Structural architecture of a dimeric class C GPCR based on co-trafficking of sweet taste receptor subunits

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Class C G protein–coupled receptors (GPCRs) are obligatory dimers that are particularly important for neuronal responses to endogenous and environmental stimuli. Ligand recognition through large extracellular domains leads to the reorganization of transmembrane regions to activate G protein signaling. Although structures of individual domains are known, the complete architecture of a class C GPCR and the mechanism of interdomain coupling during receptor activation are unclear. By screening a mutagenesis library of the human class C sweet taste receptor subunit T1R2, we enhanced surface expression and identified a dibasic intracellular retention motif that modulates surface expression and co-trafficking with its heterodimeric partner T1R3. Using a highly expressed T1R2 variant, dimerization sites along the entire subunit within all the structural domains were identified by a comprehensive mutational scan for co-trafficking with T1R3 in human cells. The data further reveal that the C terminus of the extracellular cysteine-rich domain needs to be properly folded for T1R3 dimerization and co-trafficking, but not for surface expression of T1R2 alone. These results guided the modeling of the T1R2–T1R3 dimer in living cells, which predicts a twisted arrangement of domains around the central axis, and a continuous folded structure between transmembrane domain loops and the cysteine-rich domains. These insights have implications for how conformational changes between domains are coupled within class C GPCRs.

The perception of sweet taste guides animals to seek nutritious food. Initial molecular recognition of sweet-tasting substances, which includes sugars, artificial sweeteners, and some proteins, is by sweet taste receptors comprising taste 1 receptor member 2 (T1R2) and member 3 (T1R3) (1, 2). The sweet taste receptor is conserved in vertebrates, but there are species-specific differences in ligand specificity and sensitivity to inhibitors (3–6).

T1R2 and T1R3 belong to the class C G protein–coupled receptor (GPCR) family. This family is dominated by important neuronal receptors, including eight types of metabotropic glutamate receptor (mGluR) and the γ-aminobutyric acid type B receptor (GABA_B_R) that recognize the major excitatory and inhibitory neurotransmitters, respectively, and the umami taste receptor (T1R1–T1R3). Other class C GPCRs include the calcium-sensing receptor (CaSR) for regulating calcium homeostasis and the promiscuous androgen receptor GPRC6A. Class C GPCRs have a large N-terminal extracellular domain (ECD) genetically fused to the seven-helix transmembrane domain (TMD). The ECD is further divided into a ligand-binding domain (LBD), which adopts a bi-lobed structure that closes over a ligand similar to the action of a venus-fly trap, and, with the exception of GABA_B_R, a cysteine-rich domain (CRD) that connects to the TMD (7, 8). In the human sweet taste receptor, sugars bind and induce closure of the T1R2 LBD, whereas artificial sweeteners and sweet proteins can bind other sites dispersed across the receptor to stabilize an active conformation (3, 9–18).

Class C GPCRs are obligate dimers, and their dimeric architecture is critical for converting agonist binding at the extracellular LBDs into active G protein signaling in the cytoplasm (19). Multiple studies have determined dimeric ECD structures in ligand-bound and -unbound states and identified that ECD dimerization occurs through hydrophobic contacts between the N-terminal subdomains, referred to as lobe 1 (20–27). The relative orientation between lobes 1 and 2 can vary in both

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2 The abbreviations used are: T1R2, taste 1 receptor member 2; T1R3, taste 1 receptor member 3; GPCR, G protein–coupled receptor; mGluR, metabotropic glutamate receptor; GABA_B_R, γ-aminobutyric acid type B receptor; CaSR, calcium-sensing receptor; ECD, extracellular domain; TMD, transmembrane domain; LBD, ligand-binding domain; CRD, cysteine-rich domain; TM and H, transmembrane helix; SSM, site-saturation mutagenesis; ECL2, second extracellular loop; MD, molecular dynamics; ER, endoplasmic reticulum; HA, influenza hemagglutinin; HLA, human leukocyte antigen; HBSS, Hanks’ balanced salt solution; DMEM, Dulbecco’s modified Eagle’s medium; FBS, fetal bovine serum; RMSD, root mean square deviation; Asp, aspartate; SC, SC45647; Cycl, cyclamate; PROP, 6-n-propylthiouracil; Iso, isoproterenol; PDB, Protein Data Bank; MHC, major histocompatibility complex; EBV, Epstein-Barr virus; β2AR, β2-adrenergic receptor.
ligand-free and -bound states, but it is their closure around an agonist that ultimately generates conformational changes in the TMD for active signaling (28–33). There are two isolated TMD structures from mGluR1 and mGluR5 (34–36). Although the mGluR1–TMD crystal packing captured a possible dimer mediated by transmembrane helix 1 (TM1 or H1), cysteine cross-linking has shown that mGluR2 dimerizes via TM4/TM5 contacts in the resting state and via TM6 in the active state (35, 37). It is unclear whether the dimeric mGluR1 TMD structure is physiological or perhaps represents an alternative conformational state. Although these studies provide insights into receptor dimerization and ligand-induced conformational change, it remains unclear how individual domains are assembled into a full-length receptor. Understanding how T1R2 and T1R3 associate can therefore illuminate the mechanism of class C GPCRs generally.

In this study, we identify a conserved surface during in vitro selection of human T1R2 for associating with T1R3. Identifying the dimerization surface of the sweet taste receptor was challenging due to low surface expression (2, 38, 39), which necessitated the selection of T1R2 mutants for enhanced surface expression and co-trafficking with T1R3. A subsequent deep mutational scan of T1R2 in human cells identified surfaces on lobe 1 of the LBD, on the CRD, and in TM6 for dimerization with T1R3 as assessed by their co-trafficking to the plasma membrane. These surfaces are also conserved in natural evolution, but whereas natural sequence conservation is shaped by multiple properties such as protein stability, ligand binding, and signaling, the deep mutational scan could be focused on T1R2–T1R3 co-trafficking to identify residues under explicit conservation for dimerization. We consequently propose a model of the T1R2–T1R3 heterodimer consistent with the saturation mutagenesis and previously published studies.

Results
Selection of T1R2 mutants with enhanced surface expression

Surface localization of human T1R3 depends on human T1R2 co-expression, suggesting the two subunits must form a heterodimeric complex for appropriate trafficking of a functional sweet taste receptor (39). We focus on the mutagenesis and selection of T1R2 to tease apart the effects of mutations on subunit stability (based on the independent trafficking of T1R2 to the cell surface) and dimerization (based on T1R2–T1R3 co-trafficking).

