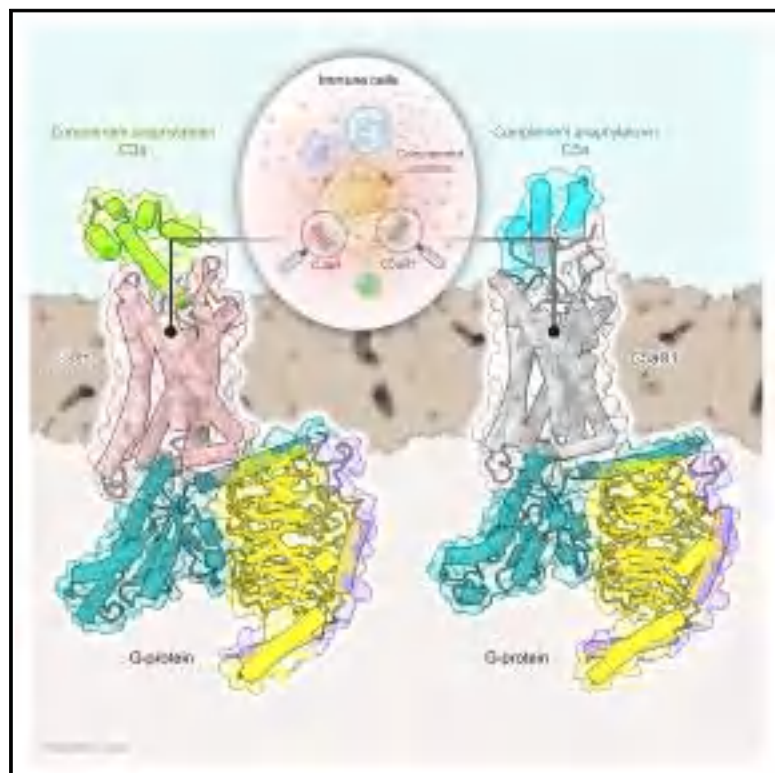


LIST OF PUBLICATIONS:

- Molecular insights into ligand promiscuity and atypical dimerization of the C-X-C chemokine receptor, CXCR2
S Saha, FK Sano, S Sharma, M Ganguly, S Saha, H Akasaka, T Kobayashi, N Zaidi, S Mishra, A Dalal, S Mohapatra, M Yadav, Y Itoh, A Chevigne, R Banerjee, W Shihoya, O Nureki, AK Shukla (***Under revision***) (***Joint first author***)
- Molecular insights into dual-agonism and biased-signaling at the C-X-C type chemokine receptors CXCR3 and CXCR7
S Saha, FK Sano, S Sharma, P Sarma, M Ganguly, H Akasaka, T Kobayashi, N Zaidi, S Mishra, A Dalal, Y Itoh, R Leurs, GD Jhingan, R Banerjee, W Shihoya, O Nureki, AK Shukla (***Under revision***) (***Joint first author***)
- Molecular mechanism of distinct chemokine engagement and functional divergence of the human Duffy antigen receptor
S Saha, B Khanppnavar, J Maharana, H Kim, CMC Carino, C Daly, S Houston, P Kumari, PN Yadav, B Plouffe, A Inoue, KY Chung, R Banerjee, VM Korkhov, AK Shukla (***Joint first author***)
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- Molecular basis of anaphylatoxin binding, activation, and signaling bias at complement receptors
MK Yadav, J Maharana, R Yadav, **S Saha**, P Sarma, C Soni, V Singh, S Saha, M Ganguly, XX Li, S Mohapatra, S Mishra, H Khant, M Chami, T Woodruff, R Banerjee, AK Shukla, C Gati (***Joint first author***)
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Molecular Cell 80 (1), 3-5, 2020
- The Inside Story: Crystal Structure of the Chemokine Receptor CCR7 with an Intracellular Allosteric Antagonist
S Saha, AK Shukla
Biochemistry 59 (1), 12-14, 2019

Molecular basis of anaphylatoxin binding, activation, and signaling bias at complement receptors

Graphical abstract



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In brief

A series of cryo-EM structures offer insights into how anaphylatoxins C3a and C5a engage complement receptors, illustrating agonist binding modes, pathways of receptor activation, and signaling bias that should facilitate the discovery of ligands and therapeutic candidates targeting these receptors.

Highlights

- Cryo-EM structures of complement receptors C5aR1 and C3aR in complex with G proteins
- Molecular mechanism of ligand binding, selectivity, activation, and signaling bias
- Structural basis of reduced efficacy of C5a^{des-Arg} through proteolytic cleavage
- Discovery and structural mechanism of a subtype-specific biased agonist of C3aR



Article

Molecular basis of anaphylatoxin binding, activation, and signaling bias at complement receptors

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SUMMARY

The complement system is a critical part of our innate immune response, and the terminal products of this cascade, anaphylatoxins C3a and C5a, exert their physiological and pathophysiological responses primarily via two GPCRs, C3aR and C5aR1. However, the molecular mechanism of ligand recognition, activation, and signaling bias of these receptors remains mostly elusive. Here, we present nine cryo-EM structures of C3aR and C5aR1 activated by their natural and synthetic agonists, which reveal distinct binding pocket topologies of complement anaphylatoxins and provide key insights into receptor activation and transducer coupling. We also uncover the structural basis of a naturally occurring mechanism to dampen the inflammatory response of C5a via proteolytic cleavage of the terminal arginine and the G-protein signaling bias elicited by a peptide agonist of C3aR identified here. In summary, our study elucidates the innerworkings of the complement anaphylatoxin receptors and should facilitate structure-guided drug discovery to target these receptors in a spectrum of disorders.

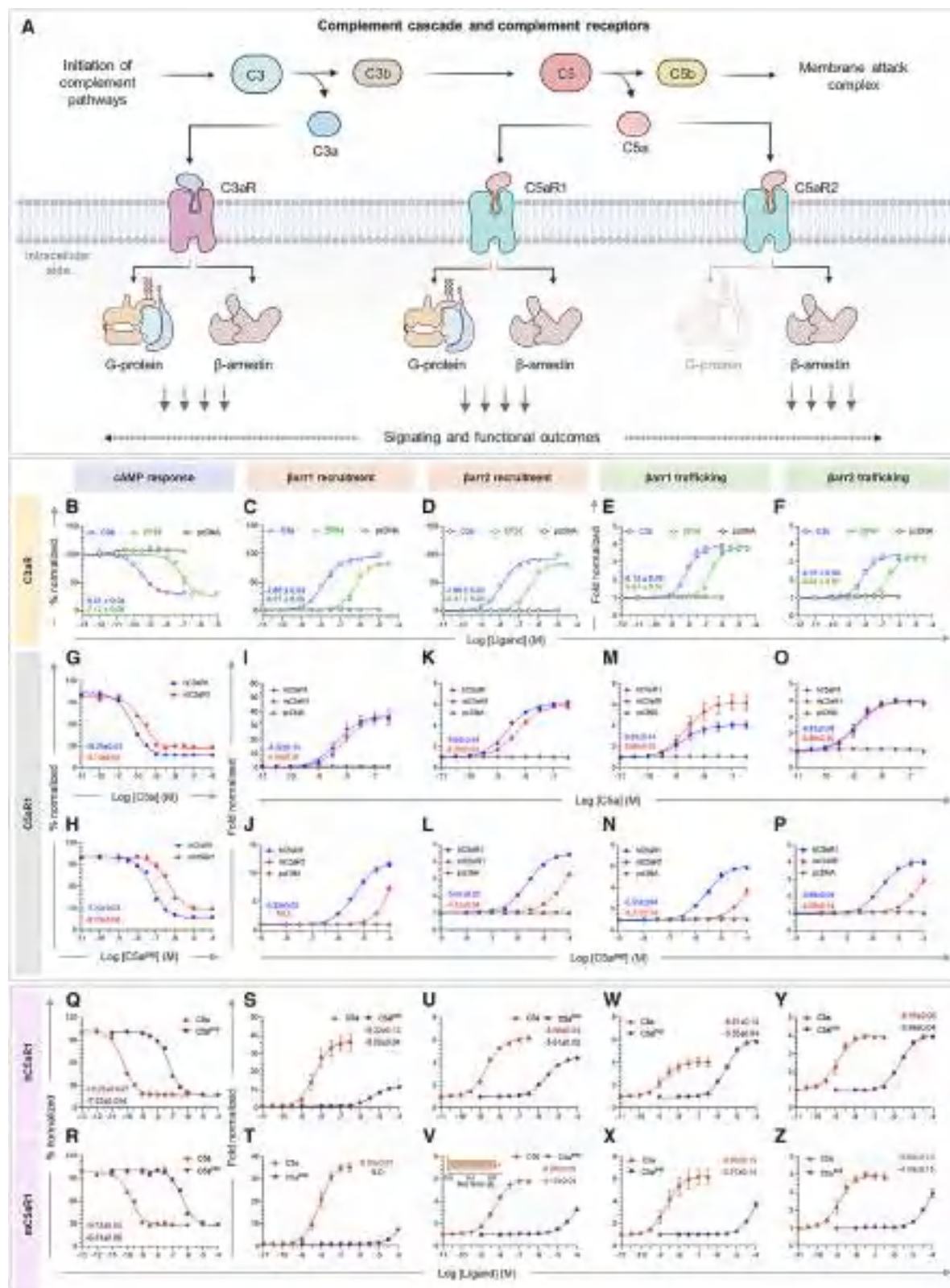
INTRODUCTION

One of the key mechanisms through which the immune system combats pathogenic infections is the activation of the complement cascade.^{1–4} It is an intricate network of plasma proteins including inflammatory peptides, proteases, and integral membrane receptors that work in a concerted fashion.^{1–4} Once activated, it plays a vital role in the efficient elimination of microbial agents through the formation of the membrane attack complex and associated mechanisms.^{3,4} Complement activation results in the generation of several peptide fragments by the action of different proteases, and these complement factors subsequently exert their functions through their corresponding receptors and effectors.^{1–4} Abnormal activation of the complement system is directly linked with multiple disease conditions, including immunodeficiency, autoimmune disorders such as rheumatoid arthritis, hematological and vascular disease, ocular disease,

neurodegenerative and neuropsychiatric disease, renal disease, and inflammatory bowel disease.^{5–7} Complement fragments C3a and C5a generated from the proteolytic cleavage of complement C3 and C5, respectively, also known as anaphylatoxins, play a central role in priming and amplifying the immune response by recruiting immune cells, such as leukocytes, and triggering the secretion of pro-inflammatory molecules, such as cytokines.^{1–4}

C3a and C5a bind to distinct seven transmembrane receptors (7TMRs), with C3a being selective for C3aR, whereas C5a can bind to two different receptors, namely, C5aR1 and C5aR2^{8–11} (Figure 1A). C3aR is a prototypical G-protein-coupled receptor (GPCR), which exhibits primary and secondary coupling to Gi and Gq subtypes of heterotrimeric G proteins and also recruits β -arrestins (β arrestins) upon activation^{11–14} (Figure 1A). On the other hand, although C5aR1 couples to both Gi and β arrestins upon activation by C5a, C5aR2 solely signals via β arrestins without any measurable G-protein activation and hence is also referred to as an arrestin-





(legend on next page)

coupled receptor (ACR)^{15–17} (Figure 1A). C3aR and C5aR1 are expressed in multiple types of immune cells, including mast cells, neutrophils, and monocytes/macrophages, and their aberrant signaling is linked to numerous inflammatory disorders, such as sepsis, vasculitis, pulmonary fibrosis, arthritis, asthma, and lupus,^{5,6,18–21} making them important drug targets. Moreover, the interaction of C5a with C5aR1 and ensuing downstream signaling responses have been implicated in the disease severity of COVID-19 patients, including a potential chemoattractant role leading to infiltration of neutrophils and monocytes in the broncho-alveolar lavage fluid (BALF) of patients.²² In addition, a monoclonal antibody that targets complement C5a and thereby blocks its interaction with C5aR1, has recently been approved for use in hospitalized patients of COVID-19.²³ However, despite its fundamental importance, our current understanding of complement anaphylatoxin recognition by complement receptors remains limited and is based predominantly on biochemical studies.

Human C3a and C5a contain 77 and 74 amino acids, respectively, and exhibit a four-helix bundle architecture.^{24,25} Previous studies have suggested a two-site binding mechanism to the corresponding receptors involving the N terminus, the 2nd extracellular loop (ECL2), and the transmembrane core of the receptor.^{26–29} Interestingly, peptides derived from and modified based on the carboxyl terminus of both C3a and C5a have been identified as agonists of C3aR and C5aR1, although their binding affinity and potency differ significantly.^{16,30–38} For example, EP54 and EP67, two decapeptides derived from the carboxyl terminus of C5a, and several peptides derived from the carboxyl terminus of C3a exhibit full agonism at C3aR in ERK1/2 mitogen-activated protein (MAP) kinase phosphorylation assay.³³ Moreover, a previous study has also reported that EP54 and EP67 can induce cytokine production and, to a lesser extent, enzyme release from neutrophils.³⁹ Similarly, a hexapeptide designed based on C5a, referred to as C5a^{pep}, behaves as a G-protein-biased agonist for C5aR1.⁴⁰ These studies underscore the

critical contribution of the carboxyl terminus of C3a and C5a in eliciting transducer coupling and downstream functional responses. Finally, the activity of C3a and C5a is regulated by a physiological mechanism where the terminal arginine residues are cleaved off by the action of carboxypeptidases, and the resulting fragments, referred to as C3a^{des-Arg} and C5a^{des-Arg} exhibit a significantly reduced functional responses through the cognate receptors.^{41–44} However, the direct structural visualization of agonist binding to C3aR and C5aR1 and ensuing mechanism of receptor activation and transducer coupling still remain elusive. This represents a major knowledge gap in our current understanding of complement receptor activation and signaling that restricts the possibility of structure-guided design of ligands and therapeutics targeting these receptors.

In this manuscript, we present nine cryoelectron microscopy (cryo-EM) structures of agonist-bound C3aR and C5aR1 in complex with heterotrimeric G proteins, which elucidate intricate molecular details of complement recognition, receptor activation, and downstream signaling. In particular, these structures uncover previously unanticipated distinct binding modes for C3a and C5a on their respective receptors while maintaining a converged positioning of their carboxyl terminus in the binding pocket. The structural insights help rationalize the subtype selectivity of C3a and C5a, a large body of data on agonism and cross-reactivity of C3a/C5a-derived peptides, and allow us to identify a subtype-selective G-protein-biased agonist at C3aR. Together with biochemical and pharmacological data, the structural snapshots presented here unravel the molecular mechanism driving signaling bias in the complement receptor system and the natural mechanism to dampen the inflammatory responses of complement anaphylatoxins through the removal of the terminal arginine residue. Collectively, this study offers a previously lacking platform to facilitate structure-guided drug discovery at the complement receptors with enhanced subtype selectivity and biased agonism.

Figure 1. Activation of complement receptors and downstream functional outcomes

(A) Triggering of the various complement pathways leads to the generation of complement peptides and subsequent activation of cognate complement receptors. An overview of the activation of C3aR, C5aR1, and C5aR2 and their signaling has been illustrated.

(B) To study G α i activation, forskolin-elevated decrease in cAMP level is measured using GloSensor assay downstream of C3aR in response to indicated ligands (mean \pm SEM; n = 4; normalized with starting value for each ligand as 100%).

(C and D) β arr1/2 recruitment to C3aR in response to indicated ligands as measured by NanoBIT assay (receptor-SmBIT + LgBIT- β arr1/2), respectively (mean \pm SEM; n = 4; normalized with the luminescence signal at maximal ligand dose of C3a as 100%).

(E and F) β arr1/2 trafficking to the endosomes downstream of C3aR in response to indicated ligands as measured by NanoBIT assay (receptor + SmBIT- β arr1/2 + LgBIT-FYVE) (mean \pm SEM; n = 4; normalized with the luminescence signal at minimal ligand dose of each condition as 1).

(G and H) C5a (top) and C5a^{pep} (bottom) driven G α i-mediated second messenger response as measured by agonist-dependent decrease in forskolin-induced cytosolic cAMP levels downstream to C5aR1. Respective logEC50 values are mentioned in the inset. Data (mean \pm SEM) represent four independent experiments, normalized with respect to the highest signal (measured as 100%) for each receptor.

(I–P) C5a/C5a^{pep} induced β arr1/2 recruitment and trafficking as measured by NanoBIT assay. Respective logEC50 values are mentioned in the inset. Data (mean \pm SEM) represent four independent experiments, fold normalized with respect to luminescence observed at the lowest dose (measured as 1) for each receptor.

(Q and R) Comparison of C5a/C5a^{pep}-mediated cAMP response downstream of human (top) and mouse (bottom) C5aR1 reveals reduced potency of C5a^{pep} as compared with C5a. Respective logEC50 values are mentioned in the inset. Data (mean \pm SEM) represent four independent experiments, normalized with respect to the highest signal (measured as 100%) in response to each ligand.

(S–Z) Measurement of β arr1/2 recruitment and trafficking to human (top) and mouse (bottom) C5aR1 upon stimulation with C5a and C5a^{pep}. Respective logEC50 values are mentioned in the inset. Data (mean \pm SEM) represent four independent experiments, fold normalized with respect to luminescence observed at the lowest dose (measured as 1) for each ligand. Bias factor (β value) determined by taking C5a as reference elucidates the G-protein-biased nature of C5a^{pep} that has been provided in insets.

Plots explaining the functional bias (Q–Z) of C5a^{pep} are from the data shown in (G)–(P) and presented separately to highlight the effects compared with C5a in both human and mouse C5aR1.

See also Figure S1.

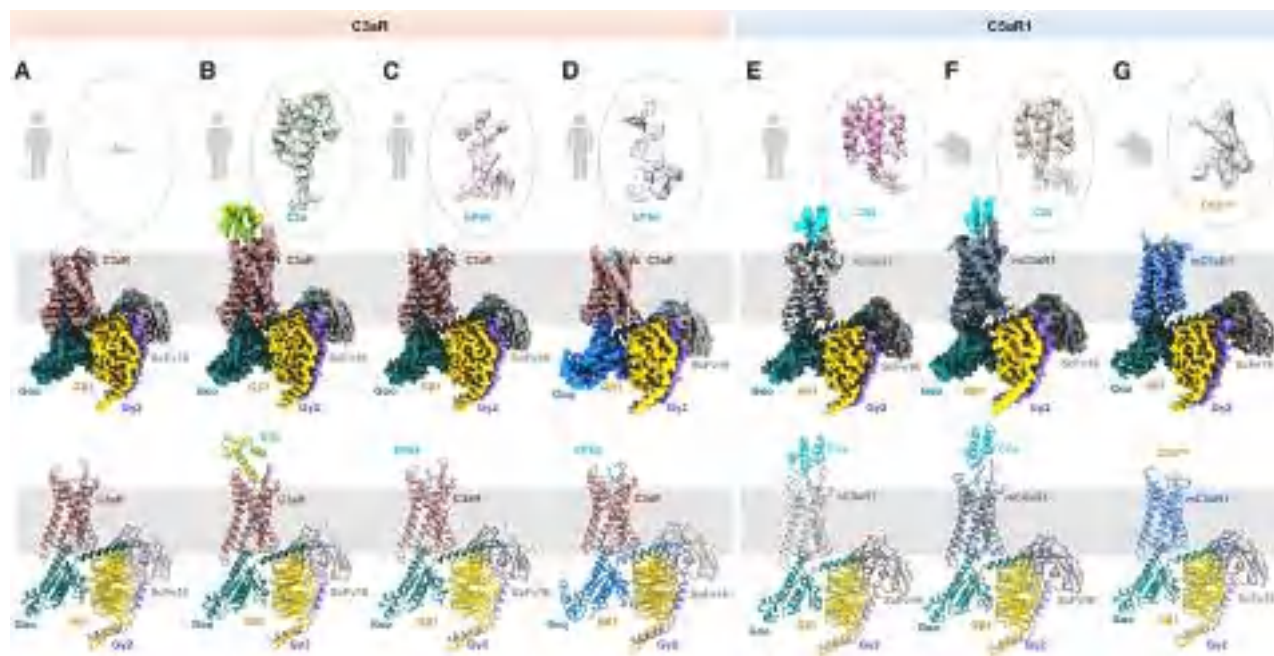


Figure 2. Structures of complement receptor signaling complexes

(A–G) Cryo-EM density maps (top) and corresponding models (bottom) of Apo-C3aR-Go (Glacios), C3a-C3aR-Go, EP54-C3aR-Go, EP54-C3aR-Gq, C5a-hC5aR1-Go, C5a-mC5aR1-Go, and C5a^{pep}-mC5aR1-Go. Cryo-EM density maps of respective ligands have been shown in gray dotted circles (top corner). (Rosy brown, C3aR; green, C3a; pale blue, EP54; light gray, hC5aR1; slate gray and blue, mC5aR1; cyan, C5a; yellow, C5a^{pep}; teal, Gαo; deep blue, Gαq; gold, Gβ1; purple, Gβ2; dark gray, ScFv16.)

See also Table S1 and Methods S1.

RESULTS

Ligand pharmacology at the complement receptors

In order to study agonist binding and activation of C3aR and C5aR1, we focused our efforts on both natural and synthetic agonists of these two receptors. For C3aR, we selected C3a and a decaemic peptide agonist EP54. We first measured the pharmacological profile of EP54 with C3a as a reference in G-protein and β arr assays (Figures 1B–1F and S1A–S1C). We observed that EP54 acts as a full agonist for Gαi coupling, as measured using cyclic AMP (cAMP) response in a GloSensor assay, and β arr1/2 recruitment and endosomal trafficking as measured using NanoBIT-based assays, albeit with lower potency (Figures 1B–1F). These findings make EP54 a suitable candidate, together with C3a, to decipher the structural determinants of C3a recognition and agonism at C3aR. In order to further our understanding of complement receptor activation, we next turned our attention to the complement C5a receptor, C5aR1. Previous studies have suggested that selected ligands exhibit species-specific differences in their pharmacology at C5aR1.²⁰ Therefore, we first compared the pharmacology of human C5a and C5a^{pep}, a synthetic peptide designed based on the carboxyl-terminal sequence of C5a, on the human and mouse C5aR1, referred to as hC5aR1 and mC5aR1, respectively, in G-protein and β arr assays. We observed that both C5a and C5a^{pep} behave as full agonists on mC5aR1 with slightly lower potency compared with hC5aR1 in terms of G-protein-

mediated cAMP response (Figures 1G and 1H). On the other hand, although C5a exhibits full agonism for β arr recruitment and endosomal trafficking on mC5aR1 (Figures 1I, 1K, 1M, and 1O), C5a^{pep} displays partial agonism at mC5aR1 compared with hC5aR1 in these assays (Figures 1J, 1L, 1N, and 1P). Moreover, C5a^{pep} acts as G-protein-biased agonist at both hC5aR1 and mC5aR1 (Figures 1Q–1Z). The mC5aR1 and hC5aR1 were expressed at comparable levels in these assays (Figures S1D–S1M). These findings suggest that structural visualization of agonist-bound human and mouse C5aR1 may yield interesting insights into species-specific agonist pharmacology. We also note that in some assays, C5a^{pep} dose-response curves do not reach complete saturation phase due to limitations associated with its relatively lower affinity and poor solubility at higher concentrations, and therefore, we have not calculated bias factor for the corresponding datasets.

Structure determination of agonist-C3aR/C5aR1-G-protein complexes

Taking the lead from the agonist pharmacology data presented in Figure 1, we determined seven different cryo-EM structures of G-protein-bound C3aR and C5aR1 occupied with natural and synthetic agonists (Figures 2A–2G). First, we reconstituted C3a-C3aR-Gαoβ1γ2 complex stabilized by ScFv16³⁰ (Methods S1) and determined the structure at ~3.2 Å (Methods S1). However, we did not observe any discernible density for C3a in the complex (Figure 2A). We, therefore, refer to this as

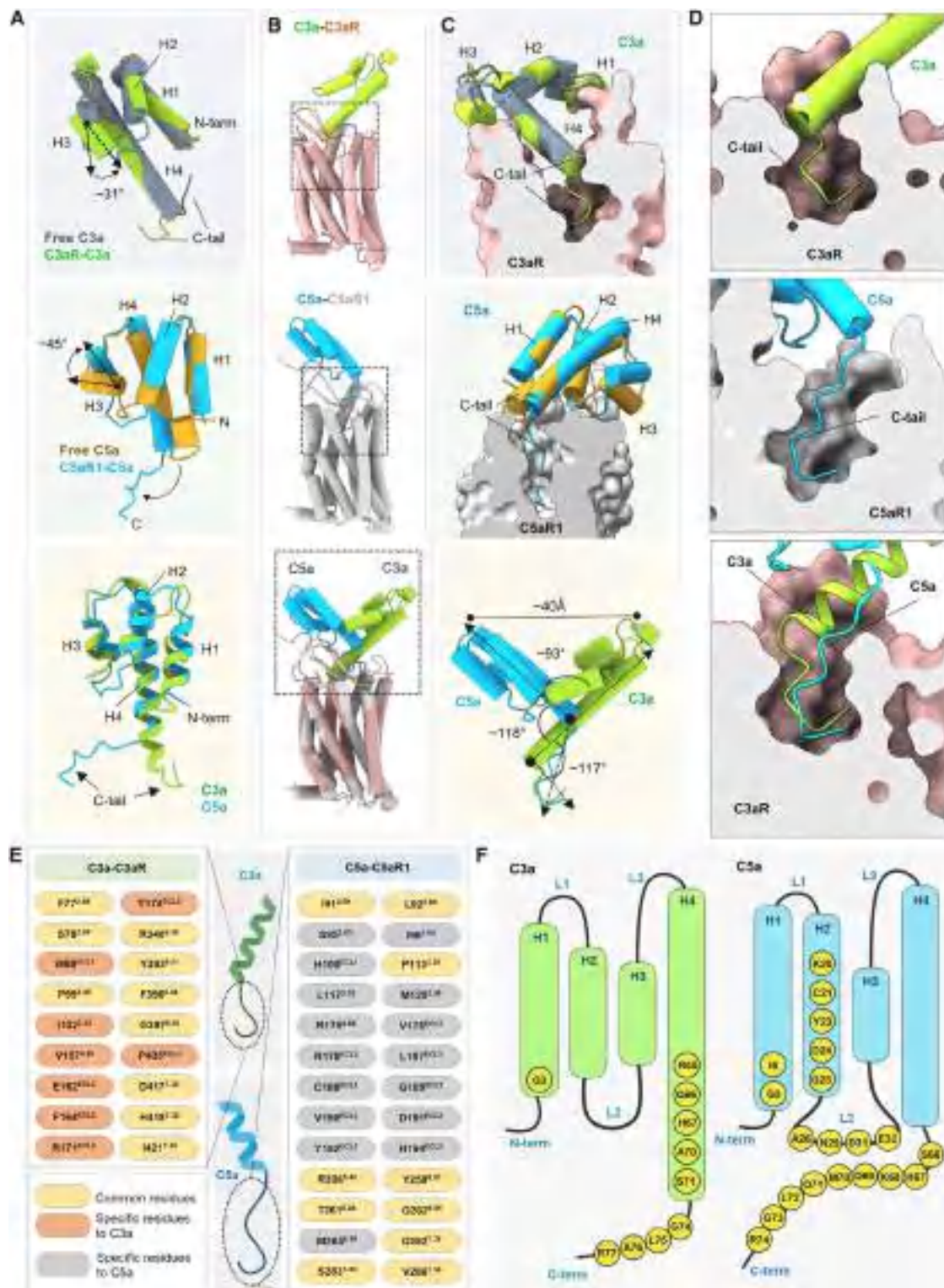


Figure 3. Complement peptide binding to complement receptors

(A) Structure of C3a (top) and C5a (middle), showing four-helix bundle with a short C-terminal tail. Free C3a (PDB: 4HW5) and free C5a (PDB: 1KJS) have been superimposed with C3aR and C5aR, respectively. Structural alignment of C3a and C5a in receptor-bound forms (bottom).

(B) Side view of C3aR (top), C5aR1 (middle), and both (bottom) bound to their endogenous ligands.

(C) The C termini of the ligands change their conformation upon binding to their corresponding receptors compared with the basal states.

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apo-C3aR-Go complex in subsequent discussion. Considering the potential dissociation of C3a during purification as a plausible reason for the empty ligand-binding pocket, we supplemented the reconstituted C3a-C3aR-G $\alpha\beta\gamma$ 2 complex with additional C3a immediately before freezing the samples for cryo-EM. This strategy allowed us to obtain a C3a-C3aR-G $\alpha\beta\gamma$ 2 complex structure at 3.2 Å with a clear C3a density (Figure 2B; Methods S1). Next, we also reconstituted EP54-C3aR-G $\alpha\beta\gamma$ 2 and EP54-C3aR-G $\alpha\beta\gamma$ 2 complexes (Methods S1) and determined their structures at 2.9 and 3.6 Å, respectively (Methods S1), and we observed clear densities for EP54 in these complexes (Figures 2C and 2D). Finally, we reconstituted C5a-hC5aR1-G $\alpha\beta\gamma$ 2, C5a-mC5aR1-G $\alpha\beta\gamma$ 2, and C5a^{pep}-mC5aR1-G $\alpha\beta\gamma$ 2 complexes and determined their structures at 3.3, 3.9, and 3.4 Å, respectively (Figures 2E–2G; Methods S1). Despite the modest resolution of the mC5aR1 complexes, our generated cryo-EM maps allowed unambiguous modeling of the secondary structures of all components, including C5a and C5a^{pep} (Figures 2F and 2G). The overall data collection and refinement statistics, the density maps of the key segments in the complexes, including the ligands, and the description of the residues that are resolved in these structures are presented in Tables S1 and S2 and Methods S1.

Recognition of complement anaphylatoxins by C3aR and C5aR1

C3a and C5a share about 35% sequence identity, and they both adopt a four-helix bundle architecture stabilized by multiple disulfide bridges and a terminal arginine located at the end of helix 4 (H4)^{24,25} (Arg⁷⁷ in C3a and Arg⁷⁴ in C5a) (Figure 3A). Site-directed mutagenesis studies, coupled with ligand binding and second messenger assays, have proposed a two-site model for C3a-C3aR and C5a-C5aR1 interaction, with the first site involving the ECL2 of the receptor, whereas the second site engages multiple residues from the extracellular side of the transmembrane domain (TMD).^{26,27} In addition, the N terminus of C5aR1 has also been proposed to be involved in C5a binding.⁹ It is noteworthy that, although ECL2 of C3aR is extraordinarily long, with more than 150 amino acids, functional studies have demonstrated that a large part of it is dispensable for the binding of C3a, as measured by calcium response assays.²⁷ In the structures determined here, the stretch from Lys175^{ECL2} to Pro330^{5,32} is not resolved, likely due to inherent flexibility in this region. We observe that both C3a and C5a retain their helical fold upon binding to their respective receptors (Figure 3A), although there are some structural changes compared with their basal states, which are more pronounced in C5a than in C3a (Figure 3A). For example, H3 in C3a and C5a tilts by about 30° and 45°, respectively, upon binding to their corresponding receptors (Figure 3A). Moreover, the distal carboxyl terminus in C3a undergoes a significant rotation upon binding to C3aR

(Figure 3A, upper), whereas that in C5a transitions from a short α -helical turn in the basal state to an extended conformation upon binding to C5aR1 (Figure 3A, middle). Surprisingly, there is a dramatic difference in the overall positioning of C3a on C3aR and C5a on C5aR1 (Figures 3B and 3C). The globular domains of C3a and C5a are tilted at an angle of about 120° with respect to their carboxyl terminus (Figure 3C, lower). Moreover, the globular domains of C3a and C5a are oriented in opposite directions on the extracellular side of the receptor at an angle of about 90° (Figure 3C, lower). Notably, however, the carboxyl terminus of both C3a and C5a adopt a hook-like conformation (Figure 3D) and position themselves in an analogous binding pocket on the respective receptors with a buried surface area of 1,720 and 1,252 Å², respectively (Figure 3D, lower). It is worth noting that the residues from Arg⁶⁵ to Arg⁷⁷ in C3a form extensive interactions with ECL1–ECL3 and TM2–7 in C3aR (Figures 3E and 3F; Table S2). The only other residue from C3a that interacts with C3aR is Gln³, which engages with Asp167 and Thr166 in ECL2 through polar interactions (Figure S1S). On the contrary, we observed a significantly broader interface between C5a and C5aR1, which not only includes Ser⁶⁶ to Arg⁷⁴ at the distal carboxyl terminus but also multiple residues from the loop between H2 and H3, H2, and H1 (Figures 3E, 3F, and S1T). On the receptor side, C3a and C5a engage several analogous residues from TM2–3 and TM5–7, although they also display several specific interactions with their cognate receptors (Figure 3E; Table S2). Importantly, although we observe a clear density for the N terminus of C5aR1 and its interaction with C5a in the C5a-C5aR1 structures, the N terminus of C3aR is not resolved in any of the structures. Although the absence of clearly resolved density for the N terminus of C3aR may indicate its inherent flexibility and a lesser contribution in agonist binding as indicated earlier,²⁷ future studies are required to probe this further. Taken together, the potential differential involvement of the N terminus and distinct orientation of C3a and C5a provide the structural basis of subtype selectivity between these two receptors. On the other hand, an overall analogous binding pocket and positioning of distal carboxyl terminus of C3a and C5a may help rationalize the cross-reactivity of peptide agonists described earlier.²⁸ The structure of C5a-bound mouse C5aR1 exhibits an overall similar binding pose of C5a as the human C5aR1, including the extended helical conformation of its distal carboxyl terminus (Figure S1U), extensive interactions with the receptor, including the N terminus and ECL2, and a critical involvement of its terminal Arg⁷⁴ (Table S2).

