Multiple PLC β signaling pathways in taste receptor cells contribute to the detection of bitter, sweet and umami stimuli

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Summary: This study identifies a new role for PLC β 3 in the transduction of bitter, sweet and umami taste stimuli and demonstrates that multiple taste cell populations respond to these stimuli.

1 ABSTRACT

Taste receptor cells use multiple signaling pathways to detect chemicals. While 2 salty and sour stimuli bind ion channels, bitter, sweet and umami stimuli activate a 3 GPCR/PLCβ2/IP₃R3 signaling pathway. Current thinking is that all bitter, sweet and 4 umami stimuli are detected by Type II cells via a PLCβ2/IP₃R3 pathway. Using live cell 5 calcium (Ca²⁺) imaging on isolated taste receptor cells from mice lacking IP₃R3 in Type 6 7 Il cells, we find many taste cells still robustly respond to bitter, sweet and umami stimuli, including a subset of Type III taste cells. Immunohistochemical analysis reveals that 8 PLC_{B3} is not expressed in Type II cells but is present in other taste cell populations, 9 including a subset of Type III cells. Loss of either IP₃R3 or PLCβ3 in taste receptor cells 10 cause comparable reductions in c-Fos labeling in the nucleus of the solitary tract 11 following oral infusion of quinine. Short term behavioral assays also demonstrate that 12 loss of either IP₃R3 from Type II cells or loss of PLCβ3 causes similar taste deficits, 13 indicating that both molecules are required for normal taste transduction. Our data 14 identify a critical, and previously unknown, role for PLC_{B3} in taste transduction and 15 demonstrate that input from multiple PLC^β pathways/taste receptor cell populations are 16 required to transduce bitter, sweet and umami stimuli. 17

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24 SIGNIFICANCE STATEMENT

It is currently thought that the transduction of bitter, sweet and umami stimuli in 25 taste cells depends solely on GPCR/PLCB2/IP₃R3 signaling in Type II taste receptor 26 cells. However, in the absence of either PLCB2 or IP₃R3, mice have reduced, but not 27 abolished, ability to detect these stimuli, suggesting that other taste cells also contribute 28 to taste transduction. When IP₃R3 is absent, Type II cells do not generate a Ca²⁺ signal 29 to these taste stimuli; however, other taste receptor cells respond to bitter, sweet and 30 umami stimuli, including a subset of Type III cells. We identified a PLC_{β3} pathway that 31 is expressed in non-Type II taste cells, including a subset of Type III cells. Loss of 32 PLCB3 significantly reduced taste cell responses, cFos-labeling in the nucleus of the 33 solitary tract, and caused significant behavioral deficits. We conclude that PLCB3 has 34 an important role in taste transduction. 35

Chemicals in the oral cavity are detected by taste receptor cells which are 37 grouped together in taste buds found in epithelial specializations in the tongue called 38 papillae. There are three main papillae: Circumvallate (CV), foliate (Fol), and fungiform 39 (Fun) which differ in their localization on the tongue and innervation by the cranial 40 nerves. Despite these differences, taste papillae all house taste buds that are 41 42 comprised of distinct cell types that vary in their signaling functions within the bud (1). The current view of taste transduction is that taste buds are comprised of Type I, 43 Type II and Type III taste receptor cells (2). Type I cells are thought to primarily function 44 as support cells and share some characteristics of glia cells (3), while Type II cells 45 detect bitter, sweet or umami stimuli through the activation of specific GPCRs for each 46 of these taste qualities. The subsequent transduction of these taste qualities within the 47 cells are due to a single signaling pathway that is comprised of phospholipase C β 2 48 49 (PLCβ2) that activates the inositol 1,4,5-trisphosphate receptor type 3 (IP₃R3) on the endoplasmic reticulum to cause Ca^{2+} release (4-6). This Ca^{2+} release activates the 50 51 transient receptor potential cation channel subfamily M members 4 and 5 (TRPM4 and TRPM5) which depolarize the cell sufficiently to activate the release of ATP through the 52 calcium homeostasis modulator 1 (CALHM1) channel (5, 7-11). The expression of 53 PLCβ2, IP₃R3 and TRPM5 is restricted to Type II taste cells within the bud and these 54 cells lack both voltage-gated calcium channels (VGCCs) and conventional chemical 55 synapses (model shown in Figure 1A)(12-16). Type III cells detect sour and salt stimuli 56 through ionotropic receptors that depolarize the cell to activate VGCCs and cause 57 vesicular neurotransmitter release (17-23) (see Figure 1A). It is currently thought that 58 59 Type III cells do not respond to bitter, sweet or umami stimuli.