T1R2 and T1R3 were expressed with extracellular N-terminal tags for surface detection either independently or together in Expi293F cells, a suspension culture derivative of HEK293 that has advantages for sorting large mutant libraries. Based on flow cytometry of cells stained in nonpermeable and permeable conditions (Fig. 1, A–D), we found low levels of T1R2 and barely detectable levels of T1R3 on the cell surface, with most protein remaining intracellular. T1R3 surface expression was substantially increased by co-expression with T1R2 but not with the unrelated GPCR CXCR4 (Fig. 1, B and D, and Fig. S1). T1R2 surface levels, however, were decreased by co-expression with T1R3, due partially to decreased total expression (Fig. 1, A and C). Using a brighter fluorescent antibody showed that although T1R2 surface expression is reduced, some T1R2 nonetheless remains surface-localized when co-expressed with T1R3 (Fig. S1).

To identify T1R2 mutants with enhanced surface expression that may be more amenable to biochemical studies, we used fluorescence-activated cell sorting (FACS) to select high-expressing T1R2 variants. A random mutagenesis library of FLAG-tagged T1R2 was transfected into Expi293F cells using conditions that typically yield no more than one sequence variant per cell, providing a tight link between genotype and phenotype (40). The highest T1R2-expressing cells were enriched by two rounds of FACS (Fig. 1E). 61 individual clones were isolated and tested, and the 18 clones with higher T1R2 surface expression were sequenced. All were unique, and yet despite the lack of convergence, two kinds of mutations were over-represented; seven clones had a serine to asparagine substitution at position 212, and 10 clones had frameshift mutations that result in extended C termini or premature termination after TM1 (Fig. 1F and Fig. S2). We named clones with only missense mutations as M1 to M7 and clones with frameshift mutations as FS1 to FS10. An 18th mutant combined premature termination after TM3 with 9 amino acid substitutions, but because it lacked similarities to other clones, it was excluded from further analysis. In summary, a diverse library of T1R2 was successfully evolved in vitro for higher surface expression.

Single missense mutations that enhance surface expression of T1R2 and T1R3

T1R2 variants with enhanced surface expression had one or more missense mutations in the extracellular domain, with the exception of clone FS1 that only had a frameshift (Fig. 1F and Fig. S2). To determine which of the missense mutations are responsible for higher surface expression, we used targeted mutagenesis to introduce the mutations individually into T1R2. Most enhanced T1R2 surface expression, with small increases in total expression compared with WT (Fig. 2A and Fig. S3A). Mutation S212N, which was found in seven of the clones, had the largest effect. I132S, R230H, and D231G also caused substantial increases in surface expression, whereas other mutations had lesser effects. With few exceptions, mutations could generally be combined for even higher T1R2 surface expression (Fig. 2A).

Co-expression with T1R3 decreased surface T1R2 by half or more for all the missense mutations (Fig. 2A), due in part to decreased total expression (Fig. S3A). Surprisingly, T1R3 surface expression was only improved by co-expression with a subset of the T1R2 variants containing mutations I132S, S212N, R230H, D231G, or Q265R, whereas other highly-expressed T1R2 mutants failed to bring T1R3 to the cell surface as effectively as WT and are presumably defective for dimeric associations (Fig. 2B and Fig. S3B).

The five most effective T1R2 mutations–I132S, S212N, R230H, D231G, and Q265R–are located close to the T1R3-binding interface and are directed toward the solvent (Fig. 2C and Fig. S3C). I132S and S212N generate consensus glycosylation sequences (Asn<sup>130</sup>–Tyr<sup>131</sup>–Ser<sup>132</sup> and Asn<sup>212</sup>–Asp<sup>213</sup>–
Thr<sup>214</sup>). N-Linked glycosylation is often involved in folding and trafficking of membrane proteins (41). Whether glycosylation on residue 212 might interfere with ECD dimerization would depend on the relative orientations of T1R2–T1R3 lobes 1 and 2 (Fig. 2C and Fig. S3C), perhaps explaining why S212N caused a large increase in T1R2 surface expression but a smaller increase for T1R3 compared with other mutations (Fig. 2A and B). R230H and D231G, which can cap and break the C terminus of the second helix in lobe 2, may facilitate folding of the loop that forms a critical disulfide (Cys<sup>233</sup>–Cys<sup>513</sup>) between the LBD and CRD (Fig. S3D). Finally, Q265R makes favorable contacts to nearby backbone atoms (Fig. S3E).

**TMDs and cytosolic tails of T1R2 and T1R3 modulate surface expression of the dimeric sweet taste receptor**

Ten of the T1R2 clones had frameshifts generating premature termination after TM1 or an extended C terminus of altered sequence after TM7 (Fig. 1F and Fig. S2). We therefore hypothesized that a C-terminal motif in the transmembrane domain or cytosolic tail of T1R2 inhibits surface expression. To test this, we characterized truncated variants of T1R2 and T1R3 in which the extracellular domains alone (R2ECD and R3ECD, respectively) were fused to a canonical transmembrane helix from MHC class I (R2ECD–MHC and R3ECD–MHC) (Fig. 3A). Surface expression of both R2ECD–MHC and R3ECD–MHC was indeed improved, whereas total expression was similar to the WT proteins (Fig. 3B and C, and Fig. S4, A and B). We further tested two T1R2 frameshift mutants with extended (R2FS1) or pre-matured (R2FS10) cytosolic tails. Surface expression of R2FS1 and R2FS10 was substantially increased, yet these T1R2 variants had relatively similar total expression (Fig. 3B and Fig. S4 A). Cells expressing R2FS1 responded to sweet compounds similarly to cells expressing T1R2, indicating that the mutant is functional (Fig. 3 D–F). The T1R2 transmembrane domain and cytosolic tail therefore retain T1R2 intracellularly, and the transmembrane domain and cytosolic tail of T1R3 couples T1R3 escape from intracellular compartments to the presence of T1R2.

T1R2 and T1R3 variants were then co-expressed to determine whether ECD interactions are sufficient to promote T1R3 surface trafficking. Indeed, surface localization of WT T1R3 or R3ECD–MHC was enhanced by co-expression with R2FS10 and R2ECD–MHC, as well as by co-expression with soluble R2ECD that lacks any transmembrane anchor (Fig. 3 G and H).
Compared with WT T1R2, the R2FS1 variant has increased surface expression and differs by a short random extension three residues in from the native C terminus (Fig. 3, B and J). Because the cytosolic tail of T1R2 is also highly conserved (Fig. 3J), we hypothesized that the cytosolic tail regulates trafficking. To narrow down the location of a trafficking motif, we truncated the last five residues (R2A835–839), as well as substituted the highly conserved terminal basic residues at positions 837 and 838 to glycine (R2-RR837–8GG), as found in R2FS1. Surface T1R2 expression was increased by both modifications, whereas total expression was similar to WT T1R2 (Fig. 3K). However, surface T1R3 failed to increase beyond levels achieved by co-expression with WT T1R2, similar to the previously characterized R2FS1 construct (Fig. S4, F and G). T1R1, a homolog of T1R2 that also forms a heterodimer with T1R3 for the recognition of umami taste, similarly has two basic residues at its C terminus, but their substitution did not increase T1R1 surface expression, and it likely has different trafficking mechanisms (Fig. S4, H and J). We also examined the subcellular localization of T1R2 and R2FS1 using confocal microscopy and found both proteins co-localize intracellularly with GM130, suggesting the checkpoint for trafficking to the cell surface is exit from the Golgi (Fig. S5). In summary, a terminal dibasic motif within the T1R2 cytosolic tail inhibits T1R2 surface localization and favors the plasma membrane trafficking of a T1R2–T1R3 heterodimer based on co-expression analysis.

