T2Rs Function as Bitter Taste Receptors

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Introduction

Mammals can perceive and distinguish between sweet, sour, bitter, and salty tastes (Kinnamon and Cummings, 1992; Lindemann, 1996a; Stewart et al., 1997). Of these four modalities, bitter perception has a particularly important role: many naturally poisonous substances taste bitter to humans, and virtually all animal species show an aversive response to such tastants (Garca and Hankins, 1975; Glendenning, 1994; Glendenning et al., 1999), suggesting that bitter transduction evolved as a key defense mechanism against the ingestion of harmful substances.

The biology of bitter perception is very poorly understood; neither the sensory receptor cells nor the receptor molecules have been physiologically or molecularly defined (Lindemann, 1996b). However, several biochemical and physiological studies have suggested that bitter transduction in mammalian taste receptor cells is mediated by G proteins and G protein-coupled receptors (GPCRs) (Lindemann, 1996a; Wong et al., 1996). Because the universe of chemical compounds that evoke a bitter taste is structurally diverse, we reasoned that bitter receptors might encompass a large GPCR family with significant sequence variation. In the accompanying paper (Adler et al., 2000 [this issue of Cell]), we described the isolation of a novel family of 40–80 divergent GPCRs, T2Rs, selectively expressed in subsets of taste receptor cells of the tongue and palate epithelium. T2Rs in humans and mice are genetically linked to loci associated with bitter perception (Conneally et al., 1976; Capeless et al., 1992; Reed et al., 1999), and are selectively expressed in taste receptor cells that contain gustducin, a G protein α subunit implicated in bitter transduction (Wong et al., 1996; Ming et al., 1998). While the genetics, expression profile, and diversity of the T2R family support the proposal that T2Rs are taste receptors, rigorous demonstration of their role in taste transduction requires functional validation. Here we use a heterologous expression system to demonstrate that T2Rs function as receptors for bitter tastants. We analyzed mouse strains that differ in their recognition of various bitter compounds and show that mice that do not perceive low concentrations of cycloheximide contain missense mutations in the mT2R-5 gene. These amino acid changes significantly reduce the sensitivity of the mT2R-5 receptor to cycloheximide. Mice strains deficient in their ability to detect cycloheximide have amino acid substitutions in the amino acid changes significantly reduce the sensitivity shift measured in cell-based assays closely mirrors the behavioral phenotype of the Cyx-deficient mice (Lush and Holland, 1988). The discovery of mammalian bitter receptors will help understand the biology of bitter perception, from transduction pathways in receptor cells to coding of bitter signals through the afferent sensory pathway.

Results and Discussion

Functional Expression of T2Rs

A difficulty in generating a cell-based reporter system to measure T2R activity is our poor understanding of the native signaling pathway. We therefore expressed T2Rs with Gα15, a G protein α subunit that has been shown to couple a wide range of receptors to phospholipase Cβ (Offermanns and Simon, 1995; Krautwurst et al., 1998). In this system, receptor activation leads to increases in intracellular calcium [Ca^{2+}], which can be monitored at the single cell level using the FURA-2 calcium-indicator dye (Tsien et al., 1985). To test and optimize Gα15 coupling, we used two different GPCRs, a Gαi-coupled μ opioid receptor (Reisine, 1995) and a Gαq-coupled mGlur1 receptor (Masu et al., 1991). Transfection of these receptors into HEK-293 cells produced robust, agonist-selective, and Gα15-dependent Ca^{2+} responses (Figure 1). To assay T2R function, we initially generated four expression constructs containing epil
Figure 1. \(G_{\alpha}15\) Couples Activation of \(\mu\) Opioid Receptor and mGluR1 Receptor to Release of Intracellular Calcium

HEK-293 cells were transiently transfected with the \(G_{\alpha}i\)-coupled \(\mu\) opioid receptor or the \(G_{\alpha}q\)-coupled mGluR1 receptor. Transfected cells containing \(G_{\alpha}15\) were assayed for increases in \([Ca^{2+}]_i\) before (a and b) and after (c and d) the addition of receptor agonists: (c) 10 \(\mu\)M DAMGO and (d) 20 \(\mu\)M trans (-)-1-amino-1,3 cyclopentane dicarboxylic acid (ACPD). Ligand- and receptor-dependent increases in \([Ca^{2+}]_i\) were dependent on \(G_{\alpha}15\) (e and f). Scales indicate \([Ca^{2+}]_i\) (nM) determined from FURA-2 emission ratios.

to peptide-tagged hT2R-3, hT2R-5, hT2R-10, and hT2R-16 (see Adler et al., 2000). However, none of the receptors was efficiently targeted to the plasma membrane.

