STRUCTURAL BIOLOGY

Structures of neurokinin 1 receptor in complex with G_q and G_s proteins reveal substance P binding mode and unique activation features

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The neurokinin 1 receptor (NK₁R) is involved in inflammation and pain transmission. This pathophysiologically important G protein–coupled receptor is predominantly activated by its cognate agonist substance P (SP) but also by the closely related neurokinins A and B. Here, we report cryo–electron microscopy structures of SP-bound NK₁R in complex with its primary downstream signal mediators, G_q and G_s . Our structures reveal how a polar network at the extracellular, solvent-exposed receptor surface shapes the orthosteric pocket and that NK₁R adopts a noncanonical active-state conformation with an interface for G protein binding, which is distinct from previously reported structures. Detailed comparisons with antagonist-bound NK₁R crystal structures reveal that insurmountable antagonists induce a distinct and long-lasting receptor conformation that sterically blocks SP binding. Together, our structures provide important structural insights into ligand and G protein promiscuity, the lack of basal signaling, and agonist- and antagonist-induced conformations in the neurokinin receptor family.

INTRODUCTION

The neurokinin receptor (NKR) family and its endogenous ligands are one of the most thoroughly studied neuropeptide signaling systems (1–4). Initially found in the early 1930s, to date, the human NKR family is known to consist of three members (NK₁R, NK₂R, and NK₃R), which belong to the superfamily of G proteincoupled receptors (GPCRs). NK₁R represents the most abundant and widely expressed member of the family, and this receptor has been found in neuronal, muscle, and endothelial cells of diverse tissues, ranging from the central and peripheral nervous system (5, 6) to the digestive (7) and respiratory (8) system, as well as on T cells in the immune synapse (9). Accordingly, NK₁R is implicated in major pathophysiological processes, such as inflammation (10), analgesia (11), depression (12), pruritus (13), and possibly asthma (14).

In the clinic, NK₁R is today targeted by the two marketed antagonists, aprepitant and netupitant, for the treatment of chemotherapyinduced nausea and vomiting (CINV) (15-17). Nonetheless, because of their widespread role, NK₁R ligands may find additional clinical uses, justifying a mechanistic study of this receptor. Recently, we reported crystal structures of the NK₁R bound to antagonists from different development stages, which allowed us to propose a molecular mechanistic model for the observed insurmountable antagonism underlying the long-lasting effect of the marketed drugs, which is not shown by other tight-binding tool compounds (18-21). Nevertheless, to verify this hypothesis, a structure of the agonist-bound active state would be needed.

The antagonist-bound structures also provided a mechanistic model for the lack of basal signaling of NK_1R : In contrast to other class A GPCRs where an allosteric sodium ion stabilizes the inactive

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conformation, the NK₁R-specific residue E78^{2.50} [number in superscript corresponds to Ballesteros-Weinstein numbering (22)] takes up a position in the receptor core, which stabilizes this state and, hence, prevents basal activity (23). However, in the absence of an agonist-bound structure, these studies could not explain the molecular determinants of receptor activation.

Activation of NK₁R is mediated predominantly through substance P (SP), an 11-amino acid-long peptide with the sequence RPKPQQFFGLM, comprising an amidated C terminus. However, two other neurokinins, A (NKA) and B (NKB), are also capable of binding to NK₁R, albeit with lower affinity compared to SP (24). NKA and NKB, in turn, are the main endogenous agonists of the two other NKR family members, NK₂R and NK₃R, respectively. In their C-terminal part, which mediates receptor activation, SP, NKA, and NKB share a high sequence homology with the three terminal amino acids being identical. Numerous pharmacological studies have located the determinants for ligand selectivity to the N-terminal portion of the peptide (25-28). On this basis, a message-address model has been proposed for the endogenous NKR agonists, dividing the ligand in two functionally distinct parts, the C-terminal region inducing receptor activation (message) and the N-terminal region providing receptor subtype selectivity (address) (29). Nonetheless, the precise molecular determinants of ligand binding and receptor activation have remained largely unknown because of the lack of structural information on the agonist-bound receptor in an active, G protein-bound state.

Last, the NK₁R has also been shown to couple to different heterotrimeric G protein subtypes, predominantly G_q and G_s , for intracellular signal transduction. Various previous studies suggested a coupling preference (2- to 20-fold difference in median effective concentration, EC₅₀) of the receptor to G_q over G_s (23, 30–32). Here, we elucidated the binding mode of neurokinin peptide ligands exemplified by SP and investigated the mechanism for the reported G protein–coupling preference. We used cryo–electron microscopy (cryo-EM) to determine the structures of SP-bound NK₁R complexed with G_q and G_s heterotrimers.

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RESULTS

Structure determination

For the preparation of stable, active-state complexes of SP-bound wild-type human NK₁R with G_q and G_s, we used the previously engineered G protein versions mini-G α_q and mini-G α_s (33, 34) as heterotrimers with G $\beta_1\gamma_2$. To permit binding of a complex-stabilizing single-chain antibody variable fragment (scFv16) distant from the GPCR/G protein interface (35), the N terminus of mini-G α_q and mini-G α_s was replaced with the N-terminal 18 amino acids of G α_{i1} . The resulting heterotrimeric G proteins are hereafter, for simplicity, referred to as G_q and G_s. Purified complexes of these G proteins with SP-bound NK₁R and scFv16 (fig. S1) were subjected to cryo-EM data acquisition. After iterative rounds of two-dimensional (2D) and 3D classification, the two complexes of NK₁R with G_q and G_s were resolved to 2.7 and 2.9 Å, respectively (Fig. 1, Table 1, and figs. S2 to S6).

In the structures of both complexes, the receptor region of highest resolution was observed in the transmembrane core, whereas the extracellular and intracellular transmembrane helix tips as well as their connecting flexible loops represented regions of lower local resolution (figs. S4 and S5). Intracellular loop 3 (ICL3) was not resolved in either of the complexes. In contrast to the previous crystal structures of antagonist-bound NK₁R, both SP-bound NK₁R complexes display a resolved receptor N terminus, starting at P22^{N-term}, as well as a resolved extracellular loop 3 (ECL3), due to direct contact of these regions with the ligand (*19*). ECL2 is packed against ECL1 and adopts the previously observed extended β -hairpin, which is tethered to the extracellular tip of transmembrane helix III through a disulfide between C180^{ECL2} and C105^{3.25}. ECL3 adopts short

 α -helical fold between D276^{ECL3} and L279^{ECL3}. The receptor N terminus, together with all ECLs, establishes the structural framework for extracellular SP binding. The C terminus of the ligand is located within the orthosteric pocket, where it binds deep into the receptor core in an elongated conformation.

On the intracellular portion of NK₁R, we detected an additional cholesterol density at the outer interface of helices II and IV in both the G_q -bound and the G_s -bound complexes (fig. S7). The observed binding area overlaps with a previously described cholesterol consensus motif present in the sequence of 21% of all human class A GPCRs (*36*). The presence of a cholesterol molecule in structures of the active NK₁R state may explain the observed cholesterol dependence of NK₁R activation (*37*). Furthermore, a functionally important cholesterol molecule was observed at a similar binding interface in two recently reported active-state complexes of the cholecystokinine A receptor (CCK₁R) with G_q and G_s (*38*).

Despite the high resolutions obtained and the monoolein matrix being supplemented with 10% (w/w) cholesterol, cholesterol has not been observed in any of the antagonist-bound NK₁R crystal structures (18–20). The presence of this cholesterol molecule may thus be specific to the active state of the receptor, similar to state-dependent lipid interactions of the adenosine A_{2A} receptor ($A_{2A}R$) (39, 40).

