Functional solubilization of the β2-adrenoceptor using diisobutylene maleic acid

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Highlights

DIBMA can be used to extract the human β2AR from mammalian cells

DIBMALP-β2AR retains ligand binding ability and shows improved stability

TR-FRET-based ligand binding methods avoid purification for DIBMALP-β2AR characterization

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SUMMARY
The β2-adrenoceptor (β2AR) is a well-established target in asthma and a prototypical G protein-coupled receptor for biophysical studies. Solubilization of membrane proteins has classically involved the use of detergents. However, the detergent environment differs from the native membrane environment and often destabilizes membrane proteins. Use of amphiphilic copolymers is a promising strategy to solubilize membrane proteins within their native lipid environment in the complete absence of detergents. Here we show the isolation of the β2AR in the polymer diisobutylene maleic acid (DIBMA). We demonstrate that β2AR remains functional in the DIBMA lipid particle and shows improved thermal stability compared with the n-dodecyl-β-D-maltopyranoside detergent-solubilized β2AR. This unique method of extracting β2AR offers significant advantages over previous methods routinely employed such as the introduction of thermolabile mutations and the use of detergents, particularly for functional biophysical studies.

INTRODUCTION
G protein-coupled receptors (GPCRs) are the largest family of membrane proteins within the human genome and are responsible for modulating a broad range of hormonal, neurological, and immune responses. It is well established that GPCRs have a large therapeutic potential. Indeed, GPCRs currently represent 34% of all US food and drug administration-approved drugs, with 475 drugs targeting over 100 diverse receptors (Hauser et al., 2017). The β2-adrenoceptor (β2AR) is a rhodopsin-like family GPCR (Schiøth and Fredriksson 2005) and member of the adrenoceptor family, which signals primarily through coupling the heterotrimeric Gs protein. It is a well-established target for asthma and has become one of the most studied GPCRs with several structural (Wacker et al., 2010; Rasmussen et al., 2011; Bang and Choi 2015) and detailed biophysical studies (Manglik et al., 2015; Gregorio et al., 2017) into its activation mechanism.

A prerequisite for completion of biophysical and structural studies is the extraction and isolation of the β2AR from its cellular environment. Classically, this has involved the use of detergents; in the case of the β2AR and other GPCRs, n-dodecyl-β-D-maltopyranoside (DDM) is most often used (Munk et al., 2019). However, it is well established that detergent micelles do not recapitulate the environment of the cell membrane and, as such, protein stability is compromised. Moreover, there is strong evidence that phospholipid composition affects β2AR function (Dawaliby et al., 2016). Cholesterol in particular appears associated with the β2AR in crystal structures (Cherezov et al., 2007) and improves β2AR stability (Zucher et al., 2012) and function (Paila et al., 2011). Multiple studies (Leitz et al., 2006; Whorton et al., 2007) have mimicked the native membrane environment and improved protein stability through reconstitution of membrane proteins in liposomes, amphipols, or synthetic nanodiscs; however, these all require initial use of detergents to extract the membrane protein from the membrane.

Recently, it was discovered that styrene maleic acid (SMA) copolymer directly incorporates into biological membranes and self-assembles into native nanoparticles, known as Styrene Maleic Acid Lipid Particles (SMALPs) (Knowles et al., 2009) (Stroud et al., 2018), avoiding the use of detergents at all stages. This has provided a novel method for the solubilization of membrane proteins with their native receptor-associated phospholipids, although some preferential extraction of native lipids occurs (Barniol-Xicota and Verhelst 2021).
SMA has been used to solubilize a range of membrane proteins (Dorr et al., 2014; Gulati et al., 2014; Sun et al., 2018) including GPCRs (Jamshad et al., 2015; Bada Juarez et al., 2020) for both structural and biophysical studies. Such studies either improved protein stability compared with detergent or have allowed extraction of membrane proteins that were previously unstable in detergents. There is, however, evidence that the conformational flexibility of GPCRs within SMALPs is restricted (Mosslehy et al., 2019; Routledge et al., 2020), therefore differing from the native state of the protein. Furthermore, the high absorbance of SMA copolymer in the far-UV region makes optical spectroscopic studies of membrane proteins that are encapsulated within SMALPs challenging (Gulamhussein et al., 2019).

An alternative to SMA is diisobutylene maleic acid (DIBMA), a copolymer that was developed specifically for the extraction of membrane proteins from the intact bilayer (Oluwole et al., 2017). Compared with SMALPs, DIBMALPs are believed to have only a mild effect on lipid packing, be compatible with optical spectroscopy in the far UV range, and tolerate low millimolar concentrations of divalent cations (Oluwole et al., 2017). This makes DIBMALPs far more amenable for functional biophysical studies. DIBMALPs have been shown to contain lipids of the cell membranes using lipidomic approaches (Barniol-Xicota and Verhelst 2021). Despite the natural polydispersity in length of polymer molecules, DIBMALPs form a monodisperse in size population (Oluwole et al., 2017; Gulamhussein et al., 2019; Gulamhussein et al., 2020). Inclusion of integral membrane proteins such as OmpLA and α-synuclein did not affect their size distribution (Oluwole et al., 2017; Adão et al., 2020).

In this study we demonstrate isolation of the functional β2AR from the mammalian cell membrane using DIBMA, with improved thermal stability compared with conventional detergent-based methods.