**Deep mutational scan of T1R2 for surface expression and dimerization with T1R3**

T1R3 is barely detected on the cell surface when expressed alone, yet its surface expression dramatically increases when co-expressed with T1R2 (Figs. 1–3). Furthermore, T1R3 surface expression is correlated with the various surface levels observed for different missense mutants of T1R2, strongly suggesting that T1R2 and T1R3 physically associate and traffic together (Fig. 2). Hence, using co-trafficking of T1R3 to the cell surface as an indirect measure of heterodimer interactions, we screened T1R2 mutations for T1R3 association to define the dimerization surface, using deep mutational scanning in human cells as described previously for the class A GPCRs CXCR4 and CCR5 (40). Single site-saturation mutagenesis (SSM) was applied to R2FS1 rather than WT T1R2 for three reasons. First, because R2FS1 had higher surface expression (Fig. 3G), positive cells could be well resolved for consistent FACS gating conditions. Second, we had already shown that the T1R2 C-terminal tail regulates trafficking of a T1R2–T1R3 heterodimer, but now we wanted to explore all the other potential sites on the LBD, CRD, and TMD that may mediate subunit interactions. Third, R2FS1 is functionally competent for signaling (Fig. 3, D–F). The deep mutational scan is therefore in the R2FS1 background, and from hereon we simply refer to it as T1R2.

Three SSM libraries of FLAG-tagged T1R2 were generated spanning residues 22–285 (ECD1 library), 286–549 (ECD2 library), and 550–836 (TMD library) that together cover nearly all single amino acid substitutions. Splitting mutagenesis across three libraries that are sorted independently increases sampling of each mutation. The T1R2 SSM libraries were transfected in...
Expi293F cells stably expressing c-Myc–tagged T1R3. Cells were stained and sorted for surface expression of T1R2 to assess the impact of mutations on subunit stability and escape from the ER quality control machinery, or were sorted for both T1R2 and T1R3 surface expression to indirectly assess the impact of mutations on heterodimerization (Fig. S6). Naïve plasmid libraries and cDNA from sorted cells were deep-sequenced, and enrichment ratios for all amino acid substitutions were calculated; beneficial mutations are enriched, whereas deleterious mutations are depleted (Fig. 4)(42). Residue conservation scores, which define conserved sites for the activity under selection, are calculated by averaging log₂ enrichment ratios of all mutations at a specific position (40). Both enrichment ratios and conservation scores for the three SSM libraries were closely correlated between independent replicates (Fig. S7).

In the T1R2 sequence–activity landscape, mutations in secondary structure tend to be deleterious for surface expression, whereas the connecting loops are more tolerant of amino acid substitutions. Two long loops in lobe 1 (residues 40–65 and 328–384) were highly tolerant of mutations, as were the loops connecting the seven transmembrane helices, whereas polar substitutions within transmembrane helices were depleted. As expected, stop codons at most positions were depleted, with the notable exception that premature termination after TM1 is tol-
erated for T1R2 surface expression, although such mutants are highly deleterious for "bringing" T1R3 to the cell surface, immediately indicating there are important interactions between the TMDs and cytosolic tails. These results are consistent with the T1R2 random mutagenesis screen.

T1R2 has eight asparagine residues that are putative sites of N-glycosylation. All eight are exposed to solvent on the T1R2 surface and are highly conserved in the in vitro selection except Asn248 (Fig. S8). Four (Asn84, Asn292, Asn368, and Asn428) are known N-glycosylation sites in fish T1R2 (25).

**Conserved surface on the T1R2 ECD for dimerization with T1R3**

When the T1R2 libraries were selected for T1R2–T1R3 surface co-expression, many more mutations were deleterious, and the T1R2 sequence becomes more conserved (Fig. 4E). This is especially apparent in lobe 1 of the LBD. To highlight T1R2 residues preferentially conserved for heterodimerization, we calculate the difference between conservation scores from the selections for T1R2 surface expression (which primarily capture sequence constraints for subunit folding) from the conservation scores for surface expression of T1R3. A surface patch of T1R2 lobe 1, which we refer to as site I, is under strong selective pressure for associating with T1R3 (Fig. 5, A and B; residues at predicted contact sites are listed in Table S1). Hydrophathy-weighted averages of enrichment ratios also highlight that site I is biased toward hydrophilic residues for T1R2 surface expression alone, yet it is restricted to hydrophobic residues for dimerizing and trafficking with T1R3 (Fig. 5, C and D). This is consistent with crystal structures showing that the ECDs of class C GPCRs dimerize mainly through hydrophobic interactions between lobe 1 of each subunit. The mutational scan also indicated that the tip of the CRD most distal from the membrane, which we refer to as site II, is preferentially conserved for dimeric interactions. Furthermore, a large surface on the CRD (referred to as site III) is weakly conserved and biased toward hydrophobic residues.

A dimerization interface between sites I, II, and III of two ECDs closely matches the dimeric structure of agonist-bound CaSR (21). This was surprising, because we had anticipated the mutational scan would define the dimerization interface for the sweet taste receptor in a resting conformation with the LBDs unbound and open. However, we note that the sugar-binding site at the cleft between lobes 1 and 2 of T1R2 is highly conserved in the sequence–activity landscape, and the receptor could bind one of the many sugars or sugar derivatives in the ER lumen. The data are overall highly consistent with a dimeric
T1R2–T1R3 extracellular domain conformation resembling that of ligand-bound CaSR.

Crystal structures show class C GPCRs share structural and dimerization features, and it is therefore likely that the dimerization surface is evolutionarily conserved. Natural sequence conservation for T1R2, T1R3, CaSR, or mGluR3 was calculated and mapped to the crystal structures of fish T1R2–LBD, fish T1R3–LBD, CaSR–ECD, and mGluR3–ECD, respectively. The putative dimerization surface based on the mutational scan is also conserved in natural history (Fig. 5, G–J). An advantage of the experimental mutational scan using two different selection regimes is that it filters out residue conservation for the folding and surface expression of T1R2 alone, thereby focusing attention on residues under explicit preferential conservation for T1R2–T1R3 association and co-trafficking.