Binding of SP to NK₁R

Over the past decades, the binding mode of SP on NK₁R has been investigated in numerous mutational and pharmacological studies (41-46). Despite these extensive efforts, the precise molecular details of endogenous agonist interactions have remained unknown. The



Fig. 1. Cryo-EM structures of NK₁R:SP:G_q:scFv16 and NK₁R:SP:G_s:scFv16. (A) Cryo-EM map of the NK₁R:SP:G_q:scFv16 complex resolved to 2.71 Å (FSC 0.143) as viewed from the side and from the top. (B) Cryo-EM map of the NK₁R:SP:G_s:scFv16 complex resolved to 2.87 Å (FSC 0.143) as viewed from the side and from the top. (C) Molecular model of the NK₁R:SP:G_q:scFv16 complex with SP and cholesterol shown in stick representation. (D) Molecular model of the NK₁R:SP:G_s:scFv16 complex with SP and cholesterol shown in stick representation.

	NK1R:SP:Gq:scFv16 (PDB ID: 7P00)	NK1R:SP:Gs:scFv16 (PDB ID: 7P02)
Data collection		
Microscope	Titan Krios G3i	Titan Krios G3i
Detector	Gatan K3	Gatan K3
Energy filter slit width (eV)	20	20
Magnification	130,000	130,000
Voltage (kV)	300	300
Electron exposure (e ⁻ /Å ²)	62.51	63.51
Defocus range (µm)	0.8–2.4	0.8–2.4
Pixel size (Å)	0.65	0.65
Symmetry imposed	C1	C1
Number of micrographs	5291	6341
Initial particle images (n)	3,511,990	4,227,825
Final particle images (n)	558,058	395,052
Map resolution (Å)	2.71	2.87
FSC threshold	0.143	0.143
Refinement		
Number of atoms	.	
All	9110	9061
Protein	8987	8957
Ligand	95	76
Cholesterol	28	28
Model validation		
CC map versus model (%)	79	78
RMSD		
Bond lengths (Å)	0.012	0.012
Bond angles (°)	1.679	1.510
Ramachandran statistics		
Favored regions (%)	96.63	95.36
Allowed regions (%)	3.37	4.64
Outliers (%)	0.00	0.00
Rotamer outliers (%)	0.81	1.12
Cβ deviations (%)	0.74	0.00
MolProbity overall score	4.75	6.05

Table 1. Cryo-EM processing and modeling statistics.

obtained resolution now allows us to describe the precise interaction mode of SP with NK_1R in the fully active receptor state. To also characterize the observed interactions in a quantitative manner, we performed whole-cell ligand binding assays on wild-type human NK_1R and single mutants of structurally identified key interaction residues using fluorescently labeled SP (table S1). Overall, the ligand binding pose of the endogenous agonist SP is highly similar in the G_s and the G_q complexes of NK₁R (fig. S8). The interactions of SP with NK₁R can broadly be divided into two groups: (i) those of the C terminus (residues F_7 - M_{11} , subscript indicates SP residue number), which are highly conserved between the endogenous agonists SP, NKA, and NKB, with the transmembrane helical core of the receptor, and (ii) those of the nonconserved N terminus (residues R_1 - Q_6) with the extracellular surface of the receptor (Fig. 2, A and B).

Interactions of the C terminus of SP

The interactions of the conserved SP C terminus with NK₁R are well resolved in both cryo-EM density maps of the G_q and the G_s complexes, allowing to model all side chains of the peptide (Fig. 2B). In contrast, at its N terminus, density can only be assigned to the peptide backbone. Furthermore, in the G_s -bound structure of NK₁R, we could not resolve EM densities for the N-terminal residues R_1 - P_2 and have thus modeled only K₃- M_{11} of SP (Fig. 2B). Therefore, because of the higher resolution obtained and the completely resolved ligand, we refer to the G_q -bound structure for the description of the interactions of SP with NK₁R.

The pocket to which the C-terminal part of SP binds is lined by transmembrane helices II, III, VI, and VII of NK₁R (Fig. 2A). Residues F_7 - M_{11} of SP are oriented along a hydrophobic interface established by helices VI and VII in a well-defined conformation, as evidenced by the clear EM density for all ligand side chains in this region (Fig. 2B).

Within the receptor core, the interactions of the agonist can again be divided into two main groups: (i) a hydrogen-bonding network centered at the C-terminal amide of the agonist, spanning helices II, III, and VII (Fig. 2D), and (ii) a hydrophobic contact surface lined mainly by residues from helices VI and VII (Fig. 2E).

C-terminal amidation is a central feature of tachykinins occurring in all vertebrates and invertebrates (2), and its removal has been demonstrated to be detrimental to agonist activity of SP (47, 48). In the structures of SP-bound NK₁R, we now find the C-terminal amide of the agonist positioned in close hydrogen-bonding distance (N-O distances of 2.7 and 2.9 Å) to N85^{2.57}. Accordingly, we observe a 10-fold decrease in agonist binding affinity upon changing the intermolecular distance of the interacting amides (mutation N85^{2.57}Q) and an 8- and 10-fold decrease upon either removal of the side chain or changing its electrostatics (mutations N85^{2.57}A and N85^{2.57}D, respectively) (Fig. 2D), underlining the importance of this interaction.

The binding pose of the C-terminal portion of SP is further stabilized through hydrogen bonding of the backbone oxygen and nitrogen of the agonist residue L_{10} with N89^{2.61} and Y287^{7.35}, respectively. The importance of these H bonds is underlined by a 36- to 111-fold decrease in SP binding affinity upon mutation of N89^{2.61} to A, D, or Q, and a 31-fold decrease upon removal of the side chain in Y287^{7.35}A (Fig. 2D). In addition to these hydrophilic contacts, we find His108^{3.28} in hydrogen-bonding distance to both N85^{2.57} and N89^{2.61}, potentially further stabilizing the rotamer orientation of these two residues that bind SP.

On the opposite side of this hydrogen-bonding network at the bottom of the orthosteric pocket, the side chain of M_{11} is found in a hydrophobic subpocket. This subpocket is lined by residues from helices III to VII, including I113^{3,33}, F117^{3,37}, Q165^{4,60}, H197^{5,39}, F264^{6,51}, F268^{6,55}, and M291^{7,39}, and removal of the hydrophobic



Fig. 2. Interactions of the endogenous agonist SP with NK₁R. (A) SP (stick representation, carbons depicted in pink) establishes a host of interactions with the receptor transmembrane core and the extracellular receptor surface (receptor-interacting residues depicted as sticks with green carbons). (B) Processed cryo-EM maps of SP in the NK₁R:G_q complex (colored in pink) and the NK₁R:G_s complex (colored in yellow) contoured at 4.5 σ. Bottom: The amino acid sequences (numbers on top correspond to position in SP) of the three neurokinins SP, NKA, and NKB are indicated by residues of the conserved ligand C terminus highlighted in red, and the C-terminal amide is shown. Negatively and positively charged residues are highlighted in orange and blue, respectively. (**C**) Schematic representation of the polar network shaping the extracellular receptor surface for SP binding, indicating the hydrogen bonds detected in the structure. Bottom: Alignment of residues involved in specific side-chain interactions of the polar network are shown, comparing NK₁R with NK₂R and NK₃R. Framework residues of ECL3 only interact via backbone hydrogen bonding and are therefore not included in the alignment. (**D** to **F**) Close-up on the interactions of the amidated SP C terminus at the bottom of the orthosteric binding pocket (D), the interactions of the neurokinin-conserved C-terminal portion of SP with the hydrophobic interaction surface of helices VI and VII of NK₁R (E), and the polar network at the extracellular receptor surface (F). Bottom: Agonist affinity profiles of selected mutants compared to wild-type NK₁R are shown. pK_D values were derived from whole-cell saturation binding experiments (table S1). Data are shown as means ± SEM from three to six independent experiments performed in duplicate.

moieties from residues F117^{3.37}, Q165^{4.60}, F264^{6.51}, F268^{6.55}, and M291^{7.39} by mutation to an alanine leads to a 6- to 28-fold reduction in SP binding affinity (Fig. 2D).