**RESULTS**

**Extraction of the β2AR from membranes using DIBMA**

The β2AR was extracted from the membrane of mammalian (T-Rex-293) cells using either 1% DDM or 3% DIBMA (Figure 1A). Figure 1B shows a comparison of the solubilization efficiency of 1% DDM and 3% DIBMA as 90 ± 11% and 32 ± 7%, respectively. Figure 1C shows fluorescence size exclusion chromatography (FSEC) of these β2ARs. Figure 1C shows a peak at 1.6–1.8 mL, roughly 75 kDa, which corresponds to DDM-β2AR or DIBMALP-β2AR. In addition, there was a higher-molecular-weight peak for the DIBMALP-β2AR and two higher-molecular-weight peaks for DDM-β2AR. These peaks are presumed to correspond to protein aggregates. Peaks were confirmed by in-gel fluorescence (Figures S2A and S2B).

**DIBMALP-β2AR retains its pharmacology**

A time-resolved fluorescence resonance energy transfer (TR-FRET)-based ligand binding assay was established to investigate if the β2AR remained functional when extracted from the HEK cell membranes into DIBMALPs. Förster resonance energy transfer (FRET) is the non-radiative transfer of energy from an excited donor fluorophore to a ground state acceptor fluorophore. Energy transfer will only occur when the fluorescent emission spectrum of the donor overlaps with the excitation spectrum of the acceptor fluorophore and these fluorophores are within ~10 nm of each other. In this study, the SNAP tag on the N terminus of the β2AR
was labeled with donor fluorophore terbium cryptate (Lumi4-Tb). Excitation of terbium cryptate using a laser allowed proximity of the β2AR to acceptor fluorophores fluorescent propranolol and the BIODIPY F-L cysteine dye to be quantified for ligand binding and thermostability assays, respectively. The specific labeling of the SNAP tag meant that it was not necessary to purify the β2AR in these studies. TR-FRET is becoming an increasingly used technique for ligand binding studies (Emami-Nemini et al., 2013).

Figure 2 shows saturation binding experiments for the fluorescent antagonist S-propranolol-red-630/650 (F-propranolol) binding membrane-β2AR, DDM-β2AR, and DIBMALP-β2AR. The β2AR retained ligand binding ability when extracted from the membrane using both DDM and using the copolymer DIBMA. These data showed comparable affinities for F-propranolol binding to the β2AR in membranes (pKd = 7.50 ± 0.05), DDM (pKd = 7.10 ± 0.08), and DIBMA pKd = 7.00 ± 0.13, although with slightly reduced affinity in DIBMA compared with membranes (P = 0.02, one-way ANOVA and Tukey’s multiple comparison).

In order to better understand if the conformational state of the receptor or its ability to adopt different states in DIBMALPs was affected we investigated its pharmacology using the full agonist isoprenaline, the antagonist propranolol, and the inverse agonist ICI 118,551 in equilibrium competition binding assays using F-propranolol as the tracer (Figure 3). Increasing concentrations of each competing ligand produced a reduction in the specific binding of F-propranolol to the β2AR in membranes, DDM, and DIBMALPs with largely comparable pKi values (Table 1). The only statistically significant difference was between isoprenaline binding to the β2AR found in membranes versus the DDM-solubilized β2AR (p = 0.03) (one-way ANOVA and Tukey’s post hoc). The slopes of all curves were similar to 1.

**DIBMALP-β2AR shows improved stability**

Next, we investigated the thermostability of the DIBMALP-β2AR using a novel ThermoFRET assay (Figure 4, Table 2). Labeling of the SNAP tag on the N terminus of the receptor with Lumi4-Tb allowed thermostability to be investigated without purifying the receptor. β2AR unfolding was initially measured by quantifying TR-FRET between Lumi4-Tb and BODIPY F-L-Cystine that covalently reacted with cysteines that become exposed as the receptor unfolded (Tippett et al., 2020).
Figure 4B shows the Tm of DDM-solubilized β2AR as 35.2 ± 2.4°C. Ligand-induced shifts in thermostability were seen when the DDM-solubilized β2AR was incubated with F-propranolol (Tm = 37.8 ± 0.4°C, p > 0.05) and cyanopindolol (Tm = 41.9 ± 0.1°C, p = 0.04) (one-way ANOVA and Tukey’s multiple comparison test).

Figure 4A shows the Tm of β2AR in the membrane environment as 62.42 ± 0.1°C. No ligand-induced shift was observed when β2AR membranes were pre-incubated with F-propranolol or cyanopindolol; this suggests the unfolding of the receptor itself is not directly measurable and perhaps that these data show the disintegration of the membrane itself. Figure 4C shows TR-FRET thermostability data for the DIBMALP-β2AR; these data did not fit a Boltzmann sigmoidal curve as the top end of the temperature range did not plateau. No effect on any part of the curve was observed with the addition of F-propranolol or cyanopindolol. Therefore, as was the case in membranes, the observed thermostability changes in DIBMALPs likely reflect the melting of the lipid particles as opposed to the receptor itself.

