thank Dr. Thomas Sakmar (The Rockefeller University, New York, NY, USA) for providing the plasmid for ZAP1-expression. We acknowledge Dr. Rosalie Crouch (Medical University of South Carolina, Charleston, SC, USA) for a generous gift of 11-cis retinal through financial support from NEI. We thank Dr. Vincent Lemieux (Radboud University, Nijmegen, The Netherlands) for support with the DLS assay. This research was supported by funds to WJD from the Council for Chemical Sciences of the Netherlands Organization for Scientific Research (NWO-CW project 700.54.008).

References


**Supplemental Material:** The online version of this article (DOI: 10.1515/hsz-2015-0100) offers supplementary material, available to authorized users.