From the bottom of the orthosteric ligand binding pocket toward the extracellular opening, the peptide agonist packs with residues F_7 and G_9 against the hydrophobic surface of helices VI, VII, and ECL3, thereby enforcing an elongated conformation of the SP peptide backbone (Fig. 2E). Thus, F_8 and L_{10} of SP are facing away from helices VI and VII and pack against ECL1 and ECL2 of NK₁R. At the extracellular boundary of the orthosteric ligand binding pocket. SP is held in position through hydrophobic interactions with V179^{ECL3} and Y278^{ECL3} as well as with Y92^{2.64}, A93^{2.65}, I283^{7.31}, and Q284^{7.32} from the helical tips of helices II and VII. In line with the hydrophobic nature of these interactions, SP binding affinity is reduced by a factor of 2 to 16 upon removing interaction surface by mutation of these residues to alanine, while removal of the hydroxy headgroup of Y92^{2.64} by mutation Y92^{2.64}F leaves SP binding affinity unchanged (Fig. 2E).

 F_8 of SP, being the first amino acid in C- to N-terminal direction, which is not conserved between SP, NKA, and NKB, is not involved

in a direct interaction with the transmembrane domain of NK₁R but rather facing toward the open extracellular space. This orientation explains the relative ease of replacing this residue of the agonist with different amino acids or labeling it with radioactive tracers, as was done in earlier pharmacological studies (49, 50).

Interactions of the N terminus of SP

In the N-terminal portion (residues R_1 - Q_6), the binding of SP to NK₁R is governed by several interactions of the receptor with the peptide backbone of the ligand (Fig. 2F and fig. S9). The residue side chains in this region of the agonist are not resolved (except for Q_6 , which packs against ECL3), likely due to higher flexibility in the receptor:ligand complex (Fig. 2B). Specifically, the backbone carbonyl oxygen of Y278^{ECL3} engages in a hydrogen bond to the backbone nitrogen of F₇ of the ligand. Furthermore, R177^{ECL2}, located at the tip of the β -turn in ECL2, establishes two hydrogen bonds to the backbone carbonyls of Q_6 and P_4 of SP. R177^{ECL2} is also within hydrogen-bonding distance of N96^{2.68} at the start of ECL1 and N23^{N-term}. Through its interaction with N23^{N-term}, R177^{ECL2} helps to position the receptor N terminus from the tip of helix I toward

the peptide agonist, enabling the formation of two additional hydrogen bonds of Q24^{N-term} with the backbone carbonyl of Q₅ of SP and with the backbone carbonyl of L279^{ECL3}. Hence, R177^{ECL2} acts as a central residue, which engages a polar network that links all ECLs with the receptor N terminus, thereby shaping the entire extracellular surface for ligand binding (Fig. 2, C and F, and fig. S9). The importance of this polar network for SP binding to NK₁R is exemplified by the 7- to 46-fold loss in ligand binding affinity upon mutation of either of the involved residues to alanine, with mutation of the central R177^{ECL2} solely emerge from its positive charge, we furthermore substituted R177^{ECL2} with lysine, resulting in a 60-fold reduced binding affinity.

The preorientation of the NK₁R N terminus through this polar network enables direct interaction of F25^{N-term} with F₇ of the ligand (Fig. 2E), a residue conserved in the neurokinin peptide family (Fig. 2B), thereby introducing a pronounced kink in the receptor N terminus (Fig. 2F). The importance of this specific interaction for SP binding is underlined by the 16-fold reduced binding affinity observed for the mutation F25^{N-term}A, removing the hydrophobic moiety, while the mutation F25^{N-term}Y retaining the aromatic properties resulted only in a subtle change in binding affinity (Fig. 2E). Together, the participation of N23^{N-term} and Q24^{N-term} in the observed polar network at the extracellular receptor surface and the direct interaction between F25^{N-term} and the conserved F₇ of the peptide agonist provide a structural basis for the previously described importance of the receptor N terminus for ligand binding (42).

Implications on ligand subtype selectivity

As early as the late 1970s, a message-address hypothesis was proposed for several neuropeptide receptors (29, 51, 52), which broadly divides the endogenous neurokinin peptide agonists in two functionally distinct domains: a "message" domain responsible for receptor activation and an "address" domain establishing NKR subtype selectivity. The observation of numerous well-resolved interactions of the hydrophobic C terminus of SP with the core of the receptor and the observed effects of mutations are consistent with a major contribution of the neurokinin family–conserved peptide C terminus to the overall binding energy.

Unexpectedly, no defined short-range interactions between side chains of the ligand and NK1R are observed for the charged N terminus of SP, which is divergent in both length and residue composition between the different neurokinin peptides. Instead, it appears that binding interactions are predominantly established through interactions of NK1R with the peptide backbone of SP (Fig. 2F and fig. S9). This finding implies that there are several more subtle components contributing to the preference of SP binding to NK1R over NKA and NKB. The N terminus of SP has a double-positive charge in this region, while NKA is zwitterionic and NKB carries a doublenegative charge (Fig. 2B). These differences in charge distribution of the neurokinin peptides favor interactions between the positively charged N terminus of SP and the negatively charged extracellular receptor surface formed by the solvent-exposed residues $E172^{ECL2}$, $E183^{ECL2}_{ECL2}$, $E186^{ECL2}$, and $D276^{ECL3}$ (fig. S10). Notably, mutation of E172^{ECL2}, which is found closest to R_1 - K_3 of SP, to a positively charged lysine, as found in NK₃R, increases affinity of NKB, presumably because of the interaction with the NKB N-terminal D₁ (corresponding to P_2 for SP) (41, 53). Furthermore, the presence of two prolines in SP may alter the preferred conformers of the flexible N terminus,

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compared to NKA and NKB. Therefore, the subtype specificity does not seem to be achieved through specific short-range side-chain interactions but rather through a mechanism of electrostatic and conformational selection of the N terminus.

Last, the polar network established by residue $R177^{ECL2}$ points toward an additional layer of complexity to subtype-specific ligand binding in NKRs: The receptor residues involved in establishing this polar network, which structures the entire extracellular receptor surface for ligand binding, are only partially conserved between NK₁R, NK₂R, and NK₃R (Fig. 2C). Because of the geometry of the observed interactions, the exact identity of all of these residues is pivotal to this network, and hence, it can only be established in this form by NK₁R, but not by its close relatives NK₂R and NK₃R.

Activation of NK₁R

The comparison of the active, G protein–bound NK₁R:SP complexes with the inactive-state structure of NK₁R in complex with the antagonist CP-99,994 (19) enables us to identify the molecular changes underlying receptor activation. In the canonical activation mechanism of class A receptors, binding of the agonist to the orthosteric pocket of the receptor induces a cascade of conformational changes in several conserved motifs and culminates in a pronounced outward swing of helix VI to accommodate the C-terminal C α 5 helix of the G protein (54). Unexpectedly, NK₁R adopts most but not all of the canonical rearrangements observed for active-state class A GPCRs, and we use the closely related neuropeptide orexin-B receptor (OX₂R) to exemplify the specific differences (Fig. 3) (55).