We then investigated the thermostability of the β2AR by measuring the reduction in TR-FRET binding of F-propranolol over an increasing temperature range (Figure 4D). This gave the Tm values of 60.1 ± 0.6°C for membrane-β2AR (60.1 ± 0.6°C) and DDM-β2AR (36.0 ± 0.6°C) similar to those obtained using BODIPY FL L-Cystine in the presence of F-propranolol. Unpaired two-tailed t tests showed no statistically significant differences between membrane-β2AR + F-propranolol or DDM-β2AR + F-propranolol Tm values obtained using ThermoFRET versus that obtained by measuring the decrease in F-propranolol ligands binding.

Thermostability of DIBMALP-β2AR measured by the decrease in F-propranolol binding gave a curve that could be fitted to a Boltzmann with a Tm value of 46.8 ± 2.1°C. This Tm value is statistically significant from that of membrane-β2AR (p = 0.0002) and DDM-β2AR (p = 0.0009) obtained by the same method (one-way ANOVA and Tukey’s multiple comparison test). Therefore, the DIBMALP-β2AR shows approximately 10°C improved stability over the conventional DDM-β2AR. We also observed differences in the slopes of DIBMALP-β2AR and DDM-β2AR thermostability curves obtained by this method; these were −3.2 and −2.7, respectively. In addition, we investigated the thermostability of another rhodopsin-like GPCR, the adenosine 2A receptor (A2AR), when solubilized into a DIBMALP using fluorescent adenosine receptor antagonist (F-XAC) (Hello Bio, UK). Measuring the reduction in F-XAC bound to A2AR over an increased temperature range gave a Tm value of 44.8°C ± 0.7, which was not statistically significantly different from that of the DIBMALP-β2AR.

Table 1. Ligand binding parameters of different preparations of β2AR

|                | Membranes | DDM          | DIBMA         |
|----------------|-----------|--------------|---------------|
|                | pIC50     | pKi          | Slope         | pIC50     | pKi          | Slope         | pIC50     | pKi          | Slope         |
| Propranolol    | 8.7 ± 0.13| 9.5 ± 0.03   | 1.0 ± 0.02    | 9.0 ± 0.04  | 9.5 ± 0.03   | 1.2 ± 0.04    | 9.1 ± 0.10  | 9.6 ± 0.10   | 0.8 ± 0.30    |
| ICI 118,551    | 8.5 ± 0.10| 9.3 ± 0.15   | 1.1 ± 0.22    | 8.5 ± 0.02  | 8.9 ± 0.10   | 1.0 ± 0.06    | 8.3 ± 0.15  | 9.1 ± 0.06   | 1.3 ± 0.23    |
| Isoprenaline   | 4.7 ± 0.12| 5.5 ± 0.20   | 1.1 ± 0.11    | 5.8 ± 0.06  | 6.3 ± 0.13   | 1.1 ± 0.09    | 5.1 ± 0.18  | 5.8 ± 0.10   | 1.1 ± 0.15    |

Values shown are mean of n = 3 individually fitted curves ± SEM, as determined by TR-FRET competition binding assays.
DISCUSSION

The β2AR has become the prototypical GPCR for understanding GPCR structure and the molecular basis of signaling (Bang and Choi 2015; Gregorio et al., 2017); these studies have all required the use of detergents to extract the β2AR from the plasma membrane. Detergents do not recapitulate the complexity of the native membrane environment and are known to damage membrane proteins. Here, we demonstrate that the polymer DIBMA can be used to extract the β2AR from the plasma membrane, together with its native phospholipids, avoiding the use of detergents at any stage.

Similarly to Gulamhussein et al. (2020), we show that DIBMA can be used to extract GPCRs from cell membranes and that the solubilization efficiency of DIBMA is lower than that of SMA or in our case the detergent DDM. We then used TR-FRET ligand binding studies to show that the β2AR remained functional inside the DIBMALP (Figures 2 and 3). Ligand binding data showed comparable affinity (pKd/Ki) values for the β2AR binding F-propranolol, propranolol, ICI 118,551, and isoprenaline solubilized in DIBMA compared with membranes. Although the difference in pKd values for F-propranolol binding membranes-β2AR (7.5 ± 0.05) and DIBMALP-β2AR (7.0 ± 0.13) was statistically different (P = 0.02), this is only a 3-fold difference,

| Table 2. Thermal stability of different preparations of β2AR |
|---------------------------------------------------------------|
| **Tm (°C) ThermoFRET** | **Tm (°C) F-ligand** |
| Membrane β2AR | 62.4 ± 0.2 | – |
| Membrane β2AR + F-propranolol | 61.6 ± 0.4 | 60.1 ± 0.6a |
| Membrane β2AR + cyanopindolol | 63.0 ± 0.4 | – |
| DDM β2AR | 35.2 ± 2.4 | – |
| DDM β2AR + F-propranolol | 37.8 ± 0.4 | 36.0 ± 0.6a |
| DDM β2AR + cyanopindolol | 41.9 ± 0.1 | – |
| DIBMALP β2AR | – | 46.8 ± 2.1a |
| DIBMALP A2A + F-XAC | – | 44.8 ± 0.7b |

Reported error values are SEM.

αβ2AR thermostability was measured using F-propranolol dissociation.

βDIBMALP-A2A thermostability was measured using F-XAC dissociation.
and the pharmacological importance of this remains to be seen. There was no statistical difference between F-propranolol pKᵢ values for DDM-β₂AR and DIBMALP-β₂AR. All ligand binding curves showed one phase binding and a slope of 1 indicating no co-operativity of ligand binding.