A number of studies have shown that many GPCRs, in particular sensory receptors, require specific "chaperones" for maturation and targeting through the secretory pathway (Baker et al., 1994; Dwyer et al., 1998). Recently, Krautwurst et al. (1998) generated chimeric receptors consisting of the first 20 amino acids of rhodopsin and various rodent olfactory receptors. These were targeted to the plasma membrane and functioned as odorant receptors in HEK-293 cells. We constructed rhodopsin-T2R chimeras (rho-T2Rs) and determined that the first 39 amino acids of bovine rhodopsin are very effective in targeting T2Rs to the plasma membrane of HEK-293 cells (Figure 2). Similar results were obtained with 11 human and 16 rodent T2Rs (see below). Inclusion of this N-terminal sequence also increased membrane expression of control mGluR1 receptors, and significantly augmented their \(G_{\alpha}15\)-mediated responses (data not shown). To further enhance the level of T2R expression, rho-T2Rs were placed under the control of a strong EF-
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Figure 3. T2R Receptors Are Stimulated by Bitter Compounds
HEK-293 cells were transfected with rho-mT2R-5 (a, d, and g), rho-hT2R-4 (b, e, and h), and rho-mT2R-8 (c, f, and i). Cells expressing mT2R-5 were stimulated using 1.5 mM cycloheximide (d and g) and those expressing hT2R-4 and mT2R-8 with 1.5 mM denatonium (e, f, h, and i). No increase in \([\text{Ca}^{2+}]_i\) was observed in the absence of \(G_{\alpha15}\) (g–i); in contrast robust \(G_{\alpha15}\)-dependent responses were observed in the presence of tastants (d–f); scales indicate \([\text{Ca}^{2+}]_i\) (nM) determined from FURA-2 emission ratios. Line traces (j–l) show the kinetics of the \([\text{Ca}^{2+}]_i\) changes for representative cells from panels (d–f); arrows indicate addition of tastants.

1× promoter and introduced as episomal plasmids into modified HEK-293 cells expressing \(G_{\alpha15}\) (PEAKmod cells; see Experimental Procedures).

We employed two parallel strategies to identify ligands for T2Rs. In one, we chose a random set of human, rat, and mouse T2R receptors, and individually tested them against a collection of 55 bitter and sweet tastants (see Experimental Procedures). We expected functional coupling to meet four criteria: tastant selectivity, temporal specificity, and receptor- and G protein-dependence. In the other, we used data on the genetics of bitter perception in mice to link candidate receptors with specific tastants.