At the extracellular receptor portion, SP binding causes a contraction of the orthosteric pocket through an inward movement of helices VI and VII, and ECL3 toward the central axis of the receptor (Fig. 3A). These rearrangements reposition F264^{6.51} at the bottom of the orthosteric pocket to trigger the transmission switch W261^{6.48}, leading to the canonical outward move of F257^{6.44} of the PIF motif together with L209^{5.51} and an inward repositioning of I120^{3.40} (Fig. 3B). On the same horizontal plane through the receptor, an extensive water-mediated hydrogen bonding network, clustered around a central sodium ion that links helices II, III, VI, and VII, was described to stabilize class A GPCRs in the inactive state, involving interactions with the highly (98% of class A GPCRs) conserved $D^{2.50}$ (56). Upon receptor activation, this sodium ion is expelled, and the conserved $D^{2.50}$ is repositioned to establish a direct hydrogen bond to N^{7.49} of the NPxxY motif in helix VII (57). NK₁R, however, has a glutamate $E78^{2.50}$ at the position of the conserved $D^{2.50}$, a sequence variation that has been linked to the unusual lack of constitutive signaling of NK₁R in functional studies (23). In the inactive state of the antagonist CP-99,994-bound receptor, the longer side chain of E78^{2.50} enables the establishment of a direct hydrogen bond to N301^{7.49} without the need of a bridging sodium ion. In the activestate structure, we now find that the entire helix VII is pushed downward and toward the intracellular space owing to the longer side chain of E78^{2.50} (Fig. 3C).

This NK₁R-specific translocation restricts the canonical inward shift of helix VII at the intracellular receptor surface upon receptor activation and prevents Y305^{7.53} of the conserved NPxxY motif from adopting the expected inward-facing orientation toward R130^{3.50} of the DRY motif (Fig. 3, C and D) (58). As a consequence, the outward swing of helix VI is less pronounced in NK₁R, and M249^{6.36} is rotated further into the receptor core instead of adopting an outward-facing orientation, while the ratchet-like rearrangement of M250^{6.37}



Fig. 3. Molecular characteristics of NK₁R activation by SP. (A) Superposition of the active-state NK₁R:SP:G_q (green) with the inactive-state crystal structure of NK₁R in complex with the antagonist CP-99,994 (PDB ID: 6HLL) (blue) and the active-state OX₂R:G_q complex (PDB ID: 7L1U) (brown). Relative changes from inactive to active NK₁R are indicated by black arrows. Left: extracellular view, right: intracellular view, middle: side view with approximate position of conserved motifs involved in receptor activation indicated, and sections of the close-up views in (B) to (D) highlighted by dotted boxes. (**B** to **D**) Close-up views on conserved motifs involved in receptor activation. Repositioning of side chains from the inactive to the active state are indicated by arrows.

enables the canonical packing against Y243^{5.58} (Fig. 3D). In summary, NK₁R adopts an active conformation with smaller rearrangements of the intracellular tips of helices VI and VII, thus presenting a distinctly shaped intracellular binding surface for the C-terminal C α 5 helix of the G α protein (Fig. 4, B and C).

Coupling of NK₁R to different G proteins

Promiscuity for different G proteins has been described for many GPCRs including NK₁R, which mainly signals through the G_{s} - and G_{q} -coupled pathways (59). Because of the lack of structural information on the signaling-active complexes of NK₁R with either of the two G proteins, the molecular details of receptor–G protein interaction and the differences between G_{s} - and G_{q} -bound NK₁R have remained unknown. Our structures of SP-bound NK₁R in complex with both G_{s} and G_{q} now reveal detailed information on these interactions.

Overall, in both resolved complexes, G_s and G_q adopt a highly similar conformation relative to NK₁R, where the C-terminal C α 5 of G_q is shifted 0.9 Å deeper into the intracellular crevice created by the activated receptor (Fig. 4A). Unexpectedly, in the NK₁R:G_s complex, we observe a relative conformation of the two interacting proteins toward each other that is highly distinct from those observed in other class A GPCR:G_s complex structures (Fig. 4B). In the NK₁R complex, the tip of C α 5 of G_s is oriented further away from helices V and VI and more toward helix II, when compared to previously reported G_s complexes: melanocortin 4 receptor (MC₄R) (60), A₂AR

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(61), β_1 -adrenergic receptor (β_1AR), and β_2 -adrenergic receptor (β_2AR) (62, 63). The distinct orientation of C α 5 of G_s in NK₁R can be attributed to the less-pronounced outward swing of helix VI that is stabilized through the reduced inward movement of helix VII and the movement of helix V toward helix VI. Thus, the NK₁R:G_s intracellular helical movements more closely resemble those generally observed for G_q complexes, including the NK₁R:G_q structure we determined (Fig. 4C). These observations are consistent with the ability of the receptor to couple to both G_q and G_s.

To date, only few structures of class A GPCRs in complex with G_q have been determined, including those of the CCK₁R (38), OX₂R (55), and 5-hydroxytryptamine receptor 2A (HTR_{2A}) (64). Common to all these complexes is a less-pronounced outward swing of the intracellular tip of helix VI, compared to G_s complexes, along with a C α 5 positioning closer to helix II. Although overall very similar, C α 5 in NK1R:G_q is bound more deeply in the receptor cavity than in the other G_q -bound complex structures. This deeper binding mode is enabled by the NK₁R-specific reduced inward shift of helix VII and the shift of helices V and VI toward helix VII (Fig. 4C). In summary, the NK₁R-specific intracellular positioning of transmembrane helices V, VI, and VII, resulting from the unexpected inactive-state orientation of Y305^{7.53} in the conserved NPxxY motif, translates into distinct C α 5 positions in both active-state complexes of NK₁R:G_g and NK₁R:G_g.

G protein interactions in GPCRs are mainly governed by contacts established through the C-terminal α 5 helix of the G α protein

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Fig. 4. Interactions of NK₁R with Ga_s and Ga_q. (**A**) Structural superposition of NK₁R:G_s and NK₁R:G_q with a close-up of the C-terminal tip of Ca⁵, highlighting the 0.9 Å deeper binding of Ga_q compared to Ga_s. Gβ₁γ₂ and scFv16 are omitted for clarity. (**B**) Structural superposition of NK₁R:G_s with class A GPCR:G_s complexes viewed from the cytoplasmic side, highlighting the distinct Ca⁵ binding angle in NK₁R and differing positions of helices V, VI, and VII. A2A:G_s (PDB ID: 6GDG), MC4R:G_s (PDB ID: 7AUE), β₂AR:G_s (PDB ID: 7BZ2), and β₁AR:G_s (PDB ID: 7JJO). The arrows indicate the relative helical position from the group of class A GPCR:G_s structures toward NK₁R:G_s. (**C**) Structural superposition of NK₁R:G_q with class A GPCR:G_q shown from the cytoplasmic side highlighting deeper insertion position of the Ca⁵ the NK₁R cavity and differing positions of helices V, VI, and VII. OX₂R:G_q (PDB ID: 7L1U), HTR_{2A}:G_q (PDB ID: 6WHA), and CCK₁R:G_q (PDB ID: 7MBY). The arrows indicate the relative helical position from the ca⁵ the IX₁R:G_s and Ga_q subunits with H-bonded and hydrophobic contacts indicated by a dashed and solid line, respectively. Residues differing between Ca⁵ of Ga_s and Ga_q are indicated by a lighter color. G protein residues are labeled according to CGN numbering beneath each residue. (**E** and **F**) Interactions of Ca⁵ of Ga_s (**E**) and Ga_q (**F**) with NK₁R. Hydrogen bonds are indicated by black, dotted lines.