Although the pKᵢ values for different preparations of the receptor were comparable, the signal amplitude obtained for F-propranolol binding DIBMALP-β₂AR in TR-FRET experiments was 3-fold lower than for membranes-β₂AR. This reduction in signal amplitude could be due to an effect of the DIBMA polymer on the TR-FRET, for example, fluorescence quenching. Alternatively, it could reflect that a lower fraction of the ligand binding capable β₂AR receptors are present compared with the amount of Tb3+ labeled receptor molecules. However, it should be noted that the assay window for DDM-β₂AR was higher than that of membranes, whereas it would be expected that less β₂AR is functional, suggesting that the solubilization environment can influence the observed signal amplitude. Although the concentration of β₂AR used in each experimental condition was quantified using 620 nm emission of Lum4-Tb, it was not possible to account for difference in Lum4-Tb quantum yield in the membrane, DDM and DIBMALP environments.

It has been shown that the conformational changes of another class A GPCR, Rhodopsin II, in response to agonist (isoprenaline), antagonist (propranolol), and inverse agonist (ICI 118,551) to be able to ascertain if conformational states of the β₂AR differed in a membrane, DDM micelle, or DIBMALP environment. A substantial increase or decrease in pKᵢ value would demonstrate an increase or decrease in the population of the receptors in the conformational state stabilized by the ligand, and therefore a difference in the conformational landscape of the receptor. As there was no statistically significant difference in pKᵢ values between membrane-β₂AR and DIBMALP-β₂AR, it can be concluded that the DIBMALP-β₂AR represents the native conformational landscape of the β₂AR. The difference in pKᵢ values between DDM-β₂AR (6.3 ± 0.13) and membrane-β₂AR (5.5 ± 0.2) for isoprenaline was statistically significant (p = 0.03); this may indicate a change in the conformational state of β₂AR in the DDM micelle compared with its native conformational state. Propranolol, ICI 118,551, and isoprenaline pKᵢ values obtained in this study are in line with the previous studies that investigated the affinity of these compounds for the β₂AR (Baker 2005; Sykes et al., 2014).

Furthermore, we employed a ThermoFRET-based thermostability assay to investigate the stability of the DIBMALP-β₂AR compared with the DDM-β₂AR. We show that the thermostability of DIBMALP-β₂AR is 10°C higher than that of the DDM-β₂AR. It was not possible to find any thermostability data for the β₂AR in synthetic nanodiscs; however, the only other method to show a similar (11°C) increase in thermostability for β₂AR is that of thermostabilizing mutations (Serrano-Vega and Tate, 2009). Since these mutations also lead to a shift in the β₂AR’s conformational landscape to the antagonist-bound and inactive form, the DIBMALP-β₂AR offers a clear advantage for study of receptor function.

Moreover, thermostability data for DIBMALP-β₂AR using F-propranolol showed a Tm value that was very similar to the Tm of DIBMALP-A₂AR. In addition, no shift in thermostability was observed for DIBMALP-β₂AR preincubated with F-propranolol or the high-affinity antagonist cyanopindolol. It therefore seems likely that this Tm value of ~45°C for DIBMALP-β₂AR corresponds to the melting temperature of DIBMALP. Interestingly, this Tm value of ~45°C is lower than that of ~50°C reported for SMALP-A₂AR extracted from yeast membranes using a radioligand-based thermostability assay but slightly higher than ~42°C A₂AR extracted from mammalian (HEK293) membranes (Jamshad et al., 2015). The Tm of 60.2 ± 0.2°C seen for β₂AR in membranes was also unaffected by the presence of F-propranolol and cyanopindolol. As this Tm of 60.2 ± 0.2°C is statistically significant from that of the DIBMALPs it seems likely that the Tm of ~45°C corresponds to disruption of the protein-lipid-polymer particles, whereas the Tm of 60.2 ± 0.2°C corresponds to the melting or disruption of the membrane itself. We also noted a shallower slope for DIBMALP-β₂AR (~3.2) compared with DDM-β₂AR (~2.7); this broader transition may reflect the more heterogeneous nature of DIBMALPs compared with the detergent micelle.

In summary, here we show the utility of the copolymer DIBMA to solubilize the β₂AR in a functional form. We show that this method offers improved stability over the use of the conventional detergent DDM and has allowed us to maintain the native environment and ligand binding activity of the β₂AR. This could therefore provide an improved solubilization method for structural and biophysical studies. Moreover, we demonstrate this using novel TR-FRET ligand binding-based methods that should allow for easier screening of membrane protein solubilization conditions and anticipate that this approach could be applied to other GPCRs.
Limitations of the study
As discussed above, one limitation of the study is that we cannot be certain about the cause of the decreased signal window in the ligand binding assays for DIBMALP-β2AR versus membrane-β2AR and DDM-β2AR. This could be indicative of less DIBMALP-β2AR being functional compared with in membranes, although there could be a number of other reasons such as decreased quantum yield of either donor or acceptor, as discussed. Furthermore, although we showed that DIBMALP-β2AR retains ligand binding, we have not tested other functionality of the receptor such as its ability to activate G proteins or recruit arrestins. Such studies would require the purification of G proteins, which was beyond the scope of this study.

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SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.103362.

