

HHS Public Access

Author manuscript Nature. Author manuscript; available in PMC 2015 February 14.

Published in final edited form as:

Nature. 2014 August 14; 512(7513): 218-222. doi:10.1038/nature13430.

Visualization of arrestin recruitment by a G Protein-Coupled Receptor

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Author information: VW is deceased. The authors declare no competing financial interests.

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Authors contribution: AKS designed and optimized the procedures for forming and purifying the complex, executed and optimized "on-column" cross-linking protocol and provided the preparations of complex used for EM, HDXMS and cross-link mapping experiments with assistance from PTS. RIR and LYH performed biochemical and pharmacological characterization of the complex. GHW performed EM analysis assisted by MS, ANO and AMD and supervised by GS. KHX performed the HDXMS experiments assisted by SL, JQ, AWK and AB, performed the cross-link mapping experiments assisted by JQ and AWK and designed the disulphide trapping experiments carried out by MC. VLW supervised the initial phase of the HDXMS experiments. CRL, LLG, JMS and XC synthesized the high affinity agonist BI-167107. RH and SSS provided the linker sequence, vector and advice on ScFv conversion and expression. XJY and BUK contributed in assessing various methods of complex formation. PAP provided advice on implementation of ISAC²⁸. SK and AAK provided the phage display library and protocols for Fab selection, expression and purification. BKK conceived the "on-column" cross-linking strategy, advised AKS on its execution and optimization, assisted with molecular modeling of the complex and participated in supervision of the project. GS directly supervised the EM studies, performed the molecular modeling of the complex and supervised overall project execution. RJL supervised overall project design and execution. AKS, GHW, KHX, GS, BKK and RJL participated in data analysis and interpretation. AKS, GHW, KHX, GS, BKK and RJL wrote the manuscript. All authors have seen and commented on the manuscript.

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Abstract

G Protein Coupled Receptors (GPCRs) are critically regulated by β -arrestins (β arrs), which not only desensitize G protein signaling but also initiate a G protein independent wave of signaling¹⁻⁵. A recent surge of structural data on a number of GPCRs, including the β_2 adrenergic receptor $(\beta_2 AR)$ -G protein complex, has provided novel insights into the structural basis of receptor activation⁶⁻¹¹. Lacking however has been complementary information on recruitment of β arrs to activated GPCRs primarily due to challenges in obtaining stable receptor-Barr complexes for structural studies. Here, we devised a strategy for forming and purifying a functional β_2 AR- β_3 arr1 complex that allowed us to visualize its architecture by single particle negative stain electron microscopy (EM) and to characterize the interactions between β_2AR and β_{arr1} using hydrogendeuterium exchange mass spectrometry (HDXMS) and chemical cross-linking. EM 2D averages and 3D reconstructions reveal bimodal binding of $\beta arr1$ to the $\beta_2 AR$, involving two separate sets of interactions, one with the phosphorylated carboxy-terminus of the receptor and the other with its seven-transmembrane core. Areas of reduced HDX together with identification of cross-linked residues suggest engagement of the finger loop of β arr1 with the seven-transmembrane core of the receptor. In contrast, focal areas of increased HDX indicate regions of increased dynamics in both N and C domains of $\beta arr1$ when coupled to the $\beta_2 AR$. A molecular model of the $\beta_2 AR$ - βarr signaling complex was made by docking activated βarr1 and β₂AR crystal structures into the EM map densities with constraints provided by HDXMS and cross-linking, allowing us to obtain valuable insights into the overall architecture of a receptor-arrestin complex. The dynamic and structural information presented herein provides a framework for better understanding the basis of GPCR regulation by arrestins.

To facilitate the isolation of a stable β_2 AR- β_{arr} complex, we utilized a modified β_2 AR construct with its carboxy-terminus replaced by that of the arginine vasopressin type 2 receptor (AVPR₂). This chimeric receptor (β_2 V₂R) maintains pharmacological properties identical to the β_2 AR, but it binds β_{arrs} with higher affinity compared to wild-type β_2 AR¹². We co-expressed the β_2 V₂R, β_{arr1} (1-393) and GRK2^{CAAX} in insect cells followed by agonist stimulation and affinity purification through the FLAG-tagged receptor (Fig. 1a) However, since the isolation of a stable complex was still not feasible (Fig. 1b, lanes1,2), we explored enhancing its stability by adding Fab30, an antibody fragment that we previously reported to selectively recognize and stabilize the active conformation of β_{arr1}^{13} . Indeed, incubation of He β_2 V₂R- β_{arr1} complex (Fig. 1b, lanes 5,6), whereas a non-specific Fab (referred to as Fab1) did not support complex stabilization (Fig.1b, lanes 3,4). Complex

isolation was only possible in response to an agonist (BI-167107) and not an inverse agonist (ICI-118551) (Fig. 1b, lanes 5,6). Furthermore, the efficiency of complex purification using this approach directly mirrors the pharmacological efficacy of the ligand used to stimulate the cells (Fig. 1c). While stimulation of cells with inverse agonists does not yield significant co-purification of β arr1, agonists robustly stabilize the complex and partial agonists yield co-purification of βarr1 at moderate levels. Moreover, the efficiency of complex formation also corresponds to the ligand occupancy of the receptor as reflected by the increasing amount of β arr1 co-purification with increasing agonist concentrations (Extended Data Fig. 1a,b). The direct correlation of ligand efficacy and occupancy with purification efficiency reflects that this approach yields a complex that depends on both activated receptor conformation and receptor phosphorylation. The purified $\beta_2 V_2 R$ - $\beta_3 rr 1$ -Fab30 complex also exhibited robust interaction with purified clathrin terminal domain compared to βarr1 alone, suggesting that Barr1 in this complex is in a physiologically relevant and functional conformation (Extended Data Fig. 2)¹⁴⁻¹⁶. Importantly, this strategy allowed preparative scale purification of a highly stable $\beta_2 V_2 R$ - $\beta arr1$ -Fab30 complex as assessed by analytical size exclusion chromatography (Fig. 1a, last panel, green trace and Extended Data Fig. 1c). In addition to the Fab30 stabilized $\beta_2 V_2 R$ - $\beta arr1$ complex, we were also able to obtain equally stable $\beta_2 V_2 R$ - $\beta_3 rr1$ complexes using the single chain variable fragment of Fab30 (ScFv30) (Fig. 1a, last panel, blue trace).