(Fig. 4, D to F, and tables S2 and S3). Owing to the similar overall conformation of NK₁R:G_s and NK₁R:G_q, the interactions at the contact surface of receptor and G protein are also closely related, whereas the differences are due to the differing residues between G_s and G_q within the respective C α 5 (Fig. 4D). Furthermore, G_q engages in a larger number of hydrogen-bonding interactions (seven in G_q compared to six in G_s) with NK₁R due to the deeper binding of C α 5 in the crevice formed by the receptor.

In NK₁R:G_s, G protein coupling involves contacts with receptor transmembrane helices II, III, V, VI, and VII; helix VIII; and ICL2, with most contacts in G_s being established by residues L376^{H5.25}, L377^{H5.26}, and Y374^{H5.23} [common G α numbering for G proteins indicated in superscript (65)]. Enabled by the distinct orientation of C α 5, NK₁R establishes an additional hydrogen-bonding contact between T67^{2.39} at the intracellular tip of transmembrane helix II and Q373^{H5.22} of the G protein, which had not been observed in previous G_s complexes (Fig. 4E). A similar interaction, also not observed in previous GPCR complexes with G_q, is established between T67^{2.39} and E373^{H5.22} of G_q (Fig. 4F). In this region of the receptor, we furthermore find two hydrogen bonds of N375^{H5.24} with N68^{2.40} and N309^{8.47} and, accordingly, a central role of N375^{H5.24} in G_q binding, as demonstrated for HTR_{2A} (64). Together, the deeper binding of C α 5 of G_q in NK₁R, compared to G_s, and the larger number of interactions may provide a structural rationale for the reported preference of NK₁R signaling via G_q over G_s (30).

SP binding is incompatible with the insurmountable antagonist-bound NK₁R conformation

The clinically used NK_1R antagonists aprepitant and netupitant behave as pseudo-irreversible antagonists, establishing long-lasting inhibition of receptor activation by SP despite their competitive binding to the orthosteric pocket (fig. S11) (21). This so-called insurmountable antagonism and the markedly prolonged in vivo efficacy of these drugs could only in part be explained by slower dissociation kinetics alone.

In our previous structural studies of NK1R, we found a distinct conformation of NK1R when in complex with aprepitant and netupitant, compared to the tight-binding, but surmountable progenitor antagonist CP-99,994 (19), and this was backed by clear evidence in the electron density. On the basis of this structural evidence, we were able to pinpoint a hydrogen-bonding network, cross-linking the extracellular tips of transmembrane helices V and VI as a molecular determinant of insurmountable antagonism. Furthermore, in the insurmountable antagonist-bound structures, we observed a cis configuration of the peptide bond preceding P175^{ECL2} at the tip of the β-hairpin loop of ECL2, which was absent in the CP-99,994bound state of the surmountable antagonist. Comparison of the active-state NK1R:SP structures with the inactive-state crystal structures of NK1R:aprepitant and NK1R:CP-99,994 now provides evidence for the insurmountable antagonist-induced state to directly interfere with SP binding.

The active-state NK₁R adopts a conformation more closely related to the surmountable antagonist complex with CP-99,994 with a root mean square deviation (RMSD) over all atoms of 1.65 Å and 1.77 Å, compared to NK₁R:CP-99,994 and NK₁R:aprepitant (insurmountable), respectively. Looking more closely at the extracellular portion of the receptor, the most dominant differences between the active state and both of the antagonist-bound structures are rearrangements of ECL2 together with ECL1 and the inward shift of the tip of helices VI and VII due to direct contacts of the agonist SP with residues from these helices (Fig. 5, A and B). However, the rearrangements of ECL2 are different for the two types of antagonists.

In the surmountable antagonist CP-99,994–bound NK₁R, ECL2 adopts a conformation similar to what we now revealed for the active state receptor yet with ECL2 positioned farther away from the central axis of the receptor (Fig. 5A). In contrast, in the insurmountable antagonist-bound NK₁R structures, the *cis* configuration of the peptide bond preceding P175^{ECL2} causes a twist of the loop toward the central axis of the receptor, where the agonist is located in the SP-bound structures (Fig. 5B). Thus, in addition to an inward



Fig. 5. The insurmountable antagonist-bound NK₁R state prohibits SP binding. (A) Superposition of agonist-bound NK₁R:SP:G_q (green) with the surmountable antagonist CP-99,994–bound NK₁R (PDB ID: 6HLL) (blue) reveals a slightly outward-moved but similar ECL2 conformation for the antagonist conformation. Note the small movement of R177 ^{ECL2}. **(B)** Superposition of agonist-bound NK₁R:SP:G_q (with the insurmountable antagonist aprepitant-bound NK₁R (PDB ID: 6HLO) reveals an inward shifted and twisted ECL2 in the insurmountable conformation that clashes at M174^{ECL2} with P₄ of the endogenous agonist and also poses R177^{ECL2} away from its central position in the polar network at the extracellular receptor surface. **(C)** Superposition of agonist-bound NK₁R:SP:G_q, CP-99,994- and aprepitant-bound NK₁R showing the hydrogen bonds (indicated as orange, dotted lines) that lock helices V and VI in the insurmountable conformation. The rotamer orientation of H197^{5.39} in the insurmountable conformation of a new hydrogen bond between H197^{5.39} with the backbone carbonyl oxygen of F268^{6.55} (indicated as green, dotted line) in the active state.

shift of the entire ECL2, the *cis* configuration of P175^{ECL2} positions M174^{ECL2} in an orientation where its side chain clashes with P_4 of SP and thus directly interferes with agonist binding.

Potentially even more important, the *cis* configuration of P175^{ECL2} orients R177^{ECL2} away from the ligand-binding interface, thereby preventing this residue from establishing its central role in the hydrophilic network connecting ECL1, ECL2, and ECL3 and the receptor N terminus for optimal agonist binding (Fig. 5B). The direct clash of M174^{ECL2} with SP and the prevention of R177^{ECL2} to engage in shaping the entire extracellular receptor surface necessary for agonist binding provide an additional structural rationale for the long-lasting effects of insurmountable antagonists on NK₁R.

Furthermore, in the active-state structures, SP establishes contacts to F264^{6.51} and F268^{6.55} of transmembrane helix VI. The associated inward shift of helix VI enables the formation of a hydrogen bond between the backbone carbonyl oxygen of F268^{6.55} and H197^{5.39}, thus further stabilizing the active conformation (Fig. 5C). However, H197^{5.39} is the central residue of a hydrogen-bonding network crosslinking helices V and VI in the insurmountable antagonist-bound NK₁R crystal structures (19). In the structural superposition, we now find that the orientation of H197^{5.39} in the insurmountable conformation interferes with the inward repositioning of helix VI in the active conformation (C-C distance of H197^{5.39} and F268^{6.55} backbone of 2 Å) (Fig. 5C). The formation of the hydrogen-bonding network between Y272^{6.59}, H197^{5.39}, and T205^{5.43} in the insurmountable antagonist-bound conformation may thus directly prevent the receptor from adopting an active conformation by sterically preventing the necessary translocation of helix VI. In summary, SP binding to NK1R is precluded in the insurmountable antagonist-bound receptor conformation by direct steric incompatibility of ECL2 and helix VI with agonist binding as well as by preventing the formation of a critical polar network established by R177^{ECL2} of ECL2.