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AUTHOR CONTRIBUTIONS
C.R.H. performed the experiments. C.R.H. wrote the manuscript and it was edited by D.A.S., D.R.P., S.J.B., and D.B.V. D.A.S. gave technical advice and generated reagents. R.U. and D.R.P. gave technical advice. B.L.H. and F.M.H. generated reagents. C.R.H., S.J.B., and D.B.V. conceived the idea. S.J.B. and D.B.V. supervised the project.

DECLARATION OF INTERESTS
D.A.S. and D.B.V. are founding directors of Z7 Biotech Ltd, an early-stage drug discovery company. All other authors declare no conflict of interest.

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Bastos, M. (2020). DIBMA nanodiscs keep low-expressing GPCR from a human cell line. Biochim. Biophys. Acta Biomembr. Acta Biomembr. 1862, 183315.

Bang, I., and Choi, H.J. (2015). Structural features of beta2 adrenergic receptor. crystal structures and beyond. Mol. Cells 38, 105–111.

Barniol-Xicota, M., and Verhelst, S.H.L. (2021). Lipidomic and in-gel analysis of maleic acid copolymer nanodiscs reveals differences in composition of solubilized membranes. Commun. Biol. 4, 218.

Cherezov, V., Rosenbaum, D.M., Hanson, M.A., Rasmussen, S.G., Thian, F.S., Koblika, T.S., Choi, H.J., Kuhn, P., Weiss, W.I., Koblika, B.K., and Stevens, R.C. (2007). High-resolution crystal structure of an engineered human beta2-adrenergic G protein-coupled receptor. Science 318, 1258–1265.

Comeo, E., Kindon, N.D., Soave, M., Stoddart, L.A., Kilpatrick, L.E., Scammells, P.J., Hill, S.J., and Kellam, B. (2009). Subtype-selective fluorescent ligands as pharmacological research tools for the human adenosine A2A receptor. J. Med. Chem. 52, 2656–2672.

Dawaliby, R., Trubbia, C., Delporte, C., Masurel, M., Van Antwerpen, P., Koblika, B.K., and Govaerts, C. (2016). Allosteric regulation of G protein-coupled receptor activity by phospholipids. Nat. Chem. Biol. 12, 35–39.

Dorr, J.M., Koorengelvel, M.C., Schafer, M., Prokopenko, A.V., Scheidelaa, S., van der Cuijsten, E.A., Daftorn, T.R., Baldus, M., and Killian, J.A. (2014). Detergent-free isolation, characterization, and functional reconstitution of a tetrameric K+ channel. The power of native nanodics. Proc. Natl. Acad. Sci. U S A 111, 18607–18612.

Emami-Nemini, A., Roux, T., Leblay, M., Bourrier, E., Lamarque, L., Trinquet, E., and Lohe, M.J. (2013). Time-resolved fluorescence ligand binding for G protein-coupled receptors. Nat. Protoc. 8, 1307–1320.

Gregorio, G.G., Masureel, M., Hilger, D., Terry, D.S., Juette, M., Zhao, H., Zhou, Z., Perez-Aguilar, J.M., Hauge, M., Mathiasen, S., et al. (2017). Single-molecule analysis of ligand efficacy in beta2AR-G protein activation. Nature 547, 68–73.

Gulamhussein, A.A., Uddin, R., Tighe, B.J., Poyner, D.R., and Rothney, A.J. (2020). A comparison of SMA (styrne maleic acid) and DIBMA (di-isobutylene maleic acid) for membrane protein purification. Biochim. Biophys. Acta Biomembr. Acta Biomembr. 1862, 183291.

Gulati, S., Jamshad, M., Knowles, T.J., Morrison, K.A., Downing, R., Cant, N., Collins, R., Koenderink, J.B., Ford, R.C., Overdun, M., et al. (2014). Detergent-free purification of ABC (ATP-binding-cassette) transporters. Biochem. J. 461, 269–278.

Hauser, A.S., Attwood, M.M., Rask-Andersen, M., Schoth, H.B., and Gloriam, D.E. (2017). Trends in GPCR drug discovery: New agents, targets and indications. Nat. Rev. Drug Discov. 16, 829–842.

Heydenreich, F., Miljus, T., Milic, D., and Veprintsev, D. (2020). High-throughput site-directed scanning mutagenesis using a two fragment PCR approach. Bio Protoc. 10.

Heydenreich, F.M., Miljus, T., Jaussi, R., Benoit, R., Milic, D., and Veprintsev, D.B. (2017). High-throughput mutagenesis using a two-fragment PCR approach. Sci. Rep. 7, 6787.

Jamshad, M., Charlton, J., Lin, Y.P., Routledge, S.J., Bawa, Z., Knowles, T.J., Overdun, M., Dekker, N., Daftorn, T.R., Bill, R.M., et al. (2015). G-protein coupled receptor solubilization and purification for biophysical and functional studies, in the total absence of detergent. Biosci. Rep. 35, e00188.

Knowles, T.J., Finka, R., Smith, C., Lin, Y.P., Daftorn, T.R., and Overdun, M. (2009). Membrane proteins solubilized intact in lipid containing nanoparticles bounded by styrene maleic acid copolymer. J. Am. Chem. Soc. 131, 7458–7485.

Leitz, A.J., Bayburt, T.H., Barnakov, A.N., Springer, B.A., and Sigal, S.G. (2006). Functional reconstitution of Beta2-adrenergic receptors utilizing self-assembling nanodisc technology. Biotechniques 40, 601–602, 604, 606, passim.