The interaction of β arrs with activated GPCRs is proposed to involve two sequential steps¹⁷. First, the phosphorylated carboxy-terminus of activated GPCRs is thought to engage the N-domain of β arrs, a high affinity charge-charge interaction primarily mediated between the phosphates on the receptor tail and basic residues on β arrs^{13,17}. This first engagement is hypothesized to facilitate activating conformational changes in β arr, leading in turn to additional interactions with the transmembrane core of the receptor¹⁷. To obtain dynamic structural information on the receptor- β arr complex, we carried out hydrogen-deuterium exchange mass spectrometry (HDXMS) analysis on the purified assembly^{18,19}. In addition to the $\beta_2 V_2 R$ - β arr1-Fab30 complex, we used the arginine vasopressin type 2 receptor carboxy-terminal phosphopeptide (V₂Rpp)- β arr1-Fab30 complex as a reference to extract specific information about the core interaction between the receptor and β arr1.

We observed a reduction in the hydrogen-deuterium exchange rate in the three major loops, the finger loop (55%), the middle loop (16%) and lariat loop (23%) of β arr1 when we compared the HDXMS profile of the $\beta_2 V_2 R$ - β arr1-Fab30 complex with that of $V_2 R p$ - β arr1-Fab30 complex (Fig. 2 and Extended Data Figure 3a). Thus, these regions, and especially the finger loop, are likely to be buried (or have reduced solvent exposure) in the $\beta_2 V_2 R$ - β arr1-Fab30 complex, most likely through an intricate engagement with the transmembrane receptor core. This finding is consistent with previous EPR studies on rhodopsin-arrestin interactions, which revealed a crucial involvement of the finger loop of arrestin with the core of rhodopsin^{17,20-22}. Interestingly, several regions in both the N- and the C-domains of β arr1, in contrast, reveal enhanced hydrogen-deuterium exchange rates indicating that they become more dynamic upon interaction of β arr with the agonist bound phosphorylated receptor. This observation suggests that the core interaction between β arr1 and $\beta_2 V_2 R$ likely has long-range effects in β arr1 structure. Previous studies mapping interactions between GPCRs and arrestins suggested that receptors may also interact with

the broad concave surfaces of the N- and C-domains of arrestins^{21,23-25}. However, peptides representing these surfaces are not fully represented in our HDXMS studies, thus limiting our ability to detect these interactions. We also note that our previously published high affinity agonist radioligand binding data on the T4L- β_2 V₂R- β_{arr1} -Fab30 complex in membranes, which provides a readout of the fully engaged β_{arr} conformation, suggested that approximately 32% of the receptor is in a high affinity agonist binding state¹³. This indicates that our HDXMS data represents an average of two mixed complex populations, one with fully engaged β_{arr1} with the receptor and the other displaying partially engaged β_{arr1} .

Our previous crystal structure of V₂Rpp bound activated βarr1 revealed a significant repositioning of the finger loop compared to the inactive βarr1, presumably primed to engage with the transmembrane core of the activated receptor¹³. To test this we carried out mass-spectrometry based mapping of the T4L- $\beta_2 V_2 R$ - β_{arr1} interface using the homobifunctional, primary amine reactive chemical cross-linker DiSuccinimidyl Adipate (DSA). We found that Lys^{77} on $\beta arr1$ (towards the distal end of the finger loop) cross-links with Lys²³⁵ in the 3rd intracellular loop of the β_2AR (Extended Data Fig. 3b-e). These findings are in line with previously published biochemical and biophysical data suggesting an intricate interaction of the receptor core and the finger loop in arrestins. As an additional control for the close proximity of these residues, we created a series of mutants with single cysteine substitutions around Lys²³⁵ in the amino terminal end of the 3rd intracellular loop of the $\beta_2 V_2 R$ (amino acids 231-236) and in the finger loop around Lys⁷⁷ of β arr1 (amino acids 75-79) and evaluated the formation of disulphide trapped complexes in pairs of receptor and ßarr1 mutants. Consistent with our chemical cross-linking data, cysteines engineered at position 235 of the receptor and at position 78 in βarr1 yielded the most robust disulphide trapped complex, suggesting a close proximity of these two residues in the complex (Extended Data Fig. 4). Taken together these findings demonstrate a direct interaction of the finger loop with the receptor core.

We next employed single particle EM to examine the architecture and conformational dynamics of $\beta_2 V_2 R$ - $\beta arr1$ complexes. Due to the asymmetric nature and small size of these complexes (~150 kDa and ~125 kDa for the Fab and ScFv complexes, respectively) characterization attempts with cryo-EM were not successful and we thus applied negative stain EM that provides adequate contrast for alignment of small particle projections. This approach also enabled a direct comparison with our earlier negative stain EM analysis of the $\beta_2 AR$ -Gas protein complex⁹. As in that work, here we used a T4 lysozyme fusion at the N-terminus of the receptor (referred as T4L- $\beta_2 V_2 R$) to provide a marker for the receptor orientation⁹. The negative stain EM visualization showed a monodisperse particle population (Fig. 3a, Extended Data Fig. 5) and we applied reference-free alignment and classification to obtain 2D averages of the complex.

The majority of averages of the $\beta_2 V_2 R$ - $\beta arr1$ -Fab30 complex revealed distinct projection profiles of an ovoid density, attributed to the receptor in partially flattened detergent micelle, with an attached T-like density attributed to the Fab30- $\beta arr1$ complex (Fig. 3b, Extended Data Fig. 6a). Comparisons with averages of the $\beta_2 V_2 R$ - $\beta arr1$ -ScFv30 complex identify the Fab30 density engaging the middle of $\beta arr1$, in agreement with our recent crystal structure

of β arr1-Fab30 co-crystallized with the V₂Rpp (Fig. 3b, Extended data Fig. 6b). In this conformation β arr1 appears to hang off the receptor via a single point interaction presumably involving only the flexible V₂Rpp fused on β_2 AR. The flexible nature of this interaction is further supported by the variable receptor orientation in these averages, as judged by the T4L domain positioning. It is possible that the "hanging" arrestin conformation based on the V₂Rpp- β arr1 interaction represents a transient intermediate step in the recruitment process that has been stabilized by Fab30. Strikingly, we also observe a significant number of class averages representing ~37% of particles where β arr1 forms a much more extensive interface with the receptor, employing roughly the opposite face of the Fab30 binding region (Fig. 3b, lower panel). The observed fraction of particles displaying the extensive interface is in agreement with our previous radioligand binding results on the T4L- β_2 V₂R- β arr1-Fab30 complex in membranes, which suggested approximately 32% of the receptor in a high affinity agonist binding state¹³. This observation also raised the possibility that β arr1 fully engages the receptor through a second set of weak interactions.