As mentioned earlier, we serendipitously determined the structure of ligand-free C3aR-G $\alpha\beta\gamma$ 2 complex, which exhibits an empty ligand-binding pocket (Figure 2A). Structural comparison between ligand-free and agonist-bound C3aR structures revealed an overall similar conformation of the ligand-binding

(D) The C-terminal tails of C3a (top) and C5a (middle) adopt a hook-like conformation upon entering deep into the orthosteric pocket of respective receptors. C3aR and C5aR1 have been shown in surface slice, and C3a/C5a as cartoon representation.

(E) The distal C-terminal portions of C3a/C5a make extensive contacts with C3aR/C5aR1. Residue at the interface between the H4 and C termini of C3a/C5a and C3aR/C5aR1 is depicted.

(F) C3a and C5a are shown in topology diagrams. Residues making contact with receptors are highlighted in yellow circles.

See also Figures S1 and S2 and Table S2.

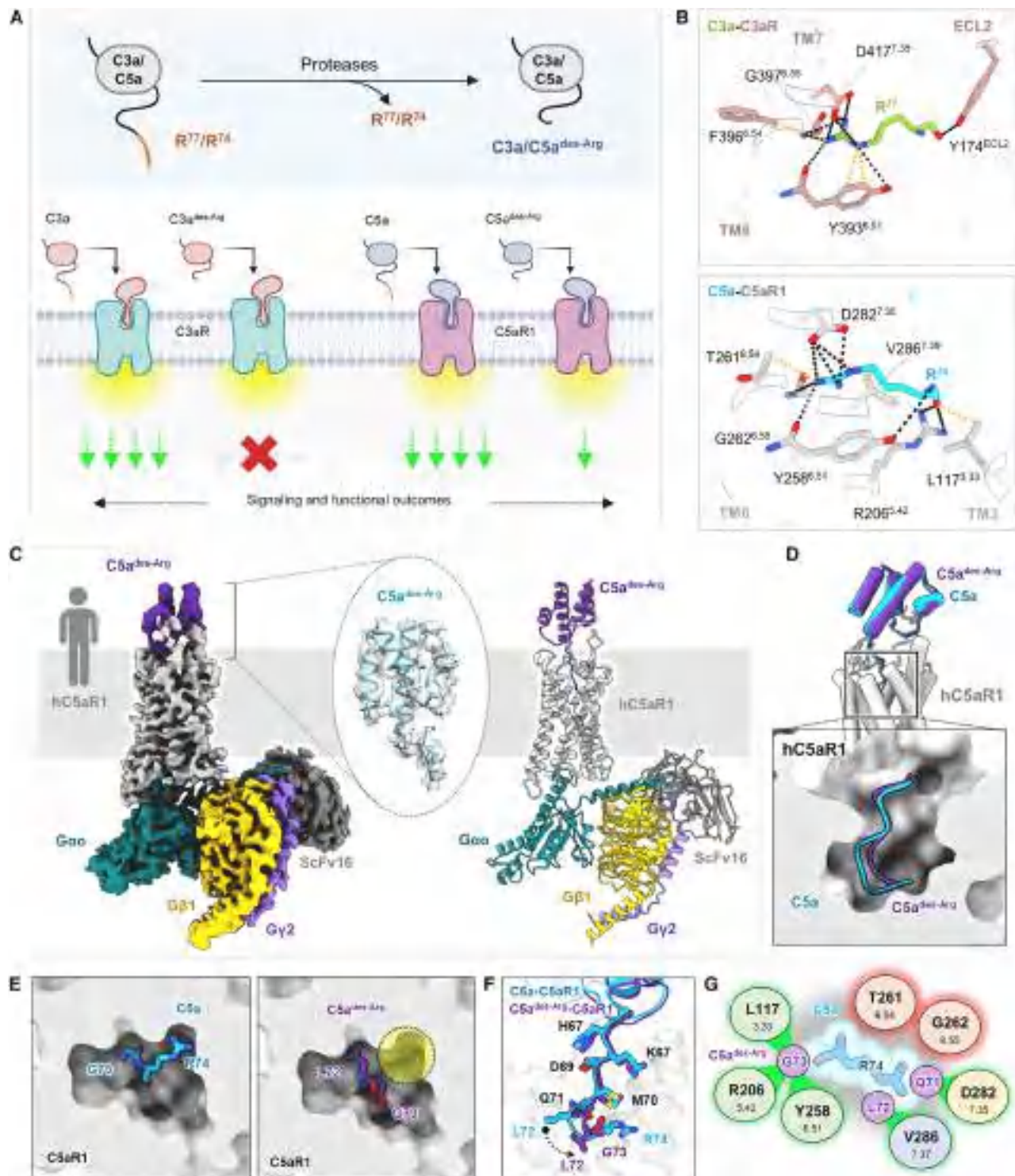


Figure 4. Structural determinants of reduced functional efficacy of C3a^{des-Arg} and C5a^{des-Arg}

(A) Proteolytic cleavage of the C-terminal Arg⁷⁷ from C3a and Arg⁷⁴ from C5a results in C3a^{des-Arg} and C5a^{des-Arg}, respectively (top). C3a^{des-Arg} is reported to elicit almost no functional outcomes upon binding to C3aR, whereas C5a^{des-Arg} retains minimal potency in driving downstream signaling via C5aR1 (bottom). (B) Interactions of Arg⁷⁷ in C3a (top) and Arg⁷⁴ in C5a with the residues of C3aR and C5aR1 have been illustrated. (C) Cryo-EM density map (left) and corresponding model (right) of C5a^{des-Arg}-hC5aR1-Go. Cryo-EM density map of C5a^{des-Arg} has been shown in gray dotted circles.

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pocket in C3aR (Figures S2A–S2D), although with a few notable differences (Figures S2E and S2F). For example, Arg340^{5,42} of C3aR, which engages with Arg⁷⁷ of C3a, displays a linear inward shift of about 5 Å, possibly due to a lack of constraints imposed by ligand binding (Figure S2E). Similarly, Arg161^{4,64} in TM4 of C3aR, which engages with Leu⁸ in EP54, exhibits a rotation of about 150° in apo-C3aR, which allows it to interact with the main chain of Lys96^{3,26} in TM3 (Figure S2F).

Structural mechanism of reduced efficacy by C3a^{des-Arg} and C5a^{des-Arg}

C3a and C5a are potent anaphylatoxins with strong inflammatory response, and there exists a natural mechanism to dampen their excessive activity in physiological context. The terminal arginine residues (Arg⁷⁷ in C3a and Arg⁷⁴ in C5a) are cleaved off by carboxypeptidases, resulting in C3a^{des-Arg} and C5a^{des-Arg} (Figure 4A).⁶ Although C5a^{des-Arg} retains about one-tenth of the activity compared with C5a, C3a^{des-Arg} appears to completely lose the pro-inflammatory activity (Figure 4A).^{8,41–43} However, a molecular basis to explain this mechanism operating *in vivo* has remained elusive, and our structural snapshots now provide a potential rationale for this. It is particularly striking that Arg⁷⁷ in C3a and Arg⁷⁴ in C5a form multiple polar interactions and non-bonded contacts with the residues in TMDs and ECL2 of C3aR and C5aR1, respectively (Figure 4B). This is also supported by site-directed mutagenesis studies where Arg³⁴⁰Ala and Asp⁴¹⁷Ala mutations in C3aR, which are critical interaction partners of Arg⁷⁷ in C3a, diminish agonist binding and receptor activation.²⁶ In order to directly elucidate the molecular mechanism of dramatically altered functional efficacy of C5a^{des-Arg}, we reconstituted and determined the structure of C5a^{des-Arg}-C5aR1-Go complex (Figures 4C, S2G, S4D, and S5; Tables S1 and S2). We observed that the binding pose of C5a^{des-Arg} on C5aR1 superimposes well with C5a, and its carboxy terminus adopts a similar hook-like conformation in the orthosteric binding pocket on the receptor (Figure 4D). Strikingly, however, a minor sub-pocket occupied by Arg⁷⁴ is empty in case of C5a^{des-Arg} (Figure 4E), and three terminal residues, namely Gln⁷¹, Leu⁷², and Gly⁷³, display spatial repositioning compared to C5a (Figure 4F). Although this repositioning in C5a^{des-Arg} compensates for some of the interactions made by C5a, two critical interactions with Thr261^{6,54} and Gly262^{6,55} in TM6 are absent (Figure 4G). Arg⁷⁴ in C5a is involved in most extensive interactions in the binding pocket, and therefore, C5a^{des-Arg} is able to maintain at least some level of transducer coupling through compensating interactions but not at the same level as C5a due to lack of engagement of the Arg⁷⁴ sub-pocket and key missing interactions with TM6 residues.

Structural basis of agonism exhibited by carboxyl-terminal C3a/C5a peptides

In order to gain insights into the mechanism of receptor agonism exhibited by C3a and C5a carboxyl terminus peptides, we analyzed the EP54-C3aR-Go and C5a^{pep}-mC5aR1-Go structures. We observed that similar to the carboxyl terminus of C3a and C5a, EP54 and C5a^{pep} also adopt hook-like conformations and position themselves in a similar binding pocket on the corresponding receptors (Figures 5A and 5B). Specifically, the binding pocket of EP54 and C5a^{pep} on C3aR and C5aR1 are primarily similar to that of C3a and C5a, with buried surface area of 1,640 and 1,140 Å², respectively, and a major involvement of TM2–3, TM5–7, ECL2, and ECL3 (Figures 4C and 4D; Table S2). Moreover, the terminal Arg¹⁰ of EP54 and d-Arg⁶ in C5a^{pep} are positioned in a similar fashion as Arg⁷⁷ and Arg⁷⁴ in C3a and C5a and make similar contacts with key residues in the binding pocket (Figures S1V–S1W). These observations provide a structural rationale for the ability of peptides derived from the carboxyl terminus of C3a and C5a to act as potent receptor agonists and, therefore, also identify ligand-receptor interactions that are critical and sufficient for driving downstream responses. It is also important to note that there are a few interactions that are specific to either EP54 vs. C3a, and C5a^{pep} vs. C5a (Figures 5C and 5D), and it is tempting to speculate that they are responsible for driving lower potency of EP54 and C5a^{pep} compared with C3a and C5a as observed in functional assays (Figures 1B–1P). Finally, the overall binding pose and interactions of EP54 in the C3aR-Gq complex were similar to that in the C3aR-Go complex (Figures S3A–S3E).

Structure-guided reversal of species-specific pharmacology at C5aR1

As presented earlier, we observed only a small difference between human and mouse C5aR1 for C5a-induced G-protein coupling, whereas β arr interaction and trafficking were identical. On the other hand, β arr recruitment and trafficking were dramatically different between the human and mouse receptors upon C5a^{pep} stimulation. Structural superimposition of the human and mouse C5aR1 with respect to C5a- and C5a^{pep}-interacting residues provides the potential structural basis for this observation (Figure 5E). Although the binding poses of C5a^{pep}/C5a on mC5aR1 and C5a on hC5aR1 are similar and the carboxyl terminus hooks occupy similar orientations within the orthosteric pocket, minor rearrangements can be seen in the core helices of C5a (Figure 5F). These helical rearrangements result in significant variation in the docking poses of the N terminus onto C5a in both mouse and human C5aR1 (Figure 5F). Although there are some differences in C5a-interacting residues between the human and mouse receptors, they appear to be minor (Figure 5G). On the other hand, the differences are more

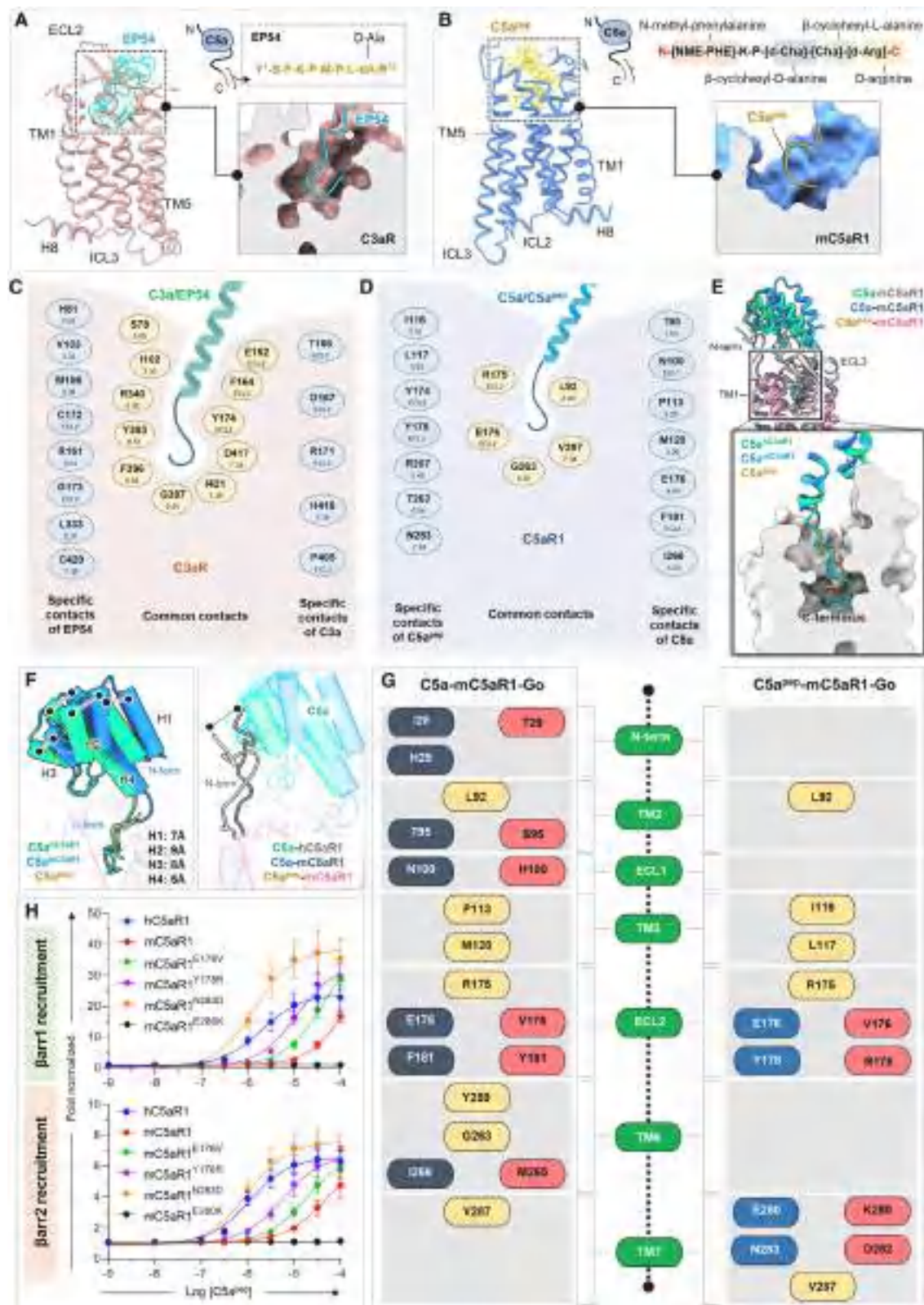
(D) Structural superimposition of C5a- and C5a^{des-Arg}-bound C5aR1 highlighting a similar binding pose. The hook-like C termini of both the ligands are shown within the ligand-binding pocket.

(E) The sub-pocket in C5aR1 occupied by the guanidinium side chain of Arg⁷⁴ is empty in case of C5a^{des-Arg}.

(F) Conformational switching of Leu⁷² and Gly⁷³ located toward the distal end of C5a^{des-Arg} is shown.

(G) Residue contacts by Arg⁷⁴ of C5a in C5aR1 have been shown in circles. Interactions of Gln⁷¹, Leu⁷², and Gly⁷³ compensate for the missing terminal Arg in C5a^{des-Arg} are shown as green circles. Residues highlighted in red do not make any interactions in C5a^{des-Arg}-bound C5aR1 structure.

See also Figures S1 and S7, Table S2, and Methods S1.



(legend on next page)

pronounced in case of C5a^{pep} interacting residues (Figure 5G). For example, Glu176^{ECL2}, Tyr178^{ECL2}, Glu280^{7,31}, and Asn283^{7,34} in mouse C5aR1 are substituted with Val176^{ECL2}, Arg178^{ECL2}, Lys280^{7,31}, and Asp282^{7,34}, respectively, in human C5aR1 (Figure 5G). In the structure of C5a^{pep}-mC5aR1, Tyr178^{ECL2} interacts with d-Cha⁴ in the ligand, and substitution of Tyr with Arg would reverse the polarity in these positions. Likewise, substitution of Glu280^{7,31} and Asn283^{7,34} in mouse C5aR1 with Lys and Asp, respectively, would alter the individual polarity patterns and possibly allow differential interactions in these sites. These alterations between the amino acid sequence of mouse and human C5aR1 may account for the difference observed in β arr recruitment and trafficking between the human and mouse receptors upon C5a^{pep} stimulation (Figures 1J–1P). In fact, mutating these residues in mouse C5aR1 to the corresponding human C5aR1 residues enhances the potency and efficacy of C5a^{pep} in β arr1/2 recruitment, which supports the structural interpretation outlined above (Figures 5H, S1N, and S1O). Although additional studies are required to correlate the structural observations with functional differences, it provides a plausible structural explanation for species-specific differences in the pharmacology of C5aR1.

Agonist-induced activation and G-protein coupling of the complement receptors

The structural snapshots determined in this study provide comprehensive insights into agonist-induced complement receptor activation. Considering that the only inactive-state structure of the complement receptor family is the antagonist-bound C5aR1 structure,^{45,46} we used the same to identify activation-dependent structural features in both C3aR and C5aR1 (Figures 6A, 6B, and S4A). Superimposition of the C3aR structures determined here with antagonist-bound C5aR1 reveals significant conformational changes on the intracellular side of the receptor, including an outward movement of TM6 by approximately 9 Å and an inward shift of TM7 by about 11 Å (Figure 6A). Similar to C3aR, C5aR1 also exhibits the major hallmarks of GPCR activation, including a large outward movement of TM6 by approximately 8 Å, and an inward movement of TM7 by about 10 Å (Figure 6B). Interestingly, in the inactive-state structure of C5aR1, H8 exhibits an inverted orientation and is sandwiched

between TM1 and TM7, whereas in both C3aR and C5aR1, H8 undergoes a dramatic movement of nearly 180° compared with the inactive C5aR1 structure, which potentially facilitates the opening of the transducer-coupling interface on the receptor (Figures 6A and 6B). However, we note that the dramatic reorientation of H8 observed here compared with the crystal structure in the inactive state requires further investigation in future studies, preferably in cellular context to establish its functional significance. In addition, the conserved microswitches in C3aR and C5aR1, including DRY, NPxxY, CWxP, and PIF motifs, exhibit significant structural rearrangements upon activation, as outlined in Figures 6C and 6D. We also note that the overall movement of TM6 and TM7, the rotation of H8, and the activation-dependent conformational changes in the microswitches are conserved in all the structures (Figures S4B–S4H and S5). Taken together, these interactions promote the opening of a cavity toward the cytoplasmic face of the receptor, capable of accommodating G protein and the core interaction with β arrs (Figure S4I).

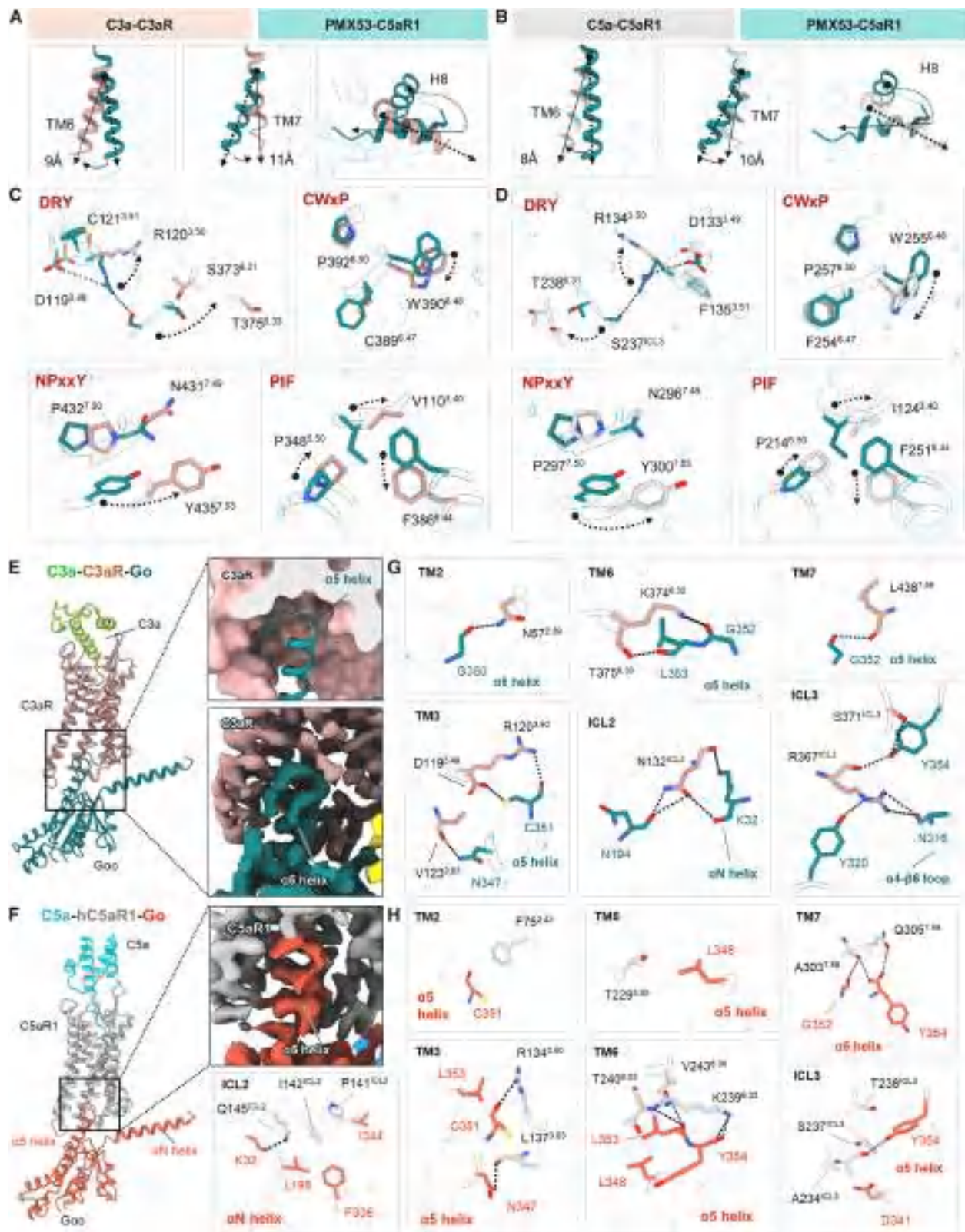
The overall interaction interface between the complement receptors and G α o is very similar in all structures (Figures 6E, 6F, and S6). For example, the carboxyl terminus of the α 5 helix in G α o adopts a loop conformation and inserts into the cytoplasmic core of the receptor with an approximate buried surface area of 2,200 and 1,800 Å² for C3aR and C5aR1, respectively. The key interaction interfaces are composed of TM2, TM3, TM6, TM7, ICL2, and ICL3 in the receptors and α 5 helix, α N helix, and α N- β 1 loop of the G α o subunit (Figures 6G, 6H, and S6). Moreover, the overall engagement and interaction interface of C3aR with Gq is similar to that observed for C3aR-G α o complex, although there are some differences, as well (Figures S3E and S6). A comprehensive list of all the receptor-G-protein interfaces is included in Table S2, and the EM densities of the critical residue-level interactions are presented in Figure S7.

Discovery of a G-protein-biased agonist at C3aR

Based on the structural insights wherein the terminal carboxyl segments of the C3a and C5a appear to be most critical for receptor binding and activation, we tested a set of peptide ligands including C3a and C5a on C3aR and C5aR1 in G-protein-coupling and β arr-recruitment assays (Figures 7A and S1P–S1R). Interestingly, although C3a did not exhibit any measurable

Figure 5. Binding poses of C3a- and C5a-derived peptides on C3aR and C5aR1 and species-specific insights into ligand binding of C5aR1

- (A) Sequence of EP54 derived from the C terminus of C5a (top right). Side view of EP54 (transparent surface) binding to C3aR (ribbon) (left). EP54 docks into C3aR and forms a hook-like structure (bottom right).
- (B) Sequence of C5a^{pep} derived from the C terminus of C5a (top right). Side view of C5a^{pep} (transparent surface) binding to C5aR1 (ribbon) (left). C5a^{pep} docks into C5aR1 and forms a hook-like structure (bottom right).
- (C and D) Comparative analysis of common and specific interactions of C3a/EP54 with C3aR and C5a/C5a^{pep} with C5aR1.
- (E) Superimposition of agonist-C5aR1 structures. C5a and C5a^{pep} in the ligand-binding pocket are shown. (hC5aR1 in surface slice and ligands in cartoon representation.)
- (F) Helical shifts in the core domain of C5a upon binding to mouse and human C5aR1 are shown (left). Conformational changes in the N termini of mouse and human C5aR1 upon interaction with C5a (right).
- (G) Schematic representation of residue contacts between C5a and C5a^{pep} with mC5aR1. Residues present at the orthosteric pocket of ligand binding are depicted. Residues from C5a-hC5aR1 structure shown in red boxes, C5a-mC5aR1 structure in dark gray, C5a^{pep}-mC5aR1 in blue, and common residue contacts at the interface have been shown in yellow boxes. Different regions from the receptors have been highlighted in green boxes.
- (H) Measuring β arr1/2 recruitment in response to C5a^{pep} downstream to a series of mC5aR1 mutants mimicking the corresponding hC5aR1 residues show dramatic increase in both potency and efficacy of β arr1/2 recruitment compared with the wild-type mouse receptor. Data (mean \pm SEM) represent six independent experiments, fold normalized with respect to luminescence observed at the lowest dose (measured as 1) for each receptor.
- See also Figures S1 and S3 and Table S2.



(legend on next page)

transducer-coupling response at C5aR1, C5a exhibited some cross-reactivity with C3aR, albeit only at moderately high concentration (i.e., 1 μ M) in the G-protein-coupling assay (Figure 7A). More importantly, a pentadecameric peptide referred to as EP141 that is derived from and modified based on the carboxyl terminus sequence of C3a (Figure S7E) showed full efficacy for C3aR in G-protein-coupling assay, although it has lower potency compared with C3a (EC50 for C3a vs. EP141, \sim 1.5 vs. 90 nM). On the other hand, it displays only partial efficacy in β arr-recruitment assay compared with C3a (Emax for EP141, \sim 55%–65% of C3a) and lower potency (EC50 for C3a vs. EP141, \sim 10–20 nM vs. \sim 300–500 nM) (Figure 7A). In addition, although EP54 binds to both C3aR and C5aR1, EP141 is selective for C3aR (Figure 7A). The analysis of EP141 responses in G-protein and β arr1/2-recruitment assays yields a bias factor of \sim 0.5 \pm 0.05 (Figure 7A). We also tested EP141 using human monocyte-derived macrophages (HMDMs) and observed that it is a full agonist in terms of intracellular Ca^{2+} response, but unlike C3a, it does not attenuate lipopolysaccharide (LPS)-induced tumor necrosis factor alpha (TNF- α) and interleukin (IL)-6 secretion (Figure 7B). Interestingly, EP141 stimulation of HMDMs leads to an even higher ERK1/2 MAP kinase response than C3a (Figure 7B). This is particularly intriguing as previous studies have suggested that, unlike other GPCRs, β arrs play an inhibitory role in C3aR-induced ERK1/2 phosphorylation,^{12,13} and therefore, lower efficacy of EP141 in terms of β arr recruitment may underlie the enhanced ERK response observed. In order to understand the molecular basis of G-protein bias exhibited by EP141, we reconstituted and determined the structure of EP141-C3aR-G α complex (Figures 7C, S2H, S4E, and S5; Tables S1 and S2). We observed that similar to the carboxyl terminus of C3a and EP54, EP141 also adopts a hook-like conformation to position itself in the orthosteric binding pocket, albeit it is aligned a little differently than EP54 (Figure 7D). Although EP141 exhibits extensive interactions in the orthosteric pocket, similar to those displayed by the analogous residues of C3a and EP54, including the contacts made by terminal Arg⁷⁴, it also displays additional interactions through Arg⁶⁹ that were not previously observed. As presented in Figure 7E, Arg⁶⁹ in EP141 forms a hydrogen bond network with Ser400^{6,58}, Thr403^{6,61}, and Asp404^{ECL3} of the C3aR. These interactions result in an outward movement of the extracellular portions of TM6 and TM7 in C3aR by approximately 1.5 and 2.8 Å, respectively, and also a shift in ECL3 compared with C3a-bound C3aR (Figure 7F). It is interesting to note that Arg⁶⁹ in C3a is a part of an α helix (H4), and thus, perhaps it is structurally constrained and unable to engage in

hydrogen bond networks in a fashion similar to EP141. On the other hand, EP54 contains a lysine at the corresponding position that attains an alternative rotameric conformation, which precludes the possibility of it engaging in the same hydrogen bonding network as EP141. Considering the partial efficacy of EP141 in β arr assays compared with EP54 while maintaining similar potency and efficacy in terms of cAMP response, the structural observations presented here now lay a foundation to rationally design and optimize signaling-biased peptide agonists of C3aR. Taken together, these observations identify EP141 as a C3aR-selective, G-protein-biased agonist and provide a framework for designing additional biased agonists at C3aR. Considering the striking selectivity of EP141 for C3aR, even more than that displayed by C5a, it should allow the delineation of receptor subtype-selective cellular and physiological responses observed upon complement activation, and therefore, it may help refine the precise therapeutic window of selectively blocking the complement receptors. Moreover, these findings also underscore EP141 as a valuable tool compound to decipher the intricate details of biased agonism at C3aR and investigate the physiological outcomes of biased signaling at C3aR that has remained mostly unexplored.