Nearly 30 years ago, it was first reported that various inbred strains of mice differ in their sensitivity to the bitter compound sucrose-octaacetate (Warren and Lewis, 1970). Subsequently, a number of studies demonstrated that this strain difference was due to allelic variation at a single genetic locus (Soa) (Whitney and Harder, 1986; Capeless et al., 1992). These findings were extended to additional loci influencing sensitivity to various bitter tastants, including raffinose undecaacetate (Rua), cycloheximide (Cyx), copper glycinate (Glb), and quinine (Qui) (Lush, 1984, 1986; Lush and Holland, 1988). Genetic mapping experiments showed that the Soa, Rua, Cyx, Qui, and Glb loci are clustered at the distal end of chromosome 6 (Lush and Holland, 1988; Capeless et al., 1992). In the accompanying paper, we show that at least 25 mT2Rs colocalize with this mouse chromosome 6 bitter cluster (Adler et al., 2000). Therefore, we selected
Figure 4. mT2R-5 Is a Taste Receptor for Cycloheximide
(a) HEK-293 cells expressing $\alpha_{15}$ and rho-mT2R-5 were challenged with multiple pulses of 2 $\mu$M cycloheximide (CYX), 3 mM 6-n-propyl thiouracil (PROP), or 5 mM denatonium (DEN); dots and horizontal bars above the traces indicate the time and duration of tastant pulses. Cycloheximide triggers robust receptor activation. This experiment also illustrates desensitization to repeated stimulation or during sustained application of the stimulus. The data shown here were derived from 50 responding cells on a high-density plate. Equivalent results were obtained in HEK-293 cell plated at high (confluent) or low density. (b) Responses to cycloheximide are highly specific and are not observed after addition of buffer (CON) or high concentrations of other tastants. Abbreviations and concentrations used are: cycloheximide, CYX (5 $\mu$M); atropine, ATR (5 mM); brucine, BRU (5 mM); caffeic acid, CAFF (2 mM); denatonium, DEN (5 mM); epicatechin, (-)-EPI (3 mM); phenyl thiocarbamide, PTC (3 mM); 6-n-propyl thiouracil, PROP (10 mM); saccharin, SAC (10 mM); strychnine, STR (5 mM); sucrose octaacetate, SOA (3 mM). Columns represent the mean $\pm$ SE of at least six independent experiments. (c) The mT2R-5 gene from taster (DBA/2-allele) and nontaster (C57BL/6-allele) strains mediate differential $[Ca^{2+}]_{i}$ changes to pulses of cycloheximide. Horizontal bars depict the time and duration of the stimulus. We waited 200 s between stimuli to ensure that cells were not desensitized due to the successive application of cycloheximide. (d) Cycloheximide dose response of mT2R-5. Changes in $[Ca^{2+}]_{i}$ are shown as FURA-2 ($F_{340}/F_{380}$) ratios normalized to the response at 30 $\mu$M cycloheximide; points represent the mean $\pm$ SE of at least six determinations. The nontaster allele shows a marked decrease in cycloheximide sensitivity relative to the taster allele. The data shown in panels (a), (c), and (d) were obtained from measurements of $[Ca^{2+}]_{i}$ from 50 individual responding cells. Because HEK-293 cells plated at high density may form functional gap junctions, our quantitative studies were based on recordings from isolated cells (see Experimental Procedures). Qualitatively similar data was obtained in whole-field recordings.

T2R receptors from this array, constructed the corresponding rho-mT2R chimeras and individually transfected them into HEK-293 cells expressing the promiscuous $\alpha_{15}$ protein. After loading the cells with FURA-2, we assayed for responses to sucrose octaacetate, raffinose undecaacetate, copper glycinate, quinine, and cycloheximide. As controls for transfection efficiencies, we used a CMV-GFP construct, and as a control for $\alpha_{15}$ signaling a set of plates was cotransfected with rho-mGluR1 and assayed for responses to the mGluR1-agonist ACPD.

Cells expressing mT2R-5 specifically responded to cycloheximide (Figure 3). The response occurred in nearly all transfected cells and was receptor- and $\alpha_{15}$-dependent because cells lacking either of these components did not trigger $[Ca^{2+}]_{i}$ changes (Figure 3g), even at 5000-fold higher cycloheximide concentration. As expected for this coupling system, the taster-induced increase in $[Ca^{2+}]_{i}$ was due to release from internal stores, since analogous results were obtained in nominally zero $[Ca^{2+}]_{i}$ data (data not shown). The activation of mT2R-5 by cycloheximide is very selective; this receptor did not respond to any other tastants (Figures 4a and 4b), even at concentrations that far exceeded their biologically relevant range of action (Saroli, 1984; Glendinning, 1994). While cycloheximide is only moderately bitter to humans (Lush and Holland, 1988), it is strongly aversive to rodents with a sensitivity threshold of $\sim$0.25 $\mu$M (Kusano et al., 1971; Lush and Holland, 1988). In our cell-based assay, the concentration of cycloheximide required to induce half-maximal response of mT2R-5 was 0.5 $\mu$M, and the threshold was $\sim$0.2 $\mu$M (Figures
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4c and 4d). Notably, this dose response closely matches the sensitivity range of cycloheximide tasting in mice (Lush and Holland, 1988; see next section).

To examine the kinetics of the cycloheximide response, rho-mT2R-5 transfected cells were placed on a microperfusion chamber and superfused with test solutions under various conditions. Figure 4a shows robust transient responses to micromolar concentrations of cycloheximide that closely follow application of the stimulus (latency < 1 s). As expected, when the tastant was removed, \([\text{Ca}^{2+}_i]\) returned to baseline. A prolonged exposure to cycloheximide (~10 s) resulted in adaptation: a fast increase of \([\text{Ca}^{2+}_i]\) followed by a gradual, but incomplete decline to the resting level (Figure 4a). Similarly, successive applications of cycloheximide led to significantly reduced responses, indicative of desensitization (Lefkowitz et al., 1992). This is likely to occur at the level of receptor, since responses of a cotransfected mGluR1 were not altered during the period of cycloheximide desensitization (data not shown).