DISCUSSION

The neurokinin family has served as a model system in neuropeptide research since its discovery more than half a century ago, and many of the breakthroughs in this field have been associated with SP, the main endogenous agonist of NK₁R (*1*, *2*). Already early studies on NK₁R have been hampered by the promiscuity of NK₁R toward ligand interaction with NKA and NKB, the cognate agonists of NK₂R and NK₃R, and the coupling of NK₁R to different G proteins for intracellular signal transmission, and these promiscuities have therefore subsequently received much attention (*28, 30*).

Furthermore, NK₁R has been the subject of intense high-throughput screening and medicinal chemistry campaigns over the course of several decades, resulting in marketed drugs against CINV (66). While we recently found the molecular determinants for the efficiency of the marketed drugs aprepitant and netupitant in inactive-state NK₁R crystal structures by comparing them with a precursor compound (*19*), we could not fully describe the mechanism of preventing SP binding in the absence of structural information of the agonist-bound active state of the receptor. Our active-state complexes have allowed us now to advance the understanding of this receptor system in several major aspects: (i) the unexpected mechanism of agonist preference for SP, (ii) the unusual geometry of interaction with G_s and the ensuing G_q preference, (iii) the mechanism of eliminating basal activity, and (iv) the insurmountable antagonism of some clinically used drugs.

Our structures of SP-bound NK₁R in complex with G_q and G_s reveal the interactions of the endogenous agonist with its receptor in great detail. While the conserved amidated C terminus binds deep into the pocket in a well-defined orientation to trigger activation, the N terminus engages a polar network, shaping the entire extracellular receptor surface for ligand binding. In the polar N terminus of the ligand, which highly deviates between neurokinin family endogenous agonists SP, NKA, and NKB in both length and residue composition, stable binding is established through defined direct interactions of NK1R with the peptide backbone of SP but not with any of its side chains. Thus, subtype specificity does not seem to be achieved through stable short-range side-chain interactions but electrostatics and conformational fine-tuning of the N terminus. Furthermore, the residues of the extracellular polar network, which establishes the structural framework for binding of the agonist N terminus in NK₁R, are not conserved among the three NKR subtypes, implying a distinct ligand interaction mode of NK₂R and NK₃R. Therefore, for a detailed understanding of the determinants for neurokinin family subtype specificity, further studies including NK₂R, NK₃R, and their endogenous ligands NKA and NKB are needed.

Only very recently have structures of the same GPCR in complex with several different G proteins become available (*61*, *67*). Similar to CCK₁R, we cannot observe notable differences in ligand binding depending on the identity of the coupled G protein. In contrast to all other G_s-bound structures, including CCK₁R, NK₁R adopts a distinct C α 5 orientation resembling more the conformation that is generally observed in G_q complexes. We find that in the NK1R:G_q complex, C α 5 binds deeper into the intracellular receptor crevice than in any of the previously reported G_q complexes, which is consistent with the preference of G_q over G_s.

Recently, our inactive-state NK_1R crystal structures in complex with antagonists of different clinical development stages have allowed us to provide a structural rationale for two peculiarities of NK_1R : the reported lack of basal signaling in NK_1R and the molecular determinants of insurmountable antagonism (23, 50). The determined active-state structures allowed us to now extend and refine our previous hypotheses.

First, the unusual D78^{2.50}E substitution in the otherwise highly conserved allosteric sodium ion binding site enables the establishment of a direct hydrogen bond to N301^{7,49} in the inactive state. This prevents the canonical rearrangements in the conserved NPxxY motif upon receptor activation, thereby restricting the intracellular outward movement of helix VI, all in all preventing fluctuations, which would allow basal signaling.

Second, we find that the insurmountable antagonist-bound NK₁R conformation directly interferes with SP binding through steric clashes of the agonist with ECL2 and the prevention of the inward movement of the extracellular tip of helix VI necessary for receptor activation. A key component is the prevention of the formation of a critical polar network established by R177^{ECL2} of ECL2. The fact that this conformation of ECL2 contains a *cis* proline and results in a series of interdependent conformational changes, as described above, makes it plausible that it is not "surmountable" by adding agonist. Quite in contrast, in the surmountable antagonist complex structure, the ECL2 is merely positioned further away from the central axis, but otherwise found in a conformation similar to the one in the active-state receptor. In summary, our structures provide important insights into agonist preferences, antagonist mechanisms, lack of basal activity, and G protein promiscuity in the NKR family.

METHODS

Constructs

The sequences of scFv16 (*35*) and of wild-type human NK₁R (C-terminally truncated after residue 335) were cloned into a modified pFL vector (MultiBac system, Geneva Biotech), each resulting in an expression construct with a melittin signal sequence, followed by a FLAG-tag, a His₁₀-tag, and a human rhinovirus 3C protease cleavage site N-terminal to the gene of interest. To generate the G α_s and G α_q subunits, the N-terminal 20 amino acids of the engineered mini-G_{s399} and mini-G_{sq70} were replaced by the 18 N-terminal amino acids of G_{i1} (*34*, *35*). Each engineered G α protein sequence was, together with G β_1 including an N-terminal noncleavable His₁₀-tag and G γ_2 (not containing any tags), cloned into one pFL vector, each gene under the control of its own polyhedrin promoter.

Expression in insect cells

Recombinant baculovirus was generated using the MultiBac expression system. Virus production and protein expression were performed as previously described (19). The expression cassettes were integrated into the DH10EMBacY baculovirus genome, and the resulting bacmids were transfected into Sf9 cells in six-well tissue culture plates (2 ml, density of 4×10^5 cells/ml) using 8 µl of Cellfectin II Reagent (Thermo Fisher Scientific) and Sf-900 II serum-free medium (Thermo Fisher Scientific). Viral P0 stocks were harvested as the supernatant after 5 days and were subsequently amplified to obtain high-titer viral P1 stocks. For expression, 4 to 5 liters of Sf9 insect cells in Sf-900 II SFM medium were infected with P1 virus at a cell density of 3×10^6 cells/ml and a multiplicity of infection of 5. Expression was performed for 72 hours at 27°C under constant shaking. Cells expressing NK₁R or G proteins were harvested by centrifugation, washed with phosphate-buffered saline (MilliporeSigma), frozen in liquid nitrogen, and stored at -80°C. Cells secreting scFv16 were separated from the medium by centrifugation immediately before purification.

Purification of human NK₁R

Insect cells expressing the human NK₁R construct were lysed, and receptor-containing membranes were isolated by repeated Dounce homogenization in hypotonic buffer {10 mM Hepes (pH 7.5), 20 mM KCl, 10 mM MgCl₂, and protease inhibitors [Pefabloc SC (50 μ g/ml; Carl Roth) and Pepstatin A (1 μ g/ml; Carl Roth)]} and hypertonic buffer [10 mM Hepes (pH 7.5), 20 mM KCl, 10 mM MgCl₂, 1.0 M NaCl, and protease inhibitors] in the presence of deoxyribonuclease (DNase, MilliporeSigma). Purified membranes were resuspended in hypotonic buffer, frozen in liquid nitrogen, and stored at -80° C until further use.