While this is the widely accepted view of signal transduction in peripheral taste 60 cells, multiple genetic studies have reported that the deletion of individual signaling 61 proteins in Type II cells, including PLCβ2, IP₃R3 and TRPM5, does not abolish the 62 ability to detect all bitter sweet and umami stimuli (24-28). Similarly, the loss of Type II 63 64 cells leads to severely reduced, but not abolished, responses to bitter, sweet and 65 umami stimuli (29). These studies support the idea that additional signaling pathways are present in taste cells, but to date, few studies have focused on identifying these 66 67 alternatives (30, 31). We previously reported that some taste cells generate Ca²⁺ 68 responses to a bitter stimulus and cell depolarization, suggesting that taste cells with VGCCs, presumably Type III cells, respond to at least one bitter compound (32). A 69 separate report also found some taste cells with VGCCs can respond to bitter stimuli 70 71 (19). These studies led us to hypothesize that in addition to Type II cells, a separate 72 population of taste cells may contribute to the detection of bitter, sweet and umami stimuli. To test this hypothesis, we used live cell imaging in mice that lack IP₃R3 and 73 found that many taste cells still responded to bitter, sweet and umami stimuli, including 74 75 a subset of Type III cells. These cells responded to taste stimuli through a distinct PLC β which we identified as PLC β 3. We conclude that multiple populations of taste 76 cells use distinct signaling pathways to respond to bitter, sweet and umami stimuli. 77

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79 **RESULTS**

Bitter, sweet and umami taste evoked responses are still present in IP₃R3-KO
 mice

82	IP_3R3 is part of the canonical signaling pathway in Type II cells that respond to
83	bitter, sweet and umami stimuli and its expression is localized in Type II cells (4, 12, 16,
84	33). One study using single cell PCR reported mRNA levels for IP ₃ R3 in both Type II
85	cells and some SNAP25+ cells (a Type III marker) but did not evaluate protein
86	expression for these compounds (34). In contrast, another study using an IP $_3$ R3-GFP
87	mouse reported that GFP+ cells did not functionally express VGCCs, a physiological
88	marker for Type III cells (32). While these conflicting reports suggest that it is possible
89	that IP ₃ R3 expression is not restricted to Type II cells, for the purposes of our current
90	study, we are presuming that IP $_3$ R3 is only expressed in Type II cells.
91	To evaluate the taste evoked signaling in taste receptor cells that lack the ability
92	to release Ca ²⁺ via IP ₃ R3, we used a transgenic mouse in which GFP replaces the
93	coding region of IP ₃ R3. In these mice, GFP labels the cells that should express IP ₃ R3
94	but no longer do so. Initial immunohistochemical analyses found that taste cells from the
95	CV papillae of wild type mice were successfully labeled with anti-IP $_3$ R3 antibody (n=3;
96	Figure S1A), while their KO littermates lacked IP $_3$ R3 labeling but instead have GFP
97	expression (n=6; Figure S1B). Furthermore, anti-IP $_3$ R3 labels the GFP expressing cells
98	in IP ₃ R3-heterzygous mice (n=4; Figure S1C); confirming that GFP expression identifies
99	the taste cells that normally express IP $_3$ R3. To further characterize this mouse, we
100	evaluated the expression of other proteins that are part of this established signaling
101	pathway in Type II cells. The GFP expression in the KO mice co-localized with both

102 PLC β 2 (n=4; Figure S1D) and gustducin (n=3; Figure S1E) labeling. As expected,

gustducin expression was restricted to the IP3R3-GFP-KO cells but not all GFP

expressing cells had gustducin while there was complete overlap between PLCβ2

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labeling and GFP. Co-localization analyses are reported in Figures S1F. Thus the
 other components of the signaling pathway normally found in Type II cells are intact but
 these cells no longer have the capacity to release Ca²⁺ in response to bitter, sweet or
 umami stimuli.