To stabilize this weak interaction, we developed an approach whereby the $\beta_2 V_2 R$ - β_{arr1} -Fab30/ScFv30 complex is cross-linked by exposure to a glutaraldehyde containing buffer zone while migrating through a size exclusion column (Extended Data Fig. 7a). This method facilitated near complete cross-linking of preformed complexes at relatively high concentrations and simultaneously enabled the isolation of highly monodisperse sample (Extended Data Fig. 7b,c, 8 and 9).

EM classification and averaging of the cross-linked $\beta_2 V_2 R$ - β_{arr1} -Fab30/ScFv30 complexes revealed distinct views of a uniform particle architecture, suggesting that cross-linking stabilized a single complex conformer (Fig. 3c). More importantly, the averages show that arrestin interacts extensively with the receptor in a configuration that appears very similar to the one observed in the smaller fraction ($\sim 37\%$) of the native complex. The conformational stabilizing action of the cross-linking is also evidenced by the consistent position of the T4L projection profile, in contrast to its variable positioning observed in averages of the native complex. To better characterize the $\beta_2 V_2 R$ - β_{arr1} assembly, we employed the random conical-tilt approach²⁶ to calculate low resolution 3D maps (~29 Å) from selected classes of the cross-linked complex (Extended Data Fig. 10). The 3D reconstructions show distinct densities for the main complex components, in full agreement with our domain assignment in the 2D projections averages (Fig. 4a, Extended Data Fig. 10). The receptor containing region appears ovoid due to the large micelle "belt" characteristic of the LMNG detergent, as we also observed in the case of the β_2 AR-Gas complex⁹. A protrusion on one end of the receptor-micelle globular density represents the T4L domain that marks the receptor extracellular region. On the opposite side, the ßarr1 density lies longitudinally on the receptor, engaging roughly the opposite side of the Fab30 interacting region. In this configuration, both β arr domains appear to engage the receptor but one of the domains lies mostly outside the interacting zone.

The HDXMS, chemical cross-linking and disulphide trapping data allowed us to constrain the modeling of the T4L- β_2 AR and β_{arr1} -Fab30 crystal structures within the density of the EM 3D maps and generate a low-resolution model for the overall conformation of the β_2 AR- β_{arr1} complex (Fig. 4b). This model can accommodate limited rotations and translations of

the individual crystal structures, which are also expected to undergo conformational changes upon complex formation. Lys⁷⁷ of β arr1 in our model is placed in close proximity to β_2 AR Lys²³⁵ which is located at the end of a helical extension of TM5 in the β_2 AR-G α_s complex¹⁰. This prompted us to use this structure to model the β_2 AR- β arr1 complex. In our model, β arr1 forms an extensive interface with the receptor through its N-terminal domain involving interactions with the phosphorylated receptor tail and the insertion of the finger loop directly in the receptor core, involving the space between TM 3,5 and 6. We note that the finger loop insertion is likely associated with outward shifts in the positioning of TM helices 3, 5 and 6 and also helix 8. The middle and lariat loops of β arr1 are not participating in major interactions but reside close to the interface as suggested by their modest reduction in hydrogen-deuterium exchange rates observed by HDXMS (Fig. 2c). The relative positioning of these loops is also in agreement with previous EPR studies on visual arrestin in complex with activated and phosphorylated rhodopsin^{20,21}.

In regards to β_2AR , TM5 and the 3rd intracellular loop in this model locate above the concave β -sheet region of the N-terminal domain of β arr1. The placement of these receptor elements implies that the N-terminus of the V₂Rpp cannot be in the position observed in the crystal structure of V₂Rpp- β arr1-Fab30¹³, suggesting that the V₂R carboxy-terminus in the β_2V_2R chimeric receptor is mobile and repositions itself significantly upon β arr1 interaction with the receptor core. In contrast to the N-terminal domain, the C-terminal domain of β arr1 lies mostly outside the interaction zone, apart from the loop of residues 242-246 that is in interacting distance from the short α -helical segment connecting TM3 and TM4 of β_2V_2R . This observation is intriguing considering that mutation of the residues distal to the DRY motif (at the end of TM3) have been reported to directly affect β arr recruitment for a number of GPCRs including the β_2AR^{27} .

Our results suggest that arrestin likely employs a biphasic mechanism to engage the receptor (Fig. 4c). The first phase involves an interaction between the phosphorylated C-terminal tail of the receptor and the N-terminal domain of arrestin. Given the flexibility and the length of the C-terminal receptor tail, it is expected to act like a fishing line sampling a wide interaction space at a high rate. The second point of interaction appears weak and involves primarily the insertion of the finger loop within the receptor core, resulting in a longitudinal arrangement of arrestin on the receptor (Fig 4a,c). This arrangement would most certainly preclude GPCR engagement of G-protein heterotrimers, thereby blocking classical GPCR signaling and inducing desensitization. While it is not yet clear whether the single point interaction resulting in a "hanging" arrestin configuration has other physiological functions, it seems possible that these might involve recruitment and complex formation with components of the receptor endocytosis and signaling machinery such as clathrin and G $\beta\gamma$.

Online methods

Reagents

Insect cell culture media and transfection kits to generate various virus stocks were purchased from Expression systems. Superdex 200 SEC columns were purchased from GE healthcare and detergents were purchased from Anatrace. EM grade glutaraldehyde solution and other general chemicals were purchased from Sigma.