DISCUSSION

The structural snapshots presented here provide the mechanistic basis of C3a and C5a recognition by their cognate receptors, C3aR and C5aR1, and elucidate previously unanticipated distinct binding modes of these anaphylatoxins to these receptors (Figure 7G). Importantly, we also uncover the molecular mechanism that allows the carboxyl-terminal peptide fragments of C3a and C5a to bind and activate C3aR and C5aR1, including the identification of a G-protein-biased agonist at C3aR, which illuminates intricate details of the so-called effector sites on these receptors sufficient for driving downstream functional responses. Moreover, these structures also offer previously lacking templates to facilitate structure-guided ligand discovery with subtype-selective pharmacology and signaling bias, and therefore, taken together with recent studies,^{47,48} they should allow us to overcome a major limitation that has previously stunted the discovery of potential therapeutic molecules targeting the complement receptor in inflammatory disorders. It is also interesting to note that both C3aR and C5aR1 recruit β arrs; however, they appear to have distinct contributions in downstream signaling for these two receptors. For example, β arrs play an inhibitory role in agonist-induced ERK1/2 MAP kinase activation

Figure 6. Mechanism of activation and G-protein coupling at C3aR and C5aR1

(A and B) Dynamic changes in TMs of activated C3aR/C5aR1 compared with the inactive state of C5aR1. The TM6, TM7, and H8 from different receptor complexes shown are from receptors mentioned in boxes. Solid lines (active receptors) and dotted lines (inactive C5aR1) indicate direction of movement. The respective degrees of movements in corresponding TMs have also been mentioned.

(C and D) Close-up views of the conserved DRY, NPxxY, CWxP, and PIF motifs show conformational changes upon C3aR and C5aR1 activation. The names of the hallmark microswitches are noted inside respective boxes. Polar contacts are depicted as black dashed lines.

(E and F) α 5 helix of G α docks into the cytoplasmic core of C3aR and C5aR1. Only receptor and G α are shown in ribbon representations to highlight the binding pose of G proteins with receptor core. Surface slice presentations (top) and cryo-EM maps are shown in inset boxes to highlight the direct docking of G α to receptors.

(G and H) Magnified view of the interactions between TM2, TM3, TM6, TM7, ICL2, and ICL3 of C3aR and hC5aR1 with G α . Ionic bonds are depicted as black dashed lines.

See also Figures S4, S5, S6, and S7 and Table S2.



for C3aR^{12,13} but a supportive role in the case of C5aR1.⁴⁰ Additionally, the second C5a receptor, C5aR2, selectively recruits β arrs without any measurable G-protein-dependent second messenger response, which makes it an example of intrinsically β arr-biased 7TMR.⁴⁰ Still, however, the structural basis of this unique functional manifestation by C5aR2 remains elusive, and therefore, future studies focused on this receptor are required to complete the molecular understanding of complement signaling through 7TMRs (Figure 7G). Taken together, these three complement receptors make a fascinating model system to decipher the intricate details of 7TMR- β arr interaction with distinct functional outcomes.

As mentioned earlier, excessive complement activation is linked to many diseases, and therefore, it continues to remain an important therapeutic area. For example, a small-molecule inhibitor of C5aR1 has recently been approved by the United States Food and Drug Administration for anti-neutrophil cytoplasmic antibody vasculitis (ANCA-vasculitis),⁴⁹ and a monoclonal antibody, marketed under the brand name Gohibic, targeting C5a to block its interaction with C5aR1, has been approved for use in hospitalized patients of COVID-19.²³ Moreover, a cyclic peptide inhibitor of complement C3 has also been approved for paroxysmal nocturnal hemoglobinuria (PNH)⁵⁰ and age-related macular degeneration (AMD).⁵¹ However, the complement cascade also plays numerous roles in host protective mechanisms. For example, in a mouse model, C5a-C5aR1 interaction appears to contribute to the clearance of *Candida albicans* and improved host survival.⁵² Furthermore, although C3a and C5a are broadly labeled as pro-inflammatory, there is now an appreciation that, unlike C5a, C3a exerts many immune-protective activities in the host. For example, following acute organ injury or infection, complement-mediated produc-

tion of C3a confers protection in the brain, kidney, vessels, and intestine.^{53–56} The development of biased ligands for C3a and C5a receptors may therefore allow for the specific protective activities of these receptors to be separated from their pathogenic activities. Indeed, our recent study demonstrates that EP141, the G-protein-biased agonist for C3aR identified here, can protect against tissue injury in a mouse ischemia reperfusion injury model.⁵³ In addition, we have also demonstrated previously that C5a^{pep}, a G-protein-biased agonist for C5aR1, attenuates LPS-mediated cytokine production from macrophages similar to C5a while exhibiting diminished pro-inflammatory response of neutrophil migration.⁴⁰ We also note that cytokines such as TNF- α and IL-6 are linked to inflammatory diseases such as rheumatoid arthritis and Crohn's disease; however, these cytokines are also important for host defense, and current anti-TNF and anti-IL-6 therapies increase infection risk.⁵⁷ Our finding that the endogenous C3a response to these cytokines can be modulated by biased receptor ligands may assist in the development of therapies tuned for different aspects of complement receptor function. Collectively, these observations underscore the therapeutic potential for biased agonists targeting C3aR and C5aR1, and the structural and pharmacological data presented here now provides a framework for exploring this direction going forward.

Limitations of the study

We note that the rotation of H8 observed in the active state structures of C5aR1 presented here, compared with the previously determined crystal structure of antagonist-bound C5aR1, merits further studies to probe its functional significance with respect to receptor signaling and regulation. We also acknowledge that our study is focused primarily on understanding the

Figure 7. Identification of a biased agonist and schematic of complement recognition by C3aR and C5aR1

(A) G-protein activation and β arr1/2 recruitment were studied using GloSensor assay and NanoBIT-based assay (receptor-SmBIT + LgBIT- β arr1/2), respectively, first panel: forskolin-induced cAMP level downstream of C3aR in response to indicated ligands (mean \pm SEM; n = 4; normalized with the lowest ligand concentration for each ligand as 100%); second panel: β arr1 recruitment to C3aR (mean \pm SEM; n = 4); and third panel: β arr2 recruitment to C3aR (mean \pm SEM; n = 3); normalized with the highest ligand concentration of C3a as 100% (top). G-protein activation and β arr1/2 recruitment downstream of C5aR1 in response to indicated ligands, first panel: forskolin-induced cAMP level decrease downstream of C5aR1 in response to indicated ligands (mean \pm SEM; n = 5; normalized with the lowest concentration of each ligand as 100%), second panel: β arr1 recruitment to C5aR1 (mean \pm SEM; n = 5) and third panel: β arr2 recruitment to C5aR1 (mean \pm SEM; n = 4); normalized with the highest ligand concentration of C5a as 100% (bottom). Bias factors (β value) determined taking C3a as reference elucidate the G-protein-biased nature of EP141 have been shown in insets. Bias factor was calculated using <https://biasedcalculator.shinyapps.io/calc/>.

(B) Intracellular calcium response, ERK1/2 activation, cytokine release upon stimulation with C3a, and EP141 were studied in human macrophages. Top left: calcium response (mean \pm SEM; n = 9, data were normalized to the bottom [0%] and top [100%] values of the C3a dose-response curve). Bottom left: ERK1/2 activation (mean \pm SEM; n = 9, normalized to the peak C3a-induced response for that donor). For cytokine release, top right: TNF- α release (mean \pm SEM; n = 4), and bottom right: IL-6 release (mean \pm SEM; n = 5), data were normalized to the medium only (0%) and LPS (100%) triggered response, analyzed using two-way ANOVA, Tukey's multiple comparison test. The exact p values are as follows: for TNF- α release, LPS vs. LPS+C3a: (p = 0.0066), LPS only vs. LPS+EP141: (p = 0.3947). For IL6 release, LPS vs. LPS+C3a: (p = 0.0002), LPS vs. LPS+EP141 (p = 0.1048) (**p < 0.01, ***p < 0.001, ns, non-significant).

(C) Cryo-EM density map and corresponding model of EP141-C3aR-Go. Cryo-EM density map of EP141 has been shown in gray dotted circles (left). Sequence of EP141 derived from the C terminus of C3a. Side view of EP141 (surface) bound to C3aR (ribbon).

(D) The ligand binding pocket has been shown as surface slice representation to highlight the positioning of C3a, EP54, and EP141 on C3aR (right).

(E) Unique residue contacts between EP141 and C3aR at the ligand binding pocket might help explain the phenotypic behavior exhibited by EP141. Arg⁶⁹ of EP141 makes extensive interactions with residues from TM6 and ECL3 (left), which are absent in C3a-C3aR (middle) and EP54-C3aR complexes (right). Black dotted arrows represent probable movements of Arg⁶⁹ in C3a and Lys⁴ in EP54 with respect to Arg⁶⁹ in EP141.

(F) Interactions by Arg⁶⁹ of EP141 with C3aR result in outward movements in TM6, TM7, and ECL3 in C3aR.

(G) Schematic diagram showing ligand binding of the complement receptors C3aR and C5aR1. The C-terminal tail of C3a/C5a changes its conformation to form a hook-like loop upon interacting with C3aR/C5aR1. In addition to the ECLs and the extracellular side of TMs of both the receptors, C5a engages with an extra interface on N terminus of C5aR1. Absence of this interaction network on N terminus might explain the differential positioning of C3a compared with C5a. A group of charged residues on C3aR/C5aR1 interact and stabilize the critical terminal Arg with polar contacts from the C terminus of complement peptides. C5a also binds to C5aR2, but the mechanism behind this interaction has yet to be explored in high detail.

See also Figures S1 and S7, Table S2, and Methods S1.

fundamental mechanism of complement anaphylatoxin recognition by their cognate receptors, and although it paves the way for structure-guided ligand discovery, it does not directly delve into such an exercise. In addition, we also note that this study is focused primarily on C3aR and C5aR1 and not on C5aR2, which remains to be explored in future studies with respect to complement C5a binding, activation, and its intrinsic β arr bias that we have described earlier.^{15,40}

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.cell.2023.09.020>.

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AUTHOR CONTRIBUTIONS

M.K.Y. expressed and purified C3aR and hC5aR1 and reconstituted the complex with G proteins for negative staining and cryo-EM. R.Y. carried out cryo-EM screening, data collection, and structure determination on C3aR and hC5aR1 complexes. J.M. and R.B. carried out negative staining on the complexes, participated in data processing, structure determination, and analyzed the structures. Shirsha Saha and M.K.Y. expressed and purified mC5aR1 and reconstituted the complex with G proteins for negative staining and cryo-EM, and Shirsha Saha carried out the functional assays on C5aR1 with help from P.S. P.S. generated all the constructs for functional assays on C3aR and performed the cellular assays with help from C.S. and Shirsha Saha. C.S. started the expression, purification, and reconstitution of C3aR complexes including purification of G proteins and C3a. V.S. and Sayantan Saha contributed to the purification of G proteins, C3a, C5a, and ScFv16 and also assisted with reconstitution and purification of the complexes. S. Mohapatra and S. Mishra contributed to purification of ScFv16 and C5a. H.A.K. assisted with data collection on C3aR and hC5aR1 complexes. M.C. prepared the grids and collected cryo-EM data on mC5aR1 complexes. X.X.L. and T.M.W. designed and performed the HMDM assays. J.M. prepared all the figures with inputs from R.B., M.K.Y., R.Y., Shirsha Saha, P.S., and M.G. A.K.S. and C.G. supervised and managed the overall project. All authors contributed to data analysis, interpretation, and manuscript writing.

DECLARATION OF INTERESTS

The authors declare no competing interests.

INCLUSION AND DIVERSITY

We support inclusive, diverse, and equitable conduct of research.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Monoclonal ANTI-FLAG M2-HRP antibody	Sigma-Aldrich	Cat# A8592; RRID: AB_439702
Bacterial and virus strains		
<i>E. coli</i> strain BL21(DE3)	New England Biolabs	Cat# C2527H
<i>E. coli</i> strain SHuffle	New England Biolabs	Cat# C3028
<i>E. coli</i> strain Rosetta (DE3)	Sigma-Aldrich	Cat# 70954
Chemicals, peptides, and recombinant proteins		
TRIS	SRL	Cat# 71033
HEPES	SRL	Cat# 63732
NaCl	SRL	Cat# 41721
EDTA	SRL	Cat# 12070
Phenylmethanesulfonyl Fluoride (PMSF)	SRL	Cat# 84375
L-Cysteine Hydrochloride Monohydrate	Sigma Aldrich	Cat# C7880
Iodoacetamide	Sigma Aldrich	Cat# I1149
Imidazole	Sigma Aldrich	Cat# I202-500G
Benzamidine Hydrochloride	SRL	Cat# 93014 (0248255)
Lysozyme	SRL	Cat# 45822
Glycerol	SRL	Cat# 77453
Dithiothreitol	HiMedia	Cat# MB070
Lauryl Maltose Neopentyl Glycol (MNG)	Anatrace	Cat# NG310, CAS no.1257852-96-2
Dodecyl Maltoside (DDM)	Anatrace	Cat# D310A
Cholesteryl Hemisuccinate (CHS)	Sigma Aldrich	Cat# C6512
Paraformaldehyde (PFA)	Sigma Aldrich	Cat# P6148, CAS no. 30525-89-4
Poly-D-lysine	Sigma Aldrich	Cat# P0899
TMB (Tetramethylbenzidine)	Thermo Fisher Scientific	Cat# 34028
Janus Green B	Sigma Aldrich	Cat# 201677
PEI (Polyethylenimine)	Polysciences	Cat# 23966
Bovine Serum Albumin, BSA	SRL	Cat# 83803 (0140105)
FLAG peptide	GenScript	N/A
HBSS - Hank's Balanced Salt Solution	Thermo Fisher Scientific	Cat# 14065
GIBCO Fetal Bovine Serum	Thermo Fisher Scientific	Cat# 10270-106
Iscove's Modified Dulbecco's Medium	Thermo Fisher Scientific	Cat# 12440053
DMEM	Cellclone	Cat# CC3004
Phosphate-buffered saline (PBS)	Sigma Aldrich	Cat# D1283
GIBCO Penicillin-Streptomycin	Thermo Fisher Scientific	Cat# 15140122
ESF921 Insect Cell Culture Medium	Expression Systems	Cat#96-001-01
Coelenterazine	Goldbio	Cat# CZ05
D-Luciferin Sodium Salt	Goldbio	Cat# LUCNA-1G
Glyco-diosgenin (GDN)	Anatrace	GDN101
Coomassie Brilliant Blue	SRL	Cat# 64222
Uranyl formate	Polysciences	Cat# 24762-1
Apyrase	New England Biolabs	Cat# M0398S
Recombinant G α o	This study	N/A
Recombinant G α q	This study	N/A

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Recombinant Gβ1γ2 heterodimer	This study	N/A
Recombinant ScFv16	This study	N/A
Recombinant human C5a	This study	N/A
Recombinant mouse C5a	This study	N/A
Recombinant human C3a	This study	N/A
Recombinant human C5adesArg	This study	N/A
C5a ^{pep}	GenScript	N/A
EP54	GenScript	N/A
EP67	GenScript	N/A
EP141	GenScript	N/A
Recombinant human macrophage colony stimulating factor	BioLegend, San Diego, USA	Cat# 574814
Formvar/carbon coated 300 mesh copper grids	PELCO (Ted Pella)	Cat# 01753-F

Critical commercial assays

Site Directed Mutagenesis Kit	NEB	Cat# E0554
NanoBIT assay	Promega	N/A
GloSensor assay	Promega	N/A
BD OptEIA™ Human IL-6 ELISA Set	BD OptEIA	Cat# 555220
BD OptEIA™ Human TNF ELISA Set	BD OptEIA	Cat# 555221
Fluo-4 NW Calcium Assay Kit	Thermo Fisher Scientific	Cat# F36206
AlphaLISA SureFire Ultra p-ERK1/2 (Thr202/Tyr204) Assay Kit	Perkinelmer	Cat# ALSU-PERK-A-HV

Deposited data

Apo-C3aR-Go complex (Glacios)	This study	PDB: 8I97, EMD-35259
Apo-C3aR-Go complex (Titan)	This study	PDB: 8I9S, EMD-35282
C3a-C3aR-Go complex (Composite map)	This study	PDB: 8I9L, EMD-35275
C3a-C3aR-Go (C3aR-Go complex only, Original Map)	This study	EMD-35293
C3a-C3aR-Go (C3a only, Original Map)	This study	EMD-35294
C5a-hC5aR1-Go (Composite map)	This study	PDB: 8IA2, EMD-35292
C5a-hC5aR1-Go (hC5aR1-Go complex only, original map)	This study	EMD-35295
C5a-hC5aR1-Go complex (C5a only, original map)	This study	EMD-35296
C5a-mC5aR1-Go	This study	PDB: 8HQC, EMD-34947
C5a-pep-mC5aR1-Go	This study	PDB: 8HPT, EMD-34943
EP141-C3aR-Go	This study	PDB: 8J6D, EMD-36001
EP54-C3aR-Go	This study	PDB: 8I95, EMD-35257
EP54-C3aR-Gq	This study	PDB: 8I9A, EMD-35263
C5a-desArg-hC5aR1-Go	This study	PDB: 8JZZ, EMD-36755
Crystal Structure of the Human C3a anaphylatoxin	Bajic et al. ²⁴	PDB: 4HW5
Human formyl peptide receptor 2 in complex with fMLFI and Gi2	Zhu et al. ⁵⁸	PDB: 7WVV
Muscarinic acetylcholine receptor 2-Go complex	Maeda et al. ⁵⁹	PDB: 6OIK
Human c5a in complex with MEDI7814 a neutralizing antibody	Colley et al. ⁶⁰	PDB: 4UU9
Gel images and Functional assay data	This study	Mendeley data https://doi.org/10.17632/km5vft8gkd.1

Experimental models: Cell lines

Human: HEK293	ATCC	Cat# CRL-3216
Human monocyte-derived macrophages (HMDM)	Australian Red Cross LifeBlood Service, Brisbane, Australia	N/A

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
<i>Spodoptera frugiperda</i> (Sf9) Cell line	Expression Systems	Cat# 94-001F
Oligonucleotides		
mC5aR1 (wild type and mutants) cloning in pCAGGS vector_Foward: CGGGGTACCGAGGAGATCTGCCA CCATGGGCAAGACCATCATCGC	This study	N/A
mC5aR1 (wild type and mutants) cloning in pCAGGS vector_Reverse: TCCCCCGGGGACGGCCTGGGACT TGCG	This study	N/A
hC5aR1 cloning in pCAGGS vector_Foward: CGGGGTACCGAGGAGATCTGCCACCATGGGGAAG ACGATCATCGCC	This study	N/A
hC5aR1 cloning in pCAGGS vector_Reverse: TCCC CCGGGCACTGCCTGGGTC	This study	N/A
C3aR cloning in pCAGGS vector_Foward: CGGGGTACCGAGGAGATCTGCCACCATGGGGA GACGATCATCGCC	This study	N/A
C3aR cloning in pCAGGS vector_Reverse: TCCCCCGGGCACTGTGGTAGAGTTTCTC	This study	N/A
mC5aR1_E176V SDM primer Forward: CGTGACAGAggtGCTTACAAGG	This study	N/A
mC5aR1_E176V SDM primer Reverse: AATGATGGGATAGTCAGCAG	This study	N/A
mC5aR1_Y178R SDM primer Forward: CAGAGAGGCTcgcaAGGACTTCTAC	This study	N/A
mC5aR1_Y178R SDM primer Reverse: TACACGAATGATGGGATAG	This study	N/A
mC5aR1_E280K SDM primer Forward: GAAGCGCGTgaagAAGCTGAACA	This study	N/A
mC5aR1_E280K SDM primer Reverse: AGAGTAGGGGATGAGGGAG	This study	N/A
mC5aR1_N283D SDM primer Forward: GGAGAAGCTGgacAGCCTGTGCG	This study	N/A
mC5aR1_N283D SDM primer Reverse: ACGCGCTTCAGAGTAGGG	This study	N/A
Recombinant DNA		
pCDNA_3.1 (empty vector)	Dr. Arun K Shukla	N/A
pcDNA3.1_hC5aR1	Dr. Arun K Shukla	N/A
pcDNA3.1_mC5aR1	This study	N/A
pcDNA3.1_C3aR	This study	N/A
pCAGGS_hC5aR1	This study	N/A
pCAGGS_mC5aR1	This study	N/A
pCAGGS_mC5aR1 ^{E176V}	This study	N/A
pCAGGS_mC5aR1 ^{Y178R}	This study	N/A
pCAGGS_mC5aR1 ^{E280K}	This study	N/A
pCAGGS_mC5aR1 ^{N283D}	This study	N/A
pCAGGS_C3aR	This study	N/A
pCAGGS_LgBiT-βarr1	Dr. Asuka Inoue	N/A
pCAGGS_LgBiT-βarr2	Dr. Asuka Inoue	N/A
pCAGGS_SmBiT-βarr1	Dr. Asuka Inoue	N/A
pCAGGS_SmBiT-βarr2	Dr. Asuka Inoue	N/A
pCAGGS_LgBiT-FYVE	Dr. Asuka Inoue	N/A

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
pVL1393-FLAG-humanC5aR1	GenScript	N/A
pVL1393-FLAG-mouseC5aR1	GenScript	N/A
pVL1393-FLAG-C3aR	GenScript	N/A
pVL1392Dual-β1γ2 vector	GenScript	N/A
Software and algorithms		
Relion3.1.2	Zivanov et al. ^{61,62,63}	https://www3.mrc-lmb.cam.ac.uk/relion/index.php?title=Main_Page
cryoSPARC	Punjani et al. ⁶⁴	https://cryosparc.com/
UCSF Chimera X	Pettersen et al. ⁶⁵	https://www.rbvi.ucsf.edu/chimerax/
UCSF Chimera	Pettersen et al. ⁶⁶	https://www.cgl.ucsf.edu/chimera/
COOT	Casañal et al. ⁶⁷ ; Emsley et al. ⁶⁸ ; Emsley and Cowtan ⁶⁹	https://www2.mrc-lmb.cam.ac.uk/personal/pemsley/coot/
Phenix	Liebschner et al. ⁷⁰ ; Adams et al. ⁷¹	https://www.phenix-online.org/
PDBePISA webserver	Krissinel and Henrick ⁷²	https://www.ebi.ac.uk/pdbe/pisa/
PDBsum	Laskowski et al. ⁷³	http://www.ebi.ac.uk/thornton-srv/databases/pdbsum/
Graphpad Prism v8 and v9	GraphPad Software, San Diego, California USA	https://www.graphpad.com/scientific-software/prism/
Others		
100kDa Cutoff Concentrators	Cytiva	Code# 28932319
30kDa Cutoff Concentrators	Cytiva	Code# 28932317
10kDa Cutoff Concentrators	Cytiva	Code# 28932296

RESOURCE AVAILABILITY

Lead contact

Further information and requests for reagents should be addressed to the lead contact Dr. Arun K. Shukla (arshukla@iitk.ac.in).

Materials availability

ALL unique reagents described in this manuscript are available from the [lead contact](#) with a completed Materials Transfer Agreement.

Data and code availability

- All three-dimensional cryo-EM density maps, coordinates for the atomic models and local-refined maps generated in this study have been deposited and are publicly available as of the date of publication. Accession numbers (EMDB and PDB IDs) are listed in the [key resources table](#). Original gel images have been deposited to Mendeley data, and they are publicly available after publication. The DOI is listed in the [key resources table](#).
- This paper does not report any original code.
- Any additional information required to reanalyze the data reported in this paper is available from the [lead contact](#) upon request.

EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

Human cell line

HEK-293 cells were purchased from ATCC for all the cellular experiments performed in the study. The cell line was examined frequently under the microscope for proper morphology, but they were not authenticated or tested for mycoplasma contamination. They were cultured in DMEM with fetal bovine serum (FBS) at 37°C in 5% CO₂. In this study, any stable, knockout, or knockdown cell lines were not generated, and the details of previously generated cell lines are referenced in the manuscript.

Human monocyte cell-derived macrophages (HMDMs)

Human buffy coat blood from anonymous healthy donors was obtained through the Australian Red Cross LifeBlood Service (Brisbane, Australia) with the experimental approval by The University of Queensland Human Research Ethics Committee (approval reference 2020000559), and monocytes were isolated as mentioned in the subsequent method section. The isolated monocytes

were differentiated in Iscove's Modified Dulbecco's Medium supplemented with 10% FBS, 100 U mL⁻¹ penicillin, 100 µg mL⁻¹ streptomycin and 15 ng mL⁻¹ recombinant human macrophage colony stimulating factor. Further demographic information on donors was not available as per the deed agreement with Australian Red Cross Lifeblood.

Insect cell line

Sf9 cells were obtained from Expression systems, and they were routinely monitored under the microscope for proper morphology, but they were not authenticated or tested for mycoplasma contamination. These cells were maintained in a shaker incubator at 27°C with 135rpm shaking, and sub-cultured in protein-free insect cell medium purchased from Expression Systems.

Bacterial cell culture

Escherichia coli strain DH5alpha were used for plasmid DNA amplification and isolation, and they were cultured in Luria-Bertani (LB) broth at 37°C with shaking at 160rpm. For protein expression, BL21 (DE3), Rosetta (DE3), Shuffle strains of *Escherichia coli* were used, and they were cultured using Luria-Bertani (LB), Terrific Broth (TB), or 2XYT media under the indicated culture conditions (temperature and shaking) as described in the subsequent method sections.

METHOD DETAILS

General chemicals, plasmids, and cell culture

Unless otherwise stated, most standard reagents were purchased from Sigma Aldrich. Dulbecco's Modified Eagle's Medium (DMEM), Phosphate Buffer Saline (PBS), Trypsin-EDTA, Foetal-Bovine Serum (FBS), Hank's Balanced Salt Solution (HBSS), and Penicillin-Streptomycin solution were purchased from Thermo Fisher Scientific. HEK-293T cells (ATCC) were maintained in DMEM (Gibco, Cat. no: 12800-017) supplemented with 10% (v/v) FBS (Gibco, Cat. no: 10270-106) and 100 U mL⁻¹ penicillin and 100 µg mL⁻¹ streptomycin (Gibco, Cat. no: 15140122) at 37 °C under 5% CO₂. Sf9 cells were maintained in protein-free cell culture media purchased from Expression Systems (Cat. no: 96-001-01) at 27 °C with 135 rpm shaking. The cDNA coding region of C3aR was cloned in pcDNA3.1 vector with an N-terminal FLAG tag and in pVL1393 vector with an N-terminal FLAG tag followed by the N-terminal region of M4 receptor (residues 2-23). For the constructs used in NanoBiT assay to monitor C3aR/C5aR1-βarr interaction, SmBiT was fused at the C-terminus of the receptors and LgBiT was fused at the N-terminus of βarrs through sub-cloning. For endosomal localization assay, SmBiT was fused at the N-terminus of βarrs and LgBiT was fused at the N-terminus of the FYVE domain of the human Endofin protein. Additional technical details, assay establishment and optimization using these constructs have been described previously.^{74,75} All DNA constructs were verified by sequencing from Macrogen. EP54, EP67, EP141 and C5a^{pep} were synthesized from GenScript.

GloSensor-based cAMP assay

G-protein activation on agonist stimulation was quantified by GloSensor assay using cAMP level as readout as described previously.⁷⁶⁻⁷⁸ Briefly, HEK-293 cells were co-transfected either with 5 µg of FLAG-tagged C3aR along with 2 µg F22 plasmid or 3.5 µg of FLAG-tagged C5aR1 along with 3.5 µg of F22 plasmid. Post 16-18 h of transfection, cells were trypsinized and harvested, followed by resuspension in assay buffer composed of 1X HBSS, 20 mM of 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES), pH 7.4, and D-luciferin (0.5 mg mL⁻¹) (GoldBio, Cat. no: LUCNA-1G). Cells were seeded in 96-well flat bottom white plate (Corning) at a density of 200,000 cells per 100 µL and incubated at 37 °C for 90 min followed by 30 min incubation at room temperature. Basal readings were taken before ligand stimulation. To study ligand-induced Gi activation, cells were treated with forskolin at 1 µM concentration before stimulation, and readings were recorded until maximum luminescence signal was obtained. For stimulation, ligand concentrations were prepared by serial dilution in 1X HBSS, 20 mM HEPES, pH 7.4. The cells were stimulated with indicated doses of respective ligands. Changes in luminescence were recorded using a microplate reader (BMG Labtech). Data were normalized by treating the minimum agonist concentration as 100% and plotted using nonlinear regression analysis in GraphPad Prism software.

Surface expression assay

To study receptor surface expression, whole cell-based receptor surface ELISA was performed as previously discussed.⁷⁹ Briefly, cells transfected with receptor construct for respective assays were seeded in a 24-well with 0.01% poly-D-Lysine pre-coated plate at a density of 0.2 million cells well⁻¹ and incubated at 37°C for 24 h. After 24 h, the plate was removed, growth media was aspirated, and the plate was washed with ice-cold 1X TBS, followed by 20 min of fixation with 4% PFA (w/v in 1X TBS) on ice. After fixation, cells were washed thrice with 1X TBS (400 µL in each wash), followed by blocking with 1% BSA prepared in 1X TBS at room temperature for 90 min. Afterward, cells were incubated for 90 min with anti-FLAG M2-HRP (prepared in 1% BSA, 1:5000) (Sigma, Cat. no: A8592). Following antibody incubation, cells were washed thrice with 1% BSA (in 1X TBS). Thereafter, assay was developed by incubating cells with 200 µL of TMB-ELISA (Thermo Scientific, Cat no: 34028) until the light blue color appeared, followed by quenching with 100 µL of 1 M H₂SO₄ by transferring the blue-colored solution to a 96-well plate. Absorbance was measured at 450 nm using a multi-mode plate reader. Afterward, cells were washed twice with 200 µL of 1X TBS and then incubated with 0.2% Janus Green (Sigma; Cat no: 201677) w/v for 15 min. The excess stain was removed by three washes with distilled water. The stain was eluted

by adding 800 μL of 0.5 N HCl per well. 200 μL of the eluted solution was transferred to a 96-well plate, and absorbance was recorded at 595 nm. The signal intensity was normalized by calculating the ratio of A450/A595 values. For data normalization, the ratio of A450/A595 values was calculated, followed by considering pcDNA transfected cells reading as 1, and receptor expression was calculated with respect to pcDNA. Data were analyzed in GraphPad Prism software.

NanoBiT-based β arr-recruitment assay

Agonist-induced β arr1/2 recruitment to the plasma membrane downstream of C3aR and C5aR1 was measured by luminescence-based enzyme-linked complementation assay (NanoBiT-based assay) following the protocol described earlier.^{80,81} Briefly, HEK-293 cells were transfected with C3aR (5 μg) and C5aR1 (1 μg of either mC5aR1 or hC5aR1) harboring carboxyl-terminus fusion of SmBiT and β arr1/2 constructs (2 μg for C3aR and 3.5 μg for C5aR1) with N-terminal fusion of LgBiT using transfection reagent poly-ethyleneimine (PEI) at DNA:PEI ratio of 1:3. Post 16–18 h of transfection, cells were trypsinized, harvested, and resuspended in the NanoBiT assay buffer (HBSS containing 0.01% BSA, 5 mM HEPES, pH 7.4, and 10 μM coelenterazine (GoldBio, Cat no: CZ05). After resuspension, 0.1 million cells per 100 μL were seeded in the flat-bottom white 96-well plate. The plate was incubated at 37 $^{\circ}\text{C}$ for 90 min, followed by 30 min at room temperature. After 2 h of incubation, three cycles of luminescence reading were taken before ligand addition in a multi-mode plate reader (BMG Labtech). Ligand concentrations were prepared by serial dilution in 1X HBSS, 20 mM HEPES, pH 7.4. The cells were stimulated with indicated doses of respective ligands followed by measurement of luminescence signal using a multi-mode plate reader for 20 cycles, and average data of 5 cycles showing highest range of luminescence were used for analysis and presentation. For measuring β arr1/2 recruitment to mC5aR1 mutants, following amount of DNA was used for each of the FLAG-tagged receptors harbouring carboxyl-terminus fusion of SmBiT: 1 μg for hC5aR1/mC5aR1, 2.5 μg for mC5aR1^{E176V}/mC5aR1^{Y178R} and 3.5 μg for mC5aR1^{N283D}/mC5aR1^{E280K} along with 3.5 μg of either β arr1/2 harbouring N-terminal fusion of LgBiT.

Agonist-induced endosomal trafficking of β arrs

Agonist-induced β arr1/2 trafficking to the endosomes was monitored using NanoBiT assay following the same protocol as described above for β arr recruitment. HEK-293 cells were transfected with β arr1/2 tagged with SmBiT at the N-terminus, and N-terminal LgBiT-tagged FYVE constructs were used for enzyme complementation. The amount of DNA for receptor, β arr1/2 and FYVE was kept at 1 μg (for hC5aR1 and mC5aR1) or 5 μg (for C3aR), 2 μg , and 5 μg , respectively.