Manglik, A., Krn, T.H., Masureel, M., Altenbach, C., Yang, Z., Hilger, D., Lerch, M.T., Koblika, T.S., Thian, F.S., Hubbell, W.L., et al. (2015). Structural insights into the dynamic process of beta2-adrenergic receptor signaling. Cell 161, 1101–1111.

Mossley, W., Voskoboykina, N., Collasevici, A., Ricke, A., Klose, D., Klare, J.P., Muklidjanian, A.Y., and Steinhoff, H.J. (2019). Conformational dynamics of sensory rhodopsin II in nanolipoprotein and styrene-maleic acid lipid particles. Photochem. Photobiol. 95, 1195–1204.

Munk, C., Mutt, E., Isberg, V., Nikolajsen, L.F., Bibbe, J.M., Flock, T., Hanson, M.A., Stevens, R.C., Deupi, X., and Gloriam, D.E. (2019). An online resource for GPCR structure determination and analysis. Nat. Methods 16, 151–162.

Muller, D.J. (2012). Cholesterol increases kinetic, energetic, and mechanical stability of the human beta2-adrenergic receptor. Proc. Natl. Acad. Sci. U S A 109, E3463–E3472.
# STAR METHODS

## KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Bacterial and virus strains** | | |
| One Shot™ TOP10 chemically competent E.coli | Invitrogen | C404010 |
| **Chemicals, peptides, and recombinant proteins** | | |
| SNAP-Lumi4-Tb | Cis bio | SSNPTBX |
| n-Dodecyl-β-D-maltopyranoside (DDM) | Anatrace, OH, US | D3010S |
| Diisobutylene Maleic acid (DIBMA) | Anatrace, OH, US | BMA1011 |
| 5% Magstrep ‘type3’ XT magnetic bead suspension | IBA Lifesciences, DE | 2-4090-002 |
| 633/650 S-propranolol-red | CellAura, UK, supplied by Hello Bio, UK | Cat no. HB7817 |
| Fluorescent XAC CA200634 | CellAura, UK, supplied by Hello Bio, UK | Cat no. HB7814 |
| ICI, 118 551 hydrochloride | Selleckchem, US | Cat no. 58114 1217094-53-5 |
| Isoprenaline hydrochloride | Sigma | Cat no. 15627 CAS-51-30-9 |
| (s)-(-)-Propranolol hydrochloride | Tocris, UK | Cat no. 0834 CAS 4199-10-4 |
| Cyanopindolol hemifumarate | Tocris, UK | Cat no. 0993 CAS 69906-86-1 |
| BODIPY™ FL L- Cystine dye | Molecular Probes, US | B20340 |
| NuPAGE™ MOPS SDS running buffer (x20) | Invitrogen | NP0001 |
| **Experimental models: Cell lines** | | |
| T-Rex™-293 cells (parent cell line) | Invitrogen | Cat.no. R71007 |
| Stable cell line T-Rex™-293 expressing TS-SNAP- β2AR plasmid generated in this study | This study | This study, SI |
| Stable cell line T-Rex™-293 expressing TS-SNAP- A2A plasmid generated in this study | This study | This study, SI |
| **Oligonucleotides** | | |
| CMV forward sequencing primer | Genewiz, UK | CGCAAATGGGCGGTAGGCGTG |
| BGH reverse sequencing primer | Genewiz, UK | TAGAAGGACAGTCGAGG |
| **Recombinant DNA** | | |
| TS-SNAP- β2AR in pcDNA4/TO | This study | This study, SI |
| TS-SNAP- A2A in pcDNA4/TO | This study | This study, SI |
| **Software and algorithms** | | |
| GraphPad Prism 8.0.0 | GraphPad software, CA, US | www.graphpad.com |
| PHERAstar v5.41 | BMG, UK | BMG Pherastar FSX platereader |
| **Others** | | |
| Shimadzu Prominence modular HPLC system | Shimadzu, Kyoto, Japan | Prominence modular HPLC system |
| Yarra 1.8um SEC-x300 2.5mL column | Phenomenex, CA, US | 00H-4743-ED-SS |
| TruePage™ Precast Gels 4-20% | Sigma | PCG2008 |
| PHERAstar ® FSX equipped with Time Resolved Fluorescence lasers and module, and TR337/665/620 and TR337/520/620modules | BMG, UK | PHERAstar ® FSX |
| ProxiPlate-384 Plus | PerkinElmer, MA, US | 6008280 |
| OptiPlate-384 | PerkinElmer, MA, US | 6007290 |
| GE Amersham Typhoon™ | GE, US | GE Amersham Typhoon™ |
RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Dmitry Veprintsev (Dmitry.Veprintsev@nottingham.ac.uk).

Materials availability
Plasmids generated in this study are available from the lead contact upon request, subject to the MTA with the University of Nottingham.

Data and code availability
Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

No human subject or animal models were used in this study. The cell lines T-RexTM-293 cells stably expressing pcDNA4TO-TS-SNAP-β2AR or pcDNA4TO-TS-SNAP- A2A were used in this study. These cell lines were maintained in high glucose DMEM (Sigma D6429) with 10% foetal bovine serum (FBS), 5μg/μL blasticidin and 20μg/μL zeocin, at 37°C and 5% CO₂.