Conserved surface on the T1R2 TMD for dimerization with T1R3

Beyond the extracellular region, the difference map also highlighted a region in the T1R2 TMD preferentially conserved for T1R2–T1R3 surface co-expression. Although most of the TMD surface was similarly conserved for both T1R2 surface expression alone or for T1R2–T1R3 co-expression, site IV was notably much more intolerant of mutations in the latter selection (Fig. 5, K and L; Fig. S9, A–J). Site IV is composed of TM6 and the cytoplasmic base of TM5. Interestingly, this region is also extremely conserved during natural evolution based
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Figure 6. Mutational tolerance of T1R2 cysteine residues. A, plot showing average conservation scores of all cysteine residues following in vitro selection. Negative scores indicate greater conservation and intolerance to mutations. Cysteine residues that are preferentially highly conserved for T1R2–T1R3 co-trafficking are highlighted by the difference score and are colored red, B and C, cartoon representations (views are rotated by 180°) showing the positions of cysteine residues in a model of the ECD. T1R2 is gray, and T1R3 is green. Loops containing cysteines (magenta sticks) are colored cyan, and the C-terminal end of lobe 1 is colored yellow. Cys\textsuperscript{359}, which forms a disulfide to T1R3, is light green. D, topology of the CRD and TMD showing disulfides (red lines) between cysteine residues (red circles). Highly conserved cysteines based on the difference conservation score are in bold red text. One A1 module and two A1 modules are colored blue, green, and yellow, respectively, and one A1-like module is colored red. LP, linker peptide. E and F, cartoon representations showing close-up view of the boxed region in D. Cysteines and highly conserved residues in the CRD for T1R2–T1R3 co-expression are shown with sticks. Expression (E) and difference (F) conservation scores are mapped to T1R2; conservation scores are colored from $-2$ (conserved, orange) to $+2$ (blue).

on alignments of orthologous mGluR1 or T1R2 sequences, whereas the surrounding TMD surfaces are highly variable (Fig. 5, M–O; Fig. S9, K–T). Because the extracellular dimerization surface identified by the mutational scan is both well-matched to crystal structures of ECD dimers and evolutionarily conserved, we argue that site IV, which is similarly highly conserved both in natural history and our mutational scan, forms the dimerization interface between the TMDs. A previous study used cross-linking to show that TM6 helices of both subunits are in proximity in agonist-bound mGluR2 (37). This further suggests that site IV is a real dimerization surface of T1R2 and that class C GPCRs share a common dimerization mechanism for the ECDs and TMDs. Because residues in TM6 of mGluR2 are only cross-linked in the active conformation (37), the mutational scan likely informs on a ligand-bound T1R2 conformation, as also suggested by the data’s agreement with the agonist-bound CaSR–ECD crystal structure. Finally, the cytosolic surface of the TMD is conserved both in the deep mutational scan and in natural evolution, suggesting that proper structure of the G protein–binding site is important for sweet taste receptor dimerization and surface expression (Fig. S9, E, J, O, and T).

CRD and the second extracellular loop (ECL2) of the TMD are involved in T1R2–T1R3 surface co-expression

T1R2 has 19 extracellular cysteine residues, all of which form disulfide bonds. In the LBD, three cysteines were highly conserved, whereas five cysteines within or bonded to the two long variable loops were generally tolerant of substitutions (Fig. 6A and Fig. S10, A and B). The cysteines with high-mutational tolerance are Cys\textsuperscript{102} of the lobe 1 core that bonds to Cys\textsuperscript{59} in the first variable loop, Cys\textsuperscript{363} and Cys\textsuperscript{366} that form a disulfide between nearby positions within the same loop, and Cys\textsuperscript{359} that forms an intersubunit disulfide with Cys\textsuperscript{129} of T1R3. This final result was unexpected; the cysteine forming a disulfide connection between T1R2 and T1R3 is clearly unnecessary for subunit expression and dimerization, and rather the major determinants of T1R2–T1R3 association are noncovalent interactions. In metabotropic glutamate receptors, the disulfide linking the two subunits is also dispensable for expression of a functional dimer (29). Of the three highly conserved LBD cysteines in the mutational scan, Cys\textsuperscript{405} and Cys\textsuperscript{410} form a disulfide that stabilizes a sharp turn at the apex of lobe 1, and Cys\textsuperscript{233} connects the LBD and CRD to couple conformational dynamics (Fig. 6, B–D).

With the exception of the GABA\textsubscript{B} receptor, class C GPCRs have nine strictly conserved cysteines in the CRD, and all nine were also conserved in our experimental deep mutational scans for T1R2 surface expression and dimerization (Fig. S10, A and B). However, sequence constraints were tighter for the cysteines during selection for T1R2–T1R3 co-expression, and this was most apparent for cysteine residues near the CRD C terminus. In particular, the last two cysteine residues of the CRD
(Cys$^{538}$ and Cys$^{551}$) are highlighted in the difference conservation scores, indicating that the CRD C terminus must be properly structured for T1R2–T1R3 dimerization but less so for T1R2 surface expression alone (Fig. 6, A, E and F). Furthermore, ECL2 and three residues (Asn$^{541}$, Phe$^{552}$, and Arg$^{554}$) at the CRD C terminus are also highlighted in the difference map as under greater conservation for dimerization. A structural model of the heterodimeric sweet taste receptor must account for why the bases of the CRD and ECL2 are conserved in the mutational scan, despite the fact that both these regions are solvent-exposed in crystal structures of the separate parts.

**Model of T1R2–T1R3 dimerization**

The deep mutational scans indicated interaction sites on LBD lobe 1 (site I), the CRD (sites II and III), and the TMD (site IV) for co-expression of T1R2 and T1R3 on the cell surface. The data are consistent with a conformation of the T1R2–ECD resembling the crystal structure of ligand-bound CaSR–ECDs, in which the CRDs are in close proximity with contacts mediated by sites I, II, and III. The data are also consistent with contacts between TM6 of the TMD as found earlier by cross-linking agonist-bound mGlur2 subunits, in addition to contacts mediated by the base of TM5 (collectively forming site IV) (37). A model was constructed of the human sweet taste receptor in a putative ligand-bound conformation by bringing together homology models of the individual domains in an arrangement that satisfies the experimental data (Fig. 7A and Fig. S11). Side-chain and backbone conformations were minimized using the ROSETTA energy function (43), followed by all-atom molecular dynamics (MD) in a lipid bilayer with explicit solvent. Distance restraints between conserved residues predicted to be at the dimer interface (Table S2–S4) maintained a pseudo-symmetric architecture. C-terminal peptides of G proteins were modeled bound to the TMDS for the purpose of forcing an active conformation; this is not to imply that the dimeric receptor trafficking in our experimental system is necessarily bound to G proteins, and actual G protein signaling occurs through a single TMD (44).