Assays with HMDMs

HMDMs were generated and cultured as previously described,^{33,82} with experiments approved by The University of Queensland Human Research Ethics Committee. Briefly, human buffy coat blood from anonymous healthy donors was obtained through the Australian Red Cross LifeBlood Service (Brisbane, Australia). Monocytes were isolated using Lymphoprep density centrifugation (STEMCELL, Melbourne, Australia) followed by CD14⁺ MACS magnetic bead separation (Miltenyi Biotec, Sydney, Australia). The isolated monocytes were differentiated for 7 days in Iscove's Modified Dulbecco's Medium supplemented with 10% FBS, 100 U mL⁻¹ penicillin, 100 μg mL⁻¹ streptomycin (Thermo Fisher Scientific, Melbourne, Australia) and 15 ng mL⁻¹ recombinant human macrophage colony stimulating factor (BioLegend, San Diego, USA) on 10 mm square dishes (Bio-strategy, Brisbane, Australia). Non-adherent cells were removed by washing with DPBS, and the adherent differentiated HMDMs were harvested by gentle scraping.

The immunomodulatory effect of human C3a (Merck) and EP141 on lipopolysaccharide (LPS)-induced cytokine release was assessed in HMDMs as previously described.^{13,28} HMDMs were seeded in 96-well tissue culture plates (100,000 /well) for 24 h before treatment. All ligands were prepared in serum-free IMDM containing 0.1% BSA. For stimulation, cells were co-treated with 10 ng mL⁻¹ LPS and C3a (10 nM) or EP141 (100 nM) for 24 h (37 $^{\circ}\text{C}$, 5% CO₂). The supernatant was collected and stored at -20 $^{\circ}\text{C}$ till further use. The IL-6 and TNF- α levels in the supernatant were quantified using respective human enzyme-linked immunosorbent assay (ELISA) kits (BD OptEIA) as per the manufacturer's protocol.

Ligand-induced intracellular calcium mobilization was assessed using Fluo-4 NW Calcium Assay kit following the manufacturer's instructions (Thermo Fisher Scientific, Melbourne, Australia). Briefly, HMDMs were seeded (70,000/well) in black clear-bottom 96-well tissue culture plates overnight. Cells were firstly stained with the Fluo-4 dye in assay buffer (1X HBSS, 20 mM HEPES) for 45 min (37 $^{\circ}\text{C}$, 5% CO₂). C3a and EP141 dilutions were prepared in assay buffer containing 0.1% BSA. On a Flexstation 3 platform, the fluorescence (Ex/Em: 494/516 nm) was continually monitored for a total of 100 s with ligand addition performed at 16 s.

The ligand-induced phospho-ERK1/2 signaling was assessed using the AlphaLISA Surefire Ultra p-ERK1/2 (Thr202/Tyr204) kit (PerkinElmer, Melbourne, Australia) as previously described.^{33,82} Briefly, HMDMs were seeded (50,000/well) in tissue culture-treated 96-well plates for 24 h and serum-starved overnight. C3a and EP141 serial dilutions were prepared in serum-free IMDM containing 0.1% BSA. For stimulation, cells were incubated with C3a or EP141 for 10 min at room temperature and then immediately lysed using AlphaLISA lysis buffer on a microplate shaker (450 rpm, 10 min). For the detection of phospho-ERK1/2 content, cell lysate (5 μL /well) was transferred to a 384-well ProxiPlate (PerkinElmer) and added to the donor and acceptor reaction mixes (2.5 μL /well, respectively) with 2-h incubation at room temperature in the dark. The plate was read on Tecan Spark 20M following standard AlphaLISA settings.

Expression and purification of C3a and C3aR

Gene encoding human C3a was cloned in pET-32a(+) vector with a Trx-6X-His tag at the N-terminal end and purified following the previously described protocol for C5a purification with some modifications.^{24,25} Briefly, freshly transformed *E. coli* SHuffle cells were inoculated in 50 mL 2XYT media with 100 $\mu\text{g mL}^{-1}$ ampicillin for starter culture at 30°C. Overnight grown primary culture was inoculated into 1.5 L 2XYT media with 100 $\mu\text{g mL}^{-1}$ ampicillin, and the culture was allowed to grow at 30°C. At O.D. \sim 0.6, culture was induced with 1 mM IPTG and shifted to 16 °C for overnight induction. Cells were harvested and incubated with 1 mg mL⁻¹ lysozyme in 50 mM HEPES, pH 7.4, 300 mM NaCl, 20 mM Imidazole, 1 mM PMSF, and 2 mM Benzamidine for 2 h at 4°C. Cells were disrupted with ultrasonication, and cell debris was removed with high-speed centrifugation. C3a was enriched on Ni-NTA resins using gravity flow. Nonspecific proteins were removed with extensive washing (50 mM HEPES, pH 7.4, 1 M NaCl, 20 mM Imidazole), and fusion-C3a was eluted with 50 mM HEPES, pH 7.4, 150 mM NaCl, 300 mM Imidazole. Trx-his tag was cleaved with 8–10 h TEV treatment (1:20 w/w, TEV: Fusion protein) at room temperature. Purified C3a was further cleaned by cation-exchange chromatography and stored at -80°C with 10% final glycerol concentration.

Codon-optimized human C3aR was expressed in *Spodoptera frugiperda* (Sf9) cells using the baculovirus expression system with an N-terminal FLAG tag to facilitate purification. The receptor was purified as described previously. Briefly, the insect cells were harvested at 72 h post-infection and lysed by sequential douncing in a low-salt buffer (20 mM HEPES, pH 7.4, 10 mM MgCl₂, 20 mM KCl, 1 mM PMSF, and 2 mM Benzamidine), high salt buffer (20 mM HEPES, pH 7.4, 1 M NaCl, 10 mM MgCl₂, 20 mM KCl, 1 mM PMSF, and 2 mM Benzamidine), and lysis buffer (20 mM HEPES, pH 7.4, 450 mM NaCl, 2 mM CaCl₂, 1 mM PMSF, 2 mM Benzamidine and 2 mM Iodoacetamide). After lysis, the receptor was solubilized for 2 h at 4°C with continual stirring in a solution of 0.5% L-MNG (Anatrace, Cat. no: NG310) and 0.1% cholesteryl hemisuccinate (Sigma, Cat. no: C6512). Post solubilization, salt concentration was lowered to 150 mM, and the receptor was purified on M1-FLAG column. In order to remove nonspecific proteins from FLAG beads, three washes of low salt buffer (20 mM HEPES, pH 7.4, 2 mM CaCl₂, 0.01% CHS, 0.01% L-MNG) were alternated with two washes of high salt buffer (20 mM HEPES, pH 7.4, 450 mM NaCl, 2 mM CaCl₂, 0.01% L-MNG) after binding. The bound receptor was eluted with FLAG elution buffer (20 mM HEPES, pH 7.4, 150 mM NaCl, 0.01% MNG, 2 mM EDTA, and 250 $\mu\text{g mL}^{-1}$ FLAG peptide) and alkylated with iodoacetamide to prevent aggregation. The purified receptor was concentrated using a 30 kDa MWCO concentrator and stored at -80°C in 10% glycerol till further use. 100 nM of hC3a or 1 μM of EP54 and EP141 were kept in all steps of receptor purification.

Expression and purification of C5a, C5a^{des-Arg}, and C5aR1

Genes encoding human C5a and C5a^{des-Arg} were cloned in pET-32a(+) vector with a Trx-6X-His tag at the N-terminal end and purified following previously described protocol with slight modification.^{25,40} After Ni-NTA purification, we directly proceeded to TEV cleavage followed by cation-exchange chromatography. Codon-optimized human and mouse C5aR1 (hC5aR1 and mC5aR1) were expressed in *Spodoptera frugiperda* (Sf9) cells using baculovirus expression system with an N-terminal FLAG tag to facilitate purification. The receptor was purified as described previously.¹³ Briefly, 72h post-infection, insect cells were harvested and lysed by sequentially douncing in low salt buffer (20mM HEPES pH 7.4, 10mM MgCl₂, 20mM KCl, 1mM PMSF, and 2mM Benzamidine), high salt buffer (20mM HEPES pH 7.4, 1M NaCl, 10mM MgCl₂, 20mM KCl, 1mM PMSF, and 2mM Benzamidine), and lysis buffer (20mM HEPES pH 7.4, 450mM NaCl, 2mM CaCl₂, 1mM PMSF, 2mM Benzamidine and 2mM Iodoacetamide). After lysis, receptor was solubilized in 0.5% L-MNG (Anatrace, Cat. no: NG310) and 0.1% cholesteryl hemisuccinate (Sigma, Cat. no: C6512) for 2 h at 4°C, under constant stirring. Post-solubilization, salt concentration was lowered to 150mM, and the receptor was purified on M1-FLAG column. After binding, FLAG beads were washed alternately with three washes of low salt buffer (20mM HEPES pH 7.4, 2mM CaCl₂, 0.01% CHS, 0.01% L-MNG) and two washes of high salt buffer (20mM HEPES pH 7.4, 450mM NaCl, 2mM CaCl₂, 0.01% L-MNG) to remove non-specific proteins. The bound receptor was eluted with FLAG elution buffer (20mM HEPES pH 7.4, 150mM NaCl, 0.01% MNG, 2mM EDTA, and 250 $\mu\text{g mL}^{-1}$ FLAG peptide) and alkylated with iodoacetamide to prevent aggregation. The purified receptor was concentrated using a 30kDa MWCO concentrator and stored at -80°C in 10% glycerol till further use. 100 nM of C5a and C5a^{des-Arg}, or 1 μM of C5a^{dep} were kept in all steps of receptor purification.

Expression and purification of G proteins

Construct for miniG α o and miniG α q were synthesized as described previously^{83,84} with additional ScFv16 binding sequence at N-terminus of miniG α o. Briefly Genes for miniG α o1 and miniG α q subunit were cloned in pET-15b(+) vector with an in-frame 6X-His tag at the N-terminal and expressed in *E. coli* BL21(DE3) cells.^{83,84} A starter culture in LB media was grown at 37°C for 6–8 h at 220 rpm. This was followed by an overnight primary culture at 30°C with 0.2% glucose supplementation. 15 mL primary culture was inoculated in 1.5 L TB (Terrific Broth) media and induced with 50 μM IPTG at an O.D of 0.8 and cultured at 25°C for 18–20 h. Cells were lysed in lysis buffer (40 mM HEPES, pH 7.4, 100 mM NaCl, 10 mM Imidazole, 10% Glycerol, 5 mM MgCl₂, 1 mM PMSF, 2 mM Benzamidine) in the presence of 1 mg mL⁻¹ lysozyme, 50 μM GDP and 100 μM DTT. Cell debris was pelleted down by centrifuging at 18000 rpm for 30 min at 4°C. Protein was enriched on Ni-NTA bead and after washing extensively with wash buffer (20 mM HEPES, pH 7.4, 500 mM NaCl, 40 mM Imidazole, 10% Glycerol, 50 μM GDP and 1 mM MgCl₂), eluted with elution buffer (20 mM HEPES, pH 7.4, 100 mM NaCl, 10% Glycerol, 500 mM Imidazole). His tag was cleaved by overnight TEV treatment at room temperature (1:20, TEV: protein), and untagged protein was recovered by size exclusion chromatography on Hi-Load Superdex 200 PG 16/600 column (Cytiva, Cat. no. 17517501). Fractions corresponding to cleaved protein were pooled, analyzed on SDS-PAGE, and stored at -80°C with 10% glycerol. The genes encoding the G β 1 and G γ 2 subunits cloned in a bi-cistronic pVL1932-

based vector with N-terminal histidine tag at the G β 1 were expressed in *Sf9* cells using the baculovirus expression system. Cells were harvested 72 h after infection and resuspended in lysis buffer (20 mM Tris-Cl, pH 8.0, 150 mM NaCl, 10% Glycerol, 1 mM PMSF, 2 mM Benzamidine and 1 mM MgCl₂). The cells were lysed via douncing and centrifuged at 4°C for 40 min at 18000 rpm. Pellet was resuspended and dounced in solubilization buffer (20 mM Tris-Cl, pH 8.0, 150 mM NaCl, 10% Glycerol, 1% DDM, 5 mM β -ME, 10 mM Imidazole, 1 mM PMSF, and 2 mM Benzamidine) and solubilized at 4°C under constant stirring for 2 h. Cell debris was pelleted down by centrifuging at 20000 rpm for 40 min at 4°C. Protein was enriched on Ni-NTA resin, and after extensive washing with wash buffer (20 mM Tris-Cl, pH 8.0, 150 mM NaCl, 30 mM Imidazole, 0.02% DDM), the protein was eluted with 300 mM Imidazole in 20 mM Tris-Cl, pH 8.0, 150 mM NaCl, 0.01% MNG. Eluted protein was concentrated with a 10 kDa MWCO concentrator (Cytiva Cat no: GE28-9322-96) and stored at -80°C with 10% glycerol.

Expression and purification of ScFv16

Gene encoding ScFv16 was cloned in pET-42a (+) vector with an in-frame N-terminal 10X-His-MBP tag followed by a TEV cleavage site and expressed in *E. coli* Rosetta (DE3) strain.⁸⁵ Overnight primary culture was sub-cultured in 1L 2x YT media supplemented with 0.5% glucose and 5 mM MgSO₄. At O.D₆₀₀ ~0.6, culture was induced with 250 μ M IPTG for 16–18 h at 18°C. Cells were resuspended in 20 mM HEPES, pH 7.4, 200 mM NaCl, 2 mM Benzamidine, and 1 mM PMSF and incubated at 4°C for 1 h with constant stirring. Cells were disrupted by ultrasonication, and cell debris was removed by centrifugation at 18000 rpm for 40 min at 4°C. Protein was enriched on Ni-NTA resins, and non-specifically bound proteins were removed by extensive washing (20 mM HEPES, pH 7.4, 200 mM NaCl, 50 mM Imidazole). Bound protein was eluted with 300 mM Imidazole in 20 mM HEPES, pH 7.4, 200 mM NaCl. Subsequently, Ni-NTA elute was enriched on amylose resin (NEB, Cat. no: E8021L) and washed with buffer (20 mM HEPES pH 7.4, 200 mM NaCl) to remove nonspecific proteins. Protein was eluted with 10 mM maltose (prepared in 20 mM HEPES, pH 7.4, 200 mM NaCl), and the His-MBP tag was removed by overnight treatment with TEV protease. Tag-free ScFv16 was recovered by passing TEV-cleaved protein through Ni-NTA resin. Eluted protein was concentrated and cleaned by size exclusion chromatography on Hi-Load Superdex 200 PG 16/600 column (Cytiva Life sciences, Cat. no: 17517501). Purified protein was flash-frozen and stored at -80°C with 10% glycerol.

Reconstitution of the C3a/EP54/EP141-C3aR-G α /G α q-G β γ -ScFv16 complexes

Purified C3aR was incubated with 1.2 molar excess of G α , G β 1 γ 2, and ScFv16 at room temperature for 2 h in the presence of 25 mU mL⁻¹ apyrase (NEB, Cat. no: M0398S) and either C3a or EP54 or EP141 for complex formation. The G-protein complex was separated from unbound components by loading on Superose 6 increase 10/300 GL SEC column and analyzed by SDS PAGE. Complex fractions were pooled and concentrated to ~10 mg mL⁻¹ using a 100 MWCO concentrator (Cytiva, Cat. no: GE28-9323-19) and stored at -80°C until further use.

Reconstitution of the C5a/C5a^{pep}-C5aR1-G α β 1 γ 2-ScFv16 complexes

Purified m/hC5aR1 was incubated with 1.2 molar excess of G α , G β 1 γ 2, and ScFv16 at room temperature for 2 h in the presence of 25 mU mL⁻¹ apyrase (NEB, Cat. no: M0398L) and either C5a/C5a^{pep}/C5a^{des-Arg} for complex formation. The G-protein complex was separated from unbound components by loading on Superose 6 increase 10/300 GL SEC column and analyzed on SDS PAGE. Complex fractions were pooled and concentrated to ~10mg mL⁻¹ using a 100MWCO concentrator (Cytiva, Cat. no: GE28-9323-19) and stored at -80°C until further use.

Single-particle, negative-stain electron microscopy

In order to confirm homogeneity and complex formation, negative stain electron microscopy was performed on all the samples before proceeding on with grid preparation for cryo-EM data collection. The individual samples were diluted to 0.02 mg mL⁻¹ just prior to grid preparation and 3 μ L of the samples were dispensed onto the carbon side of a glow discharged carbon/formvar coated 300 mesh Cu grids (PELCO, Ted Pella). The extra protein sample was blotted off after incubation for 1 min using a filter paper. The grid with the adsorbed protein sample was then touched on a first drop of 0.75% uranyl formate stain, and immediately blotted off using a filter paper. The grid was then touched onto a second drop of stain and moved gently in a rotating fashion for 30 s to increase the efficiency of staining. The grid so prepared was then left in a desiccator or on the bench in a petri-plate for air drying. Data collection was performed with a FEI Tecnai G2 12 Twin TEM (LaB6) operating at 120 kV and equipped with a Gatan CCD camera (4k x 4k) at 30,000x magnification. Processing of the collected dataset was performed with Relion 3.1.2^{61–63} where almost 10,000 particles were autopicked and subjected to reference free 2D classification, generating the 2D class averages.

Cryo-EM grid preparation and data collection

Purified ligand-C3aR-Go complexes were applied onto glow-charged grids (EasiGlow, 20 mA current with 40 s glow and 10 s hold) at ~10 mg mL⁻¹ concentration and blotted for 5–7 s followed by plunge-freezing into liquid ethane using a Vitrobot MarkIV (Thermo Fisher Scientific, USA) operating at 100% humidity. Briefly, ligand free-C3aR-Go and EP54-C3aR-Go complexes were applied onto Quantifoil R1.2/1.3 Au 300-mesh grids and blotted for 5 s at 4°C. For C3a-C3aR-Go, EP54-C3aR-Gq, C5a-hC5aR1-Go, EP141-C3aR-Go, and C5a^{des-Arg}-hC5aR1-Go complexes, the sample was applied onto Quantifoil R1.2/1.3 Au 200-mesh grids and blotted for 5–7 s at either 4 or 22°C. Cryo-EM data were collected using a Glacios microscope operating at 200 kV with a Falcon

4 direct electron detector operating in counting mode at nominal magnification of 150,000x resulting in pixel size of 0.92 Å using EPU. An additional dataset of ligand free-C3aR-Go was collected using Titan Krios operating at 300 kV with a Gatan K3 direct electron detector operating in counting mode at 105,000 magnifications with a pixel size of 0.86 Å (2-fold binned) using SerialEM. Movies were recorded with a defocus range of -0.8 to -3.0 µm and total dose of ~50 e⁻/Å². Additional data collection parameters are listed in [Figures S4, S5, S6, S7, and S8](#). Total 4,611 movies for ligand-free-C3aR-Go (Glacios), 5,740 for ligand free-C3aR-Go (Titan), 20,051 movies for C3a-C3aR-Go, 4,614 movies for EP54-C3aR-Go, 4,445 movies for EP54-C3aR-Gq, 10,151 movies for C5a-hC5aR1-Go, 9,276 movies for C5a^{des-Arg}-hC5aR1-Go, and 15,216 movies for EP141-C3aR-Go were recorded.

For the cryo-EM data collection on mouse C5aR1 complexes, 3 µL of the purified complexes of C5a-mC5aR1-Go or C5a^{pep}-mC5aR1-Go were applied onto glow discharged Quantifoil holey carbon grids (Au, R2/1 M300) and vitrified using a Vitrobot Mark IV (Thermo Fisher Scientific, USA) operating at 10°C and maintained at 90% humidity. Data collection was performed with a Titan Krios electron microscope (ThermoFisher Scientific, USA) operating at 300 kV equipped with Gatan Energy Filter. Movies were recorded in counting mode with a Gatan K2 Summit direct electron detector DED (Gatan, USA) using the automated SerialEM software at a nominal magnification of 165,000x and a pixel size of 0.82 Å at the specimen level. Total 22,014 movie stacks for C5a-mC5aR1-Go and 24,711 movie stacks for C5a^{pep}-mC5aR1-Go consisting of 40 frames were collected with a defocus value in the range of 0.5 to 2.5 µm with a total accumulated dose of 42 e⁻/Å² and total exposure time of 4 s.

Image processing and map construction

The overall cryo-EM data processing pipeline for all structures are shown in [Figures S3 and S4](#). Data processing for EP54-C3aR-Go, C3a-C3aR-Go, Apo-C3aR-Go, EP54-C3aR-Gq, C5a-hC5aR1-Go and EP141-C3aR-Go complexes were performed in cryoSPARC version 4.0,⁶⁴ whereas that of C5a^{pep}-mC5aR1-Go and C5a-mC5aR1-Go were performed with cryoSPARC version 3.3.2.⁶⁴ Cryo-EM movie stacks were aligned using Patch motion correction (multi) followed by CTF estimation with Patch CTF estimation (multi). Micrographs were curated based on CTF resolution and selected micrographs were used for particle picking using blob-picker.

For the apo-C3aR-Go complex, two independent datasets were collected – one with Titan Krios microscope operating at 300 kV and the other with a Glacios microscope operating at 200 kV. For the Glacios dataset, 2D class averages with clear secondary features were selected to prepare a sub-set of 693,550 particles for further processing. Subsequent ab-initio reconstruction and 3D/Heterogeneous classification with C1 symmetry yielded the best class with 464,408 particles which was further refined to an overall resolution of 3.19 Å (voxel size of 1.063 Å) with NU refinement. For the apo-C3aR-Go Titan dataset, total 754,251 particles corresponding to 2D averages with clear secondary structural features were selected, re-extracted with a box size of 416 px and Fourier cropped to 256 px (pixel size 1.40 Å). This clean particle stack was used for multiclass ab-initio reconstruction followed by heterogeneous classification. Non-uniform refinement with 327,193 particles from the best 3D class yielded a map with a final estimated global resolution of 3.26 Å (voxel size of 1.40 Å).

For the C3a-C3aR-Go dataset, post-2D classification, 922,698 selected particles were extracted with a box size of 416 px and fourier cropped to 360 px (pixel size of 1.063 Å). This clean particle stack was then subjected to ab-initio reconstruction and heterogeneous refinement. The 3D class containing 418,953 particles with evident secondary features were subjected to non-uniform refinement, followed by local refinement with mask to exclude the micelle resulting in a final map at 3.18 Å resolution (voxel size of 1.063 Å). The density corresponding to C3a was noisy, therefore, local refinement was performed with a mask covering the C3a to improve the interpretability of the map in this region yielding a map with global resolution of 4.55 Å. The two local refinement maps were combined to obtain a composite map which was used for subsequent model building.

For the EP54-C3aR-Go complex dataset, a clean particle stack of 767,052 from the 2D classification step was selected and re-extracted with a box size of 360 px and fourier cropped to 256 px (pixel size of 1.495 Å). This particle stack was subjected to ab-initio reconstruction, followed by heterogeneous refinement with C1 symmetry. Total 600,173 high-quality particles with evident structural features were selected and subjected to non-uniform refinement with C1 symmetry, followed by local refinement with a mask on the micelle. This led to a reconstruction with a global estimated resolution of 2.88 Å (voxel size of 0.92 Å). Local resolution estimation was performed with the Blocres sub-program within cryoSPARC version 4.0. Maps were sharpened using the “Autosharpen” sub-program within the Phenix suite⁸² for better visualization and model building.

Initial processing of EP54-C3aR-Gq micrographs showed limited top/bottom views. Therefore, a conventional neural network-based method TOPAZ,⁸³ implemented in cryoSPARC, was used for particle picking. Briefly, Topaz was trained on 37,054 particles picked from denoised micrographs with the ResNet8 convolutional neural network model using down-sampling factor 8, and the expected number of particles per micrograph was set to 500. A total of 2,007,547 particles were picked, extracted, and subjected to iterative rounds of reference-free 2D classification followed by multiclass ab-initio reconstruction/heterogeneous refinement. The best 3D class with 101,400 particles was selected for subsequent non-uniform refinement, which yielded a map of 3.57 Å resolution.

For the C5a-hC5aR1-Go complex, 10,259,948 particle projections were selected and re-extracted with a box size of 416 px, fourier cropped to 256 px (pixel size of 1.50 Å). These extracted particles were used for generating 2 ab-initio models for subsequent heterogeneous refinement. The best class containing 292,441 particle projections were subjected to non-uniform refinement to obtain a density map with nominal resolution of 3.21 Å (voxel size of 1.06 Å). Density corresponding to C5a was poorly resolved, thus local refinement was done with a mask covering the C5a density to improve the interpretability of the map. The focused refinement against C5a yielded a map with final global resolution of 3.88 Å. The two maps were combined with “Combine focus maps” within Phenix to obtain a composite map which was used for model building.

For the C5a-mC5aR1-Go dataset, automated particle picking with blob-picker resulted in 2,601,754 particles which were extracted with a box size of 360 px and fourier cropped to box size of 64 (pixel size of 4.61). These particles were then subjected to several rounds of 2D classification and class averages with clear conformations of the complex were selected and extracted with a box size of 360 px and fourier cropped to 256 px (pixel size of 1.15). These clean set of particles were subjected to Ab-initio reconstruction and heterogeneous refinement yielding 3 models. A particle stack with 173,416 particles corresponding to the 3D class with evident secondary features were re-extracted with full box size of 416 px and fourier cropped to 360 px. This was followed by non-uniform refinement and local refinement with mask on the complex resulting in a final map at 3.89 Å resolution (voxel size of 0.9476 Å).

For C5a^{pep}-mC5aR1-Go complex, 1,886,363 particles were autopicked with the blob-picker sub-program within the cryoSPARC suite, extracted with a box size of 360 px and fourier cropped to 64 px (pixel size of 4.61) for reference free 2D classification. Several rounds of iterative 2D classification yielded class averages representing different orientations of the complex. A subset of 835,654 clean particles from the 2D classification were re-extracted with a box size of 360 px and fourier cropped to 256 px (pixel size of 1.15). This was followed by Ab-initio reconstruction and heterogeneous refinement with C1 symmetry yielding 3 models. 380,463 particles corresponding to the class with clear complex conformation were re-extracted with full box size of 416 px, fourier cropped to 360 px and subjected to non-uniform refinement followed by local refinement with mask on the complex excluding the micelle yielding a reconstruction at 3.39 Å resolution (voxel size of 0.9476 Å).

For the C5a^{des-Arg}-hC5aR1-Go dataset, 8,932 motion corrected micrographs with CTF fit resolution better than 4 Å were selected for further processing. 8,059,635 particles were autopicked with the blob-picker subprogram, extracted with a box size of 360 px (fourier cropped to 64 px) and subjected to 2D classification. 1,013,163 particles corresponding to the clean 2D class averages were selected, re-extracted with a box size of 360 px (fourier cropped to 180 px) and subjected to ab-initio reconstruction and subsequent heterogeneous refinement yielding 3 classes. 860,786 particles corresponding to the best class with clear secondary features were re-extracted with full box size of 360 px and subjected to non-uniform refinement yielding a final reconstruction with an overall resolution of 3.31 Å at 0.143 cutoff.

For the EP141-C3aR-Go dataset, 15,216 motion corrected micrographs were manually curated and 12,114 micrographs with CTF fit resolution greater than 4 Å were selected for automated particle picking with the blob-picker subprogram. 9,530,582 autopicked particles were extracted with a box size of 416 px and fourier cropped to 64 px for subsequent reference free 2D classification. 2D classes with clear secondary features were selected and re-extracted with a box size of 360 px and fourier cropped to 288 px for generating 4 ab-initio reconstructions and downstream heterogeneous refinement. 914,579 particles corresponding to the best 3D class with essential features of a GPCR-G protein complex were selected and subjected to non-uniform refinement to yield a map with an estimated resolution of 3.1 Å at 0.143 cutoff.

Model building and refinement

For the C5a^{pep}-mC5aR1-Go complex, the receptor coordinates from the cryo-EM structure of human formyl peptide receptor 2 (PDB: 7WVW)⁵⁸ and the coordinates for the G α , G β 1, G γ 2 from the cryo-EM structure of Muscarinic acetylcholine receptor 2-Go complex (PDB: 6OIK)⁵⁹ were used as an initial model to dock into the EM density using Chimera.^{65,66} This was followed by manual rebuilding of the model along with the ligand in COOT^{67–69} and iterative real space refinement in Phenix.^{70,71} This yielded a model with 96.63% in the most favored region and 3.37% in the allowed region of the Ramachandran plot.

For the C5a-mC5aR1-Go complex map, the coordinates of C5a^{pep}-mC5aR1-Go complex was used as an initial model and docked into the EM density with the “Fit in map” extension in Chimera. Similarly, the coordinates corresponding to human C5a were taken from a previously solved crystal structure of the human C5a in complex with MEDI7814, a neutralizing antibody (PDB: 4UU9).⁶⁰ The model so obtained was docked in Chimera, manually rebuilt in COOT and subjected to several rounds of real space refinement in Phenix to reach a final model with 95.87% in the favored region and 4.13% in the allowed region of the Ramachandran plot. Data collection, 3D reconstruction and refinement statistics have been included as Table S1.

Coordinates of C5aR1 receptor, G α , G β 1 and G γ 2 from the cryo-EM structure of C5a-pep-C5aR1-Go (PDB: 8HPT) were used as initial models to dock into the EM density of EP54-C3aR-Go complex using the “Fit in map” extension in Chimera.^{65,66} The fitted model along with the map was imported into COOT^{67–69} for manual rebuilding of the model along with the ligand (EP54), with iterative rounds of real space refinement in Phenix.^{70,71} This yielded a final refined model with 94.95% of the residues in the most favored region and 5.05% in the allowed region of the Ramachandran plot.

For the C3a-C3aR-G α complex map, coordinates of the receptor, G α , G β 1 and G γ 2 from EP54-C5aR1-G α complex were used as initial models. The coordinates of C3a were obtained from a previously solved crystal structure of C3a (PDB: 4HW5).²⁴ Chimera was used to dock the individual components in the cryo-EM map and obtain a merged model. The combined model was manually rebuilt in COOT^{61–63} and subjected to multiple rounds of real space refinement in Phenix. The final refined model showed good validation statistics with 95.44% in the favored region and 4.56% in the allowed region of the Ramachandran plot.

For EP54-C3aR-Gq and both the ligand free-C3aR-Go cryo-EM maps, the coordinates corresponding to the individual components were obtained from the model of EP54-C3aR-Go which were used to fit into the coulombic map with the “Fit in map” extension in Chimera. Iterative rounds of manual adjustment and building in COOT^{67–69} and refinement with real space refine in Phenix resulted in the final model with excellent validation statistics and no Ramachandran outliers. For ligand free-C3aR-Go, the model obtained from the Glacios data was used for analysis of the ligand free structure due to its higher resolution and more interpretable map. Data collection and refinement statistics have been included in the supplemental information.

For the C5a-hC5aR1-Go complex map, the coordinates for the receptor were taken from the cryo-EM structure of human formyl peptide receptor 2 (PDB: 7WVV), while the coordinates for G α o, G β 1, G γ 2 were obtained from the cryo-EM structure of C5a^{deP}-mC5aR1-Go complex. The coordinates were docked into the EM density map in Chimera and the combined model so obtained were transferred to COOT for performing subsequent mutations and model manipulations. Several rounds of real space refinement in Phenix yielded the final model with 92.90% in the favored region and 7.10% in the allowed region of the Ramachandran plot.

For the C5a^{des-Arg}-hC5aR1-Go complex, the atomic coordinates of C5a, C5aR1, G α o, G β 1 and G γ 2 were obtained from the cryo-EM structure of C5a-hC5aR1-Go structure (PDB: 8IA2), and docked into the electron microscopy density map using UCSF Chimera, followed by iterative manual adjustment and rebuilding in COOT. The coordinates were refined against the final map using phenix.real_space_refine with secondary structure and geometry restraints. The final refined model showed excellent statistics with 95.59% in the most favored region and 4.41% in the allowed region of the Ramachandran plot.