METHOD DETAILS

Molecular biology
The construct pcDNA4TO-TwinStrep (TS)-SNAP-β2AR was generated by amplification of the SNAP and β2AR sequences of the pSNAPf-ADRB2 plasmid (NEB) and insertion into pcDNA4TO-TS using Gibson assembly (Heydenreich et al., 2017; Heydenreich et al., 2020). The construct pcDNA4TO-TS-SNAP-A2A was generated by amplifying the A2A receptor from the pDNA3.1 SNAP A2A construct described in (Comeo et al., 2020) and inserting into pcDNA4TO-TS-SNAP vector using Gibson assembly. This therefore gave the construct pcDNA4TO-TS-SNAP-A2A. Both constructs used a signal peptide based on the 5HT3A receptor to increase protein folding and expression.

Transfection and mammalian cell culture
pcDNA4TO-TS-SNAP-β2AR or pcDNA4TO-TS-SNAP-A2A were stably transfected into T-RexTM-293 cells (Invitrogen) using polyethylenimine (PEI). A mixed population stable line was selected by resistance to 5μg/mL blasticidin and 20μg/mL Zeocin. Stable cell lines were maintained in high glucose DMEM (Sigma D6429) with 10% foetal bovine serum (FBS), 5μg/μL blasticidin and 20μg/μL zeocin, at 37°C and 5% CO₂. When ~70% confluent TS-SNAP-β2AR expression was induced with 1μg/mL tetracycline. Cells were left to express for 50hrs before harvesting.

Labelling TS-SNAP-β2AR with terbium cryptate or SNAP-AlexaFluor488 or 647
Media was aspirated from T175 flasks and adherent cells washed twice at room temperature with Phosphate Buffered Saline (PBS). Adherent cells were labelled with 100nM SNAP-Lumi4-Tb labelling reagent in Labmed buffer (both Cisbio, UK) for 1 hr at 37°C and 5% CO₂ or for 30mins with SNAP-AlexaFluor 488 (NEB) in cell culture media. Cells were washed twice more with PBS and detached with 5mL non enzymatic cell dissociation solution (Sigma, UK). Cells were pelleted by centrifugation for 10 min at 1000g, supernatant was removed, and cell pellets frozen at -80°C.

TS-SNAP-β2AR membrane preparation
Cell pellets were thawed on ice and resuspended in 20mL buffer B (10mM HEPES and 10mM EDTA, pH 7.4). Suspensions were homogenised using 6 x 1 sec pulses of a Polytron tissue homogeniser (Werke, Ultra-Turrax). Suspensions were centrifuged at 48,000g and 4°C for 30 min, supernatant was removed and resuspended and centrifuged again as above. Resulting pellets were resuspended in buffer C (10mM HEPES and 0.1mM EDTA, pH 7.4) and frozen at -80°C.

Solubilisation of TS-SNAP-β2AR using DDM or DIBMA
Membranes were incubated with 3% DIBMA (w/v) (Anatrace, UK) in 20mM HEPES, 10% (v/v) glycerol, and 150mM NaCl, pH 8 at room temperature or 1% DDM (w/v), 20mM HEPES, 5% (v/v) glycerol, and 150mM...
NaCl, pH 8 at 4°C for 2-3 h. Samples were clarified by ultracentrifugation at 4°C for 1 hr at 100,000xg for ligand binding assays and 16900xg at 4°C for 45 min for thermostability assays.

**Affinity purification of DDM or DIBMALP TS-SNAP-β2AR**

Solubilised DDM-TS-SNAP-β2AR and DIBMALP-β2AR samples were purified using 20μL of 5% MagStrep “type3” XT magnetic beads suspension (IBA). Beads were prepared by removal of supernatant using magnetic rack and 2x washes in 200μL solubilisation buffers before resuspension in samples. Samples were incubated with beads for 2hrs at 80RPM on a roller in cold room. Supernatant was then removed from beads using magnetic rack and beads were washed twice with wash buffer (20mM HEPES, 10% glycerol, 150mM NaCl, pH 7.5 with 0.1% DDM for DDM sample only), before resuspension in 50μL elution buffer. Elution buffer consisted of 1part 10X buffer BXT (IBA) and 9 parts wash buffer. Elution took place for 2 hours at 80RPM on a roller in cold room. Sample were then separated from beads using magnetic rack.

**Fluorescence size exclusion chromatography (FSEC)**

30μL crude lysate samples of Alexa488-DDM or DIBMALP-β2AR were run of Yarra 1.8μm SEC-x300 2.5mL column (Phenomenex, CA, US) using shimazu prominance HPLC system. Running buffer consisted of 20mM HEPES, 150mM NaCl, 5% glycerol, and 0.03% DDM for DDM-β2AR sample only. FSEC took place at a flow rate of 0.2mL/min and 0.2mL fractions collected. Samples were excited at 488nm, and emission collected at 520nm. GE HMW calibration kit was use as the standard.

**In gel fluorescence**

15μL of each FSEC fraction was run on TruPage™ 4-20% Bis-Tris 17 well gel using NuPage™ LDS sample loading buffer with 5mM Dithiothreitol (DTT) and NuPage™ MOPs running buffer. Gels were run for 50min at 200V. Samples were not boiled prior to gel electrophoresis. 5μL PageRuler™ Prestained Protein Ladder (10-140kDa). was used as the ladder. Gels were scanned on GE Typhoon scanner using Fluorstage and Cy2 or Cy5 filter sets. PMT was set to auto and pixel size to 200μm.