Frozen membranes were thawed on ice, CP-99,994 (Tocris) was added to a final concentration of 80 μ M to maximize solubilization yield, and the suspension was incubated for 30 min while turning on a wheel. Afterward, iodoacetamide (2 mg/ml; MilliporeSigma) was added to the solution followed by another 30 min of incubation. Subsequently, the receptor was solubilized in solubilization buffer [30 mM Hepes (pH 7.5), 500 mM NaCl, 10 mM KCl, 5 mM MgCl₂, 40 μ M CP-99,994, 1% (w/v) *n*-dodecyl- β -D-maltopyranoside (DDM, Anatrace), and 0.2% (w/v) cholesteryl hemisuccinate (CHS, MilliporeSigma)] for 3 hours at 4°C. Insoluble material was removed by ultracentrifugation, and the supernatant was incubated with TALON IMAC resin (Cytiva) at 4°C overnight to bind the receptor to the resin.

The receptor-bound resin was washed with 20 column volumes (CVs) of wash buffer I [50 mM Hepes (pH 7.5), 500 mM NaCl, 10 mM MgCl₂, 5 mM imidazole, 10% (v/v) glycerol, and 8 mM adenosine triphosphate (ATP, MilliporeSigma)], stepwise reducing the concentration of detergent and CP-99,994 from 1% (w/v) DDM, 0.2% (w/v) CHS, and 20 µM antagonist to 0.5% (w/v) DDM, 0.01% (w/v) CHS, and 10 µM antagonist, respectively. Subsequently, the detergent was exchanged with 16 CVs of wash buffer I supplemented with 1% (w/v) lauryl maltose neopentyl glycol (LMNG, Anatrace), 0.1% (w/v) CHS, and 10 µM CP-99,994, and the antagonist was replaced with SP (WuXi STA) with another 16 CVs of wash buffer I supplemented with 0.5% (w/v) LMNG, 0.05% (w/v) CHS, and 100 µM SP. Agonist-bound NK1R was finally washed with 25 CVs of wash buffer II [50 mM Hepes (pH 7.5), 500 mM NaCl, 15 mM imidazole, 10% (v/v) glycerol, 0.05% (w/v) LMNG, 0.005% (w/v) CHS, and 50 µM SP] and eventually eluted stepwise with 4 CVs of elution buffer [50 mM Hepes (pH 7.5), 500 mM NaCl, 250 mM imidazole, 10% (v/v) glycerol, 0.01% (w/v) LMNG, 0.001% (w/v) CHS, and 50 µM SP].

Protein-containing fractions were concentrated to 0.5 ml using a 50-kDa molecular weight cutoff (MWCO) Vivaspin 2 concentrator (Sartorius Stedim) and added to a PD MiniTrap G-25 column (Cytiva) equilibrated with G25 buffer [50 mM Hepes (pH 7.5), 150 mM NaCl, 10% (v/v) glycerol, 0.01% (w/v) LMNG, 0.001% (w/v) CHS, and 50 µM SP] to remove imidazole. The complex was treated overnight with His-tagged 3C protease and peptide N-glycosidase (PNGase) F (both prepared in-house) to remove the N-terminal affinity tags and to deglycosylate the receptor. After incubation with Ni-nitrilotriacetic acid (Ni-NTA) resin (Cytiva) for 1 hour, cleaved receptor was collected as the flow-through, then concentrated to ~3 to 5 mg/ml with a 50-kDa MWCO Vivaspin 2 concentrator, frozen in liquid nitrogen, and stored at - 80°C until further use. Protein concentrations were determined by absorbance at 280 nm on a Nanodrop 2000 spectrophotometer (Thermo Fisher Scientific). Protein purity and monodispersity were assessed by LDS-polyacrylamide gel electrophoresis and analytical size exclusion chromatography (SEC) using a Nanofilm SEC-250 column (Sepax).

Purification of heterotrimeric G proteins

Purification of the engineered heterotrimeric G proteins was similar to receptor purification. All buffers used were supplemented with 10 μ M guanosine diphosphate (MilliporeSigma) and 100 μ M tris(2-carboxyethyl)phosphine) (Thermo Fisher Scientific). In contrast to the receptor purification, NaCl concentration never exceeded 150 mM, and all buffers were devoid of any receptor ligand and iodoacetamide. Insect cells expressing the heterotrimeric mini-G proteins were lysed by sonication in hypotonic buffer in the presence of DNase I and separated from debris by centrifugation. G protein-containing membranes were washed by Dounce homogenization in physiological buffer [10 mM Hepes (pH 7.5), 150 mM NaCl, 20 mM KCl, 10 mM MgCl₂, and protease inhibitors] and directly solubilized in solubilization buffer at 4°C for 2 hours. Insoluble material was removed by ultracentrifugation, and the supernatant was incubated with TALON IMAC resin at 4°C for 1 hour.

The G protein–bound resin was washed with 20 CVs of wash buffer I, stepwise reducing the concentration of DDM and CHS from 1% (w/v) and 0.2% (w/v) to 0.5% (w/v) and 0.01% (w/v), respectively. Subsequently, the detergent was exchanged with 16 CVs of wash buffer I supplemented with 1% (w/v) LMNG and 0.01% (w/v)

CHS, and the detergent concentration was reduced with another 16 CVs of wash buffer I supplemented with 0.5% (w/v) LMNG and 0.05% (w/v) CHS. G proteins were finally washed with 25 CVs of wash buffer II and eluted stepwise with 4 CVs of elution buffer containing 750 mM imidazole. Starting from wash buffer II, all buffers contained 1 mM MgCl₂.

Protein-containing fractions were concentrated using a 100-kDa MWCO Vivaspin 2 concentrator and further purified using a Superdex 200 10/300 (Cytiva) column equilibrated with SEC buffer [25 mM Hepes (pH 7.5), 150 mM NaCl, 2% (v/v) glycerol, 0.01% (w/v) LMNG, and 0.001% (w/v) CHS]. Fractions containing monodisperse G protein were collected, concentrated to 7 mg/ml with a 100-kDa MWCO Vivaspin 2 concentrator, frozen in liquid nitrogen, and stored at -80° C until further use.

Purification of scFv16

ScFv16-containing supernatant from expression by secretion from Sf9 cells was pH-adjusted by addition of Hepes (pH 7.5). Metalchelating agents were quenched by incubation with 1 mM CoCl₂ and 5 mM CaCl₂ for 1 hour at 22°C. Resulting precipitates were removed by centrifugation, and the filtrated supernatant was loaded onto a Co-loaded HiTrap IMAC HP column (Cytiva). The column was washed with 20 CVs of buffer A [20 mM Hepes (pH 7.5), 150 mM NaCl, 1 mM MgCl₂, 4 mM ATP, and 5 mM imidazole] followed by 20 CVs of buffer B [20 mM Hepes (pH 7.5), 150 mM NaCl, and 30 mM imidazole]. The protein was eluted with buffer C [20 mM Hepes (pH 7.5), 150 mM NaCl, and 300 mM imidazole]. Monomeric fractions were pooled, concentrated using a 10-kDa MWCO Amicon Ultra concentrator (Merck), and imidazole was removed by applying the concentrate to a PD-10 desalting column (Cytiva) equilibrated with G25 buffer lacking SP. scFv16 was treated for 3 hours with His-tagged 3C protease and PNGase F for removal of affinity tags and deglycosylation. After incubation with Ni-NTA resin for 1 hour, cleaved scFv was collected as the flow-through, concentrated with a 10-kDa MWCO Amicon Ultra concentrator, and further purified using a Superdex 200 10/300 column equilibrated with SEC buffer. Monomeric fractions were pooled, concentrated to ~6 to 7 mg/ml, flash-frozen in liquid nitrogen, and stored at -80°C until further use.