For the EP141-C3aR-Go complex, the coordinates of C3aR, G α o, G β 1 and G γ 2 were obtained from EP54-C5aR1-Go complex structure and were used as initial models to dock into the coulombic map using Chimera. The merged model obtained was imported into COOT and further fitted into the density with the “Fit in map” extension. The EP141 peptide ligand was then manually built into the ligand density followed by multiple rounds of real space refinement with phenix.real_space_refine. The final model so obtained exhibited good statistics with 96.19% in the most favored region and 3.81% in the allowed region of the Ramachandran plot.

All figures were prepared either with Chimera or ChimeraX software.^{65,66} Buried surface and interface surface area have been calculated with PDBePISA webserver.⁷² Ligand-receptor and receptor-G-protein interactions were identified using PDBsum.⁷³

QUANTIFICATION AND STATISTICAL ANALYSIS

GraphPad Prism v9.5 was used to plot and analyze all the functional data presented in this manuscript, and all the relevant details such as number of replicates, data normalization, mean \pm SEM, and statistical analyses are mentioned in the corresponding figure legends.

Supplemental figures

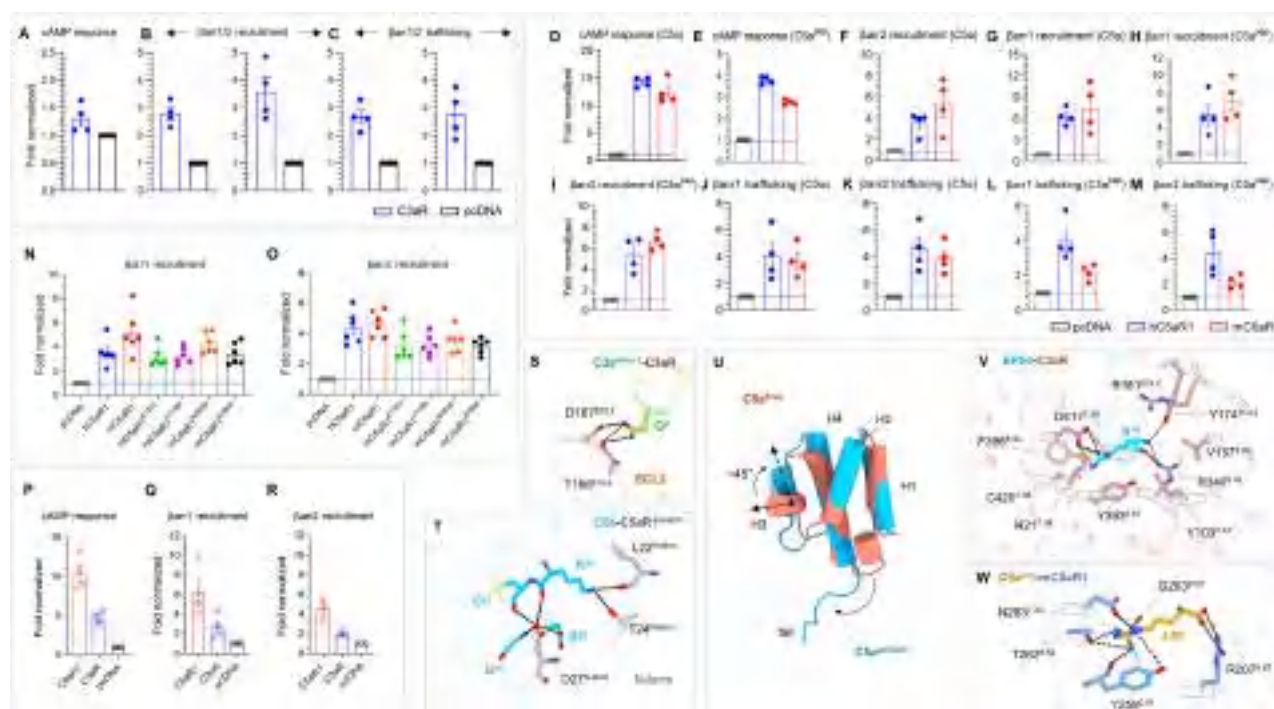
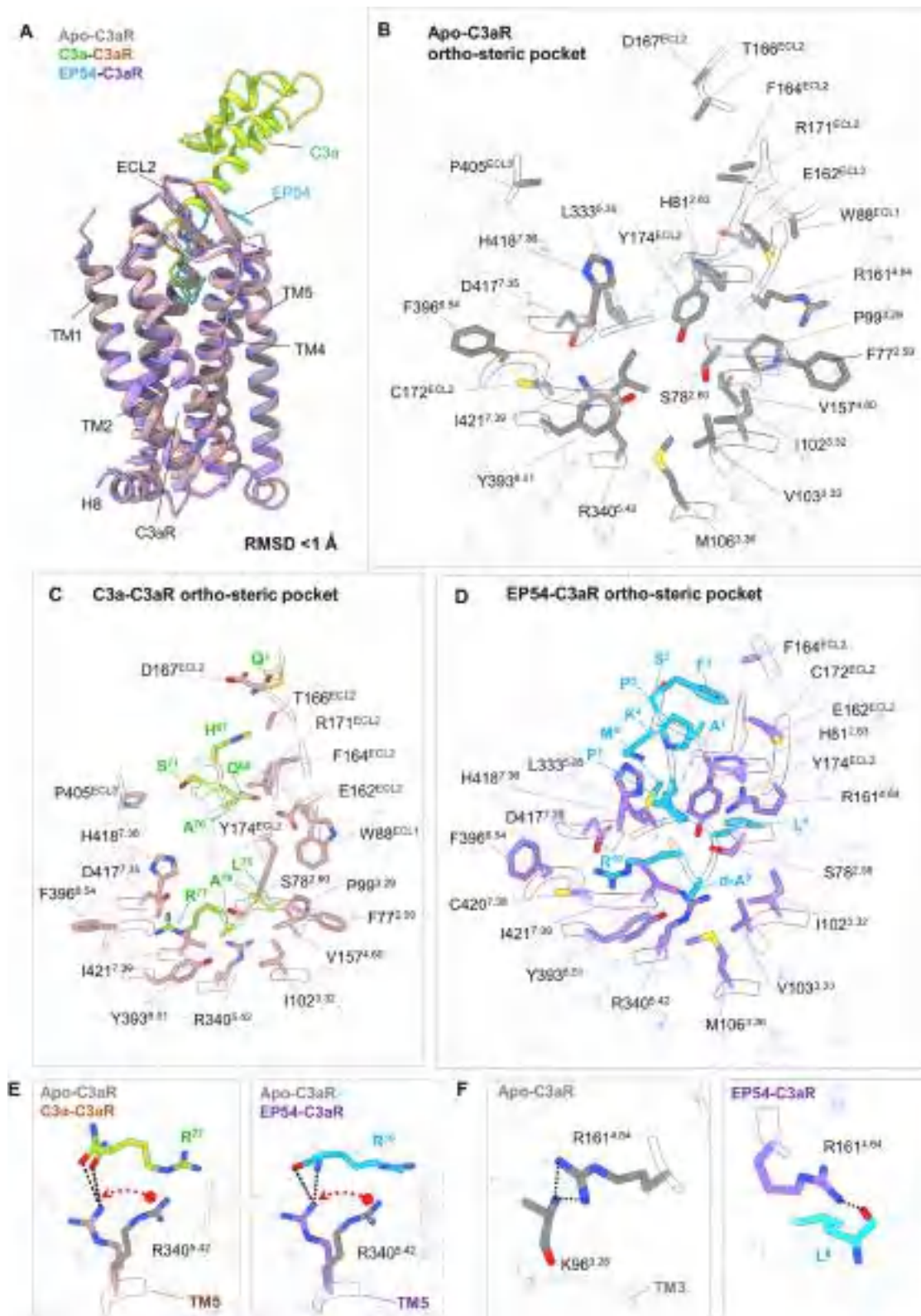


Figure S1. Surface expression profiles of C3aR and C5aR1 and critical interactions of C3a/C5a with C3aR/C5aR1, related to Figures 1, 3, and 7

(A) Surface expression of C3aR in GloSensor measured using whole-cell ELISA (mean \pm SEM; $n = 4$; normalized as fold over mock transfection).
 (B) Surface expression of C3aR in β arr1/2 recruitment assay (mean \pm SEM; $n = 4$; normalized as fold over mock transfection).
 (C) Surface expression of C3aR in β arr1/2 endosomal trafficking assay (mean \pm SEM; $n = 4$; normalized as fold over mock transfection).
 (D–M) Surface expression of indicated receptors measured using whole-cell ELISA (mean \pm SEM; $n = 4$; normalized as fold over mock transfection) in various assays.
 (N and O) Surface expression of indicated receptor constructs in β arr1 and β arr2 recruitment assays (mean \pm SEM; $n = 6$, normalized as fold over mock transfection).
 (P) Surface expression of C3aR and C5aR1 in GloSensor assay (mean \pm SEM; $n = 4$; normalized as fold over mock transfection).
 (Q and R) Surface expression of C3aR and C5aR1 in β arr1 (mean \pm SEM; $n = 4$) and β arr2 (mean \pm SEM; $n = 4$) recruitment assay (normalized as fold over mock transfection).
 (S and T) Interface between Gln³ of C3a with C3aR and C5a with N-term of C5aR1.
 (U) Structural comparison of free C5a with C5a bound to mC5aR1. H3 of C5a can be seen to exhibit a rotation of $\sim 45^\circ$ upon binding to the receptor.
 (V and W) The terminal arginine in EP54 and C5a^{PEP} makes extensive interactions with C3aR and C5aR1, respectively.



(legend on next page)

Figure S2. Comparison of the residues in the orthosteric pocket of C3aR in Apo-, EP54-, and C3a-bound states, related to Figure 3

- (A) Structural superimposition of Apo-, EP54-, and C3a-bound C3aR-Go complexes.
 (B) Conformation of all the residues present in the orthosteric pocket of C3aR involved in interaction with the ligands are shown in the Apo structure.
 (C and D) Residues of the orthosteric pocket of C3aR involved in interaction with the residues of C3a (C) and EP54 (D) are highlighted.
 (E and F) Changes in rotameric conformations of Arg340^{5,42} and Arg161^{4,64} in the Apo state as compared with the ligand-bound state.

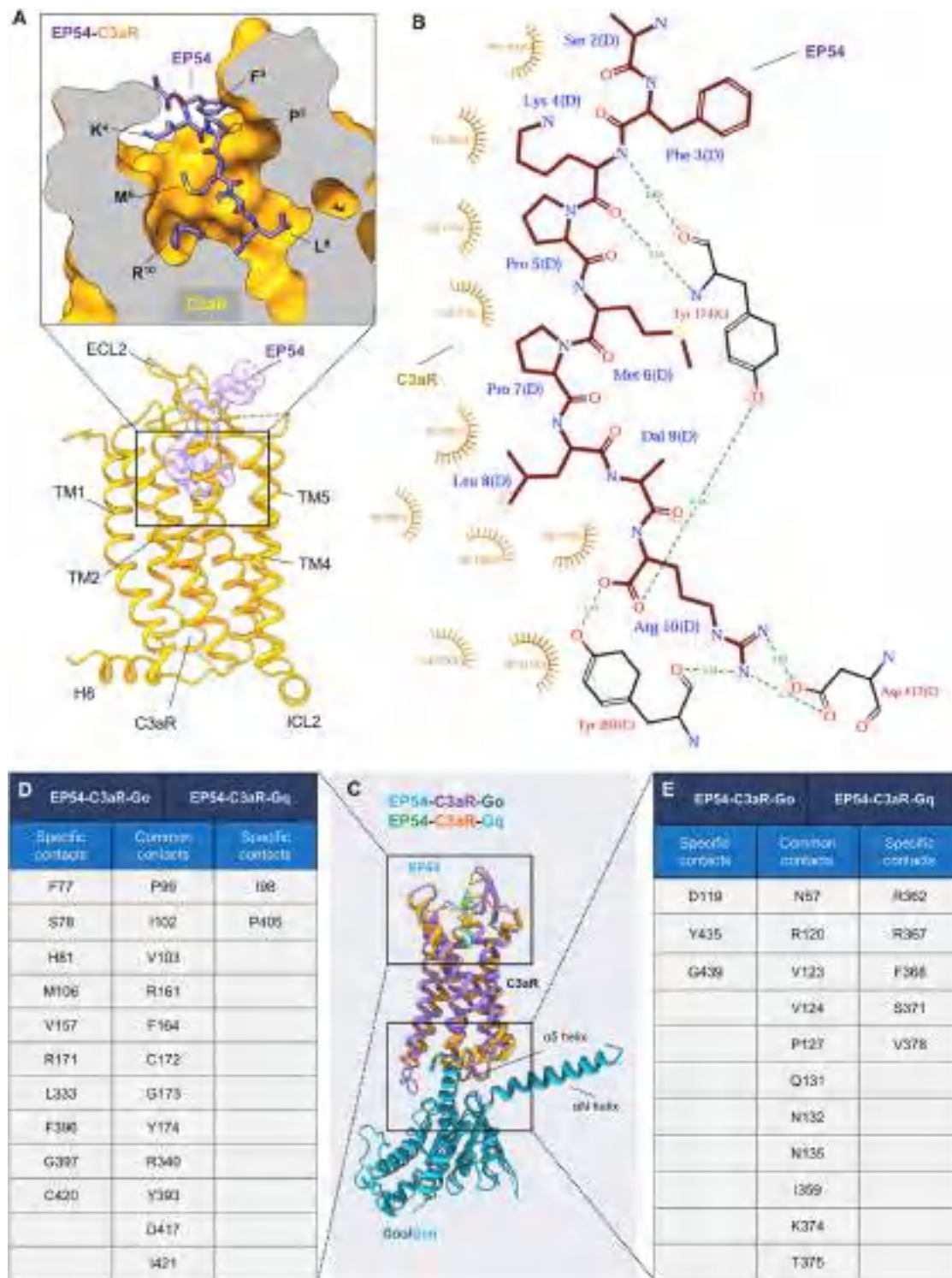


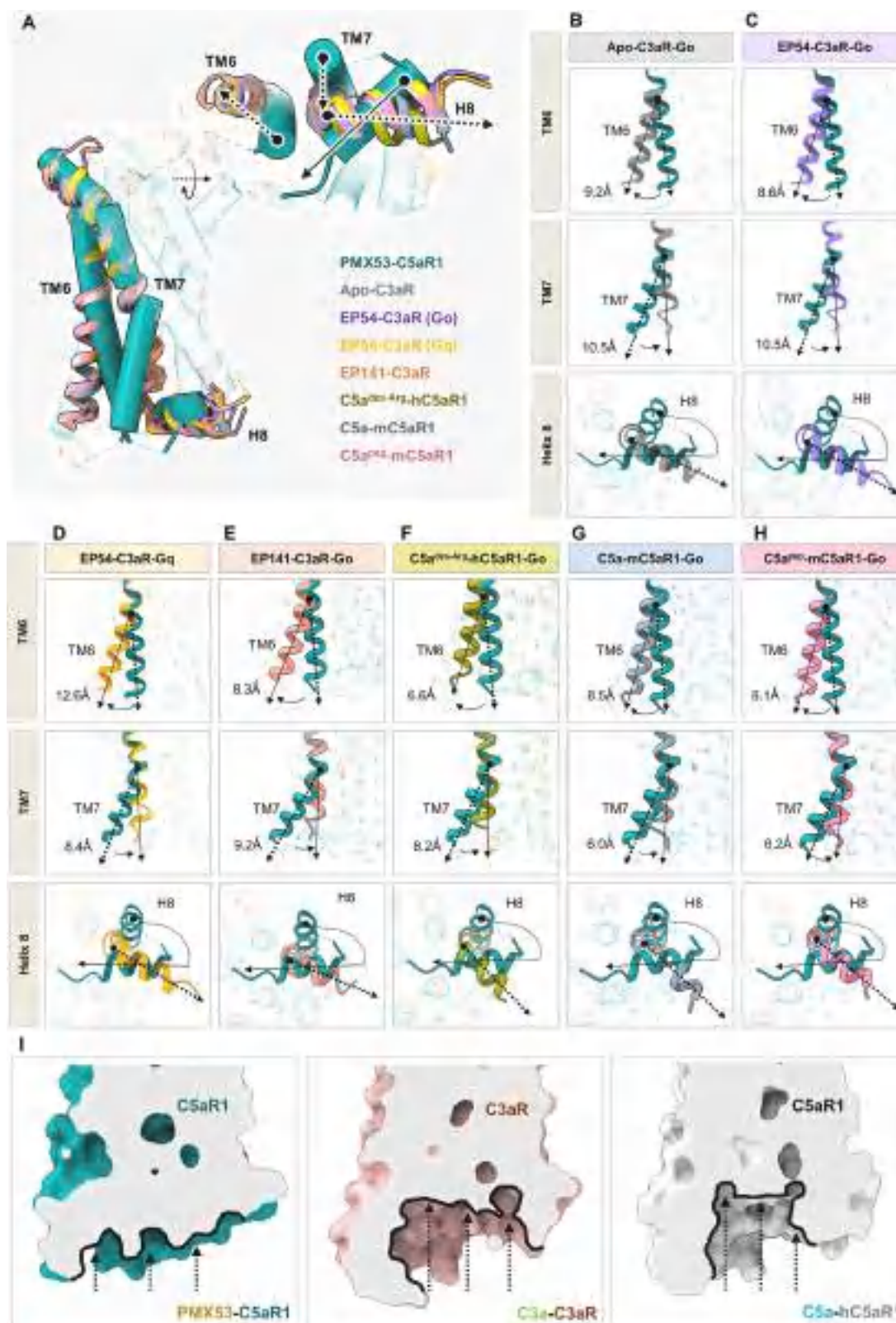
Figure S3. EP54 binding and activation of C3aR in complex with Gq and comparison with Go-bound state, related to Figures 3 and 6

(A) EP54 binding pose at the orthosteric pocket of C3aR in complex with Gq. EP54 in transparent surface (bottom) and receptor in surface slice showing side chains of EP54 in the ligand-binding pocket (top).

(B) Interaction interface between EP54 and C3aR-Gq. The interaction plot has been generated through PDBSum.

(C) Structural superposition of EP54-C3aR-Gq α o and EP54-C3aR-Gq α q complexes.

(D and E) Common and specific residue interactions between C3aR and EP54 (D) and C3aR and Gq α o/Gq α q (E), respectively.



(legend on next page)

Figure S4. Active conformations of C3aR and C5aR1, related to Figure 6

(A) Structural superimposition of C3a-bound C3aR and C5a-bound hC5aR1 with inactive C5aR1 (PDB: 6C1R). TM6, TM7, and H8 are highlighted to show the change in conformation in the receptors, and other regions are depicted in transparent ribbons.

(B–H) Dynamic changes in TMs of activated C3aR/C5aR1 compared with the inactive state of C5aR1. The TMs and H8 from different receptor complexes shown are from receptors mentioned in boxes. Solid lines (active receptors) and dotted lines (inactive C5aR1) indicate direction of movement. The respective degrees of movements in corresponding TMs have also been mentioned.

(I) The antagonist PMX53 (PDB: 6C1R) occupies an analogous binding pocket on C5aR1 and adopts a hook-like conformation similar to C5a (left). However, the cytoplasmic cavity in the inactive-state structure C5aR1 is blocked thereby preventing further transducer coupling compared with the active state C3aR and C5aR1 (middle and right).

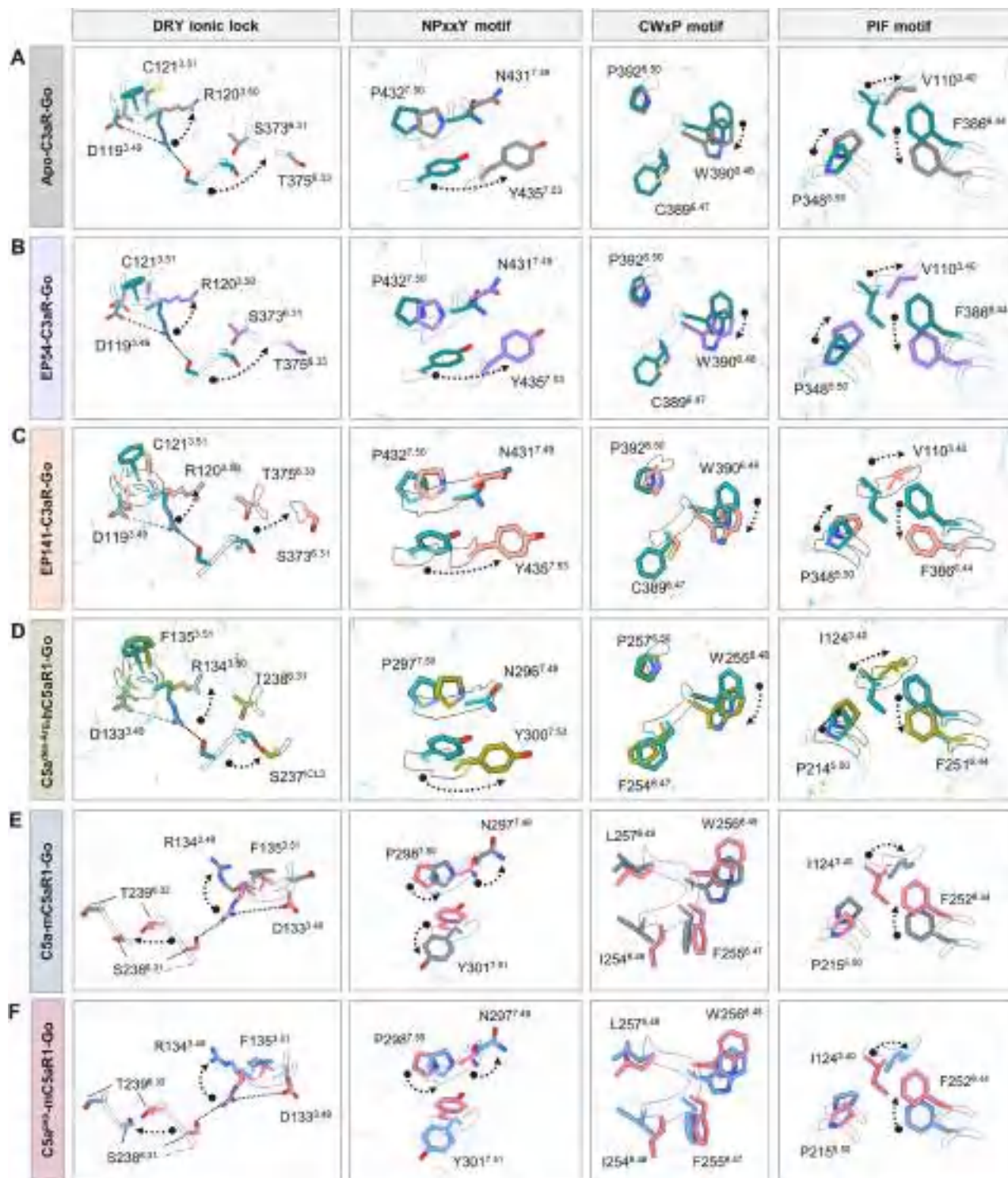


Figure S5. Conformational changes in the conserved microswitches of C3aR and C5aR1, related to Figure 6

(A–F) Conformational changes in the conserved microswitches: (DRY(F), NPxxY, C(F)WxP(L), and PIF) upon C3aR and C5aR1 activation. (Teal and pink: inactive C5aR1 for C3aR and C5aR1 structures, respectively, various colors: C3aR and C5aR1.)



Figure S6. The G-protein interface of C3aR and C5aR1, related to Figure 6

(A–G) $\alpha 5$ helix of G $\alpha o/q$ docks into the cytoplasmic core of C3aR and C5aR1. Only receptor and G α are shown in ribbon representations to highlight the binding pose of G proteins with receptor core. Surface slice presentations (top) and cryo-EM maps (bottom) have been shown in inset boxes to highlight direct docking of G α to receptors. Magnified view of the interactions between TM2, TM3, TM6, TM7, ICL2, and ICL3 of C3aR and C5aR1 with G αo . Ionic bonds are depicted as black dashed lines.

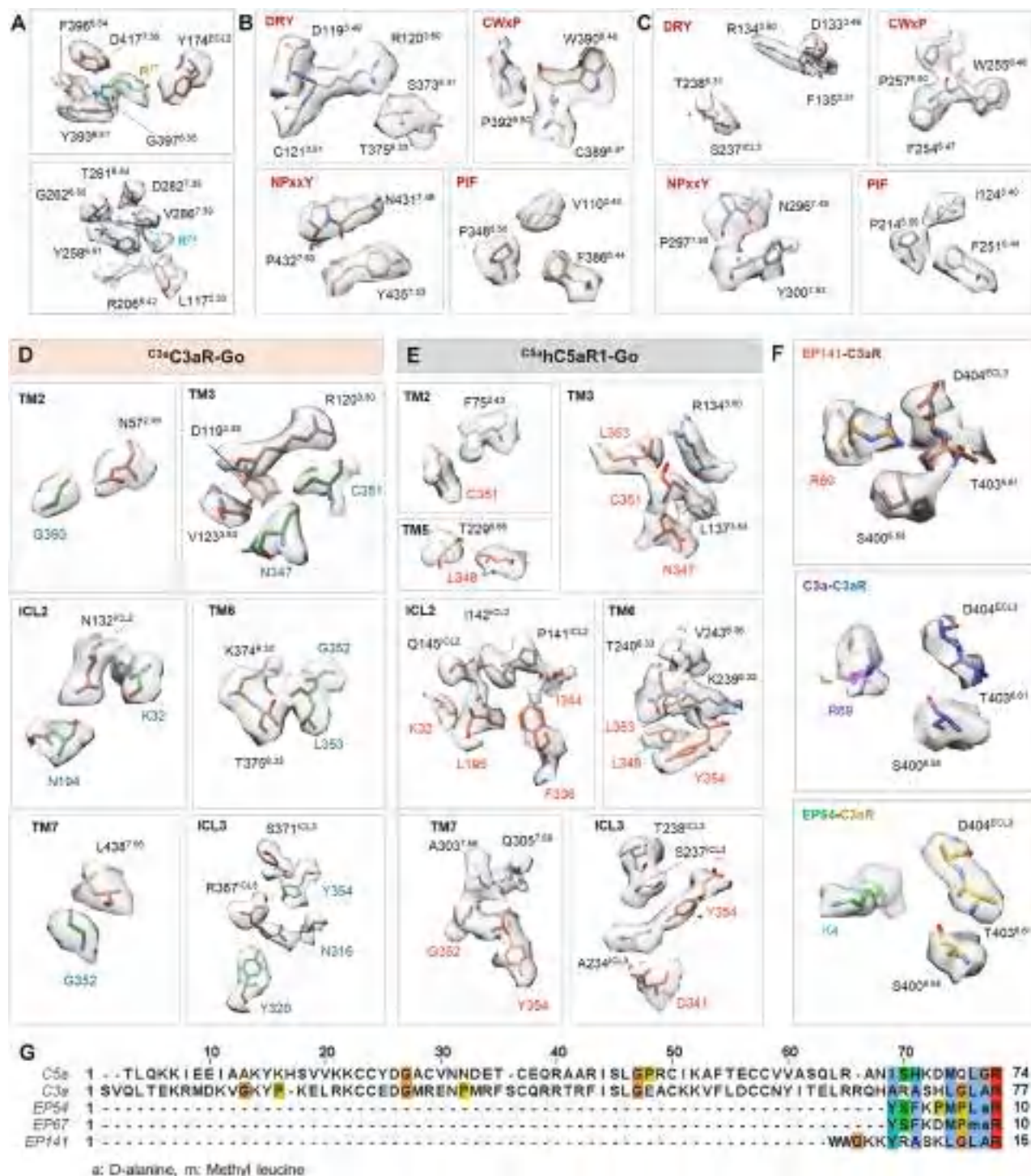
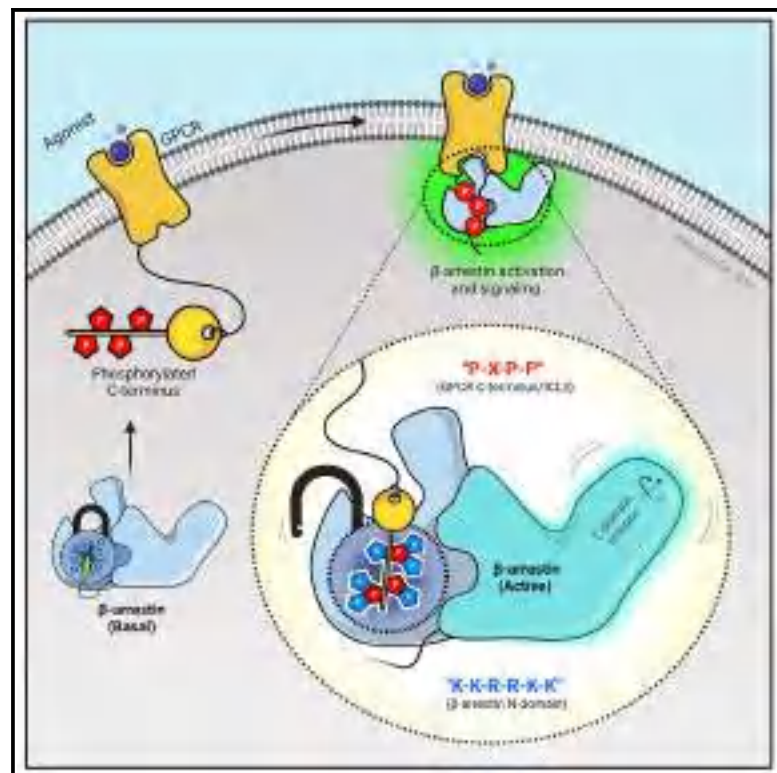


Figure S7. Cryo-EM densities of interface residues and sequence of complement peptide agonists, related to Figures 4, 6, and 7
(A–F) Densities corresponding to the amino acids shown in Figures 4B, 5C, 5D, 5F, 5H, and 7E are presented.
(G) Multiple sequence alignment of complement peptide agonists.

Structural snapshots uncover a key phosphorylation motif in GPCRs driving β -arrestin activation

Graphical abstract



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In brief

Maharana et al. determine multiple structures of activated β -arrestins in complex with the carboxyl terminus phosphopeptides of different GPCRs using cryo-EM. Based on these structural snapshots, they discover and experimentally validate a significantly conserved phosphorylation motif in GPCRs that drives β -arrestin interaction and activation.

Highlights

- Cryo-EM structure determination of β -arrestins in complex with GPCR phosphopeptides
- Identification of P-X-P-P motif in GPCRs for β -arrestin interaction and activation
- Experimental validation of P-X-P-P motif using mutagenesis and conformational sensor
- Discovery of a significantly conserved β -arrestin activation mechanism by GPCRs



Article

Structural snapshots uncover a key phosphorylation motif in GPCRs driving β -arrestin activation

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SUMMARY

Agonist-induced GPCR phosphorylation is a key determinant for the binding and activation of β -arrestins (β arrs). However, it is not entirely clear how different GPCRs harboring divergent phosphorylation patterns impart converging active conformation on β arrs leading to broadly conserved functional responses such as desensitization, endocytosis, and signaling. Here, we present multiple cryo-EM structures of activated β arrs in complex with distinct phosphorylation patterns derived from the carboxyl terminus of different GPCRs. These structures help identify a P-X-P-P type phosphorylation motif in GPCRs that interacts with a spatially organized K-K-R-R-K-K sequence in the N-domain of β arrs. Sequence analysis of the human GPCRome reveals the presence of this phosphorylation pattern in a large number of receptors, and its contribution in β arr activation is demonstrated by targeted mutagenesis experiments combined with an intra-body-based conformational sensor. Taken together, our findings provide important structural insights into the ability of distinct GPCRs to activate β arrs through a significantly conserved mechanism.