**TR-FRET ligand binding assays**

TR-FRET between the donor Lumi4-Tb and the fluorescent acceptors 633/650 S-propranolol-red (CellAura, UK, supplied by Hello Bio, UK, cat no. HB7817) (F-propranolol) was measured by exciting at 337nm and quantifying emission at 665nm and 620nm using a PheraStar FSX (BMG Labtech) and HTRF 337 665/620 module (BMG Labtech). Assay buffer consisted of 20mM HEPES, 5% glycerol, 150mM NaCl, and 0.5% Bovine Serum Albumin (BSA), pH 8.0 for DDM solubilised samples 0.1% DDM was used. All binding assays used a final concentration of 300μM DMSO. BMD was used to determine non-specific binding (NSB). Receptors were added to plates last, and the plates were incubated at room temperature for 45 mins prior to reading. For competition binding assays 100nM of F-propranolol was used for membrane and DDM samples and 200nM F-propranolol for DIBMA samples.

**ThermoFRET thermostability assays**

Solubilised Lumi4-Tb labelled β2AR was incubated with 10μM BODIPY™ FL L-Cystine dye (Molecular Probes, U.S) with or without 200mM F-propranolol or 100μM cyanopindolol, for 15 mins on ice in 20mM HEPES, 150mM NaCl, 5% glycerol, 0.5% BSA, pH8. For DDM samples 0.1% DDM was used. 20μL samples were added to each well of a 96-well PCR plate and incubated for 30 min over a temperature gradient of 20-78°C across the plate using alpha cycler 2 PCR machine (PCRmax, U.K). Samples were transferred to a 384-well proxiplate (PerkinElmer, US) and 3μM cyanopindolol was used to determine non-specific binding (NSB). Receptors were added to plates last, and the plates were incubated at room temperature for 45 mins prior to reading.

**Data analysis**

**TR-FRET ligand binding data.** Total and NSB for F-propranolol binding to the β2AR was fitted to one-site models in GraphPad Prism 8 according to Equations 1 and 2.

\[
\text{Total binding} = \frac{B_{\text{max}} \times X}{(K_d + X)} + [\text{NS} \times X + \text{background}]
\]  
(Equation 1)
Equation 1

Where:

- \( NS \) = slope of linear nonspecific binding
- \( \text{Background} \) = \( Y \) when \( X \) is 0
- \( B_{\text{max}} \) = the maximum specific binding
- \( K_d \) = the equilibrium dissociation constant
- \( Y \) = specific binding
- \( X \) = concentration of tracer

\[
\text{NSB} = \frac{[NS \times X + \text{background}]}{C_{30}}
\]  
(Equation 2)

Equation 2

Specific binding of F-propranolol to the \( \beta_2 \)AR was fitted to the one site specific binding model in GraphPad Prism 8 according to Equation 3. Final \( K_d \) values were taken as an average of \( K_d \) values from individual specific curve fits.

\[
Y = \frac{B_{\text{max}} \times X}{(K_d + X)}
\]  
(Equation 3)

Equation 3

Where:

- \( Y \) = specific binding
- \( K_d \) = the equilibrium dissociation constant of the labelled ligand

Equilibrium competition binding data was fitted to the One site Fit \( K_i \) model in GraphPad Prism 8 according to Equation 4 and 5. Final \( K_i \) values were taken as an average of individual curve fits.

\[
Y = \frac{(\text{Top} - \text{Bottom})}{\left(1 + 10^{\frac{\text{LogIC}_{50} - X}{\text{LogIC}_{50}}}ight) + \text{Bottom}}
\]  
(Equation 4)

Equation 4

Where:

- \( Y \) = binding of tracer
- \( \text{IC}_{50} \) = the concentration of competing ligand which displaces 50% of radioligand specific binding.

\[
K_i = \frac{\text{IC}_{50}}{1 + \left(\frac{[L]}{K_d}\right)}
\]  
(Equation 5)

Equation 5

Where:

- \( K_i \) = the inhibition constant of the unlabelled ligand
- \([L]\) = concentration of labelled ligand
- \( K_d \) = the equilibrium dissociation constant of the labelled ligand.
ThermoFRET thermostability curves

All ThermoFRET thermostability data from each experiment was fitted to a Boltzmann sigmoidal curve using GraphPad Prism 8 according to Equation 6 to obtain a melting temperature (Tm) value. Final Tm values were taken as an average of Tm values from individual curve fits.

\[
Y = \text{Bottom} + \frac{(\text{Top} - \text{Bottom})}{1 + \exp \left(\frac{Tm - X}{\text{Slope}}\right)}
\]  
(Equation 6)

Where:

\(Y\) = the relative concentration of proteins in the unfolded state

\(X\) = Temperature (°C)

\(Tm\) = The temperature at which half the protein of interest is unfolded

QUANTIFICATION AND STATISTICAL ANALYSIS

Comparison of \(Tm\), \(K_d\) or \(K_i\) values was made using a one-way Analysis Of Variance (ANOVA) test and Tukey’s post hoc multiple comparison test. Statistical comparison of \(Tm\) values obtained with F-propranolol Vs BODIPY™ FL L-Cystine dye was made using an unpaired t test. All statistical analysis was completed in GraphPad Prism 8 and \(p<0.05\) was considered statistically significant.