Protein expression and purification

T4L-β₂V₂R plasmid (with N-terminal fusion of T4 lysozyme) was generated by modifying the previously described T4L- $\beta_2 AR^6$. T4L- $\beta_2 V_2 R$ was co-expressed with untagged bovine βarr1 (1-393) and untagged GRK2^{CAAX} in Sf9 cells. 60-66h post-infection, cells were stimulated for 30 min (or indicated time) at 37°C with indicated ligand to induce receptor phosphorylation and ternary complex formation. Subsequently, cells were harvested, lysed in 20mM Hepes, pH 7.4, 150mM NaCl, 100nM-1uM ligand and protease inhibitors by douncing and purified Fab30/ScFv30 was added to this cell lysate. Following 1h incubation at room temperature, proteins were solubilized by adding detergent (0.5% LMNG or 1% DDM) and gentle stirring at room temperature for 1h. Solubilized material was purified on a FLAG M1 resin essentially using the protocol described for $\beta_2 AR$ purification previously¹¹. In order to avoid potential protein aggregation at high concentration, free cysteines in the eluted proteins were blocked using iodoacetamide as described previously¹¹. Proteins were concentrated using a 100 kDa Vivaspin or Millipore concentrator and loaded on to a preparative size exclusion column (Superdex 200, 16/600). The column was run at 0.3 ml/min and the column running buffer consists of 20mM Hepes, pH 7.4, 150mM NaCl, 100nM BI-167107, 0.01% LMNG, 100µM TCEP. Elution fractions corresponding to the complex peak were either pooled or used individually for further studies. Selection, characterization and purification of this Fab30 has been described previously¹³.

Biochemical and functional characterization of the complex

In order to pharmacologically validate this novel procedure, we stimulated the cells with multiple ligands (10 μ M) of varying efficacies (ranging from inverse agonists, partial agonists to full agonists). Subsequently, the cells were lysed, incubated with Fab30 and the complex was purified using FLAG M1 affinity resin. Purification of complex was assessed by the extent of β arr1 co-purified with the receptor. In order to further investigate agonist dependence of the complex formation, we assessed whether formation of this ternary complex through Fab30 stabilization follows ligand occupancy of the receptor. Cells were stimulated with varying doses of the agonist BI-167107 or Inverse agonist ICI-118551 at 37°C for 30 min. Subsequently, purification of the complex was performed as described above. Similarly, a time course of agonist and inverse agonist stimulation was carried out to identify the optimal time point to achieve maximal complex formation.

For stability analysis, the purified T4L- β_2 V₂R- β arr1-Fab30 complex (1mg/mL) was either analyzed fresh after purification (reference sample, day 0) or incubated in the cold room and at room temperature for 4 days. Subsequently, the samples were analyzed by analytical size exclusion chromatography using a Superdex 200 (10/300) column. The column was run at 0.3 ml/min and the column running buffer consists of 20mM Hepes, pH 7.4, 150mM NaCl, 100nM BI-167,107, 0.01% LMNG, 100µM TCEP.

In order to validate the active conformation of β arr1 in the T4L- β_2 V₂R- β arr1-Fab30 complex, we evaluated the ability of this complex to interact with purified clathrin terminal domain (heavy chain). The GST-clathrin-TD expression plasmid was a kind gift from Prof. Jeffrey Benovic and Dr. DongSoo Kang (Thomas Jefferson University). GST-clathrin-TD was expressed in *E. coli* using a previously published procedure²⁹ and immobilized on

glutathione sepharose beads. Subsequently, either the purified complex (in 20mM Hepes, 150mM NaCl, 0.01% LMNG, 100nM BI) or purified βarr1 (1-393) was added to the clathrin-TD and incubated at room temperature for 1h. The amount of complex used in this experiment was normalized to contain equivalent amount of βarr1 as in βarr1 alone sample in order to directly compare the extent of clathrin interaction. Subsequently, the beads were washed 3 times with the same buffer as above and proteins were eluted using the SDS-gel loading buffer. The samples were analyzed by Western blotting using anti-βarr1 antibody and anti-clathrin antibody (BD Biosciences).

Conversion of Fab30 in to ScFv30

In order to convert the Fab30 in to an ScFv, the variable domains of the light chain and the heavy chain were joined through a flexible linker (GTTAASGSSGGSSSGA) using standard PCR amplification and cloning techniques. The resulting sequence of the ScFv30 is as follows (linker sequence is underlined and highlighted in bold):

MKKNIAFLLASMFVFSIATNAYASDIQMTQSPSSLSASVGDRVTITCRASQSVSSAV AW

YQQKPGKAPKLLIYSASSLYSGVPSRFSGSRSGTDFTLTISSLQPEDFATYYCQQYKY V

 $\mathsf{PVTFGQGTKVEIK} \underline{\mathbf{GTTAASGSSGGSSSGA}} \mathsf{EVQLVESGGGLVQPGGSLRLSCAASGF} \\ \mathsf{N}$

VYSSSIHWVRQAPGKGLEWVASISSYYGYTYYADSVKGRFTISADTSKNTAYLQMN SX RAEDTAVYYCARSRQFWYSGLDYWGQGTLVTVSSAHHHHHH.

ScFv30 was expressed using the same protocol as Fab30. It was either purified on Ni-NTA resin or used directly as periplasmic extract to stabilize the T4L- β_2 V₂R- β_3 rr1 complex.

Glutaraldehyde cross-linking of the pre-formed complex

In order to mildly cross-link the complex, an "on-column" cross-linking method was used. First, a bolus of glutaraldehyde was injected to a pre-equilibrated Superdex 200 (10/300 global in 20mM Hepes, pH 7.4, 150mM NaCl, 0.01% LMNG, 100nM BI-167107) column and run at 0.25 ml/min for 20 min (i.e. a total of 5ml buffer). Subsequently, the column flow was paused, and the injection loop was flushed using buffer followed by injection of purified complex (200µl volume, at 5-10 µM concentration). Subsequently, the column was run at 0.25 ml/min and 0.3 ml fractions were collected. Cross-linking efficiency was visualized by running the individual fractions on a 4-20% SDS gel, followed by staining with "SimplyBlue" stain. Various concentrations of glutaraldehyde were tested in order to identify an optimal concentration (0.25% in this case) that yields maximal cross-linking efficiency and minimal protein aggregation.