The model has three notable features. First, ECL2 and the linker peptide of the CRD form a fold that resembles an A1 module in distantly related cysteine-rich domains (Figs. 6D and 7, B and C) (45). This forms a continuous structure similar to the CRD of tumor necrosis factor receptor, in which aromatic residues mediate module stacking (Fig. 7C) (46, 47). Aromatic residues T1R2–Trp$^{543}$ (highly conserved in both the mutational scan and natural evolution) and T1R2–Phe$^{552}$ (conserved as an aromatic in the mutational scan) are exposed on the CRD surface but become partially buried by contacts to the ECL2 backbone and T1R2–Pro$^{713}$ (tolerates hydrophobic substitutions in the mutational scan). Just as LBD lobe 2 and the CRD have two connections (the protein backbone and a disulfide), there are possibly two connections between the CRD and TMD (the protein backbone and the CRD–ECL2 folded structure) that ensure interdomain motions are coupled and not dissipated within flexible backbone linkers. Second, the two ECL2s contribute to the dimerization interface (Fig. 7B). As a result, the CRDs are stacked both within and between subunits through aromatic and proline residues and by aliphatic motifs of lysine and arginine side chains. This can account for the high conservation of the CRD C terminus in the deep mutational scan for co-expression with T1R3 (Fig. 6). Finally, our model predicts a “twisted” or “spiral” architecture of the domains around the central rotational axis (Fig. 7A). The model may account for how ligand-dependent translational motions of the CRDs are coupled to rotational motions of the TMDS (37) to reconfigure the transmembrane dimer interface (Fig. S11).

**Discussion**

Here, we identified important regions of human T1R2 for surface expression and co-trafficking with T1R3 using mutational screening and deep mutational scanning. These results guided the modeling and “visualization” of the full-length T1R2–T1R3 dimer in living cells. Our model provides insights into the general mechanism of class C GPCRs.

Many of the T1R2 clones isolated in our mutational screen for enhanced surface expression shared a S212N missense mutation, creating a new consensus N-glycosylation site within an accessible loop of lobe 2. Native N-glycosylation is required for surface expression of other class C GPCRs (21, 48–50). Notably, with only one exception, all putative N-glycosylation...
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sites of T1R2 were also highly conserved for surface expression in the deep mutational scan.

The cytosolic C terminus of T1R2 modulates surface expression of the sweet taste receptor. Replacement of the TMDs and cytosolic tails of T1R2 or T1R3 with generic transmembrane helices increased surface expression, and although ECD interactions alone were sufficient, the native TMD and cytosolic tail of T1R2 were required for optimum co- trafficking. We further identified a dibasic motif in the cytosolic tail of T1R2 that inhibits plasma membrane localization. Mutation of the dibasic motif dramatically increased surface T1R2 but failed to substantially change T1R3 levels at the cell surface, suggesting monomeric or homodimeric T1R2 was now escaping intracellular retention. Dibasic motifs in cytosolic regions of membrane proteins can, in some cases, bind adaptors to drive retrograde translocation from the Golgi back to the ER, whereas the similar K(X/D/E) motif can retain proteins in the Golgi (51, 52). This mechanism for intracellular retention is especially important for quality control of protein complexes (53), including the GABA_ß receptor (a heterodimer of GB1 and GB2), where the GB1 subunit has a C-terminal dibasic motif that becomes masked by formation of a coiled-coil with the cytosolic tail of GB2 (54). Human T1R2 also has a C-terminal dibasic trafficking motif (sequence RRD), yet when mutated, only a fraction of the protein escapes to the surface while much remains in the Golgi (Fig. S5). There is complexity in the trafficking mechanism of the sweet taste receptor that will require further study.

Based on a comprehensive mutational scan of T1R2 co-expressing with T1R3 on the cell surface, important interaction sites were identified that guided modeling of the full sweet taste receptor. The domain orientations are constrained by the mutational data in such a way as to bring ECL2 immediately below the last module of the CRD. In GPCRs generally, ECL2 is a critical determinant of ligand interactions (55, 56). ECL2 of class C GPCRs are similar in length (less than 35 residues), and the length of the linker peptide at the end of the CRD is also important for receptor activation and surface expression (57–59). Interaction of ECL2 with the CRD linker peptide was observed in the crystal structure of inactive mGluR1 TMD and was hypothesized to be important for activation (9, 36, 59), and we now predict ECL2–CRD interactions are important for dimerization. Further studies will be required to conclusively elucidate the role of ECL2 in dimerization and activation. It is also notable that the mutational scan clearly implicates TM6 and the base of TM5 at the dimer interface, as this region undergoes the most significant conformational change between resting and activated class A GPCRs (60).

The mutational scan reveals contacts between T1R2 and T1R3 that are consistent with a ligand-bound conformation trafficking to the cell surface. We showed that the WT and expression-enhanced sweet taste receptor used in this study are functional in HEK293 cells, but we failed to detect signaling by the sweet taste receptor in the Exp293F cell line, either by microscopy or flow cytometry, even when overexpressing promiscuous G proteins (positive controls with class A GPCRs showed robust responses). We speculate that due to the rich growth medium, the sweet taste receptor binds sugars or sugar derivatives in the ER shortly after synthesis and is desensitized.

Low-glucose growth medium, which is used to prevent desensitization in other lines, was toxic to Exp293F cells. It is possible that in sensory neurons of the taste bud, a T1R2–T1R3 heterodimer in a resting conformation may instead traffic to microvilli at the taste pore. Despite this limitation of the experimental system, our results are nonetheless consistent with previous data, crystal structures, and natural evolution. We therefore argue that the dimerization surface we identify by mutational scanning in living cells is real.

Finally, late during revision of this manuscript, the cryo-EM structure of the class C GPCR mGluR5 in an active conformation was published (61). Our modeled architecture of the active sweet taste receptor, with a twisted arrangement of domains and contacts to the CRDs via ECL2 loops, perfectly matches the cryo-EM structure (Fig. S12). This highlights that our innovative use of mutational scans to accurately model proteins in living cells is valid and of high impact.

Although natural history of a protein sequence contains combined information on folding, ligand recognition, signaling, and conformational dynamics, in vitro evolution and deep mutational scanning can be focused on specific protein properties. Here, we focused selections on surface expression to identify residues important for subunit folding and complex assembly. Further deep mutational scans may offer additional insights into class C GPCR mechanisms and provide a structural basis for designing low-caloric sweeteners to address obesity and diabetes.