Are other T2Rs also activated by bitter compounds? We assayed 11 rhodopsin-tagged human T2R receptors by individually transfecting them into HEK-293 cells expressing G\(_{\alpha 15}\). Each transfected line was tested against a battery of bitter and sweet tastants, including amino acids, peptides, and other natural and synthetic compounds (see Experimental Procedures). We found that the intensely bitter tastant denatonium induced a significant transient increase in \([\text{Ca}^{2+}_i]\) in cells transfected with one of the human candidate taste receptors, hT2R-4, but not in control untransfected cells (Figure 3), or in cells transfected with other hT2Rs. The denatonium response had a strong dose dependency with a threshold of ~100 \(\mu\text{M}\). While this response met the criteria of tastant selectivity, temporal specificity, and receptor- and G\(_{\alpha 15}\)-dependency, the threshold for activation was over two orders of magnitude higher than the human psycho-physical threshold for denatonium (Saroli, 1984). This could be due to poor functioning of this receptor in the heterologous expression system, or perhaps humans express another higher affinity denatonium receptor. Interestingly, hT2R-4 displayed a limited range of promiscuity since it also responded to high concentrations of the bitter tastant 6-n-propyl-2-thiouracil (PROP; Figure 5).

If the responses of hT2R-4 reflect the in vivo function of this receptor, we hypothesized that similarly tuned receptors might be found in other species. The mouse receptor mT2R-8 is a likely ortholog of hT2R-4: they share ~70% identity, while the next closest receptor is only 40% identical; these two genes are contained in homologous genomic intervals (Adler et al., 2000). We generated a rho-mT2R-8 chimeric receptor and examined its response to a wide range of tastants. Indeed, mT2R-8, like its human counterpart, is activated by denatonium and by high concentrations of PROP (Figures 3 and 5). No other tastants elicited significant responses from cells expressing mT2R-8. Because these two receptors share only 70% identity, the similarity in their responses to bitter compounds attests to their role as orthologous bitter taste receptors.

**Figure 5. hT2R-4 and mT2R-8 Respond to Denatonium**

HEK-293 cells expressing G\(_{\alpha 15}\) were transiently transfected with hT2R-4 or mT2R-8 receptors and \([\text{Ca}^{2+}_i]\) was monitored as shown in Figure 3. (a) An increase in \([\text{Ca}^{2+}_i]\) could be induced by stimulation with denatonium but not by various other bitter compounds. Response profiles of (b) hT2R-4 and (c) mT2R-8 to a set of nine out of 55 different bitter and sweet tastants (see Experimental Procedures) are shown. CON refers to control buffer addition, NAR to 2 mM naringin and LYS to 5 mM lysine. Other abbreviations and concentrations are as reported in Figure 4. The mean FURA-2 fluorescence ratio (\(F_{340}/F_{380}\)) before and after ligand addition was obtained from 100 equal-sized areas that included all responding cells. The values represent the mean ± SE of at least six experiments.

Cycloheximide Nontaster Mice Have Mutations in the mT2R-5 Taste Receptor

Our demonstration that mT2R-5 functions as a high-affinity receptor for cycloheximide suggested that the mT2R-5 gene might correspond to the Cyx locus. If this is true, we expected that either the expression profile or sequence of mT2R-5 might differ between strains categorized as Cyx tasters (DBA/2J) and nontasters (C57BL/6J) (Lush and Holland, 1988). In situ hybridizations to tissue sections demonstrated that the expression profile of mT2R-5 is indistinguishable between taster and nontaster strains (Figure 6). To determine the linkage between mT2R-5 and the Cyx locus, we identified polymorphisms in the mT2R-5 gene and determined their distribution in a recombinant inbred panel.
Figure 6. Cycloheximide Taster and Nontaster Strains Express Different Alleles of mT2R5
(a) Predicted transmembrane topology of mT2R-5; amino acid substitutions in the allele from nontaster strains are highlighted in red. The presence of only two alleles at this locus is not unexpected because the strains that share the same polymorphisms were derived from a common founder (Beck et al., 2000).

In situ hybridization showing expression of mT2R-5 in subsets of cells in the circumvallate papilla of (b) a cycloheximide taster strain (DBA/2) and (c) a nontaster strain (C57BL/6); no strain specific differences in expression pattern were detected in taste buds from other regions of the oral cavity.