We evaluated this IP₃R3-GFP-KO mouse using Fura2-AM to measure the taste-109 evoked Ca²⁺ responses in taste receptor cells. All measurements were made in 110 isolated taste cells that were no longer associated with the taste bud to ensure that the 111 taste-evoked Ca²⁺ responses were due solely to the activity of the individual taste cells 112 113 and not cell-to-cell communication (example in Figure 1B, C). We applied multiple bitter, sweet and umami stimuli as well as 50mM KCI to isolated cells from CV, Fol or 114 Fun papillae of the wild type and IP₃R3-KO mice. Since our goal was to identify all (or 115 116 most) responsive taste cells, we used the stimulus concentration that generated a maximal Ca²⁺ signal. These concentrations were based on control experiments we 117 used to identify the lowest stimulus concentration that generated the maximal Ca²⁺ 118 signal (examples shown in Figure S2, A-C). This increased our likelihood of identifying 119 responsive taste cells but did not generate any non-specific responses (Figure S2, D, 120 E). As shown in Figure S2C, a GFP+ cell did not respond to any concentration of 121 stimulus applied. Thus, the taste cells respond to a wide range of stimulus 122 123 concentrations and the reported responses are not due to a concentration or osmotic 124 effect.

We reasoned that if the Type II cells were solely responsible for all bitter, sweet and umami taste transduction, then the loss of IP_3R3 in Type II cells would abolish all Ca^{2+} responses to bitter, sweet and umami stimuli. In the IP_3R3 -KO mice, GFP-

expressing cells lack IP₃R3 and did not respond to any taste stimuli tested (n=80, Figure 128 2B, S2C); however, a large number of non-GFP expressing taste cells responded to the 129 stimuli. Initially, the response frequencies (total number of responsive cells/total 130 number of cells stimulated) were measured for all taste cells from the wild type (WT) 131 and IP₃R3-KO mice, including both GFP+ and GFP- cells (Figure 2A). The table in 132 133 Figure 2A reports the taste response frequencies in wild type (WT) and IP₃R3-KO mice with the GFP+ cells included in the analysis. Not all cells were exposed to all stimuli 134 and some cells responded to multiple stimuli. These experimental conditions preclude 135 136 the data from being additive, so we compared the frequency of responses for a given stimulus between the wild type and IP₃R3-KO mice. These data demonstrate that the 137 overall responsiveness to bitter, sweet, and/or umami stimuli was significantly reduced 138 (p<0.0001) in the KO mice compared to WT cells. The likelihood that taste cells from 139 CV would respond to taste stimuli was significantly reduced (p=0.0003). The 140 141 responsiveness of the taste cells from the Fol and Fun papillae was not significantly affected by the loss of IP₃R3, even though the GFP+ cells were included in this 142 analysis. 143

To further characterize the response profiles of the non-Type II taste cells, we analyzed the non-GFP expressing cells and plotted the response frequencies for each papillae and stimulus (Figure 2B-E). The GFP+ cells were not included in these response analyses and descriptions of the different taste cell populations are shown in Supplementary Table 1. Figure 2B compares the overall percent responsiveness of WT and KO mice for each papillae type (without the GFP+ cells) while the graphs in Figures 2C-E show the percentage of responsive cells in each papillae type for each specific