INTRODUCTION

G protein-coupled receptors (GPCRs) are typically characterized by their conserved seven transmembrane (7TM) architecture and agonist-induced coupling to heterotrimeric G-proteins and β -arrestins (β arrs).¹ Of these, β arrs are multifunctional cytosolic proteins critically involved in regulating the signaling and trafficking of GPCRs.^{2–4} Their interaction with GPCRs involves a major contribution from receptor phosphorylation, which not only drives the affinity of receptor- β arr interaction but also imparts functionally competent active conformation in β arrs driving downstream functional outcomes^{5–12} (Figure 1A). Additional interaction of β arrs with the receptor transmembrane core and membrane lipid bilayer induces further structural changes as visualized using biophysical and direct structural approaches,^{13–20} which also fine-tune the functional capabilities of β arrs and possibly spatiotemporal aspects of their regulatory mechanisms.^{21,22} Despite a poorly conserved primary sequence of GPCRs in terms of the number and spatial positioning of the putative phosphorylation sites, the near-universal nature of β arr interaction, ensuing signaling and regulatory responses, remains to be fully understood at the molecular level. Moreover, differential receptor phosphorylation by different subtypes of

GPCR kinases (GRKs), and possibly other kinases, in cell type-specific manner adds further complexity to GPCR- β arr binding modalities and context-specific functional specialization.^{23–29}

There are two isoforms of β arrs namely, β arr1 and 2, also known as Arrestin2 and 3, respectively, and despite a high degree of sequence and structural similarity, they often exhibit a significant diversity in their functional contribution toward GPCR signaling and regulation.³⁰ There are only a very few structures of β arr1 in active conformation, either in complex with receptor-derived phosphopeptides^{31,32} or in complex with agonist-bound, phosphorylated GPCRs.^{15–20} The structural coverage for β arr2 is even more sparse with the structural snapshots limited to either a complex with CXC chemokine receptor subtype 7 (CXCR7)-derived phosphopeptide³³ or in IP6-bound state.³⁴ Although these structures provide useful information about β arrs' interaction with the receptors, there is only limited information available about the phosphorylation patterns either due to chimeric constructs used in these studies or the lack of visualization of multiple phosphorylation sites resulting from insufficient structural resolution. Thus, the quest to decipher precise molecular details of convergent β arr activation by GPCRs harboring different phosphorylation patterns remains open. This represents a major knowledge gap in our current



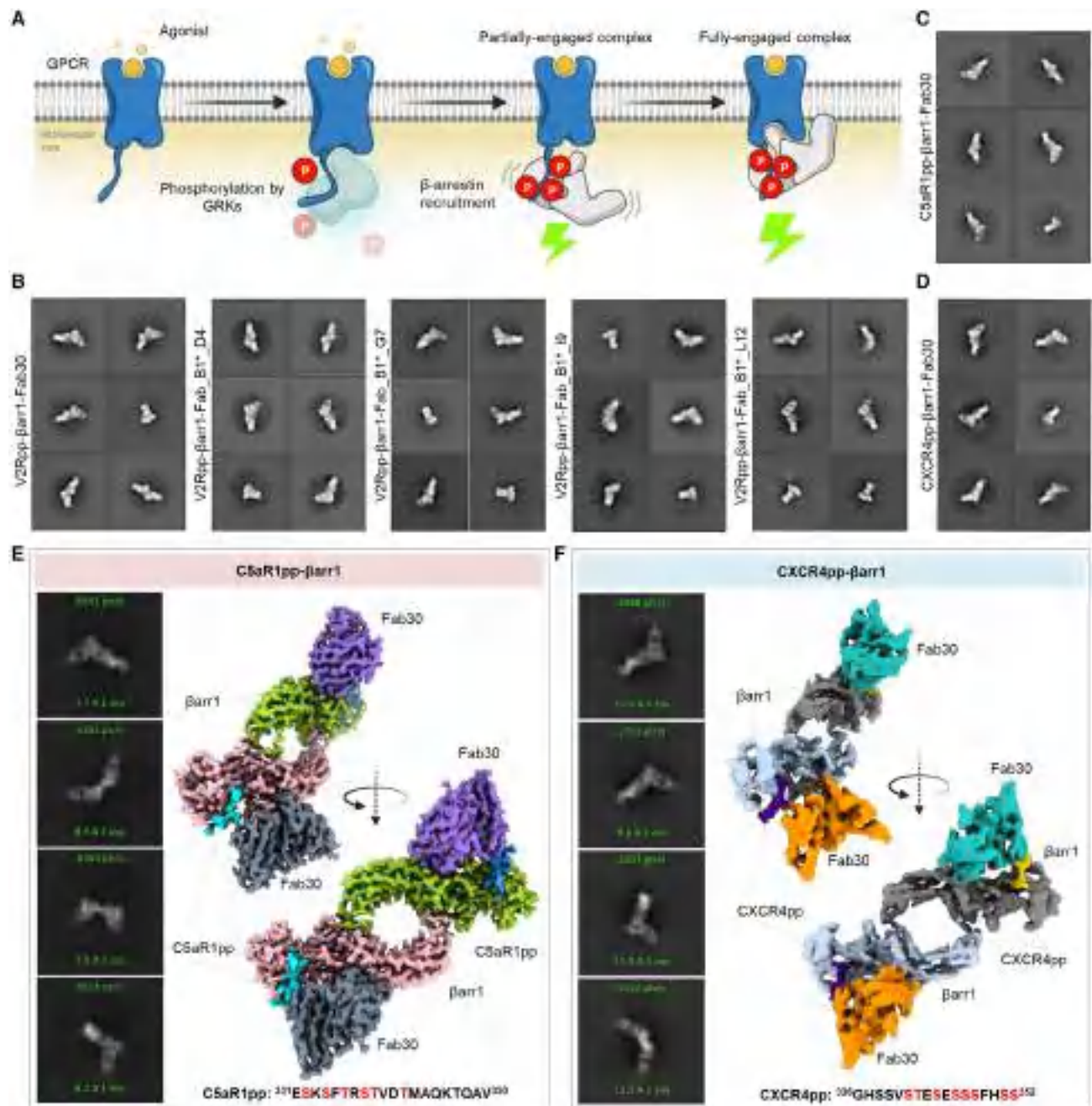


Figure 1. Reconstitution and structure determination of C5aR1pp/CXCR4pp- β arr1 complexes

(A) Agonist stimulation of GPCRs leads to receptor phosphorylation by GPCR kinases (GRKs) followed by the recruitment and activation of β arrestins governed through the phosphorylated residues and activated receptor core.

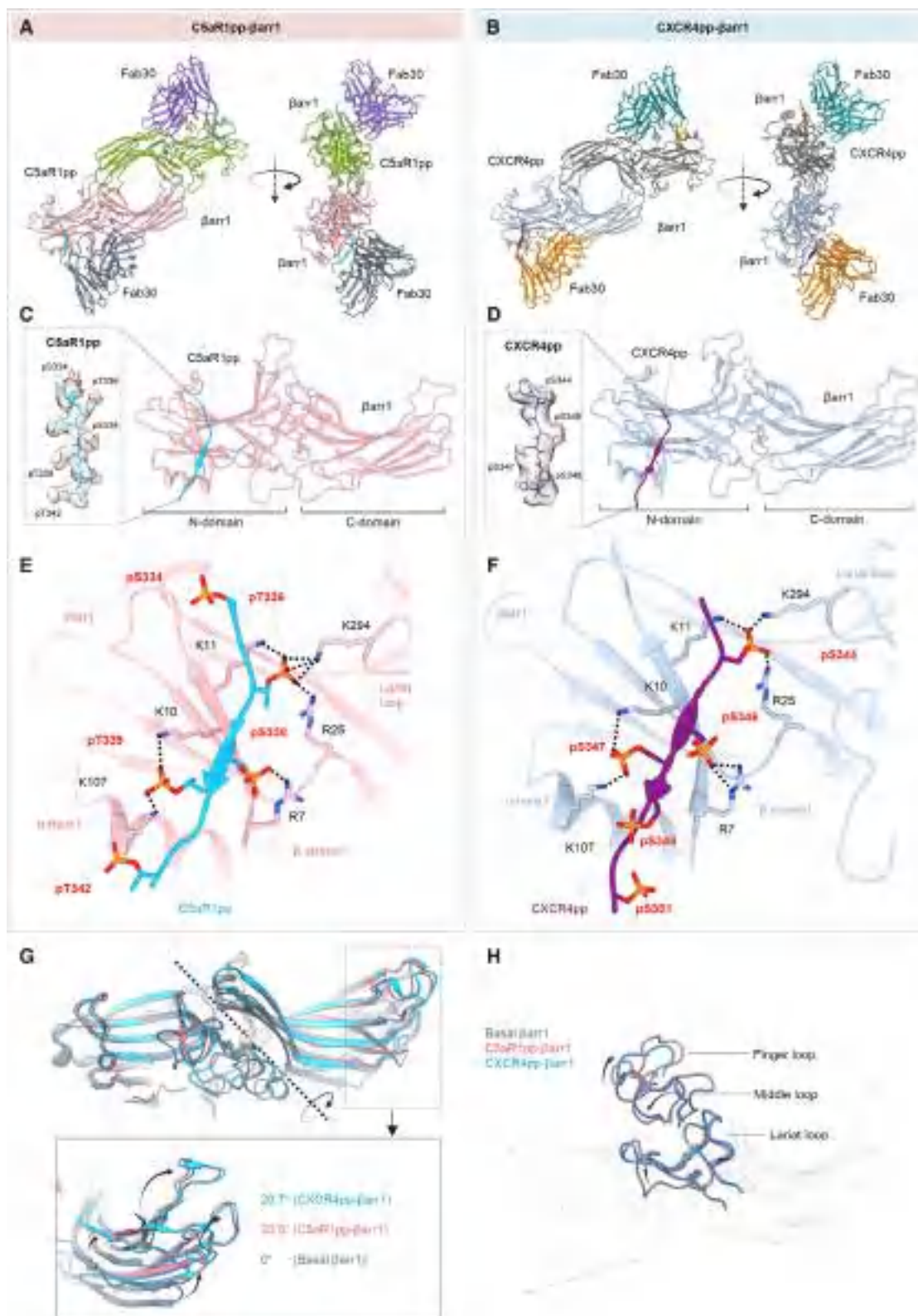
(B) Negative-staining EM-based 2D class averages of V2Rpp- β arr1 complexes stabilized by Fab30, Fab_B1*_D4, Fab_B1*_G7, Fab_B1*_I9, and Fab_B1*_L12, respectively.

(C and D) Negative-staining EM-based 2D class averages of C5aR1pp- β arr1-Fab30 and CXCR4pp- β arr1-Fab30 complexes, respectively.

(E and F) Selected 2D class averages and surface representation of C5aR1pp- β arr1-Fab30 and CXCR4pp- β arr1-Fab30 structures, respectively, determined by cryo-EM. The missing Fab30 constant domain densities were truncated during local refinement. See also Figures S1–S3 and Table 1.

understanding of GPCR signaling and regulatory paradigms governing and fine-tuning signal transduction through this versatile class of receptors.

In this backdrop, here we present cryogenic-electron microscopy (cryo-EM) structures of full-length β arr1 and 2 activated by defined phosphorylation patterns encoded in the form of



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phosphopeptides, which are derived from three different GPCRs, namely, the complement C5a receptor subtype 1 (C5aR1), the CXC chemokine receptor subtype 4 (CXCR4), and the vasopressin receptor subtype 2 (V2R). These structural snapshots reveal a P-X-P-P type pattern of phosphorylation in GPCRs that engages a K-K-R-R-K-K sequence in the N-domain of β arrs leading to β arr activation. Interestingly, a large repertoire of GPCRs encodes the P-X-P-P motif either in their carboxyl terminus or in the 3rd intracellular loop (ICL3), suggesting a broad implication of this activation mechanism. We further validate the contribution of the P-X-P-P motif with respect to β arr activation in cellular context for several GPCRs using an intrabody-based conformational biosensor and structure-guided mutagenesis studies. Collectively, our data help identify the presence of the P-X-P-P motif in GPCRs and uncover the molecular basis of its ability to activate β arrs.

RESULTS

Although cryo-EM has been used to determine the structures of GPCR- β arr1 complexes,^{15–19} all the previous structures of β arrs in basal state,^{35–38} bound to phosphopeptides^{31–33} or IP6³⁴ have been determined using X-ray crystallography. Therefore, in order to test the feasibility of structure determination of β arrs in complex with different phosphorylation patterns encoded in GPCR phosphopeptides by cryo-EM, we first reconstituted V2Rpp- β arr1 complex together with a set of conformationally selective antigen-binding fragments (Fabs), which recognize activated β arr1.^{39,40} Subsequently, we analyzed these complexes using negative-staining single particle EM, which revealed monodisperse particle distribution and 2D class averages where the densities of β arr1 and Fabs were clearly discernible (Figures 1B and S1A). Based on these observations, we synthesized and characterized a set of phosphopeptides corresponding to the carboxyl terminus of the C5aR1 and the CXCR4, and assessed their ability to activate β arr1, measured in terms of Fab30 reactivity (Figures S1B–S1E). We identified the phosphorylation patterns from C5aR1 (C5aR1pp2; referred to as C5aR1pp hereon) and CXCR4 (CXCR4pp2; referred to as CXCR4pp hereon), which elicited maximal Fab30 reactivity as a measure of β arr1 activation (Figures S1B–S1E). Subsequently, we reconstituted C5aR1pp- β arr1-Fab30 and CXCR4pp- β arr1-Fab30 complexes, validated their monodispersity and architecture using negative-staining EM (Figures 1C, 1D, S1F, and S1G), and subjected

these complexes to cryo-EM. We successfully determined the structures of C5aR1pp- β arr1-Fab30 and CXCR4pp- β arr1-Fab30 complexes at global resolutions of 3.26 and 4.45 Å, respectively (Figures 1E, 1F, and S2).

β arr1 structures in complex with C5aR1 and CXCR4 phosphorylation patterns

Both structures revealed a dimeric arrangement with the two β arr1 protomers making contacts through the C-edge loops and finger loops (Figures 2A and 2B). β arr1 protomers exhibit nearly identical overall structures with each other (root-mean-square deviation [RMSD] < 0.5 Å) with clear densities of the phosphopeptides visible in the EM map (Figures S3A and S3B). C5aR1pp and CXCR4pp are positioned in a positively charged groove on the N-domain of β arr1 (Figures 2C and 2D), and the phosphate moieties make extensive contacts with Arg/Lys residues at the binding interface (Figures 2E and 2F). Interestingly, three phosphate groups arranged in a P-X-P-P type pattern, where P is a phosphorylated residue and X is any other amino acid, in both the phosphopeptides engage a nearly identical set of Lys/Arg residues in β arr1 (Figures 2E and 2F). Specifically, the pT³³⁶-R³³⁷-pS³³⁸-pT³³⁹ pattern in C5aR1pp and pS³⁴⁴-E³⁴⁵-pS³⁴⁶-pS³⁴⁷ pattern in CXCR4pp engages K²⁹⁴-K¹¹-R²⁵-R⁷-K¹⁰-K¹⁰⁷ in β arr1. The other phosphate groups present in the phosphopeptides are either not involved in direct contact or sparsely linked with Arg/Lys or positioned outside the binding groove. The N- and C-domains of β arr1 exhibit an inter-domain rotation of approximately 20° when compared with the basal state of β arr1, in both structures, which is a hallmark of β arr activation upon binding of phosphorylated GPCRs³¹ (Figure 2G). Moreover, the three major loops in β arr1 namely, the finger, middle, and lariat loop also exhibit significant reorientation upon binding of C5aR1pp and CXCR4pp compared with the basal state, although their positioning is almost identical between the two structures (Figure 2H). Finally, the three-element interaction and polar-core network in β arr1 are also disrupted upon binding to C5aR1pp and CXCR4pp when compared with the basal state structure through the displacement of the carboxyl terminus of β arr1 from the N-domain and repositioning of the lariat loop through the interaction of K²⁹⁴ with a phosphate moiety (Figures S4A and S4B). These structural features and interaction interface are analogous to that observed in the V2Rpp- β arr1-Fab30 crystal structure determined previously,³¹ although the primary sequence and phosphorylation patterns encoded by C5aR1pp and CXCR4pp are distinct from V2Rpp (Figure S4C).

Figure 2. Overall structures and key structural features of C5aR1pp/CXCR4pp- β arr1 complexes

(A and B) Overall structures of C5aR1pp- β arr1-Fab30 and CXCR4pp- β arr1-Fab30 complexes shown with ribbon representation. The constant domains of Fab30 were masked out during refinement.

(C and D) Structure of individual C5aR1pp- β arr1 and CXCR4pp- β arr1 complex protomers shown as ribbon representation to indicate the binding of phosphopeptides on the N-domain of β arr1. Cryo-EM densities of corresponding phosphopeptides have been provided in insets.

(E and F) Stabilizing charge-charge interactions of C5aR1pp and CXCR4pp with the N-domain groove residues of β arr1 indicated as dotted lines. pS and pT refer to phospho-Ser and phospho-Thr residues, respectively.

(G) Inter-domain rotation in β arr1 upon binding of C5aR1pp (pink) and CXCR4pp (blue) is compared with the basal conformation of β arr1 determined previously (PDB: 1G4M, gray).

(H) Superimposition of C5aR1pp- and CXCR4pp-bound β arr1 structures with the basal conformation of β arr1 (PDB: 1G4M, gray) indicating the repositioning of finger, middle, and lariat loops upon β arr1 activation. See also Figure S4.

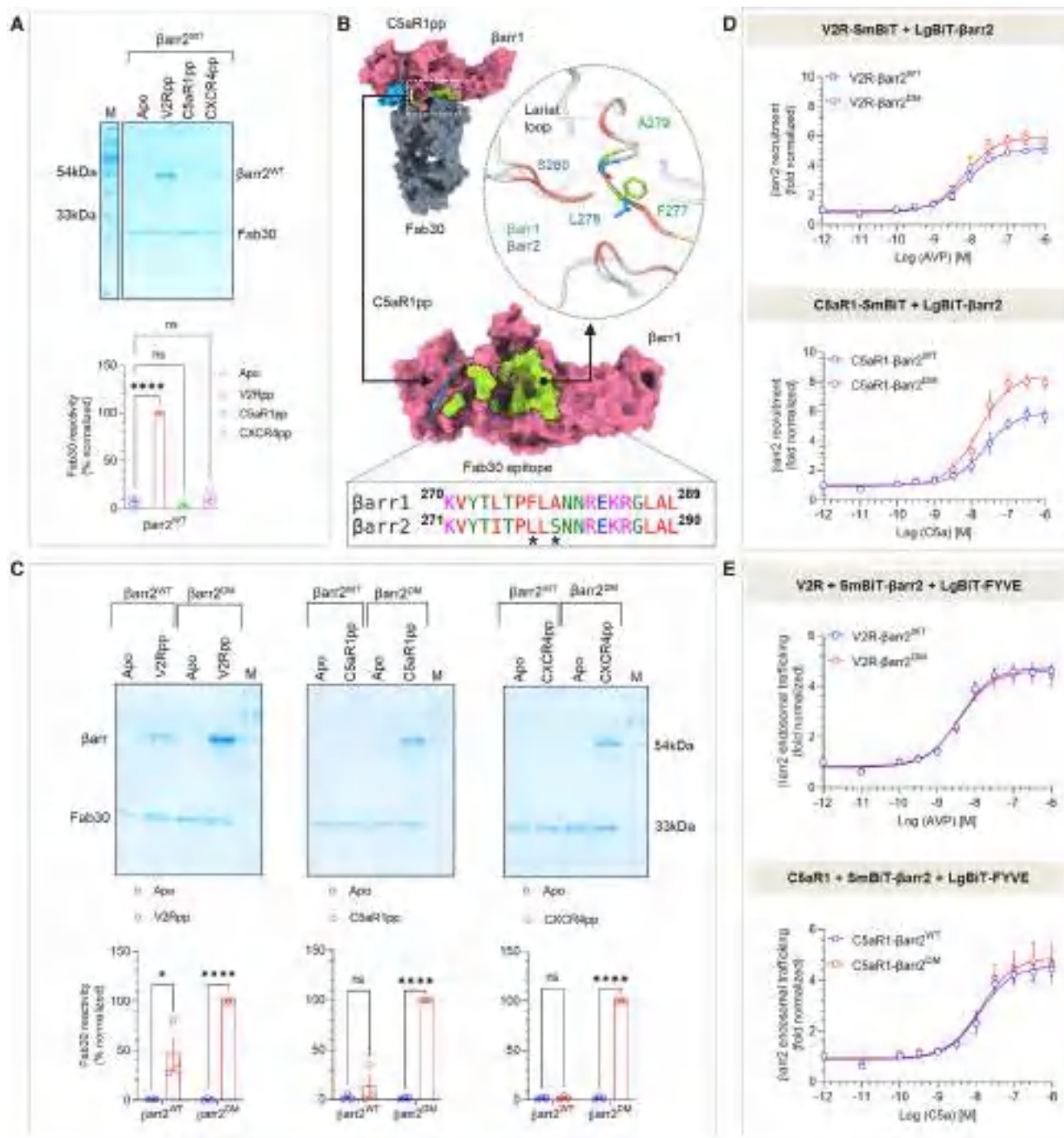


Figure 3. Generation and characterization of β arr2^{DM} for structure determination

(A) Fab30 reactivity to C5aR1pp and CXCR4pp activated β arr2^{WT} was measured by co-immunoprecipitation (coIP) assay. C5aR1pp and CXCR4pp activated β arr2^{WT} were not recognized by Fab30 (top). Densitometry-based quantification of the coIP data is presented (bottom) (mean \pm SEM; n = 3; normalized with respect to V2Rpp signal as 100%; one-way ANOVA, Dunnett's multiple comparisons test). The exact p values are as follows: Apo vs. V2Rpp (p \leq 0.0001), Apo vs. C5aR1pp (p = 0.7719), Apo vs. CXCR4pp (p = 0.7899) (****p < 0.0001; ns, non-significant).

(B) Comparison of the epitope region of Fab30 in β arr1 with β arr2 reveals that instead of F²⁷⁷ and A²⁷⁹ as in β arr1, β arr2 contains L²⁷⁸ and S²⁸⁰ in corresponding positions (indicated with an asterisk).

(C) CoIP assay showing the reactivity of Fab30 toward activated β arr2^{DM} upon binding of C5aR1pp and CXCR4pp (top). Densitometry-based quantification is presented (bottom) (mean \pm SEM; n = 3; normalized with respect to Fab30 reactivity toward activated β arr2^{DM} treated as 100%; two-way ANOVA, Sidák's multiple comparisons test). The exact p values are as follows: for β arr2^{WT}: Apo vs. V2Rpp (p = 0.018), Apo vs. C5aR1pp (p = 0.3369), Apo vs. CXCR4pp (p = 0.9338); for β arr2^{DM}: Apo vs. V2Rpp (p < 0.0001), Apo vs. C5aR1pp (p < 0.0001), Apo vs. CXCR4pp (p < 0.0001) (*p < 0.05, ****p < 0.0001).

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Structure-guided engineering yields structures of activated β arr2

As mentioned earlier, the structural coverage of active β arrs, either in complex with full GPCRs or GPCR-derived phosphopeptides, is limited primarily to β arr1. Activated structures of the other isoform i.e., β arr2 are represented only by an IP6-bound crystal structure³⁴ and a complex of truncated β arr2 with a phosphopeptide derived from a β arr-biased seven-transmembrane receptor (7TMR) (CXCR7).³³ Therefore, we set out to reconstitute and determine the structure of β arr2 in complex with the phosphopeptides derived from different receptors, i.e., C5aR1pp and CXCR4pp using cryo-EM. Surprisingly, however, we did not observe a significant Fab30 reactivity to C5aR1pp/CXCR4pp-bound β arr2 while it robustly recognized the V2Rpp- β arr2 complex (Figure 3A). Therefore, we analyzed the Fab30 interaction interface on the C5aR1pp-bound β arr1 structure to identify a potential reason for the lack of Fab30 reactivity with β arr2. Interestingly, we observed that Fab30 epitope is conserved between β arr1 and 2 with the exception of two residues, i.e., instead of F²⁷⁷ and A²⁷⁹ as in β arr1, β arr2 contains L²⁷⁸ and S²⁸⁰ in the corresponding positions, respectively (Figure 3B). Therefore, we generated a β arr2 double mutant, referred to as β arr2^{DM}, and tested its reactivity to Fab30 upon activation by distinct phosphopeptides. In line with our hypothesis, we observed a robust interaction of Fab30 with C5aR1pp- β arr2^{DM} and CXCR4pp- β arr2^{DM} complex, and we also noticed that the interaction of Fab30 with V2Rpp- β arr2^{DM} was further enhanced compared with β arr2^{WT} (Figure 3C). We also confirmed that β arr2^{DM} exhibits a similar pattern of interaction as β arr2^{WT} with V2R and C5aR1 in cellular context (Figure 3D) and shows near-identical endosomal trafficking as β arr2^{WT} upon the stimulation of V2R and C5aR1 (Figure 3E). The receptor surface expression was assessed by whole-cell-based surface ELISA assay (Figure S5A). Thus, β arr2^{DM} provides us with an excellent handle to reconstitute stable complexes with receptor phosphopeptides suitable for cryo-EM. In fact, we successfully managed to reconstitute monodisperse V2Rpp- β arr2^{DM}-Fab30 and C5aR1pp- β arr2^{DM}-Fab30 complexes (Figures S5C and S5I) and determine their cryo-EM structures at 3.96 and 4.33 Å resolution, respectively (Figures 4A, 4B, S5D–S5H, and S5J–S5N). In order to simplify the discussion, we refer to β arr2^{DM} as β arr2 from here onward unless specified otherwise.

Structures of β arr2 in complex with V2Rpp and C5aR1pp

The V2Rpp- β arr2 and C5aR1pp- β arr2 structures exhibited a trimeric assembly of β arr2 with the individual protomers arranged through N- to C-domain contacts (Figures 4A–4D and S7; Table S2). The overall structural features of the individual protomers in each structure were nearly identical as reflected by low RMSD (<0.5 Å) with the phosphopeptide densities clearly visible in the EM maps (Figures S3C and S3D). Like β arr1 structures, V2Rpp and C5aR1pp are positioned in a positively charged groove on the N-domain of β arr2 (Figures 4E and 4F),

and the phosphate moieties make extensive contacts with Arg/Lys residues at the binding interface (Figures 5A and 5B). Remarkably, we observed that three phosphate groups arranged in a P-X-P-P pattern in the phosphopeptides, engage an analogous set of Lys/Arg residues as in β arr1 (Figures 5A and 5B). Specifically, pT³⁶⁰-A³⁶¹-pS³⁶²-pS³⁶³ in V2Rpp and pT³³⁶-R³³⁷-pS³³⁸-pT³³⁹ in C5aR1pp engage K²⁹⁵-K¹²-R²⁶-R⁸-K¹¹-K¹⁰⁸ in β arr2 (Figures 5A and 5B). Similar to β arr1 structures, the other phosphate groups present in the phosphopeptides are either not involved in direct contact with Arg/Lys or positioned outside the binding groove. The N- and C-domains of β arr2 exhibit an inter-domain rotation of approximately 25° when compared with the basal state of β arr2 in both the structures, which is relatively higher from that observed in phosphopeptide-bound β arr1 (Figure 5C). Moreover, the three major loops in β arr2 namely, the finger, middle, and lariat loop also exhibit significant reorientation upon binding of V2Rpp and C5aR1pp compared with the basal state, although their positioning is almost identical between the two structures (Figure 5D). Finally, the three-element interaction and polar-core network in β arr2 are also disrupted upon binding to V2Rpp and C5aR1pp when compared with the basal state structure through the displacement of the carboxyl terminus of β arr2 from the N-domain and repositioning of the lariat loop through the interaction of K²⁹⁵ with a phosphate moiety (Figures 5E and 5F).

Identification of a key phosphorylation motif driving β arr activation

As mentioned earlier, the analysis of these structural snapshots in terms of phosphorylation sites revealed a P-X-P-P type pattern with nearly identical interactions with analogous residues in β arr1 and 2 (Figures 6A–6C). Therefore, we analyzed the primary sequence of all non-olfactory and non-orphan GPCRs in their carboxyl terminus and ICL3 to identify the occurrence of the P-X-P-P pattern in these receptors (Table S1). We observed that a large set of GPCRs harbored this motif in their carboxyl terminus sequence and several receptors also included it in their ICL3 (Figures 6F and 6G). In order to validate the functional contribution of this motif in GPCR-induced β arr activation, we employed Ib30-based conformational biosensor in cellular context using three different receptors, which possess P-X-P-P motif either in their C terminus (CXCR3), ICL3 (M2R), or lack it (CXCR7). At the level of β arr1 conformation, the Ib30 sensor reports the degree (>15°) of inter-domain rotation as a proxy of β arr activation upon its interaction with activated and phosphorylated receptors.^{40,41–43} In agreement with our hypothesis, we observed robust reactivity of Ib30 sensor with β arr1 for CXCR3 and M2R but not for CXCR7 although CXCR7 is capable of recruiting β arr1 upon agonist stimulation^{44–46} (Figure 7A). We note however that the lack of Ib30 reactivity for CXCR7 does not indicate the absence of β arr1 activation but instead, a different active conformation that is not recognized by the Ib30 sensor. Surface expression of the receptors was confirmed in

(D) A side-by-side comparison of agonist-induced β arr2^{WT} and β arr2^{DM} recruitment to V2R (top) and C5aR1 (bottom) in the NanoBIT assay (Receptor-SmBIT+LgBIT- β arr2) (mean \pm SEM; n = 4; normalized as fold over basal).

(E) A side-by-side comparison of β arr2^{WT} and β arr2^{DM} endosomal trafficking in response to agonist (arginine vasopressin peptide [AVP] for V2R and C5aR1) in the NanoBIT assay (Receptor+SmBIT- β arr2+LgBIT-FYVE) (mean \pm SEM; n = 5; normalized as fold over basal). See also Figure S5.