Hydrogen-Deuterium Exchange Mass Spectrometry (HDXMS)

Fragmentation tuning for βarr1

We first performed fragmentation tuning experiments in order to optimize experimental conditions to achieve the best peptide fragmentation pattern for $\beta arr1$. In brief, 3 μL of $\beta arr1$

protein solution was diluted with 9 μ L of a buffer containing 150mM NaCl, 8.3mM Tris-HCl, pH7.2 in H₂O and then divided into equal sized aliquots. Each aliquot was mixed individually with 18 μ L of a series of quench solutions containing 0.8% formic acid, 16.6% glycerol, and various concentrations of GuHCl (ranging from 0.05M to 4M) and TCEP (ranging from 0.015M to 0.5M). The protein-buffer-quench solution mixtures were incubated at 0°C for 1 minute, frozen on dry ice and then stored at -80 °C until subjected to on-line pepsin digestion and LC/MS analysis. Protein fragmentation maps under different conditions were generated and compared. The condition that produced the best peptide fragmentation pattern (the best coverage along the amino acid sequence, the most number of fragments, and highest number of high quality peptides) was used for all the HDXMS experiments described in this study.

Hydrogen-deuterium exchange experiments

The V₂Rpp- β arr1-Fab30 and T4L- β ₂V₂R- β arr1-Fab30 complexes were prepared as described above. Free β arr1 protein was used as a control. For each protein complex, three sets of samples were prepared: 1) non-deuterated (ND), 2) fully deuterated (FD), and 3) time-dependent on-exchange samples. A protein: buffer: quench solution ratio of 1:3:6 (volume) was used for all sample preparations. The FD sample sets were prepared by mixing protein samples with D₂O buffer (0.8% Formic acid in 100% D₂O) and incubated at room temperature for 12 hours before quenching. The ND sample sets were prepared using a similar procedure with H₂O buffer (150mM NaCl, 8.3 mM Tris-HCl, pH7.2 in H₂O) without the incubation step. The on-exchange sample sets were prepared by adding 3 volumes of D₂O buffer (150mM NaCl, 8.3 mM Tris-HCl, pH7.2 in D₂O) at 0°C and incubating for varying time-points (10s, 10²s, 10³s, 10⁴s and 10⁵s). Six volumes of ice-cold quench solution was then added to each sample, followed by snap-freezing on dry ice and stored at -80°C.

On-line pepsin digestion, LC/MS analysis and data processing

Samples prepared above were thawed right before the on-line pepsin digestion at 0° C using a cryogenic autosampler and immediately passed over an immobilized porcine pepsin column (16 μ L bed volume). Peptide fragments were collected contemporaneously on a C₁₈ trap column desalting, separated by a Magic C₁₈AQ column using a linear gradient of acetonitrile from 6.4% to 38.4% over 30 min, and followed by LC/MS analysis using an Orbitrap Elite mass spectrometer (Thermo Scientific). Both MS1 and MS2 spectra were collected using data-dependent acquisition mode. Peptide identification was performed using LC/MS data sets collected from ND samples and SEQUEST database search engine using Proteome Discoverer (Thermo Fisher Sci). The SEQUEST database search results were submitted to DXMS Explorer (Version 2.0, Sierra Analytics Inc.), filtered using several threshold parameters to create an initial peptide pool. The quality of the MS1 data for each filtered peptide was then checked by assigning an initial quality score by DXMS Explorer software, followed by a quality control process which included manual investigation of peak isotopic envelope and adjusting/improving the quality score. Only peptides with high quality in the MS1 spectra were kept in the final peptide pool. The retention times and m/z ranges of each peptide from the final peptide pool was manually verified and adjusted across all LC/MS data sets from on-exchange samples and FD samples

to ensure that DXMS Explorer had selected the correct peptide for all experiments. Results from FD samples were used to monitor the back-exchange rates during on-line pepsin digestion and LC/MS analysis. The centroids of isotopic envelopes of non-deuterated, partially deuterated and fully deuterated peptides were measured using DXMS Explorer, and then converted to deuteration level with corrections for back-exchange. A deuterium accumulation plot was created for each peptide as a further quality check and data refinement process. Rainbow maps were generated and DXMS data comparison was performed using different macros in Excel.

Chemical cross-linking and mass spectrometry

Chemical cross-linking reaction

In order to identify a potential interaction interface of the $\beta_2 V_2 R$ and β_{arr1} , we performed chemical cross-linking experiments on the pre-formed T4L- $\beta_2 V_2 R$ - β_{arr1} -Fab30 complex. To facilitate the identification of cross-linked peptides, we used an equimolar mixture of light ($^{12}C6$) and heavy ($^{13}C6$) DiSuccinimidyl Adipate (DSA, a homobifunctional amine-reactive cross-linker, spacer arm=7.7 angstrom) to cross-link the complex. The cross-linked peptides were characterized with "doublet" peak signatures in mass spectra. The T4L- $\beta_2 V_2 R$ - β_{arr1} -Fab30 complex was prepared in buffer containing 20mM HEPES, 150mM NaCl, 0.01% LMNG, 100nM BI, 100uM TCEP. Cross-linker solution was freshly prepared in DMSO at a concentration of 10 mM. For the cross-linking reaction, 100-fold excess of DSA- $^{12}C6$:DSA- $^{13}C6$ equimolar mixture was added to the T4L- $\beta_2 V_2 R$ - β_{arr1} -Fab30 complex solution and incubated for 30 min at 25 °C. Unreacted cross-linker was quenched by incubation with 100 mM ammonium bicarbonate for 20 min. The cross-linked T4L- $\beta_2 V_2 R$ - β_{arr1} -Fab30 complex was separated by SDS-PAGE, and the corresponding protein band was in-gel digested overnight at 37 °C with trypsin at a final concentration of 10 ng/µL and subjected to LC/MS analysis as described below.