151	stimulus applied. The overall responsiveness of taste cells was only significantly
152	reduced in the CV papillae when IP ₃ R3 was absent (p<0.01). Analysis of the individual
153	stimuli for each papillae type (Figures 2C-E) identified significant reductions in the
154	responsiveness to the sweet stimuli tested in the KO mice. Individual cell numbers are
155	provided in Figure S3. Therefore, removing a key protein in the canonical
156	GPCR/PLC β 2/IP ₃ R3 signaling pathway in Type II cells does not eliminate the ability of
157	all taste cells to generate Ca ²⁺ signals when stimulated by bitter, sweet or umami
158	stimuli. Based on these data, we hypothesize there are additional signaling pathways
159	that respond to these stimuli.
160	Bitter, sweet and umami stimuli are dependent on a PLC eta signaling pathway in
161	IP₃R3-KO mice
162	To identify the type of signaling pathway that generates the taste responses in
162 163	To identify the type of signaling pathway that generates the taste responses in the non-GFP cells in the IP ₃ R3-KO mice, we analyzed bitter (5mM Den), sweet (20mM
162 163 164	To identify the type of signaling pathway that generates the taste responses in the non-GFP cells in the IP ₃ R3-KO mice, we analyzed bitter (5mM Den), sweet (20mM sucrose) and umami (10mM MPG) stimuli and found they generated signals in Ca ²⁺ -free
162 163 164 165	To identify the type of signaling pathway that generates the taste responses in the non-GFP cells in the IP ₃ R3-KO mice, we analyzed bitter (5mM Den), sweet (20mM sucrose) and umami (10mM MPG) stimuli and found they generated signals in Ca ²⁺ -free Tyrode's, indicating these signals depend on Ca ²⁺ release from internal stores (Figure
162 163 164 165 166	To identify the type of signaling pathway that generates the taste responses in the non-GFP cells in the IP ₃ R3-KO mice, we analyzed bitter (5mM Den), sweet (20mM sucrose) and umami (10mM MPG) stimuli and found they generated signals in Ca ²⁺ -free Tyrode's, indicating these signals depend on Ca ²⁺ release from internal stores (Figure 3A,D,G). We confirmed these results by applying thapsigargin to disrupt the internal
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162 163 164 165 166 167 168 169 170	To identify the type of signaling pathway that generates the taste responses in the non-GFP cells in the IP ₃ R3-KO mice, we analyzed bitter (5mM Den), sweet (20mM sucrose) and umami (10mM MPG) stimuli and found they generated signals in Ca ²⁺ -free Tyrode's, indicating these signals depend on Ca ²⁺ release from internal stores (Figure 3A,D,G). We confirmed these results by applying thapsigargin to disrupt the internal Ca ²⁺ stores which abolished the taste evoked Ca ²⁺ signals (Figure 3B,E,H). These responses were also eliminated when the general PLC blocker U73122 was applied (Figure 3C,F,I), indicating that these taste evoked Ca ²⁺ responses in the IP ₃ R3-KO mice are due to the activation of another PLC signaling pathway. Individual data points for
162 163 164 165 166 167 168 169 170 171	To identify the type of signaling pathway that generates the taste responses in the non-GFP cells in the IP ₃ R3-KO mice, we analyzed bitter (5mM Den), sweet (20mM sucrose) and umami (10mM MPG) stimuli and found they generated signals in Ca ²⁺ -free Tyrode's, indicating these signals depend on Ca ²⁺ release from internal stores (Figure 3A,D,G). We confirmed these results by applying thapsigargin to disrupt the internal Ca ²⁺ stores which abolished the taste evoked Ca ²⁺ signals (Figure 3B,E,H). These responses were also eliminated when the general PLC blocker U73122 was applied (Figure 3C,F,I), indicating that these taste evoked Ca ²⁺ responses in the IP ₃ R3-KO mice are due to the activation of another PLC signaling pathway. Individual data points for each experiment are shown in Figure S4.

PLCβ3 expression in peripheral taste cells

To identify other PLC β isoforms expressed in taste receptor cells, we analyzed 173 174 previously published RNAseq data (35). We found that PLC β 2 and PLC β 3 were 175 expressed at comparable levels and were the predominant PLC β isoforms present in taste cells. Therefore, we investigated the possibility that PLC₃3 is expressed in 176 peripheral taste cells. Immunohistochemical experiments in the IP₃R3-KO mouse 177 revealed strong PLCβ3 labeling that was separate from the Type II cells (identified with 178 GFP expression) in the CV, Fol and Fun papillae (Figure 4A-C). Colocalization analysis 179 180 is shown in Figure 4D. These data were confirmed using both the TRPM5-GFP mouse 181 as well as co-labeling with PLC_β2 in C57BL/6 mice (Type II cell markers, Figure S5A, B, colocalization analysis shown in S5D). Co-labeling with the Type I taste marker 182 NTPDase2 found that PLC β 3 is also not expressed in Type I cells (Figure S5C). We 183 used qPCR to measure the relative amounts of mRNA for PLC β 3 and PLC β 2 in the 184 different papillae (Figure S5E,F) and found that both PLC_B2 and PLC_B3 are expressed 185 in CV, Fol and Fun papillae. We conclude that both PLC β 2 and PLC β 3 are highly 186 expressed in peripheral taste cells but that PLC β 3 is not expressed in either Type I or 187 Type II taste cells. Our next goal was to identify the cell population that expresses the 188 PLC_B3 signaling pathway and responds to bitter, sweet and umami stimuli. 189