Complex formation

Purified NK₁R, engineered heterotrimeric G proteins, and scFv16 were mixed in a molar ratio of 1:1.2:1.4 in complexation buffer [final concentrations: 25 mM Hepes (pH 7.5), 100 mM NaCl, 2 mM MgCl₂, 0.01% (w/v) LMNG, 0.001% (w/v) CHS, 100 μ M SP, and 100 μ m TCEP]. Apyrase (1.25 U/ml; MilliporeSigma) was added to the mixture, and the coupling reaction was allowed to proceed overnight at 4°C. The formed complex was then isolated by SEC on a Superdex 200 10/300 column equilibrated with complexation buffer with reduced detergent concentrations of 0.001% (w/v) LMNG and 0.0001% (w/v) CHS. Monomeric peak fractions were concentrated to 1 mg/ml for EM studies, using a 100-kDa MWCO Vivaspin Turbo PES concentrator.

Sample preparation for cryo-EM and image acquisition

For cryo-EM grid preparation, 3.5 μ l of purified NK₁R:SP:G_q were applied on glow-discharged holey carbon gold grids (Quantifoil R1.2/1.3, 300 mesh) and subsequently vitrified using a Vitrobot Mark IV (Thermo Fisher Scientific) operated at 100% humidity and 4°C. Cryo-EM imaging was performed on a Titan Krios G3i (Thermo Fisher Scientific), operated at 300 kV, at a nominal magnification of 130,000 using a K3 direct electron detector (Gatan) in superresolution mode, corresponding to a pixel size of 0.325 Å. A BioQuantum energy filter (Gatan) was operated with an energy slit width of 20 eV. A total of 5291 movies were obtained, with an exposure rate of ~14.8 electrons/Å² per second and a defocus range of -0.8 to $-2.4 \,\mu\text{m}$ using automatic data acquisition with EPU (Thermo Fisher Scientific). The total exposure time was 1.79 s with an accumulated dose of ~62.51 electrons/Å² and a total of 67 frames per micrograph.

The NK₁R:SP:G_s complex was imaged with the same conditions and parameters. The only difference was that a total of 6341 movies were recorded with an exposure rate of ~15.03 electrons/Å² per second for an accumulated dose of ~63.51 electrons/Å².

Cryo-EM data processing

Using MotionCor2 (68), raw movies of NK1R:SP:Gq were binned once to generate a pixel size of 0.65 Å, motion-corrected, and dose-weighted. Contrast transfer function parameters for corrected micrographs were determined by Gctf (69), implemented in cryoSPARC (version 3.0.1) (70). Subsequent cryo-EM processing steps were all performed in cryoSPARC. Initial particle selection was done using the automated blob picker on 100 micrographs, using a 100 Å minimum and 150 Å maximum particle diameter, to extract a total of 36,000 particles. The extracted particles were subjected to one round of 2D classification split into 200 classes, from which 10 classes were selected and used as a template for automatic particle picking using the template picker. A total of 3,511,990 particles were subsequently extracted from all 5291 micrographs, followed by two rounds of 2D classification split into 200 classes each. The particles from the best classes were then subjected to two rounds of ab initio 3D reconstructions split into three classes for the first round and two classes for the second round. A final dataset of 558,058 particles from the best 3D class was subjected to 3D nonuniform refinement, resulting in a 2.71 Å density map after sharpening, determined by gold standard Fourier shell correlation (FSC) using the 0.143 criterion. Local resolution estimation was performed with cryoSPARC.

The NK₁R:SP:G_s complex dataset was subjected to the same processing procedure with the following differences. A total of 4,227,825 particles were extracted from 6341 micrographs. After two rounds of 2D classification, the particles from the selected 2D classes were only subjected to one ab initio 3D reconstruction that was split into six 3D classes. A joined total of 395,052 particles from two of the 3D classes were then subjected to nonuniform refinement, resulting in a 2.87-Å density map after sharpening, determined by gold standard FSC using the 0.143 criterion. Local resolution estimation was performed with cryoSPARC.

Model building and refinement

For NK₁R:SP:G_q:scFv16, an initial model was assembled by docking of the individual complex components into the cryo-EM map using the "fit in map" routine in UCSF Chimera (71). The following components were used: NK₁R [Protein Data Bank (PDB) ID: 6HLO], $G\alpha_s$ (PDB ID: 6GDG), and $G\beta_1\gamma_2$ and scFv16 (PDB ID: 6OIJ). SP was manually built in Coot (72). The initial model was manually rebuilt in Coot, followed by several rounds of manual real-space refinement in Coot and real-space refinement with the software package Phenix.real_space_refine from Phenix (73), using information from both the processed and raw cryo-EM map. For the $NK_1R:SP:G_s:scFv16$ model, the individual components of the refined $NK_1R:SP:G_q:scFv16$ model were docked into the cryo-EM map using the "fit in map" routine in the USCF Chimera, followed by manual model building and iterative real-space refinement in Coot with the Phenix.real_space_refine routine. The quality of both models was assessed using MolProbity (74) before PDB deposition.

Whole-cell ligand binding analysis

Human embryonic kidney (HEK) 293T/17 cells (American Type Culture Collection) were cultivated in Dulbecco's modified Eagle's medium (MilliporeSigma) supplemented with penicillin (100 U/ml), streptomycin (100 μ g/ml; MilliporeSigma), and 10% (v/v) fetal calf serum (BioConcept). Cells were maintained at 37°C in a humidified atmosphere of 5% CO₂, 95% air. Transient transfections were performed with TransIT-293 (Mirus Bio) according to the manufacturer's instructions.

Ligand-binding experiments were performed on whole HEK293T cells for comparison of affinities for wild-type and receptor mutants using an homogeneous time-resolved fluorescence binding assay. HEK293T cells were seeded and transfected in poly-L-lysinecoated 384-well plates (Greiner) at a cell density of 10⁴ cells per well. Twenty-four hours after transfection, cells were labeled with 50 nM SNAP-Lumi4-Tb (Cisbio) in assay buffer [20 mM Hepes (pH 7.5), 100 mM NaCl, 3 mM MgCl₂, and 0.05% (w/v) bovine serum albumin (MilliporeSigma)] for 1.5 hours at 37°C. Cells were washed four times with ice-cold assay buffer and were then incubated for 1 hour at room temperature in assay buffer containing fluorescently labeled peptide SP-HL488 [SP labeled with HiLyte Fluor 488 at Lys-3 (Anaspec)]. Fluorescence intensities were measured on a Spark fluorescence plate reader (Tecan) with an excitation wavelength of 340 nm and emission wavelengths of 620 nm and 510 nm for Tb³⁺ and the fluorophore HiLyte Fluor 488, respectively. The ratio of fluorescence resonance energy transfer donor and acceptor fluorescence intensities was calculated (F500 nm/F620 nm). Nonspecific binding was determined in the presence of 125-fold excess of unlabeled SP. Data were analyzed by global fitting to a one-site saturation binding equation with the GraphPad Prism software (version 9.1.0, GraphPad Prism).

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at https://science.org/doi/10.1126/ sciadv.abk2872

View/request a protocol for this paper from Bio-protocol.

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NK₁R:SP:G₅ have been deposited in the Electron Microscopy Data Bank under the accession codes EMD-13140 and EMD-13141. The atomic coordinates of the NK₁R:SP:G_q and NK₁R:SP:G₅ models have been deposited in the Protein Data Bank under accession codes 7P00 and 7P02. All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials.

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