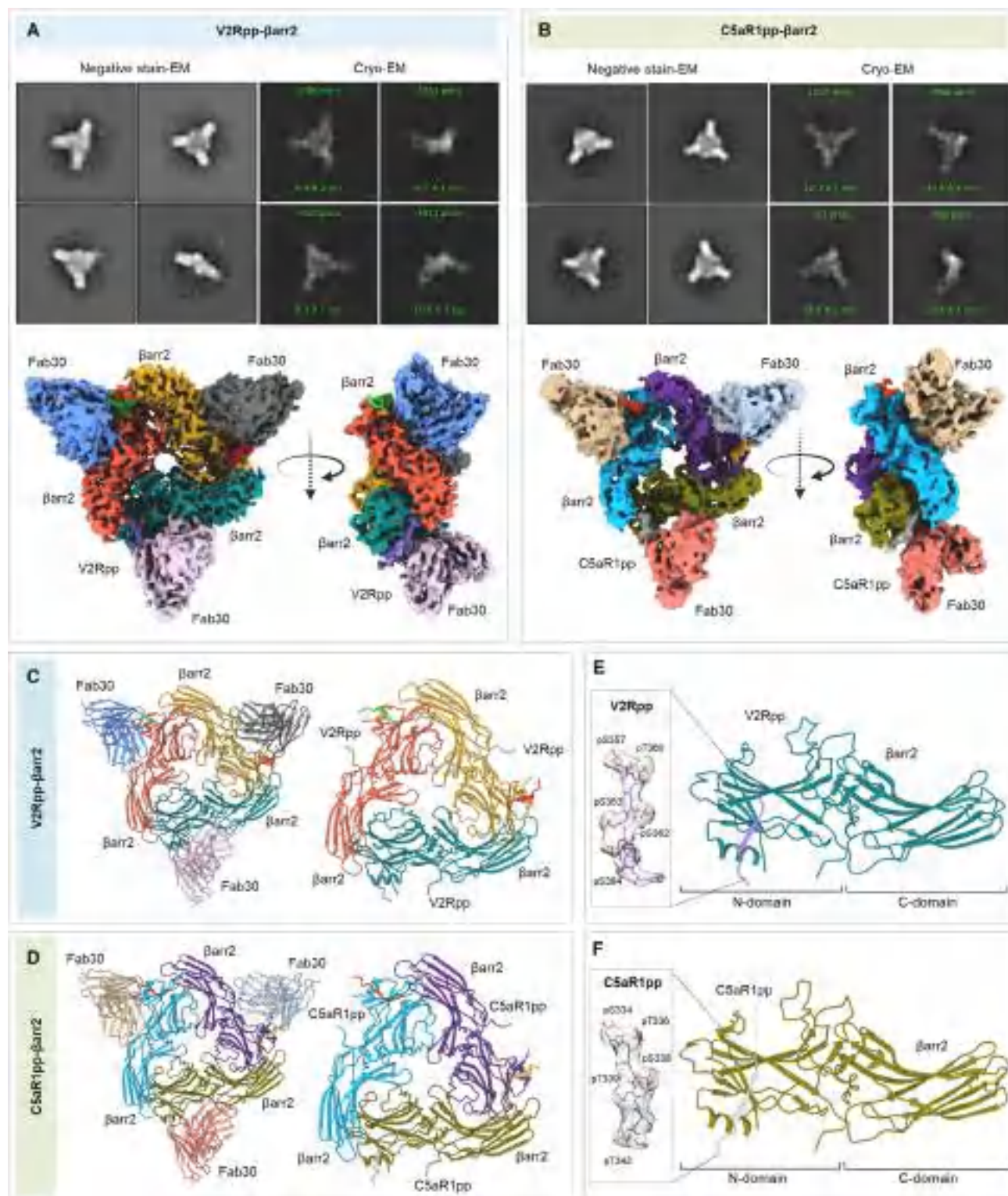


Figure 4. Structures of V2Rpp/C5aR1pp-βarr2 complexes

(A and B) Overall cryo-EM structures of V2Rpp-βarr2-Fab30 (left) and C5aR1pp-βarr2-Fab30 complexes (right), respectively, in a trimeric assembly with βarr2 and Fab30 molecules colored as individual units. Front and side views of the trimeric complex EM maps have been shown with βarr2 molecules in blue, olive green,

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these assays using the whole-cell ELISA assay (Figure S6B). To further corroborate these findings, we used two different receptors namely, the Bradykinin receptor subtype 2 (B2R) and C5aR1 for structure-based targeted mutagenesis to probe gain of function and loss of function, respectively, in terms of Ib30 reactivity pattern. As presented in Figure 7B, the activation of the wild-type B2R fails to induce an interaction between β arr1 and Ib30 as it lacks a P-X-P-P motif in its C terminus, although B2R is capable of recruiting β arrs.^{43,47} Similar to CXCR7, the lack of Ib30 reactivity for B2R likely suggests a distinct activated conformation in β arr1 that is not efficiently recognized by Ib30 sensor. Interestingly however, reconstitution of the P-X-P-P motif in B2R by double mutation ($\Delta G^{368}+L^{370}T$) results in robust Ib30 reactivity upon agonist stimulation (Figure 7B). Along the same lines, a mutant version of C5aR1pp, where the P-X-P-P motif is disrupted by the insertion of an additional arginine residue between pT³³⁶ and pS³³⁸, completely loses the ability to induce a conformation that is recognizable by Fab30 (Figure 7C). Moreover, the corresponding mutation in C5aR1 also leads to a dramatic loss of Ib30 reactivity in cellular context (Figure 7C). Finally, we also disrupted a P-X-P-P motif present in the ICL3 of M2R by site-directed mutagenesis and measured the reactivity of Ib30 upon agonist stimulation (Figure 7D). We observed that similar to C5aR1, the disruption of the P-X-P-P motif in M2R also results in a marked decrease in Ib30 reactivity (Figure 7D). Taken together these data establish that the P-X-P-P phosphorylation pattern, when present in GPCRs, serves as a critical determinant for interaction and activation of β arrs (Figure 7E). In these experiments, the wild-type and mutant receptors were expressed at comparable levels as assessed in the surface expression assay (Figures S6C–S6E).

DISCUSSION

Understanding the contribution of specific phosphorylation patterns in GPCRs for β arr recruitment, activation, and functional outcomes has been a key focus area in the field of GPCR biology, and several previous studies have provided interesting insights into this. For example, the bar-code hypothesis suggests differential contribution of distinct GPCR phosphorylation patterns in inducing specific conformations in β arrs that are linked to corresponding functional outcomes.^{28,50} Taking lead from the rhodopsin-arrestin fusion protein structure determined by X-ray free electron laser (XFEL), a previous study identified and validated PXPXP and PXXPPX type phosphorylation codes in GPCRs that appear to be critical for β arr recruitment.^{51,52} Subsequently, another study used peptide array and biophysical approaches to identify and propose a framework of the “key,” “inhibitory,” and “modulatory” phosphorylation sites in GPCRs that influence β arr binding, activation, and global conformation.⁵³ Additionally, we have also previously explored the func-

tional contribution of phosphorylation codes in several GPCRs to establish a link between different codes and ERK1/2 mitogen-activated protein (MAP) kinase activation using biochemical and cellular experiments.⁴³ This study now visualizes the molecular interaction of multiple GPCR phosphorylation patterns with β arrs and identifies a P-X-P-P motif that plays a crucial role in β arr binding and activation when present in the interacting receptor.

The structural snapshots determined here reveal that the P-X-P-P phosphorylation pattern simultaneously engages multiple elements in β arrs that are typically responsible for maintaining the inactive conformation in order to facilitate β arr activation. It is intriguing that the P-X-P-P phosphorylation pattern from different GPCR phospho-peptides utilizes a conserved set of interactions and docking interface on both isoforms of β arrs (Figures 6D and 6E). This conserved binding interface and corresponding interactions ensure the displacement of β arrs' C terminus from the N-domain and repositioning of theariat loop, leading to the release of the two major “breaks” on β arr activation namely, the three-element interaction and the polar-core network^{10,11,54} (Figure S6A). It is important to note that some GPCRs tested here such as B2R and CXCR7, which lack the P-X-P-P motif, do not exhibit Ib30 reactivity although they are capable of recruiting β arrs in functionally competent conformation as reported previously.^{43–45,47} Taken together, these data indicate that although P-X-P-P motif is not always essential for β arr binding, however, when present in GPCRs, it contributes critically in β arr interaction, and imparts an active conformation on β arrs through a structurally conserved mechanism. On the other hand, in the absence of the P-X-P-P motif, the receptors likely induce a distinct active conformation in β arrs that is also capable of directing functional responses. These scenarios underscore the structural diversity engrained in the GPCR- β arr system, which orchestrates their functional diversity and versatility. The P-X-P-P motif, when present in GPCRs, appears to be sufficient to activate β arrs as measured using Ib30 sensor; however, additional phosphorylation sites may also contribute to improve the affinity of GPCR- β arr interaction and exert modulatory activities as proposed earlier.^{52,53} Moreover, several GPCRs harbor multiple copies of P-X-P-P motif in their carboxyl terminus and ICL3 (Table S2), similar to PXPXP and PXXPPX codes as reported earlier.⁵² While it is tempting to speculate that different P-X-P-P motif may be utilized in a context-dependent fashion such as cell type-specific, GRK-specific, and ligand-specific manner, further studies are essential to systematically probe these interesting possibilities.

An intriguing observation in these structural snapshots is novel dimer and trimer assemblies of β arrs. Although β arrs have a strong propensity to adopt different oligomeric states,^{55–57} the dimer and trimer interfaces observed here differ significantly from previously reported interfaces^{34,56,57} (Figure S7). The

and purple; and Fab30 molecules in beige, red, and gray, respectively. The constant domain densities of Fab30 were removed during local refinement and not observed in the final cryo-EM maps.

(C and D) Overall trimeric arrangement of V2Rpp/C5aR1pp- β arr2 complexes in the cryo-EM structures shown here as cartoon representation (left). Domain organization of the β arr2 molecules in trimeric assembly without Fab30 shown as cartoon representation (right).

(E and F) Structure of individual β arr2 protomers in V2Rpp/C5aR1pp- β arr2 complexes showing the binding of phosphopeptides on the N-domain of β arr2. Coulombic densities of corresponding phosphopeptides have been provided in insets. See also Figures S3 and S5 and Table 1.

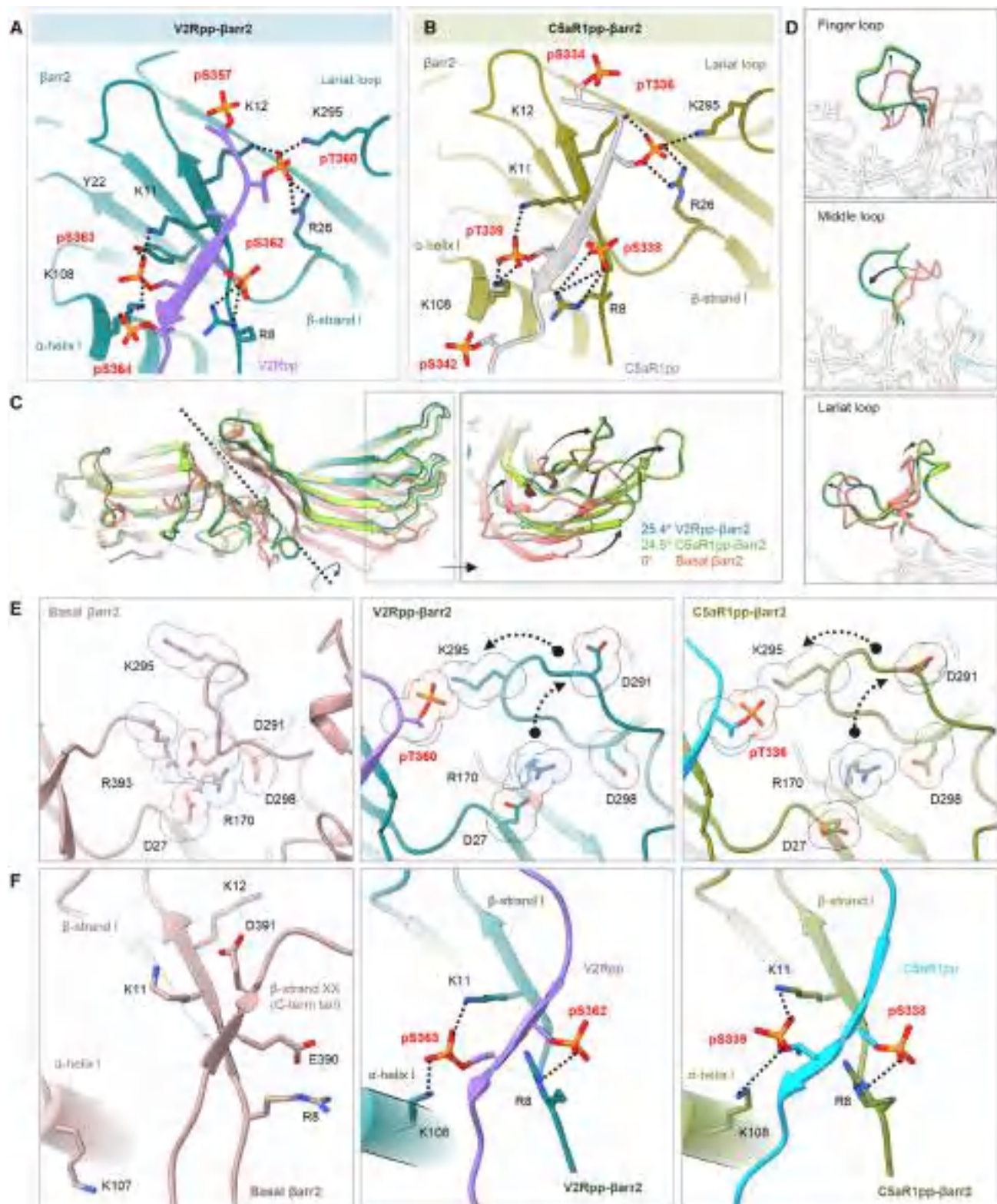


Figure 5. Active conformations of phosphopeptide-bound βarr2

(A and B) Extensive charge-charge interactions between the phosphate residues in V2Rpp/C5aR1pp with Lys/Arg in the N-domain (represented as black dotted lines) stabilize the V2Rpp and C5aR1pp into the N-domain groove of βarr2.

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overall buried surface area in dimer and trimer assemblies are approximately 1,500 and 4,500 Å², respectively, suggesting a robust and stable oligomeric arrangement. The two protomers in C5aR1pp- and CXCR4pp-bound β arr1 interface with each other through multiple hydrogen bonds, salt bridges, and non-bonded contacts in a manner where the C-edge loop residues of one protomer are positioned into the central crest of the other protomer in the proximity of the finger loop. An analogous set of interactions are also involved in the trimer arrangement of β arr2 in complex with V2Rpp and C5aR1pp. Interestingly, a previous crystal structure of β arr2 in complex with IP6 also shows a trimeric arrangement although the trimer interface is different from that observed here in phosphopeptide-bound conformations.³⁴ A comprehensive map of dimer and trimer interface with residue-level contacts is listed in Table S2. Considering the functional multiplicity of β arrs in terms of distinct signaling and regulatory outcomes and receptor-specific responses,⁵⁸ it is possible that distinct oligomeric interfaces in β arrs may be a modular mechanism to fine-tune the functional contributions by providing distinct possibilities for adaptable protein-protein interaction interfaces for binding partners. Nonetheless, the biological implications of the oligomeric assemblies observed here for activated β arrs remain to be explored further in future studies. The comparison of V2Rpp- and C5aR1pp-bound β arr1 and 2 reveal a significantly higher inter-domain rotation in β arr2 compared with β arr1 as hypothesized earlier based on cellular and biochemical studies,⁴⁰ and this may provide a plausible explanation for functional differences between the β arr isoforms as observed for multiple GPCRs.³⁰ However, structural visualization of GPCR- β arr2 complexes would be required to validate this possibility.

In summary, guided by the structural snapshots, we identify and experimentally validate a P-X-P-P motif in GPCRs that imparts an active conformation on β arrs by engaging a conserved network of interacting residues in their N-domain. Our study therefore provides an important missing link in our current conceptual framework of GPCR-mediated β arr activation and paves the way to decipher the structural and functional diversity encoded in GPCR signaling and regulatory paradigms. We also note that a companion manuscript presenting the crystal structures of β arr1 in complex with phosphopeptides derived from the chemokine receptor CCR5 converges to similar conclusions about a critical contribution of the P-X-P-P motif in GPCRs on β arr interaction and activation.⁵⁹

Limitations of the study

The structural snapshots presented here involve isolated phosphopeptides with defined phosphorylation patterns without the

transmembrane core of the receptors. As the interaction of receptor transmembrane core is also important for generating fully engaged GPCR- β arr complexes,^{60–63} it is possible that β arrs exhibit additional conformational changes in complex with activated and phosphorylated receptors. However, the ability of the P-X-P-P motif, when present in GPCRs, to engage the K-K-R-R-K-K pattern in β arrs is likely to be maintained and guide β arr activation even in the context of full-length receptors. Additionally, we cannot rule out the possibility that for some receptors, a functional P-X-P-P motif may not be generated in cellular context despite having a suitable primary sequence because all the phosphorylatable residues may not undergo efficient phosphorylation.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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- **QUANTIFICATION AND STATISTICAL ANALYSIS**

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.molcel.2023.04.025>.

(C) V2Rpp (dark green) and C5aR1pp (light green) activated β arr2 structures were superimposed with the basal state of β arr2 (PDB: 3P2D, orange), and inter-domain rotations were calculated.

(D) Conformational changes observed in the finger (top), middle (middle), and lariat loops (bottom) in the activated β arr2 compared with the basal state crystal structure of β arr2.

(E) Polar-core environment in basal β arr2 (PDB: 3P2D, left) and disruption of polar-core interactions upon binding of V2Rpp (middle) and C5aR1pp to β arr2 (right).

(F) Three-element interaction network consisting of β arr2 C-terminal β -strand XX, α -helix I, and β -strand I in the basal state of β arr2 (left). Binding of the phosphopeptides V2Rpp and C5aR1pp to β arr2 results in the displacement of the β -strand XX, and engages the phosphopeptide V2Rpp (middle) and C5aR1pp (right) into the N-domain groove of β arr2 through hydrogen bonds and polar interactions. See also Figure S6.

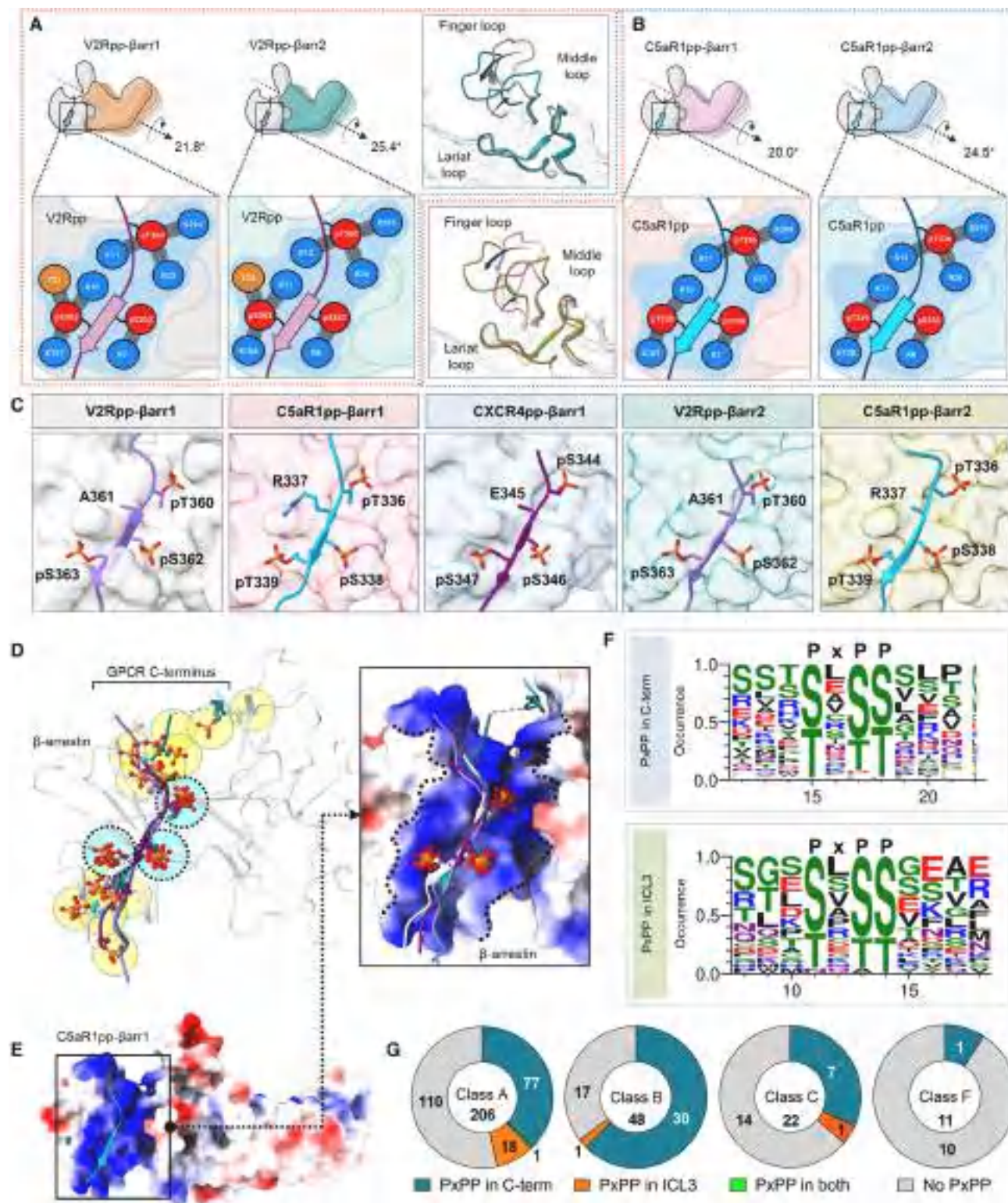


Figure 6. Identification of a key phosphorylation motif in GPCRs driving β arr activation

(A) Comparison of the V2Rpp-bound β arr1 and β arr2 structures reveals similar interactions of V2Rpp with both isoforms of β arrs although a slightly higher inter-domain rotation is observed in β arr2 (left). A schematic representation of the interface network between negatively charged phospho-residues (red) and positively

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AUTHOR CONTRIBUTIONS

J.M. and M.K.Y. prepared and characterized the β arr complexes. J.M. performed negative-staining EM with R.B. and processed the cryo-EM data with R.B. P.S. carried out all the functional assays related to β arr2^{DM} characterization and Ib30 sensor assay. M.K.Y. purified β arrs and carried out the colP experiments with V.S. and Sayantan Saha. Shirsha Saha contributed to the functional characterization of β arr2^{DM}. M.C. screened the samples and collected cryo-EM data. A.K.S. supervised and managed the overall project. All authors contributed to data analysis, interpretation, and manuscript writing.

DECLARATION OF INTERESTS

The authors declare no competing interests.

INCLUSION AND DIVERSITY

We support inclusive, diverse, and equitable conduct of research.

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charged residues (blue) of β arrs is shown (below, zoomed-in box). Although the lariat loops of the two structures align well, significant deviations can be observed for the finger and middle loops (right, inset box).

(B) Comparative analysis of C5aR1pp-bound β arr1 and β arr2 structures uncover similar interactions of C5aR1pp with both β arr isoforms, but again, a higher inter-domain rotation is observed for β arr2. A similar representation of the interface between negatively charged phospho-residues (red) and positively charged residues (blue) of β arrs is shown (below, zoomed-in box).

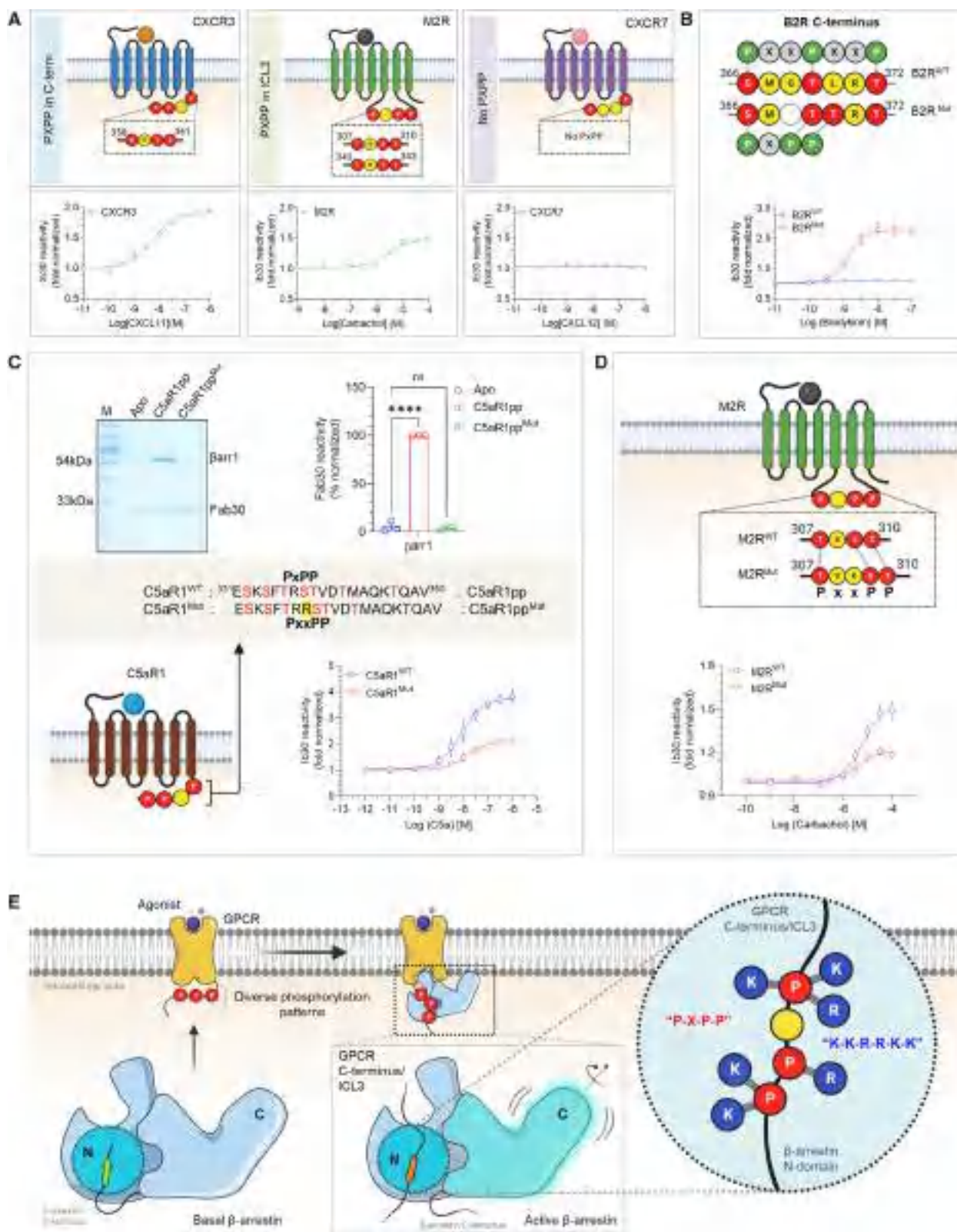
(C) In all the structures of phosphopeptide-bound β arrs, a conserved motif can be observed with respect to three phospho-residues, referred to as the P-X-P-P motif, where “P” is a phospho-Ser/Thr and “X” can be any other residue.

(D) Superposition of V2Rpp- β arr1 (PDB 4JQI), V2Rpp- β arr2, C5aR1pp- β arr1, CXCR4pp- β arr1, and C5aR1pp- β arr2 shows conservation of phosphates corresponding to P-X-P-P position, whereas other phosphates are distributed throughout the phosphopeptides.

(E) Superposition of phosphopeptides on C5aR1pp- β arr1 reveals the conserved phospho-residues on positively charged cleft present on β arrs' N-domain. β arr is shown as Coulombic-charged surface here.

(F) A sequence alignment of the C-terminal tail and ICL3 residues of non-olfactory and non-orphan class A receptors reveal the consensus sequence, “P-X-P-P” required for activation of β arrs. The consensus sequence logo was generated with the WEBLOGO tool⁴⁸ and sequence alignment was performed with Kalign.⁴⁹ A stretch of 11 amino acid residues has been shown for better representation.

(G) Proportions of GPCRs of class A, B, C, and F having P-X-P-P motif in C terminus or ICL3 have been represented as pie charts. See also Figure S6 and Table S1.



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Figure 7. A key phosphorylation motif for β arr activation

- (A) NanoBIT-based assay for assessing Ib30 reactivity to CXCR3 (left), M2R (middle), and CXCR7 (right) activated β arr1 (Receptor+SmBIT- β arr1+LgBIT-Ib30) (mean \pm SEM; n = 3; normalized as fold over basal).
- (B) Deletion of G³⁶⁸ and substitution of L³⁷⁰ to Thr in B2R engineers the P-X-P-P (referred to as B2R^{Mut}) and results in gain of function in terms of Ib30 reactivity as measured using the NanoBIT assay (Receptor+SmBIT- β arr1+LgBIT-Ib30) (mean \pm SEM; n = 3; normalized as fold over basal).
- (C) Addition of an extra Arg between positions 336 and 337 in C5aR1pp to disrupt the P-X-P-P (referred to as C5aR1pp^{Mut}) leads to a near-complete loss of Fab30 (top) reactivity as measured in colP assay (mean \pm SEM; n = 3; densitometry-based data normalized with respect to C5aR1pp signal as 100%; one-way ANOVA, Dunnett's multiple comparisons test). The exact p values are as follows: Apo vs. C5aR1pp2 (p < 0.0001), Apo vs. C5aR1pp4 (p = 0.9160) (****p < 0.0001, ns, non-significant). Corresponding mutation in C5aR1 to disrupt the P-X-P-P (referred to as C5aR1^{Mut}) results in a dramatic decrease in Ib30 reactivity (bottom) as measured using the NanoBIT assay (mean \pm SEM; n = 5; normalized as fold over basal).
- (D) Insertion of a Lys residue between positions 308 and 309 to disrupt the P-X-P-P pattern in ICL3 of M2R (referred to as M2R^{Mut}) shows reduced Ib30 reactivity compared with the wild type (M2R^{WT}) as measured using the NanoBIT assay (mean \pm SEM; n = 4; normalized as fold over basal).
- (E) Schematic representation summarizing the identification of a phosphorylation motif in GPCRs that drives β arr activation. See also Figure S6.