LC/MS analysis and cross-linked peptide identification

LC/MS analyses were performed on a LTQ Orbitrap XL mass spectrometer (Thermo Scientific) with a Finnigan Nanospray II electrospray ionization source. Tryptic peptides were injected onto a 75 μ m×150 mm BEH C18 column (particle size 1.7 μ m, Waters) and separated using a Waters nano ACOUITY Ultra Performance LCTM (UPLCTM) System (Waters, Milford, MA). The LTQ Orbitrap XL was operated in the data dependent mode using the TOP10 strategy³⁰. In brief, each scan cycle was initiated with a full MS scan of high mass accuracy 375–1800 m/z; acquired in the Orbitrap XL at 6×10^4 resolution setting and automatic gain control (AGC) target of 10⁶, which was followed by MS/MS scans (AGC target 5000; threshold 3000) in the linear ion trap on the 10 most abundant precursor ions. Selected ions were dynamically excluded for 30 s. Singly charged ions were excluded from MS/MS analysis. The LC/MS/MS data was processed and analyzed by pLink software³¹. In brief, acquired raw data from LTQ Orbitrap XL mass spectrometer was first converted to mgf files using Mascot Distiller or Proteowizard 3.0 and saved under a pLink master directory, where a composite database containing protein sequences of T4L- β_2 V₂R, βarr1 and Fab30 in FASTA format is also stored. In the pLink configuration file, both the names and paths of the mgf and database file were indicated correspondingly. The total

numbers of fixed modifications was set to 1 for cysteine carbamidomethylation. The name and number of cross-linkers used in the experiment were indicated as well in the configuration file. The filter for precursor mass accuracy was set to \pm 10 ppm. After pLink analysis, identified cross-linked peptides were manually inspected in the raw file for the appearance of a pair of doublet MS peaks.

Disulphide trapping of $\beta_2 V_2 R$ and β_{arr1} complex

Cysteine mutants of the $\beta_2 V_2 R$ and $\beta arr1$ were co-expresed in HEK-293 cells. 48h posttransfection, cells were stimulated with $\beta_2 AR$ agonist Isoproterenol (10µM) and treated with H₂O₂ (1mM) at different time points to induce the formation of disulfide bond and trap the complex. Cells were washed and lysed in lysis buffer (50mM Hepes, 250mM NaCl, pH 7.4, 2 mM EDTA, 10% glycerol, 0.5% NP40, 1mM NaV, 57mM NaF, EDTA free Complete protease inhibitor). Cell lysates were used for anti-FLAG antibody immunoprecipitation (Nterminal FLAG tagged $\beta_2 V_2$). Beads were washed and eluted proteins were separated by SDS-PAGE followed by detection using Western blotting. Expression levels of the receptor and β arr1 mutants were measured by radioligand binding and Western blotting, respectively. Densitometry analysis of the β arr1 bands was done using ImageJ software.

Specimen preparation and EM imaging of negative stained samples

T4L- β_2 V₂R- β_{arr1} -Fab30/ScFv30 complex was prepared for electron microscopy using the conventional negative staining protocol³², and imaged at room temperature with a Tecnai T12 electron microscope operated at 120 kV using low-dose procedures. Images were recorded at a magnification of 71,138x and a defocus value of ~1.5µm on a Gatan US4000 CCD camera. All images were binned (2 × 2 pixels) to obtain a pixel size of 4.16 Å on the specimen level. Particles were manually excised using Boxer [part of the EMAN 1.9 software suite] ³³ apart from tilt pairs (0° and 60°) where particles were selected using WEB³⁴.

2D classification and 3D reconstructions of negative stained T4L- β_2 V₂R- β_3 rr-Fab30(ScFv30) complex

2D reference-free alignment and classification of particle projections was performed using ISAC²⁸. 16,286 0° particle projections of native T4L- β_2 V₂R- β_{arr} -Fab30 complex and 13,703 0° projections of native T4L- β_2 V₂R- β_{arr} -ScFv30 complex were subjected to ISAC producing 186 classes accounting for 9,193 particle projections and 145 classes accounting for 8,127 projections, respectively. 14,109 0° particle projections of cross-linked T4L- β_2 V₂R- β_{arr} -ScFv30 complex were subjected to ISAC producing 111 classes accounting for 8,011 particle projections and 149 classes accounting for 8,127 projections, respectively. To determine the particle distribution of native T4L- β_2 V₂R- β_{arr} 1-Fab30 conformations, each class average was designated as "tight", "loose" or "unassigned" and the number of projections contributing to the class averages for each designation were added to calculate percentages.

For native and cross-linked Fab30 complexes the random conical tilt technique²⁶ was used to calculate a first back projection map from individual classes using the 60° tilted particle projections. After angular refinement of the 60° projections, untilted particle projections

were added to the dataset and the images were subjected to another cycle of refinement. For final reconstructions, particle projections from similar class averages were pooled together, corrected for contrast transfer function (CTF) according to local defocus values obtained by CTFTILT³⁵ and subjected to further refinement and reconstruction in FREALIGN³⁶. 2,825 particle projections from 0° and 60° images contributed to the cross-linked T4L- β_2 V₂R- β_{arr1} -Fab30 complex reconstruction. 2,196 particle projections from 0° and 60° images contributed to the native T4L- β_2 V₂R- β_{arr1} -Fab30 3D reconstruction. The resolution for each map was determined by the conventional Fourier Shell Correlation (FSC) approach at FSC=0.5 (Extended Data Figure 10).

Molecular modeling

For modeling the T4L- β_2 V₂R- β arr1-Fab30 complex we used the crystal structure of T4L- β_2 AR receptor from the T4L- β_2 AR-G α s complex (3SN6)⁶ and the crystal structure of V₂Rpp- β arr1-Fab30 (4JQI). Due to the low resolution of the EM map and the presence of the significant detergent micelle density surrounding the receptor region all docking operations within the EM maps were performed manually with visual inspection of best fit while taking into account additional constraints from cross-linking and HDXMS data. The Fab30 and T4L structures were independently shifted from the original position in the corresponding crystal structures to obtain an improved fit into the EM density map, reflecting both, the flexible nature of their association with β arr1 and β_2 AR, respectively, and a partial deformation of the complex on the carbon support of the EM grid.