From a C57BL/6J (nontaster) × DBA/2J (taster) cross. We found tight linkage between mT2R-5 and the Cyx locus but not perfect concordance in their strain distribution pattern (data not shown). We believe that this is due to the reported ambiguity in the original designation of the cycloheximide phenotype of the recombinant inbred panel progeny and parental lines (Lush and Holland, 1988). We therefore isolated the mT2R-5 gene from several additional well-characterized cycloheximide taster (CBA/Ca, BALB/c, C3H/He) and nontaster (129/Sv) strains and determined their nucleotide sequences. Indeed, as would be expected if mT2R-5 functions as the cycloheximide receptor in these strains, all the tasters share the same mT2R-5 allele as DBA/2J, while the nontasters share the C57BL/6 allele, which carries missense mutations (Figure 6), including three nonconservative amino acid substitutions (T44I, G155D and L294R).

If the mT2R-5 C57BL/6 allele is responsible for the taste deficiency of Cyx mutants, its cycloheximide dose response might recapitulate the sensitivity shift seen in Cyx mutant strains. Two-bottle preference tests have shown that Cyx taster strains avoid cycloheximide with a threshold of 0.25 μM (Lush and Holland, 1988), while nontasters avoid it strongly at 8 μM, but strongly avoid cycloheximide at 1 μM). We constructed a rhomT2R-5 fusion with the mT2R-5 gene from a nontaster strain and compared its dose response with that of the receptor from taster strains. To prevent bias due to differences in receptor numbers in the heterologous cells, we measured surface expression and assayed mT2R-5 function from the same transfection experiments (see Experimental Procedures). Remarkably, mT2R-5 from the nontaster strains displays a shift in cycloheximide sensitivity (Figure 4d) that resembles the sensitivity of these strains to this bitter tastant. Taken together, these results validate mT2R-5 as a cycloheximide receptor and strongly suggest that mT2R-5 corresponds to the Cyx locus. Formal proof that mT2R-5 is Cyx will require the knockout of this gene in taster strains, or the phenotypic rescue of nontaster animals with an mT2R-5 transgene.

T2Rs Couple to Gustducin
In the accompanying paper (Adler et al., 2000), we demonstrated that T2Rs are coexpressed with gustducin, suggesting that T2Rs may activate this G protein in response to bitter tastants. To investigate the selectivity of T2R-G protein coupling, we chose to study mT2R-5 because its activation by cycloheximide recapitulates mouse taste responses. Because of the need to assay several G proteins and the lack of a cell-based gustducin assay, we used a cell-free system. Rho-tagged mT2R-5
Concluding Remarks
To date, many putative taste receptors have been reported (Abe et al., 1993; Matsuoka et al., 1993; Ming et al., 1998; Hoon et al., 1999; Chaudhari et al., 2000). However, none have satisfied the requirements of rigorous biological verification: (1) demonstrated tissue and cell-specific expression, (2) functional validation, and (3) genetic corroboration. The T2R receptors presented in this and the accompanying paper were examined for all three criteria. First, we showed that T2Rs are selectively expressed in subsets of taste receptor cells of the tongue and palate epithelium. Second, three T2Rs (mT2R-5, hT2R-8, and mT2R-4) functioned as receptors for bitter tastants in heterologous cells. Third, polymorphisms in the mT2R-5 receptor were found to be associated with changes in bitter taste sensitivity to cycloheximide, both in vivo and in vitro. Thus, mT2R-5 is a strong candidate for Cyx. Furthermore, mT2R-5 selectively couples to gustducin, which has been implicated biochemically and genetically in taste transduction (Wong et al., 1996; Ming et al., 1998). Together, these results demonstrate that the T2R gene family contains functionally defined bitter taste receptors.

At present, we do not know what fraction of the available human and rodent receptors function in bitter transduction. However, our demonstration that all T2R-positive taste cells express multiple receptors suggests that T2R receptors may function in a similar taste modality. This is consistent with the observation that mammals can recognize a large number of bitter compounds, but do not discriminate between them (McBurney and Gent, 1979). Indeed, the two mouse receptors presented in this study (mT2R-5 and mT2R-8) respond to different bitter tastants and are expressed in combination with a number of other T2Rs in overlapping taste receptor cells (data not shown). Alternatively, if T2Rs respond to more than one modality, for example bitter and sweet then these cells would have to functionally segregate T2R receptors so as to maintain specificity and selectivity of signaling (Tsunoda et al., 1998).