Experimental procedures

Construction of plasmids

The native sequence of human T1R2, a human codon-optimized sequence of human T1R3, and an Escherichia coli codon-optimized sequence of human T1R1 were synthesized as gene fragments (Integrated DNA Technologies). Based on flow cytometric analysis of transfected cells, we found no difference in expression of native or codon-optimized human T1R3 cDNA, but due to difficulties in amplifying the native sequence for PCR-based modifications, we chose to use codon-optimized T1R3 except for the experiment described in Fig. S1, where native T1R3 was used. The native signal peptides (amino acids 1–24, 1–21, and 1–20 for T1R1, T1R2, and T1R3, respectively) were replaced with the signal peptide of influenza hemagglutinin (HA) or of HLA class I histocompatibility antigen A-2 α chain (HLA). FLAG or c-Myc tags with short flexible linker sequences were inserted after the signal peptide. T1R1 WT and mutant constructs were cloned in pcDNA3.1(+) (ThermoFisher Scientific). T1R2 and T1R3 WT and mutant constructs were cloned in pCEP4 (Invitrogen) or pcDNA3.1(+). T1R1 and T1R3 were constructed with the HLA signal peptide followed by the c-Myc tag. T1R2 constructs were fused to the HA signal peptide followed by the FLAG tag, with the exception of Fig. S1, where T1R2 was fused downstream of a HLA signal peptide and FLAG or c-Myc tag.

Cell culture and transfection

Exp293F™ cells were cultured with Exp293™ expression medium (ThermoFisher Scientific) in a 37 °C incubator with humidified atmosphere of 8% CO₂. For 1-ml transfections, up
to 1 µg of plasmid DNA and 2.7 µl of ExpiFectamine™ 293 reagent (ThermoFisher Scientific) were prepared separately in 50 µl of OptiMEM (Gibco). After 5 min of incubation at room temperature, DNA and ExpiFectamine™ 293 reagent were mixed and further incubated for 30 min. Total 100 µl of DNA/ExpiFectamine™ 293 reagent mixture was added to 1 ml of 2 x 10⁶ cells.

**Flow cytometry analysis of surface and total expression**

Total 1 µg of DNA was transfected into 1 ml of 2 x 10⁶ cells as described above. 500 ng of T1R2, T1R3, or T1R1 was co-transfected with 500 ng of empty vector. For co-expression of T1R2 and T1R3, 500 ng of each DNA was co-transfected. Transfected cells were collected after 22–27 h by centrifugation at 500 x g for 1 min at room temperature. The cell pellet was resuspended in 800 µl of cold phosphate-buffered saline (PBS) containing 0.2% BSA (PBS/BSA). Cells were collected by centrifugation and resuspended in 300 µl of PBS/BSA. 3 µl of fixable viability dye eFluor 450 (eBioscience) was added to the cells and incubated for 30 min on ice in the dark. Cells were pelleted and resuspended in 800 µl of PBS/BSA. The cells were then split into two tubes for staining surface and whole-cell protein. For surface staining, cells were centrifuged and resuspended in 40 µl of PBS/BSA. 0.3-µl antibody was then added and incubated for 20 min on ice in the dark. Unless otherwise noted, anti-FLAG–Cy3 (Sigma; clone M2) and anti-c-Myc–Alexa Fluor 647 (Cell Signaling; clone 9B11) antibodies were used. Cells were washed twice with 800 µl of PBS/BSA, resuspended in 300 µl of PBS/BSA, and analyzed. BD Cytofix/Cytoper™ Fixation/Permeabilization kit (BD Biosciences) was used for whole-cell protein staining. Cells were collected by centrifugation and resuspended in 250 µl of Fixation/Permeabilization solution for 20 min on ice in the dark. Cells were washed twice with 1 ml of 1 x BD Perm/Wash™ buffer or 0.5 x PBS–BD Perm/Wash™ buffer, and resuspended in 40 µl of wash buffer. 0.3 µl of antibody was then added and incubated for 30 min on ice in the dark. Cells were washed twice with 250 µl of wash buffer and resuspended in 250 µl of wash solution for analysis. Surface and total expression of T1R1, T1R2, and T1R3 were measured on a BD LSR II or BD Accuri™ C6 (BD Biosciences). The data were collected using BD FACSDiva software (BD Biosciences) and analyzed using FCS Express 6 (De Novo Software). Gates were set on negative controls with 0.5% of events falling within the gate, and the percentages of fluorescence-positive cells were calculated. Relative surface and total expression were normalized to WT.

**Random mutagenesis library generation**

T1R2 was mutagenized by error-prone PCR using goTaq (Promega), with an average error rate of two amino acid substitutions per 2.5-kb T1R2 gene. T1R2 PCR fragments were inserted into the pCEP4 vector using restriction and ligation methods. The T1R2 mutant library was transformed into 5-α electrocompetent E. coli cells (New England Biolabs) by electroporation. Transformed cells were collected after 1 h of incubation in S.O.C. medium at 37 °C, and 10 µl of serially diluted cells were plated to calculate the library size. Remaining cells were inoculated in 250 ml of LB medium and cultured overnight at 37 °C, and plasmid DNA was prepared using GeneJET plasmid maxiprep kit (ThermoFisher Scientific). The cDNA from sorted T1R2 mutant library-transfected Expi293F cells was similarly cloned into pCEP4, and plasmid DNA was purified as described above.

**Sorting the T1R2 random mutagenesis library**

30 ng of T1R2 mutant library and 60 µg of pUC18 carrier DNA were co-transfected into a 30-ml Expi293F culture. Medium was replaced 2 h post-transfection. Transfected cells were harvested 22 h post-transfection by centrifugation at 500 x g for 2 min at room temperature. The cell pellet was washed with 40 ml of PBS/BSA, resuspended in 4 ml of PBS/BSA containing 1/500 chicken anti-DYKDDDDK–FITC (Immunology Consultants Laboratory), and incubated for 30 min on ice in the dark. 40 ml of PBS/BSA was added to the cells, and unbound antibody was washed by centrifugation at 500 x g for 1 min at room temperature. The cells were washed twice more with 40 ml of PBS/BSA and resuspended in 4 ml of PBS/BSA for sorting. The 0.4% of cells with highest FITC signal were collected for 4 h on a BD FACS Aria II at the Roy J. Carver Biotechnology Center. Sorted cells were centrifuged at 500 x g for 2 min at 4 °C and stored at –80 °C for later RNA extraction.

**RNA extraction and cDNA synthesis**

Total RNA was extracted using GeneJET RNA purification kit (ThermoFisher Scientific). T1R2 cDNA fragments were synthesized from the extracted RNA by RT-PCR using AccuScript Hi-Fi (Agilent).