Acknowledgments

We thank Darrell Capel for excellent technical assistance, Victoria Ronk, Donna Addison and Quivetta Lennon for administrative support, Roger K. Sunahara for stimulating discussions and Alex R.B. Thomsen for critical reading of the manuscript. We acknowledge support from the National Institutes of Health Grants DK090165 (G.S.), NS028471 (B.K.K), GM072688 and GM087519 (A.A.K. and S.K.), HL075443 (K.X.), HL16037 and HL70631 (R.J.L.), from the Mathers Foundation (B.K.K.), GM60635 (P.A.P.) and from the Pew Scholars Program in Biomedical Sciences (G.S.). R.H. and S.S.S. were supported by a research grant from the Canadian Institutes of Health Research (MOP-93725). R.I.R is supported by a post-doctoral fellowship from Coordenação de Aperfeiçoamento de Pessoal de Nível Superior–CAPES. R.J.L. is an investigator with the Howard Hughes Medical Institute.

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Figure 1. Formation and functional characterization of a stable agonist- $\beta_2 V_2 R$ - $\beta arr1$ signaling complex

a. Schematic flowchart of a novel purification strategy to isolate $\beta_2 V_2 R$ - $\beta arr1$ -Fab30 complex and large scale production and separation of agonist- $\beta_2 V_2 R$ - $\beta arr1$ -Fab30/ScFv30 complex from the free receptor by size exclusion chromatography (Superdex 200, 16/600 prep grade). T4 lysozyme sequence is attached at the N-terminus of the $\beta_2 AR$. **b.** Isolation of $\beta_2 V_2 R$ - $\beta arr1$ complex requires Fab30 and it is agonist dependent. Cells were stimulated either with inverse agonist (ICI-118551) or agonist (BI-167107) followed by incubation with or without Fab and subsequent purification on FLAG M1 beads. **c.** Formation of $\beta_2 V_2 R$ - $\beta arr1$ -Fab30 complex follows ligand efficacy. Formation of the complex in response to inverse agonists, partial agonists and full agonists is shown. The data is representative of 7 independent experiments.



Figure 2. Hydrogen-deuterium exchange mass spectrometry (HDXMS) analysis reveals potential interface between $\beta_2 V_2 R$ and $\beta arr1$

Differential hydrogen-deuterium exchange (HDX) rates of $\beta arr1$ in the $\beta_2 V_2 R$ - $\beta arr1$ -Fab30 vs. $V_2 Rpp$ - $\beta arr1$ -Fab30 were mapped on to the $\beta arr1$ crystal structure (PDB: 4JQI). Blue and red color coding suggest the $\beta arr1$ regions which exchange slower and faster, respectively, in the $\beta_2 V_2 R$ - $\beta arr1$ -Fab30 complex when compared to the $V_2 Rpp$ - $\beta arr1$ -Fab30 complex. Some regions (boxed) with significant HDX rate changes are enlarged in the panels (a)-(c). The HDX rates of the finger loop (residues 63-75) (a), middle loop (residues 129-140) (b) and lariat loop (residues 274-300) (b) became slower, whereas those of other regions, for example, (c) the β -strand I (β II, II (β II) and X (β X) in the N-domain became faster in the $\beta_2 V_2 R$ - $\beta arr1$ -Fab30 complex when compared to the $V_2 Rpp$ - $\beta arr1$ -Fab30 complex.



Figure 3. Single particle EM analysis of the $\beta_2 V_2 R$ - $\beta arr1$ -Fab30/ScFv30 complex a. Representative raw EM image of negative stained T4L- $\beta_2 V_2 R$ - $\beta arr1$ -Fab30/ScFv30 complexes. Scale bar = 25 nm. b. Representative class averages of the native T4L- $\beta_2 V_2 R$ - $\beta arr1$ -Fab30/ScFv30 complex. Class averages of particles displaying the loose "hanging" interaction (top) and the fully engaged "tight" interaction (bottom) are presented. Scale bar = 10 nm. c. Representative class averages of the "on-column" cross-linked T4L- $\beta_2 V_2 R$ - $\beta arr1$ -Fab30/ScFv30 complex. Upon cross-linking, the majority of class averages display the "tight" (fully engaged) $\beta arr1$ conformation, similar to a fraction (~37%) of particles observed in the non-cross-linked complex.



Figure 4. Structural model of the $\beta_2 V_2 R$ - $\beta_3 rr_1$ -Fab30 complex

a. Views of the T4L- $\beta_2 V_2 R$ - β_{arr1} -Fab30 complex 3D reconstruction with modeled T4L- $\beta_2 AR$ (green-orange, pdb: 3SN6), β_{arr1} (blue, pdb: 4JQI), and Fab30 (purple, pdb: 4JQI) crystal structures. The density surrounding $\beta_2 V_2 R$ represents the LMNG detergent micelle and is marked by "m". **b.** Views of the $\beta_2 V_2 R$ - β_{arr1} interface within the dashed line square of panel (a). Areas of β_{arr1} with reduced deuterium exchange are shown in cyan. Cross-linked Lys²³⁵ of $\beta_2 V_2 R$ and Lys⁷⁷ of β_{arr1} are highlighted. **c.** Illustration of the two-step GPCR- β_{arr1} interaction using surface representations of the structures of $\beta_2 AR$ (orange), the phosphorylated C-terminal tail of V₂R (yellow) and β_{arr1} (blue). The C-terminal portion of the V₂R peptide (Glu³⁵⁵ - Asp³⁶⁷) in the right model is positioned as found in the β_{arr1} -Fab30-V₂Rpp structure (pdb:4JQI), whereas the N-terminal portion (Ala³⁴² – Pro³⁵²) was remodeled to connect to the $\beta_2 AR$ C-terminus.

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Extended Data Figure 1. Formation of the $\beta 2V2R$ - $\beta arr1$ -Fab30 complex follows agonist occupancy of the receptor and it is biochemically stable

a. Sf9 cells co-expressing the β 2V2R, β arr1 and GRK2CAAX were stimulated with varying doses of high affinity agonist BI-167107 followed by addition of Fab30 and purification of the complexes. Stimulation of cells with increasing concentration of BI-167107 results in increasing amount of β arr1 co-purification indicating a direct correlation between occupancy of the receptor with agonist and complex formation. b. Quantification of agonist dependent complex formation from seven independent experiments normalized with respect to β arr1 signal at highest agonist concentration. c. Purified T4L- β 2V2R- β arr1-Fab30 complex was stored either at 4°C or at room temperature for 4 days followed by size exclusion chromatography on a superdex 200 (10/300) column (flow rate 0.5 ml/min). No significant dissociation of the complex was detected as monitored by appearance of a peak corresponding to the receptor (13.5 ml) or β arr1 (14.5 ml).