A number of studies have shown that the oral cavity displays regional differences in sensitivity to the various taste modalities (Frank et al., 1983; Nejad, 1986; Frank, 1991). Our demonstration that T2Rs are expressed in all taste buds of circumvallate, foliate, and palate epithelium (data not shown) indicates that if there are significant differences in bitter sensitivity between these three regions, they may reflect events distal to tastant recognition.

The discovery of bitter taste receptors makes it possible to experimentally approach and elucidate critical aspects of the logic of bitter coding. For instance, it should be possible to genetically mark mT2R-expressing cells and examine their physiology and connectivity patterns. Similarly, it will be possible to knock out selective subsets of mT2R receptors and study the impact on bitter taste perception.

Taste receptor cells turn over throughout life (Beidler and Smallman, 1965). Therefore, synapses need to be continuously reestablished. It will be interesting to determine how this is achieved and whether nerve terminals provide any instructive signals for the expression of T2R receptors. The observation that taste buds degenerate when denervated and regenerate when the gustatory
epithelium is reinnervated provides a tractable experimental paradigm to address this question. Finally, the identification of human bitter receptors makes it possible to use high-throughput screening strategies to identify bitter antagonists, and in a small but significant way, eliminate bitterness from the world.

Experimental Procedures

**Generation, Expression, and Immunostaining of Chimeric Receptors**

A bridge overlap PCR extension technique was used to generate rho-T2R chimeras, which contain the first 39 amino acids of bovine rhodopsin in frame with human and rodent T2R coding sequences (Mehta and Singh, 1999). The rhodopsin segment was amplified from a bovine cDNA clone kindly provided by Dr. J. Nathans. All receptors were cloned into a pEAK10 mammalian expression vector (Edge Biosystems, MD). The rho-mGlur1 chimeras were constructed using a similar strategy.

**Modified HEX-293 cells (PEAK™ cells; Edge BioSystems, MD) were grown and maintained at 37°C in UltraCulture media (Bio Whitaker) supplemented with 5% fetal bovine serum, 100 μg/ml Gentamycin sulphate (Fisher), 1 μg/ml Amphotericin B, and 2 mM Glutamax I (LifeTechnologies). For transfection, cells were seeded onto matrigel-coated 24-well culture plates or 35 mm recording chambers. After 24 hr at 37°C, cells were washed in OptiMEM medium (LifeTechnologies) and transfected using LipofectAMINE reagent (LifeTechnologies).**

Transfection efficiencies were estimated by cotransfection of a GFP reporter plasmid and were typically >70%. Immunofluorescence staining and activity assays were performed 36-48 hr after transfection.

**For immunostaining transfected cells were grown on coated glass coverslips, fixed for 20 min in ice-cold 2% paraformaldehyde, blocked with 1% BSA, and incubated with the B-630 anti-rhodopsin antibody. Surface receptors were detected using rho-mT2R-5 was assessed by Western blot using mAb B6±30 and an anti-rhodopsin antibody. Surface receptors were detected using rho-mT2R-5 was assessed by Western blot using mAb B6±30 and an anti-rhodopsin antibody.**

**In Vitro Coupling of mT2R-5 to Gustducin**

Infectious Bacmid containing rhodopsin-tagged mT2R-5 (DBA/2-j) was produced using the Bac-to-Bac system (LifeTechnologies, MD). Insect larval cells were infected for 60 hr with recombinant Bacmid and membranes were prepared as described previously (Ryba and Tirindelli, 1995). Peripheral proteins were removed by treatment with 8 M urea and membranes were resuspended in 10 mM HEPES pH 7.5, 1 mM EDTA, and 1 mM DTT. The expression of rh-mT2R-5 was assessed by Western blot using mAb B6-30 and quantitated by comparison with known amounts of rhodopsin. Approximately 300 pmol of rho-mT2R-5 could be obtained from 2 x 10^7 infected cells. Gustducin and G_{i2y} heterodimers were isolated as described previously (Hoon et al., 1995; Ryba and Tirindelli, 1995). Recombinant G_{os}, G_{oq}, and G_{ol} and bovine brain G_{o} were generously provided by Dr. Elliott Ross. Receptor-catalyzed exchange of GDP for GTP-γ-S on gustducin and other G protein γ subunits was measured in the presence of 10 mM rho-mT2R-5, 100 μM GDP, and 20 μM G_{i2y} (Hoon et al., 1995). All measurements were made at 15 min time points and reflect the initial rate of GTP-γ-S binding.

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