Table 1. Cryo-EM data collection, processing, and refinement statistics, related to Figures 1, 4, S2, and S5

	C5aR1pp- βarr1-Fab30	C5aR1pp-βarr1- Fab30 (local refined)	V2Rpp- βarr2-Fab30	V2Rpp-βarr2-Fab30 (local refined)	C5aR1pp- βarr2-Fab30	C5aR1pp-βarr2- Fab30 (local refined)	CXCR4pp- βarr1-Fab30	CXCR4pp-βarr1- Fab30 (local refined)
Code	PDB: 8GO8 EMD 34173	PDB: 8I0N EMD 35104	PDB: 8GOC EMD 34175	PDB: 8I10 EMD 35115	PDB: 8GOO EMD 34178	PDB: 8I0Z EMD 35114	PDB: 8GP3 EMD 34188	PDB: 8I0Q EMD 35106
Microscope	Titan Krios	Titan Krios	Titan Krios	Titan Krios	Glacios	Glacios	Glacios	Glacios
Camera	GIF/K2	GIF/K2	GIF/K2	GIF/K2	Gatan K3	Gatan K3	Gatan K3	Gatan K3
Magnification	165,000	165,000	165,000	165,000	46,000	46,000	46,000	46,000
Voltage (kV)	300	300	300	300	200	200	200	200
Defocus range (μm)	0.5–2.5	0.5–2.5	0.5–2.5	0.5–2.5	0.5–2.5	0.5–2.5	0.5–2.5	0.5–2.5
Exposure time (s)	5	5	4	4	3	3	2.9	2.9
Total dose (e ⁻ /Å ²)	49	49	48.7	48.7	51	51	49.38	49.38
Number of frames	40	40	40	40	40	40	40	40
Pixel size (Å)	0.82	0.82	0.82	0.82	0.878	0.878	0.878	0.878
Micrographs (no.)	6,212	6,212	9,720	9,720	8,614	8,614	5,637	5,637
Initial particles (no.)	4,304,237	4,304,237	2,444,407	2,444,407	4,012,616	4,012,616	3,236,193	3,236,193
Symmetry imposed	C2	C2	C3	C3	C3	C3	C2	C2
Final particles (no.)	80,437	80,437	92,018	92,018	38,206	38,206	86,525	86,525
FSC threshold	0.143	0.143	0.143	0.143	0.143	0.143	0.143	0.143
Map resolution (Å)	3.41	3.26	4.18	3.96	4.41	4.33	4.81	4.45
Refinement								
Initial code (PDB)	4JQI	8GO8	5TV1	8GOC	8GOC	8GOO	8GO8	8GP3
Model resolution (Å)	3.5	3.4	4.7	4.1	4.7	4.7	5.3	5.7
FSC threshold	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5
Model composition								
Non-hydrogen atoms	11,519	9,141	16,942	13,205	16,966	13,163	11,494	8,842
Protein residues	1,522	1,190	2,223	1,729	2,223	1,725	1,522	1,192
Ligand atoms	0	0	0	0	0	0	0	0
RMSD								
Bond length (Å)	0.004	0.005	0.003	0.007	0.004	0.006	0.005	0.004
Bond angle (°)	0.942	1.001	0.716	1.256	0.665	1.128	0.982	0.967
Validation								
Favored (%)	97.23	95.43	95.05	93.63	94.90	93.24	96.62	94.92
Allowed (%)	2.77	4.57	4.76	6.37	4.96	6.76	3.38	5.08
Disallowed (%)	0	0	0.18	0	0.14	0	0	0
MolProbity score	1.41	1.46	1.97	1.88	1.91	1.77	1.77	1.85
Clash score	5.23	3.49	13.07	8.60	10.83	6.12	10.81	9.44

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Monoclonal ANTI-FLAG M2-HRP antibody	Sigma-Aldrich	Cat# A8592; RRID: AB_439702
Chemicals, peptides, and recombinant proteins		
TRIS	SRL	Cat# 71033
HEPES	SRL	Cat# 63732
NaCl	SRL	Cat# 41721
EDTA	SRL	Cat# 12070
Phenylmethanesulfonyl Fluoride (PMSF)	SRL	Cat# 84375 (84375)
Benzamidine Hydrochloride	SRL	Cat# 93014 (0248255)
Lysozyme	SRL	Cat# 45822
Glycerol	SRL	Cat# 77453
Dithiothreitol	HiMedia	Cat# MB070
Lauryl Maltose Neopentyl Glycol (MNG)	Anatrace	Cat# NG310, CAS no.1257852-96-2
Paraformaldehyde (PFA)	Sigma Aldrich	Cat# P6148, CAS no. 30525-89-4
Poly-D-lysine	Sigma Aldrich	Cat# P0899
TMB (Tetramethylbenzidine)	Thermo Fisher Scientific	Cat# 34028
Janus Green B	Sigma Aldrich	Cat# 201677
PEI (Polyethylenimine)	Polysciences	Cat# 23966
Bovine Serum Albumin, BSA	SRL	Cat# 83803 (0140105)
HBSS - Hank's Balanced Salt Solution	Thermo Fisher Scientific	Cat# 14065
GIBCO Fetal Bovine Serum	Thermo Fisher Scientific	Cat# 10270-106
DMEM	Cellclone	Cat# CC3004
Phosphate-buffered saline (PBS)	Sigma Aldrich	Cat# D1283
GIBCO Penicillin-Streptomycin	Thermo Fisher Scientific	Cat# 15140122
Coelenterazine	Goldbio	Cat# CZ05
Glyco-diosgenin (GDN)	Anatrace	GDN101
Cholesteryl Hemisuccinate	Sigma	C6512
Coomassie brilliant Blue	SRL	Cat# 64222
Uranyl formate	Polysciences	Cat# 24762-1
Recombinant rat β -arrestin1	Purified	N/A
Recombinant bovine β -arrestin2	Purified	N/A
C5aR1pp1	Chemically synthesized	N/A
C5aR1pp2	Chemically synthesized	N/A
C5aR1pp3	Chemically synthesized	N/A
V2Rpp	Chemically synthesized	N/A
CXCR4pp1	Chemically synthesized	N/A
CXCR4pp2	Chemically synthesized	N/A
CXCR4pp3	Chemically synthesized	N/A
CXCR4pp4	Chemically synthesized	N/A
Recombinant human C5a	Purified	N/A
Bradykinin	Genscript	N/A
Arginine Vasopressin Peptide (AVP)	Genscript	N/A
Uranyl formate	Polysciences	Cat# 24762-1

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Formvar/carbon coated 300 mesh copper grids	PELCO (Ted Pella)	Cat# 01753-F
Critical commercial assays		
Site directed mutagenesis kit	NEB	Cat# E0554
NanoBiT assay	Promega	N/A
Deposited data		
C5aR1pp- β arr1-Fab30	This study	PDB: 8GO8, EMD-34173
V2Rpp- β arr2-Fab30	This study	PDB: 8GOC, EMD-34175
C5aR1pp- β arr2-Fab30	This study	PDB: 8GOO, EMD-34178
CXCR4pp- β arr1-Fab30	This study	PDB: 8GP3, EMD-34188
C5aR1pp- β arr1-Fab30-Local-refine	This study	PDB: 8I0N, EMD-35104
V2Rpp- β arr2-Fab30-Local-refine	This study	PDB: 8I10, EMD-35115
C5aR1pp- β arr2-Fab30-Local-refine	This study	PDB: 8I0Z, EMD-35114
CXCR4pp- β arr1-Fab30-Local-refine	This study	PDB: 8I0Q, EMD-35106
Experimental models: Cell lines		
Human: HEK293	ATCC	Cat# CRL-3216
Oligonucleotides		
C5aR1_R-insertion SDM primer Forward: CGCTCCACAGTGGACACTATGG	This study	N/A
C5aR1_R-insertion SDM primer Reverse: GCGCGTGAATGACTTGCT	This study	N/A
B2R ^{mut} Forward: AACACGGACCTCCATCTCCGTG	This study	N/A
B2R ^{mut} Reverse: GTCATGGAGTTCTCCATCTGAATGGG	This study	N/A
M2R ^{mut} Forward: AAGTCTACTTCACTGGGCCAC	This study	N/A
M2R ^{mut} Reverse: GACGGTGTTTCGTCCTG	This study	N/A
β -arrestin2 Double mutant (β arr2DM) Fw: ggcgAACAACCGTGAAAAACGTG	This study	N/A
β -arrestin2 Double mutant (β arr2DM) Rv: aggaaCGGGGTGATGGGTGTAAC	This study	N/A
Recombinant DNA		
PcDNA3.1 (empty vector)	Dr. Arun K Shukla	N/A
pcDNA3.1_V2R-WT	Dr. Arun K Shukla	N/A
pcDNA3.1_C5aR1-WT	Dr. Arun K Shukla	N/A
pcDNA3.1_CXCR3-WT	Dr. Arun K Shukla	N/A
pcDNA3.1_CXCR7-WT	Dr. Arun K Shukla	N/A
pcDNA3.1_B2R-WT	Dr. Arun K Shukla	N/A
pcDNA3.1_M2R-WT	Dr. Arun K Shukla	N/A
pcDNA3.1_C5aR1-mut ^{TRRST}	This study	N/A
pcDNA3.1_B2R-mut ^{ΔG368/L370T}	Dr. Arun K Shukla	N/A
pcDNA3.1_M2R-mut ^{TVKST}	This study	N/A
pCAGGS_V2R-WT-SmBiT	Dr. Arun K Shukla	N/A
pCAGGS_C5aR1-SmBiT	Dr. Arun K Shukla	N/A
pCAGGS_SmBiT- β arr2 ^{WT}	Dr. Asuka Inoue	N/A
pCAGGS_SmBiT- β arr2 ^{DM}	This study	N/A
pCAGGS_LgBiT- β arr2 ^{WT}	Dr. Asuka Inoue	N/A
pCAGGS_LgBiT- β arr2 ^{DM}	This study	N/A

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
pCAGGS_LgBiT-FYVE	Dr. Asuka Inoue	N/A
pCAGGS_LgBiT-Ib30	Dr. Asuka Inoue	N/A
Software and algorithms		
UCSF Chimera	Pettersen et al. ⁶⁴	https://www.cgl.ucsf.edu/chimera/
UCSF Chimera X	Pettersen et al. ⁶⁵	https://www.rbvi.ucsf.edu/chimerax/
COOT	Emsley et al. ⁶⁶	https://www2.mrc-lmb.cam.ac.uk/personal/pemsley/coot/
cryoSPARC	Punjani et al. ⁶⁷	https://cryosparc.com/
PDBsum	Laskowski et al. ⁶⁸	http://www.ebi.ac.uk/thornton-srv/databases/pdbsum/
Phenix	Liebschner et al. ⁶⁹	https://www.phenix-online.org/
PyMol	Schrodinger	https://pymol.org/2/
Prism 8	GraphPad Software	https://www.graphpad.com/scientific-software/prism/
Relion3.1.2	Zivanov et al. ⁷⁰	https://www3.mrc-lmb.cam.ac.uk/relion/index.php?title=Main_Page
SerialEM	Mastronarde ⁷¹	https://bio3d.colorado.edu/SerialEM/
Graphpad Prism 9	GraphPad Software, San Diego, California USA	https://www.graphpad.com/scientific-software/prism/
ImageJ	Schneider et al. ⁷²	https://imagej.nih.gov/ij/download.html

RESOURCE AVAILABILITY

Lead contact

Further information and requests for reagents should be addressed to the lead contact, Dr. Arun K. Shukla (arshukla@iitk.ac.in).

Materials availability

Reagents described in this manuscript are available upon reasonable request from the [lead contact](#).

Data and code availability

- All three-dimensional cryo-EM density maps, coordinates for the atomic models and local-refined maps generated in this study have been deposited and are publicly available as of the date of publication. Accession numbers (EMDB and PDB IDs) are listed in the [key resources table](#). Original gel images have been deposited to Mendeley data, and they are publicly available after publication. The DOI is listed in the [key resources table](#).
- This paper does not report any original code.
- 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

Human cell lines

HEK-293 cells were purchased from ATCC for all the cellular experiments performed in the study. The cell line was examined frequently under the microscope for proper morphology, but they were not authenticated. They were cultured in DMEM with fetal bovine serum (FBS) at 37°C in 5% CO₂. In this study, any stable, knockout, or knockdown cell lines were not generated, and the details of previously generated cell lines are referenced in the manuscript.

Insect cells

Sf9 cells were obtained from Expression systems, and they were routinely monitored under the microscope for proper morphology. These cells were maintained in a shaker incubator at 27°C with 135rpm shaking, and sub-cultured in protein-free insect cell medium purchased from Expression Systems.

METHOD DETAILS

General reagents, plasmids, and cell culture

Most of the general reagents were purchased from Sigma Aldrich unless mentioned otherwise. Dulbecco's Modified Eagle's Medium (DMEM), Dulbecco's Phosphate buffer saline (PBS), Fetal-Bovine Serum (FBS), Trypsin-EDTA, Hank's balanced salt solution (HBSS), and penicillin-streptomycin solution were purchased from Thermo Fisher Scientific. HEK-293 cells were obtained from ATCC and maintained in DMEM (Gibco, Cat no. 12800-017) supplemented with 10% FBS (Gibco, Cat no. 10270-106) and 100U ml⁻¹ penicillin (Gibco, Cat no. 15140122) and 100μg ml⁻¹ streptomycin (Gibco, Cat no. 15140-122) at 37°C under 5% CO₂. The cDNA coding region for the mentioned receptors namely, V2R, C5aR1, B2R, M2R, CXCR3, and CXCR7 were cloned in pcDNA3.1 consist of HA signal sequence followed by FLAG tag at the N-terminus of the receptor. The mutants generated for the study are as follows: deletion of G368 and a substitution L370T in B2R; insertion of an Arg residue between R337 and S338 in C5aR1; insertion of a Lys residue between V308 and S309 in M2R; using Q5 site-directed mutagenesis kit (NEB, Cat. no. E0554S). For the NanoBiT assay, receptors harboring SmBiT at the C-terminus were generated by subcloning in the lab, and other constructs have been described previously.^{41,43,73} All the constructs were verified by DNA sequencing (Macrogen).

Expression and purification of βarrs

Full length rat βarr1, βarr2^{WT} and bovine βarr2^{DM} were cloned into pGEX-4T3 vector with thrombin cleavage site between GST tag and βarr. Similar protocol was followed for purifying all three forms of βarr. βarrs were expressed in *E. coli* BL21 cells and grown in Terrific broth media supplemented with 100μg ml⁻¹ ampicillin. A primary culture of 50ml volume was inoculated with an isolated colony from freshly transformed LB-amp plate. Primary culture was grown till a cell optical density at 600nm (OD₆₀₀) of 0.8-1 and further inoculated into a secondary culture of TB-Amp of 1.5L volume till OD₆₀₀ 0.8-1. The expression of βarrs was then induced with 25μM IPTG concentration and cells were allowed to grow till 16h at 18°C. Cultures were harvested and stored at -80°C until further use. Harvested pellets were of 12-15g in mass.

For purification, cells were lysed by sonication in lysis buffer; 25mM Tris, pH 8.5, 150mM NaCl, 1mM PMSF (phenylmethylsulfonyl fluoride), 2mM Benzamidine, 1mM EDTA (Ethylenediaminetetraacetic acid), 5% Glycerol, 2mM Dithiothreitol (DTT) and 1mg ml⁻¹ Lysozyme. The lysate was centrifuged at 18,000-20,000rpm at 4°C and supernatant was allowed to bind to Glutathione resin (GS resin) (Glutathione Sepharose™ 4 Fast Flow, GE Healthcare Cat. no. 17-5132-02) in a batch binding mode for overnight at 4°C. GS-resin bound GST-βarr was transferred into Econo columns (Biorad, Cat. no. 7372512) and washed rigorously with wash buffer (25mM Tris, pH 8.5, 150mM NaCl, 2mM DTT and 0.02% n-dodecyl-β-D-maltopyranoside (DDM). Afterward, on-column cleavage was set up by adding thrombin to 1:1 resin:buffer slurry at room temperature for 2h. βarrs were then eluted with gravity flow and further with buffer 25mM Tris, pH 8.5, 350mM NaCl and 0.02% DDM and 2mM DTT. Eluted proteins were concentrated and further purified on a HiLoad 16/600 Superdex column in buffer 25mM Tris, pH 8.5, 350mM NaCl, 2mM DTT and 0.02% DDM. Fractions corresponding to pure βarr were flash frozen with 10% glycerol and stored at -80°C until further use.

Expression and purification of Fabs

A similar protocol for expression and purification was followed for all the Fabs and they were purified as previously mentioned.³⁹ Briefly, Fabs were expressed in the periplasmic fraction of *E. coli* M55244 cells (ATCC) and purified using Protein L resin (GE Healthcare Cat. no. 17547802) with gravity flow affinity chromatography. Cells transformed with Fab plasmid were grown in 50ml 2xYT media and allowed to grow overnight at 30°C. 1L 2xYT media was inoculated with 5% volume of initial inoculum and grown for an additional 8h at 30°C. Cells were collected and resuspended in an equal volume of CRAP medium supplemented with 100μg ml⁻¹ ampicillin, and grown further for 16h at 30°C. For purification, cells were lysed in lysis buffer (50mM HEPES-Na⁺, pH 8.0, 0.5M NaCl, 0.5% (v/v) Triton X-100, 0.5mM MgCl₂) by sonication. Cell lysate was heated in a 65°C water bath for 30min and cooled immediately on ice for 5min. Lysate was centrifuged at 20,000rpm and passed through pre-equilibrated Protein L resin packed gravity flow affinity columns. Binding was performed at room temperature and beads were washed extensively with wash buffer (50mM HEPES-Na⁺, pH 8.0, 0.5M NaCl). Fabs were eluted with 100mM acetic acid into tubes containing 10% volume of 1M HEPES, pH 8.0 for neutralization. Eluted samples were desalted into buffer (20mM HEPES-Na⁺, pH 8.0, 0.1M NaCl) using a pre-packed PD-10 column (GE Healthcare Cat. no. 17085101). Purified Fabs were flash-frozen and stored at -80°C supplemented with 10% (v/v) glycerol until further use.

Co-immunoprecipitation assay

For co-immunoprecipitation assay, 2.5μg of β-arrestins were incubated with different phosphopeptides at 10-fold molar excess in binding buffers (20mM HEPES, pH 7.4, 150mM NaCl) for 1h at room temperature for activation. Post peptide-induced activation, 5μg Fab30 was added, and reaction was incubated for an additional 1h at room temperature. After 1h, 25μl of pre-equilibrated protein L beads (Capto™ L resin, GE Healthcare Cat. no. 17547802) was added and reaction was incubated for 90min at room temperature, followed by five washes with binding buffer containing 0.01% LMNG. Bound protein was eluted with 2X SDS loading buffer and 15μl sample was analyzed on 12% SDS-PAGE. For statistical analyses, protein bands were quantified using ImageJ software suite⁷² and the values were plotted using GraphPad Prism software v 9.5. The data were normalized with respect to their respective experimental control and appropriate statistical analyses were performed as indicated in the corresponding figure legend.

Reconstitution of phosphopeptide- β arr-Fab complexes

A previously published protocol was followed for complex purification with minor modifications.³¹ Briefly, β arrs were activated with corresponding phosphopeptides at a 1:3 molar ratios of β arr:phosphopeptide for 30–40 min at room temperature. Respective Fabs were added to the phosphopeptide- β arr mixture at 1:1.5 molar ratio of β arr:Fab and incubated for 1 h at room temperature. To remove excess Fabs, the phosphopeptide- β arr-Fab complexes were concentrated with 30,000 MWCO concentrators (Vivaspin, Cytiva Cat. no. 28932361) and injected into Superose 6 Increase 10/300 GL (Cytiva Cat. no. 29091596) gel-filtration column. Fractions were further analyzed on SDS-PAGE and selected fractions were pooled and concentrated for structural studies.

Negative-staining EM

Complex formation, homogeneity, and particle quality of the samples were judged through negative staining of the samples prior to data collection under cryogenic conditions for high resolution reconstructions. Negative staining of the samples was performed with uranyl formate in accordance with the previously published protocols.⁷⁴ For imaging, 3.5 μ l of the samples were dispensed on glow discharged carbon/formvar coated 300 mesh Cu grids (PELCO, Ted Pella) and allowed to adsorb for 1 min, followed by blotting off the sample using a filter paper. The grid was then touched on a first drop of freshly prepared 0.75% (w/v) uranyl formate stain and immediately blotted off, followed by staining for 30 sec on a second drop of stain. Imaging of the negatively stained samples were performed on a FEI Tecnai G2 12 Twin TEM (LaB6) operating at 120 kV and equipped with a Gatan CCD camera (4k x 4k) at 30,000x magnification. Data processing of the collected micrographs for the individual samples were performed with Relion 3.1.2.⁷⁰ Approximately 10,000 particles were autopicked using the gaussian blob picker within Relion and the extracted particles were subjected to reference free 2D classification.

Cryo-EM sample preparation and data acquisition

Quantifoil holey carbon grids (Cu or Au, R2/1 or R2/2) were glow discharged for 45 sec with a Glocube glow discharge system (Quorum technologies Ltd, UK). 3 μ l of the complex was dispensed on the glow discharged grid, blotted for 3 sec with a Whatman paper filter no. 1 at 10°C and maintained at 90% humidity and then plunge frozen into liquid ethane (−180 °C) using a Leica GP plunger (Leica Microsystems, Austria).

For C5aR1pp- β arr1-Fab30 complex, cryo-EM data collection was performed on R2/2 Cu 300 mesh grid using a Titan Krios electron microscope (ThermoFisher Scientific, USA) operating at 300 kV equipped with the Gatan Energy Filter. Movies were recorded in counting mode with a Gatan K2 Summit DED (Gatan, USA) using the automated SerialEM software⁷¹ at a nominal magnification of 165,000x and a pixel size of 0.82 Å at sample level. 6,212 movie stacks consisting of 40 frames were recorded over a defocus range of 0.5 to 2.5 μ m with a total dose of 49 e[−]/Å² and total exposure time of 5 sec.

For CXCR4pp- β arr1-Fab30 complex, cryo-EM data collection was performed on R2/2 Cu 300 mesh grid using a Glacios electron microscope (ThermoFisher Scientific, USA) operating at 200 kV. Movies were recorded in counting mode with a Gatan K3 DED (Gatan, USA) using the automated SerialEM software at a nominal magnification of 46,000x and a pixel size of 0.878 Å at sample level. 5,637 movie stacks consisting of 40 frames were recorded over a defocus range of 0.5 to 2.5 μ m with a total dose of 49.3 e[−]/Å² and total exposure time of 2.9 sec.

For V2Rpp- β arr2-Fab30 complex, cryo-EM data collection was performed on R2/2 Au 200 mesh grid using a Titan Krios electron microscope (ThermoFisher Scientific, USA) operating at 300 kV equipped with the Gatan Energy Filter. Movies were recorded in counting mode with a Gatan K2 Summit DED (Gatan, USA) using the automated SerialEM software at a nominal magnification of 165,000x and a pixel size of 0.82 Å at sample level. 9,720 movie stacks consisting of 40 frames were recorded over a defocus range of 0.5 to 2.5 μ m with a total dose of 48.7 e[−]/Å² and total exposure time of 4 sec.

For C5aR1pp- β arr2-Fab30 complex, cryo-EM data collection was performed on R2/2 Cu 300 mesh grid using a Glacios electron microscope (ThermoFisher Scientific, USA) operating at 200 kV. Movies were recorded in counting mode with a Gatan K3 DED (Gatan, USA) using the automated SerialEM software at a nominal magnification of 46,000x and a pixel size of 0.878 Å at sample level. 8,614 movie stacks consisting of 40 frames were recorded over a defocus range of 0.5 to 2.5 μ m with a total dose of 51 e[−]/Å² and total exposure time of 3 sec.

Cryo-EM data processing and model building

All image processing steps were performed in cryoSPARC version 3.3.2⁶⁷ unless otherwise stated. In brief, for the C5aR1pp- β arr1-Fab30 complex, 6,212 movie stacks were subjected to patch motion correction (multi), followed by CTF refinement with patch CTF multi. 5,790 motion corrected micrographs with CTF fit resolution better than 4.5 Å were selected for further processing. 4,304,237 particle projections were automatically picked with blob picker, extracted with a box size of 480 pixels and fourier cropped to 64 pixels. The particle stack so obtained was subjected to multiple rounds of 2D classification. The class averages with clear secondary structural features were selected and re-extracted with a box size of 480 pixels and fourier cropped to 256 pixels resulting in a pixel size of 1.5375 Å. 295,922 re-extracted particles were then subjected to Ab-initio reconstruction and 3D classification/Heterogeneous refinement with C1 symmetry yielding 4 models. 80,437 particles corresponding to a dimer and containing 47.9% of the total particles were subjected to non-uniform refinement with C2 symmetry to yield a map with an estimated resolution of 3.41 Å (voxel size of 1.5375 Å). Further, local refinement was performed by masking out the variable domains of Fab30 resulting into an estimated resolution of 3.26 Å. Local resolution of all reconstructions was estimated using the Blocres within cryoSPARC version 3.3.2.

For the CXCR4pp- β arr1-Fab30 complex data set, 5,637 movies were motion corrected using a patch of 5x5 patch within patch motion correction (multi). Following CTF estimation, 5,236 motion corrected micrographs with CTF fit resolution better than 6Å were curated for further processing. 3,236,193 particles were automatically picked using the blob-picker sub-program and subsequently extracted with a box size of 480pixels and fourier cropped to 64pixels. The extracted particles were subjected to several rounds of 2D classification to remove junk particles. 104,707 particles corresponding to the clean class averages were selected, re-extracted with a box size of 480pixels and fourier cropped to 256pixels (pixel size of 1.65) and used to produce two ab-initio models. The particles corresponding to the two ab-initio models were subjected to heterogenous refinement/3D classification which produced a 3D class with clear dimeric conformation and a particle count of 86,525. This particle set was re-extracted with full box size of 480pixels (pixel size of 0.878) and subjected to non-uniform refinement with C2 symmetry which converge to a map with 4.81Å resolution as estimated using the gold standard Fourier Shell Correlation (GFSC) using the 0.143 criterion. Local refinement was performed by masking out the variable domains of Fab30 which resulted into an estimated resolution of 4.45Å.

For the V2Rpp- β arr2-Fab30 complex, 9,720 movies were motion corrected with 5x5 patches followed by CTF estimation with patch CTF (multi). Following CTF refinement, 8,295 movies with CTF fit resolution better than 4.5Å were used for further processing. Particle picking from the curated micrographs was performed automatically with the blob picker sub-program to obtain an initial stack of 2,444,407 particles. The particles were then extracted with a box size of 512pixels and fourier cropped to a box size of 64pixels. The extracted particles were subjected to several rounds of reference free 2D classification. 2D class averages with evident secondary features containing 161,436 particles were extracted with a box size of 512pixels and fourier cropped to a box size of 256 (pixel size of 1.64). This sub-set of particles was used for ab-initio reconstruction and subsequent rounds of 3D/Heterogeneous classification with C1 symmetry to obtain 2 models. 92,018 particles corresponding to a trimer were re-extracted with full box size of 512pixels which refined to an overall resolution of 4.18Å (voxel size of 0.82Å) with NU refinement (C3 symmetry) according to the gold standard Fourier shell correlation (FSC) criterion of 0.143. Subsequently, local refinement was performed on β arr and the variable domains of Fab30 and resulted into an estimated resolution of 3.96Å.

For the C5aR1pp- β arr2-Fab30 complex data set, 8,614 movies were motion corrected using patch motion correction (multi) and subsequent CTF estimation was performed through patch CTF (multi). 8,157 micrographs with CTF fit resolution better than 6Å were curated for particle picking using the blob picker sub-program. 4,012,616 particles were automatically picked and extracted with a box size of 512pixels and fourier cropped to 64pixels. Reasonable class averages after several rounds of reference free 2D classification yielded a particle set containing 54,193 particle projections, which was re-extracted with a box size of 512pixels and fourier cropped to 360pixels (pixel size of 1.2487Å) for subsequent used for ab-initio reconstruction generating two ab-initio models. Following heterogenous refinement/3D classification, the 3D class with evident features of a trimer and containing 38,206 particles was subjected to non-uniform refinement with C3 symmetry to yield a reconstruction at 4.41Å (final voxel size of 1.2487Å) as determined by gold standard Fourier Shell Correlation (FSC) using the 0.143 criterion. Local refinement was performed masking out the variable domains of Fab30 which improved the resolution to 4.33Å.

Model building and refinement

Coordinates from a previously solved V2Rpp bound β arr1 structure (PDB 4JQI) was used to dock the model into the EM density map of C5aR1pp- β arr1-Fab30 using Chimera.⁶⁴ The EM map was then used for manual rebuilding of the β arr1 residues and placing the phosphopeptide in COOT.⁶⁶ The rebuilt model was subjected to real space refinement in Phenix⁶⁹ to obtain a model with 97.23% of the residues in most favored region and 2.77% in the allowed region of the Ramachandran plot.

The protomeric structure from the IP6- β arr2 (PDB 5TV1) complex solved in a previous study was used as an initial model to dock into the density map of V2Rpp- β arr2-Fab30 complex and regenerate the trimeric complex with C3 symmetry. The rigid body fitted trimeric model and the phosphopeptides were then rebuilt manually into the EM density map. The rebuilt trimeric coordinates with the phosphopeptides were subsequently subjected to real space refinement in Phenix to reach a final model with 95.05% in the favored region and 4.76% in the allowed region of the Ramachandran plot.

For model building into the 4.33Å C5aR1pp- β arr2-Fab30 coulombic map, the co-ordinates corresponding to V2Rpp peptide were deleted from the trimeric co-ordinates of V2Rpp- β arr2-Fab30 complex (PDB 8GOC), and the resulting model was docked into the EM map in Chimera. The “all atom refine” sub-module within the “refine” module in COOT was used for initial fitting of the model into the EM map, followed by manual rebuilding of the phosphopeptides. Multiple rounds of Phenix real space refinement combined with iterative model building yielded a model with 94.9% of the residues residing in the most favored region of the Ramachandran plot.

The dimeric co-ordinates from the cryo-EM structure of C5aR1pp- β arr1-Fab30 (PDB 8GO8) without the phosphopeptide was used as an initial model to dock into the CXCR4pp- β arr1-Fab30 EM map using Chimera. The docked model along with the coulombic map were imported into COOT and the model was subjected to “all atom refine” for fitting the atoms into the density. The phosphopeptide was manually built into the density to yield a complete model, which was subsequently used to refine the model against the EM map with Phenix real space refinement. The final refined model had 96.62% residues in the most favored regions and 3.38% in the allowed regions of the Ramachandran plot. For model building into the locally refined maps, coordinates of the individual full-length structures were used to dock into the corresponding maps followed by iterative rounds of manual adjustments in COOT and real space refinement in Phenix.

All the refined models were validated using “Comprehensive Validation (cryo-EM)” sub-module in Phenix. 3D reconstruction and model refinement statistics for both full and local-refined structures are provided as Table 1. Figures in the manuscript have been

prepared with Chimera⁶⁴ and ChimeraX⁶⁵ software. Domain rotation analysis was performed with PyMOL.⁷⁵ The interaction interface of β arr oligomers in the cryo-EM structures were identified using PDBSum.⁶⁸

NanoBiT assay for β arr2^{WT} and β arr2^{DM} recruitment

β arr2^{WT} and β arr2^{DM} recruitment downstream of V2R and C5aR1 in response to AVP and C5a, respectively, was measured using NanoBiT (Enzyme linked complementation-based assay) assay following the protocol described earlier.⁷⁶ Receptor constructs were tagged with SmBiT at the carboxyl-terminus, and β arr2 constructs were N-terminally tagged with LgBiT. Briefly, HEK-293 cells were transfected with indicated receptor constructs (3.5 μ g) and β arr2 (β arr2^{WT/DM}) constructs (3.5 μ g) using polyethylenimine (PEI) linear (Polysciences, Cat. no. 19850) at a ratio of 1:3 (DNA:PEI linear). After 16–18 h of transfection, cells were trypsinized, harvested, and resuspended in assay buffer (1X HBSS, 5 mM HEPES, pH 7.4, 0.01% BSA) containing 10 μ M coelenterazine (GoldBio, Cat. no. CZ05). Resuspended cells were seeded in a white flat bottom 96-well plate (100 μ l well⁻¹). After 2 h of incubation (90 min at 37°C and 30 min at room temperature), basal luminescence was recorded using a multimode plate reader (FLUOstar Omega, BMG Labtech). Later, cells were stimulated with varying doses of indicated ligands followed by measurement of luminescence signal for 20 cycles. For data analysis, ligand induced change in signals were taken and normalized with the lowest ligand dose luminescence value, and fold normalized data was plotted using nonlinear regression three-parameter sigmoidal concentration-response curve in GraphPad Prism v 9.5 software.

NanoBiT assay for β arr trafficking

Agonist-induced β arr2^{WT} and β arr2^{DM} endosomal trafficking downstream of the receptors mentioned above was studied using NanoBiT assay as described in the recruitment experiment. The only exception from the recruitment assay was that the receptor constructs were not tagged with SmBiT, but rather β arr2 (β arr2^{WT/DM}) and FYVE constructs N-terminally fused with SmBiT and LgBiT respectively were used for enzyme complementation. For each experiment, 3 μ g of indicated receptors, 2 μ g of SmBiT- β arr2^{WT/DM}, and 5 μ g of LgBiT-FYVE were used. Fold normalized change in signals were plotted using nonlinear regression three-parameter sigmoidal concentration-response curve in GraphPad Prism v 9.5 software.

NanoBiT assay for Ib30 reactivity

To assess Ib30 reactivity in response to an agonist for the mentioned receptors, NanoBiT assay was used following the same protocol as discussed in the β arr2^{WT} and β arr2^{DM} recruitment assay.⁷⁷ For enzyme complementation, N-terminally SmBiT fused β arr1 and N-terminally LgBiT fused Ib30 were used. For transfection, 3 μ g receptor except for M2R, CXCR7 (5 μ g), and CXCR3 (7 μ g), 2 μ g SmBiT- β arr1, and 5 μ g LgBiT-Ib30 were used. Transfected cells were stimulated with varying doses of respective ligands (mentioned in corresponding figures). Fold normalized change in luminescence were plotted using nonlinear regression three/four-parameter sigmoidal concentration-response curve in GraphPad Prism v 9.5 software.

Receptor surface expression

Receptor surface expression in various assays was measured using a previously described whole cell-based surface ELISA assay.⁷⁸ To study the surface expression of the receptor, cells transfected with a particular receptor were seeded into a 0.01% poly-D-Lysine pre-coated 24-well plate at a density of 2 \times 10⁵ cells well⁻¹. Post 24 h of seeding cells were washed once with ice-cold 1XTBS, fixed with 4% PFA (w/v in 1XTBS) on ice for 20 min, washed again three times with 1XTBS, and blocked with 1% BSA (prepared in 1XTBS) at room temperature for 1.5 h. Afterward, cells were incubated with anti-FLAG M2-HRP antibody at 1:5000 dilution (Sigma, Cat. no. A8592) for 1.5 h, which was followed by three washes in 1% BSA. Subsequently, incubated with TMB-ELISA substrate (Thermo Fisher Scientific, Cat. no. 34028) until a light blue color appeared. To quench the reaction, 100 μ l of the colored solution was transferred to another 96-well plate containing 100 μ l of 1M H₂SO₄, and the absorbance was measured at 450 nm. Afterward, the TMB substrate was removed, washed twice with 1XTBS, and incubated with 0.2% (w/v) Janus Green (Sigma; Cat. no. 201677) for 15 min at room temperature. Later, cells were washed with water to remove the excess stain, followed by the addition of 800 μ l of 0.5N HCl in each well. Thereupon, the colored solution was transferred to a 96-well plate for measuring the absorbance at 595 nm. The signal intensity was normalized by calculating the ratio of A450/A595 values followed by quantifying fold increase with respect to the A450/A595 nm value of negative control (mock transfection) and plotted using the GraphPad Prism v 9.5.

QUANTIFICATION AND STATISTICAL ANALYSIS

GraphPad Prism v9.5 was used to plot and analyze all the functional data presented in this manuscript, and all the relevant details such as number of replicates, data normalization, mean \pm sem, and statistical analyses are mentioned in the corresponding figure legends.