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Extended Data Figure 2. Functionally relevant conformation of $\beta arr1$ in the T4L- β 2V2R- β arr1-Fab30 complex as revealed by enhanced clathrin -terminal domain (clathrin TD) interaction Purified GST (glutathione S-transferase) tagged clathrin-TD was added to the purified complex or equivalent amount of β arr1 alone. Interaction of clathrin-TD with the complex or β arr1 was measured by subsequent co-immunoprecipitation and Western blot analysis. Quantification of four independent experiments shown as a bar graph. The relative intensities of the β arr1 bands are normalized with respect to β arr1 alone (set as 1). A coomassie stained gel indicating comparable amounts of β arr1 for complex vs. β arr1 alone conditions in clathrin-TD coimmunoprecipitation experiments is shown on the left. Error bar shows SEM. p<0.05 for paired t-test.

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Extended Data Figure 3. HDXMS analysis and mass-spectrometry based mapping of the crosslinking site in T4L-β2V2R-βarr1-Fab30 complex.

a. The differential H/D exchange between the T4L β 2V2R- β arr1-Fab30 complex and the V2Rpp- β arr1-Fab30 complex are mapped on the sequence of β arr1. b. Disuccinimidyl adipate (DSA), a homobifunctional amine-reactive crosslinker) was used to cross-link the pre-formed T4L- β 2V2R- β arr1-Fab30 complex. c. A representative SDS-PAGE showing the DSA cross-linking efficiency of the pre-formed complex. d. The cross-linked peptides were characterized with "doublet" peak signatures in mass spectra as described in the methods section and revelaed a cross-link between K235 of the β 2V2R and K77 at the distal end of

the finger loop in $\beta arr1$. e. Structural model of the $\beta 2V2R$ - $\beta arr1complex$ highlighting the cross-linking site.

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Extended Data Figure 4. Disulphide trapping strategy reveals close proximity of residue 235 of the β 2V2R and residue 78 at the distal end of the finger loop in β arr1.

a. Structural model of the β 2V2R- β arr1 complex depicting the proximity of K235 on the β 2V2R and D78 on β arr1. b. Single cysteine insertion mutants of the β 2V2R (covering residues 231- 236) and β arr1D78C were co-transfected in HEK-293 cells and complex formation was induced by stimulating the cells with an oxidizing agent H2O2 and agonist (Isoproterenol; Iso). Subsequently, a co-immunoprecipitation assay was performed using FLAG M2 beads (FLAG- β arr1). Formation of disulphide trapped complex was visualized by

Western blotting. c. Quantification of β arr1 in S-S trapped complex from three indepdent experiments with standard error of the mean.



Extended Data Figure 5. Raw EM images of negative stained native T4L-β2V2R-βarr1-Fab 30(ScFv30) complex.

a) Raw EM image of T4L- β 2V2R- β arr1-Fab30 complex.b) Raw EM image of T4L- β 2V2R- β arr1-ScFv30 complex. Scale bar = 100 nm.

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Extended Data Figure 6. 2D classifications of the T4L-β2V2R-βarr1-Fab30(ScFv30) complex. Reference-free 2D class averages were obtained using ISAC. a) 2D classification of the T4L- β 2V2R - β arr1-Fab30 comple x. b) 2D classification of the

T4L- β 2V2R- β arr1-ScFv30 complex. Scale bar = 10 nm.

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Extended Data Figure 7. "On-column" glutaraldehyde cross-linking of the pre-formed complex. a. Schematic representation of the "on-column" cross-linking strategy. A glutaraldehyde solution is injected to a size exclusion chromatography column, followed by injection of the purified complex protein. As the complex protein passes through the glutaraldehyde bolus, the receptor and the β arr components of the complex are cross-linked through proximal primary amine groups. This procedure allows only brief exposure of the complex to glutaraldehyde and serves as an "in-line" purification of homogenously cross-linked protein from any aggregation that may arise from non-specific cross-linking. b. "On-column" cross-

linking of the T4L- β 2V2R- β arr1-ScFv30 complex. Purified complex (approximately 20 μ M) was injected on to a 24mL Superdex 200 gel filtration column after a pre-injection of 200 μ L of 0.25% glutaraldehyde bolus. Individual fractions were collected and analyzed by "Simplyblue" stained SDS-PAGE. c. "On-column" cross-linking of the T4L- β 2V2R- β arr1-Fab30 complex performed as described for the ScFv complex above.



Extended Data Figure 8. Raw EM images of negative stained cross-linked T4L- $\beta 2V2R$ - β arr1-Fab30(ScFv30) complex.

a) Raw EM image of T4L- β 2V2R- β arr1-Fab30 complex. b) Raw EM image of T4L- β 2V2R- β arr1-ScFv30 complex. Scale bar = 100 nm.

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 $Extended \ Data \ Figure \ 9. \ 2D \ classifications \ of \ cross-linked \ T4L-\beta 2V2R-\beta arr1-Fab30 (ScFv30) \ complex.$

Reference-free 2D class averages were obtained using ISAC. a) 2D classification of cross-linked T4L- β 2V2R- β arr1-Fab30 complex. b) 2D classification of cross-linked T4L- β 2V2R- β arr1- ScFv30 complex. Scale bar = 10 nm.

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Extended Data Figure 10. 3D EM reconstructions and resolution indications by Fourier Shell Correlation (FSC) curves.

The top panel shows the 3D map from particles representing the fully engaged β 2V2R- β arr1 conformation of the T4L- β 2V2R- β arr1- Fab30 complex. The bottom panel shows the 3D reconstruction from particles displaying the loose, hanging arrestin, conformation of the same complex. Representative 2D averages of particles used for the calculation of initial models by the random conical tilt method are shown on the left of each respective 3D map.