190 Some Type III cells respond to bitter, sweet and umami stimuli

The only other identified taste receptor cell type is the Type III cell. Type III cells are the only taste cell population that express VGCCs (36) and can be functionally identified in live cell imaging by the ability to respond to cell depolarization with a Ca²⁺ influx through the opening of VGCCs (19, 37). Earlier studies have reported that Type III cells are sour sensitive and that a subset of the sour-sensitive Type III cells is also salt

sensitive (17, 19, 38). To functionally identify Type III cells, we applied 50mM KCI to 196 cause a cell depolarization which generated a Ca²⁺ signal if the cell expresses VGCCs. 197 Therefore, if a cell responded to 50mM KCI with a Ca²⁺ signal, we concluded that it 198 expresses VGCCs and identified it as a Type III cell. We found that some Type III taste 199 cells in the IP₃R3-KO mouse responded to bitter, sweet and umami stimuli (Figure 5A-200 201 C). To confirm that the taste cells that responded to 50mM KCl and bitter, sweet or umami stimuli were Type III cells, we tested a subset of cells with sour (citric acid) and 202 salt (NaCI) stimuli (Figure 5D-F). All of the taste cells that responded to 50mM KCI 203 responded to citric acid with a Ca²⁺ signal (CA) while some of these cells also 204 responded to bitter, sweet and/or umami stimuli (Figure 5D, F). Interestingly, a 205 separate subset of 50mM KCI/CA cells responded to NaCI with a Ca²⁺ signal but were 206 207 not sensitive to the bitter, sweet, umami stimuli (Figure 5E, F). This indicates that the sour/salt sensitive Type III cells are separate from the sour sensitive Type III cells that 208 respond to bitter, sweet and/or umami stimuli with a Ca²⁺ signal. Thus a subset of Type 209 III cells are dual responsive (DR) because they express VGCCs and are sour sensitive 210 but also respond to bitter, sweet and/or umami stimuli. 211

To determine if the loss of IP₃R3 in Type II cells affected the activity of DR cells, we compared the percentage of cells that responded to any stimulus (bitter, sweet, umami or 50mM KCI) in wild type and IP₃R3-KO mice. These data include all taste cells from all three papillae taken from Figure 2A. As shown before, the overall responsiveness to taste stimuli in the IP₃R3-KO mice was significantly reduced compared to wild type (Figure 5G); however, the percentage of DR cells in the IP₃R3-KO was not different from wild type mice (Figure 5G). Therefore, the ability of the DR cells to respond to taste stimuli does not depend on the presence of IP₃R3 in Type II
cells (Figure 5G, H).

We also performed control experiments using the GAD67-GFP mice as a cellular marker to identify Type III taste cells (39) and found that both GAD67-GFP positive and negative taste cells responded to taste stimuli + 50mM KCI with Ca²⁺ signals (Figure S6A-B). DR cells were also identified in C57BL/6 mice (Figure S6C). A recent study reported that some Type III cells from C57BL/6 mice responded to bitter stimuli (19) in agreement with our findings. Thus, DR Type III taste cells are present in taste cells in multiple mouse lines.