**Reconstruction of sorted T1R2 random mutagenesis library after sorts**

Because of inefficiencies of high-fidelity reverse transcription, full-length T1R2 DNA was not amplified from the cDNA. Instead, three separate fragments were amplified using Phusion (ThermoFisher Scientific). Fragment 1 was amplified from cDNA primed with an internal gene-specific primer (TGGACCTTGTCACCAATG) using PCR primers pCEP4_MCS-F (GATCTCTAGAAGCTGGGTACC) and NativeR2_775_seqR (TGGAGCTTGCCACCATG). Fragment 2 was amplified from cDNA primed with the EBV-reverse primer using PCR primers NativeR2_616_seqF (ATCATTGTCGTTGAAGCAG) and natR2_1335_seqR (TGAGAACTTCGAAGTCCAGA). Fragment 3 DNA was amplified from cDNA primed with the EBV-reverse primer using PCR primers NativeR2_1275_seqF (GAAGGTCAACTTCACTCTCC) and pCEP4_MCS-R (CAATGTATCTTTCTATCTTG). The full-length product was then assembled from the fragments by overlap extension PCR and cloned into pCEP4 for a second round of sorting.

To reassemble the T1R2 library after the second round of sorting, fragment 2 was amplified using PCR primers natR2_691F_seqF (GACATCTGCATGCGCTTTCCA) and natR2_1417_seqR (TGAGATTTCTCAGCTTGCG). Fragments 1 and 3 were amplified as described above. The full-length DNA library was amplified and cloned into pCEP4 as described above.
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Functional expression

HEK293 cells were kindly provided by Dr. Makoto Tominaga (Okaaki Institute for Integrative Bioscience). These cells were cultured at 37 °C under a humidified atmosphere containing 5% CO2 in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS). For calcium imaging experiments, cells were seeded onto a 35-mm recording chamber. After 24 h at 37 °C, plasmid DNAs were transiently co-transfected into HEK293 cells using Lipofectamine2000 reagent (Life Technologies, Inc.) (2.5 μl per 0.9 μg of DNA). TIR2 or R2FS1 was co-transfected with TIR3 and Ga16-gust44 using 0.9 μg of plasmids. Ca2+ imaging assays were performed 24 h after transfection.

Single-cell Ca2+ imaging

As described previously (62, 63), a bath perfusion system was used for determination of the kinetics of activation. Transfected cells in 35-mm recording chambers were washed in Hanks’ balanced salt solution (HBSS) (Life Technologies, Inc.) containing 10 mM HEPES (pH 7.4) and loaded with 3 mM fluo-4 acetoxymethyl ester (Life Technologies, Inc.) for 30 min at 37 °C. Taste solutions diluted in HBSS containing 10 mM HEPES were applied sequentially to the cells for 30 s with a peristaltic pump at a flow rate of 1.5 ml/min, and fluorescence images were obtained using a S Fluor 620/0.75 objective lens (Nikon) via a cooled-CCD camera (C6790, Hamamatsu Photonics) fitted to a TE300 microscope (Nikon). AquaCosmos software (version 1.3, Hamamatsu Photonics) was used to acquire and analyze fluorescence images. A 5-min interval was maintained between each tastant application to ensure that the cells were not desensitized. Responses were measured from individual responding cells. Isoproterenol (Iso, 10 μM) was used as positive control, which stimulates endogenous β-adrenergic receptors, showing that the Ga16-dependent signal transduction cascade was functional. SC45647 (0.3 mM), aspartame (10 mM), and sodium cyclamate (30 mM) were used as sweet taste stimuli. Human-specific sweet receptor inhibitor lactisole (1 mM) and 6-n-propylthiouracil (PROP: T2R38 agonist) were used as negative controls.

Ca2+ imaging data analysis

In the analysis of single cell responses, changes in [Ca2+]i were monitored as changes in fluo-4 fluorescence. Fluorometric signals are expressed as relative fluorescence changes: ΔF/F0 = (F − F0)/F0 where F0 denotes the baseline fluorescence level. The magnitude of the calcium increases from 10 to 30 s after stimulus onset was measured and averaged.

SSM library generation

Expression-enhanced variant R2FS1 was used as the template. An internal XhoI site was silenced, and silent HindIII, Pfl23II, and XhoI sites were inserted by targeted mutagenesis. Three SSM libraries covering TIR2 residues 22–285 (ECD1 library), 286–549 (ECD2 library), and 550–936 (TMD library) were generated by overlap extension PCR (64) and ligated into the Nhel-XhoI, HindIII-Pfl23II, and XhoI-BamHI sites of HA-FLAG-R2FS1 in pCEP4. The ligations were transformed into 5-α electrocompetent E. coli cells, and plasmid DNA was prepared as described above, ensuring the number of transformants was at least 10 times the possible library diversity. The three deep sequenced plasmid libraries covered 16,177 out of 16,300 possible single amino acid substitutions based on a minimum frequency of 5 × 10^-6.

c-Myc–TIR3 stable cell line generation

HLA-c-Myc–TIR3 in pcDNA3.1(+) was linearized by restriction with PvuI. 4 μg of linearized DNA was transfected in 8 ml of Expi293F cells as described above. G418 was added 24 h post-transfection at 100 μg/ml. Cells were passaged, and medium supplemented with G418 was replaced every 2–3 days. The G418 concentration was reduced to 50 μg/ml 3 weeks post-transfection. TIR3-positive cells were enriched by FACS as described above with some changes. R2FS1 cloned in pcDNA3.1(+) was used as an indicator to select cells expressing TIR3; unlike pCEP4, pcDNA3.1(+) lacks the ability to replicate extra-chromosomally and therefore will be lost with passaging unless integrated into the genome. Cells were transfected with R2FS1 to promote surface trafficking of TIR3 and stained with anti-FLAG–FITC and anti-c-Myc–Alexa 647. Propidium iodide was added for gating and exclusion of dead cells. The top 8% of Alexa 647–positive cells were collected and recovered in fresh medium supplemented with penicillin–streptomycin.

Sorting TIR2 SSM libraries

Plasmid DNA libraries were transfected into TIR3 stable Expi293F cells using conditions that typically yield no more than one coding sequence per cell (40). 1 ng/ml library DNA and 1.5 μg/ml pCEP4–ΔCMV (in which the fragment between Sall sites of pCEP4 is removed) were transfected into TIR3 stable cells at 2 × 10^6 cells/ml. Medium was replaced 2 h post-transfection, and cells were harvested 22–24 h post-transfection by centrifugation at 500 × g for 2 min at 4 °C. Cells were washed with 30 ml of PBS/BSA, and stained with 30 μl of anti-FLAG–Cy3 and anti-c-Myc–Alexa 647 in 4 ml of PBS/BSA for 30 min in the dark. Cells were washed twice with 30 ml of ice-cold PBS/BSA and resuspended in PBS/BSA for sorting on a BD FACS Aria II at the Roy J. Carver Biotechnology Center. Propidium iodide was added to the sample immediately prior to sorting, and dead cells were gated in the PE/Texas Red channel and excluded. Auto-fluorescent cells in the AmCyan channel were also removed during gating. The highest 55% (ECD1 and ECD2 libraries) and 100% (TMD library) of Cy3-positive cells were sorted for surface expression of TIR2. For co-trafficking of TIR3, the top 20% of APC-positive cells were sorted after gating 100 or 55% of Cy3-positive cells as just described. Sorted cells were collected and frozen at −80 °C. To maintain viability, prepared samples were sorted for no more than 4 h. Libraries were sorted on multiple occasions to collect greater numbers of cells, and the sorted cell pellets were pooled during RNA extraction.

Deep sequencing

RNA was extracted as described above. First-strand cDNA was synthesized with Accuscript using internal gene-specific primers to offset the poor efficiency of accurately reverse-
transcribing long messages. For the ECD1 library, reverse transcription was primed with natT1R2_882_ECD1_R (AGT-GAAGTCTGGCCGACG); for the ECD2 library with natT1R2_1682_ECD2_R (CATTCAGAAGACCAGACG); and for the TMD library with pCEP4_EBV_Rev (GTGGTTTGGTCCAACATCT). In a first round of PCR (18 thermocycles), primer overhangs added sequences complementary to the Illumina sequencing primers, and then barcodes and adaptor sequences were added in a second round of PCR amplification (15 thermocycles) for annealing to the Illumina flow cell. Two overlapping fragments were amplified for both ECD1 and ECD2 libraries, and three overlapping fragments were amplified for the TMD library to achieve full sequencing coverage. PCR fragments were then sequenced at the University of Illinois at Urbana-Champaign Roy J. Carver Biotechnology Center on an Illumina HiSeq 2500 (2 × 250-nucleotide kit). Deep sequencing data were analyzed with Enrich (65). Normalized log₂ enrichment ratios were calculated by subtracting the log₂ enrichment of the WT sequence. Commands are available in the data deposition with NCBI’s Gene Expression Omnibus (66) under series accession number GSE115751.

Estimating evolutionary conservation scores

Evolutionary conservation scores for amino acid positions of T1R2, T1R3, CaSR, mGluR1, and mGluR3 were calculated using the ConSurf server (67, 68). The search for homologs was based on the amino acid sequence of T1R2 TMD or on protein structures in other cases (69). Homologous sequences were searched using HMMER from UNIREF-90 with a number of iterations of 1 and E-value cutoff of 0.001. Orthologous sequences were manually selected for LBD of T1R2 (103 orthologous) and T1R3 (147 orthologous), ECD of CaS receptor (45 orthologous) and mGluR3 (59 orthologous), and TMD of T1R2 (132 orthologous) and mGluR1 (156 orthologous). The list of sequences used in each analysis is available upon request. Multiple sequence alignments were built using MAFFT-L-INS-I, and conservation scores were computed using Bayesian algorithm of substitution.

Structural modeling

The sequences of human T1R2 and T1R3 were threaded on the crystal structures of fish T1R2–T1R3 LBDs (PDB 5X2N), and the TMDs were manually positioned so as to bury site IV identified in the mutational scans while maintaining pseudo-C₂ symmetry. The full dimeric structure was fast-relaxed with coordinate restraints on all atoms, followed by fast-relax with distance constraints holding four of the most conserved T1R2 residues predicted to be at the dimer interface, and the equivalent T1R3 residues, in proximity of the central axis (Table S2). This prevented the extracellular domains “bending over” at flexible connections, thereby maintaining pseudo-C₂ symmetry. The lowest energy ROSETTA model was chosen from 300 decoys.

The ROSETTA model was further relaxed by molecular dynamics (MD). The MD system was built using Tleap module in AMBER14 (73). The transmembrane domains were embedded in 1-palmitoyl-2-oleoyl-phosphatidylcholine lipid bilayer, and the full receptor was solvated using TIP3P water molecules (74). The MD system was neutralized using 0.15 M NaCl. The AMBERff14SB force field (75) was used for simulation, and the protonation states were assigned to the titratable residues at physiological pH 7. The final MD system contained ~193,000 atoms. The MD system was subjected to minimization for 20,000 cycles and then heated at constant volume from 0 to 300 K over 3 ns. The system was equilibrated for 20 ns at constant pressure and temperature to 1 bar and 300 K. The backbone atoms of the receptor dimer were restrained with a force constant of 10 kcal/mol/Å² during the minimization, heating, and equilibration protocol. The hydrogen bonds were constrained using SHAKE algorithm (76), and the long-range electrostatics were treated with Particle Mesh Ewald method (77). The nonbonded interaction cutoff was set to 10 Å, and the integration time step was 2 fs. The final productions runs were performed over a period of 50 ns. During the productions runs, the combination of relaxation protocols was applied over the length of the simulation time. For the initial 10 ns, the Cα atoms of all protein residues were restrained. For the next 10 ns, the Cα atoms of residues at the domain interface were restrained (Table S3). Finally, distance restraints were applied for 30 ns between a small subset of residues at the domain interface (Table S4). A force constant of 20 kcal/mol/Å² was used in the production runs. The root mean square deviation (RMSD) plots of extracellular and transmembrane domains (Fig. S11E) show that the structures are converged with a mean RMSD of 2.5 Å. The flexible cysteine-rich loop regions are excluded from the RMSD analysis.

Confocal microscopy

HEK293T cells were seeded onto µ-slide 8-well (ibidi) at least a day before transfection and cultured with DMEM supplemented with 10% FBS at 37 °C with 5% CO₂. 50–60% confluent cells were transfected. 300 ng of plasmid DNA and 0.9 ng of linear polyethyleneimine (Polyscience, M, 250,000) were mixed in 30 μl of OptiMEM and incubated for 20 min at room temperature. The DNA/polyethyleneimine mixture was added to a well. Cells were washed with PBS after 24 h, fixed, and permeabilized with the Fixation/Permeabilization solution of BD Cytofix/CytopermTM Fixation/Permeabilization kit for 20 min. Cells were further washed and incubated with BD Perm/Wash™ buffer for 15 min. Cells were then incubated with anti-FLAG–Cy3 (1:200) and anti-GM130–Alexa Fluor 488...
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(TermoFisher Scientific, 1:400) or anti-calnexin–Alexa Fluor 488 (TermoFisher Scientific, 1:200) for 1 h in the dark. Cells were washed three times with PBS. Images were collected on Zeiss LSM 700 (Carl Zeiss) using ×63 oil-immersion objective. The collected images were processed using Fiji (ImageJ).

Data and plasmid availability

Raw deep sequencing data and calculated enrichment ratios are deposited with NCBI’s Gene Expression Omnibus (GEO) under series accession number GSE115751. Plasmids have been deposited with Addgene under accession numbers 113944–113962. Structure coordinates are available upon request.3

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