228 PLCβ3 contributes to taste evoked signaling in a subset of Type III cells

229 So far our data have identified that 1) there is a population of taste cells that do 230 not depend on IP₃R3 (in Type II cells) to respond to bitter, sweet and umami, 2) PLC β 3 is expressed in taste cells, and 3) some Type III cells are DR and can respond to bitter. 231 sweet and umami stimuli. We hypothesized that PLC₃3 is involved in the signaling of 232 the DR cells. We next performed live cell imaging on isolated cells from CV, Fol, and 233 Fun papillae of PLC β 3-KO and wild type mice to evaluate how the loss of PLC β 3 affects 234 taste cell responsiveness, in particular the responsiveness of the DR cells. Loss of 235 PLCB3 resulted in significant reductions in the overall responsiveness of the PLCB3-KO 236 Fol and Fun cells but not the CV cells (Figure 6A,B). Knocking out PLCB3 had a two-237 fold effect, it reduced the number of responses to bitter, sweet or umami stimuli and it 238 significantly reduced the number of DR taste cells in CV, Fol and Fun taste cells (Figure 239 6C-E). Control experiments found no change in the ability of Type III cells to respond to 240 241 sour or salt stimuli (Figure S7A-C). Type II cells were also still functional in PLCβ3-KO

242	mice (Figure S7D-F). Individual cell numbers are shown in Figure S8. These data
243	indicate that PLC β 3 is contributing to the activity of the DR cells.
244	We next determined if PLC eta 3 is expressed in Type III cells using SNAP-25 which
245	is a synaptic protein that is a cellular marker for Type III taste cells (16, 40).
246	Immunohistochemical analysis found some co-expression of PLC eta 3 and SNAP25 in
247	wild type mice (Figure S9A,B,E). There was no loss of SNAP-25 expression in the
248	PLC β 3-KO mouse compared to WT (Figure S9C,D). Thus, PLC β 3 is expressed in some
249	Type III cells and the loss of this protein did not affect the expression of SNAP25 in
250	Type III cells.
251	The significant reduction in the number of taste evoked Ca ²⁺ responses in the DR
252	cells from PLC β 3-KO mice suggests that PLC β 3 is responsible for transducing most of
253	the taste-evoked responses in these cells. To determine if the loss of both IP_3R3 and
254	PLC β 3 would completely abolish the ability to respond to bitter, sweet and umami
255	stimuli, we attempted to generate a double knock out mouse by breeding the IP ₃ R3-KO
256	and PLC β 3-KO mice but the mice were not viable.
257	Loss of PLC β 3 causes deficits in the taste-evoked activity in the nucleus of the
258	solitary tract.
259	To determine if the loss of IP ₃ R3 or PLC β 3 in taste cells affected the taste signal

that is sent to the brain, we measured the neural activity in the nucleus of the solitary tract (NTS) using c-Fos labeling as a marker for activated neurons. This approach is commonly used to identify recently activated neurons, including neurons in the taste pathway (41-45). The NTS receives input from the gustatory neurons and is the first synaptic relay from the peripheral taste system. Mice (n=3 for each condition) were

orally infused with guinine (5mM) or water for 30 min. The level of c-Fos labeling for 265 both IP₃R3-KO and PLCβ3-KO mice were compared to WT and water controls. Loss of 266 either of these proteins resulted in a significant reduction in c-Fos activity to a level that 267 268 was comparable to water (Figure 7). Control experiments demonstrate that neither IP₃R3 nor PLC β 3 are expressed in this region of the NTS (Figure S10). 269 270 Loss of PLCB3 significantly affects taste-driven licking for bitter, sweet and umami 271 272 We then performed brief-access licking experiments to determine if the loss of taste-evoked signals in the PLC₃-KO mice correlated with loss of taste sensitivity. 273 Measurements were also made with IP₃R3-KO mice for comparison. Since the PLC_β3-274 KO mouse has a mixed genetic background while the IP₃R3-KO mice are in a C57BL/6 275 background, wild type littermates for both mouse lines were used as controls (Figure 8). 276 The wild type mice both demonstrated concentration dependent decreases in licking to 277 denatonium while PLC_β3-KO and IP₃R3-KO mice treated the solution similarly to water 278 279 except for the highest concentration (20mM Den, Figure 8A). Wild type mice treated MSG (+ 10µM amiloride) solutions as more palatable than water up to 400mM and then 280 decreased licking at the highest concentrations (600 and 800mM, Figure 8B). Both of 281 the KO mice treated MSG like water. In the brief access test for the artificial sweetener, 282 Acesulfame K (AceK), wild type mice increased licking up to 2 mM and then decreased 283

dependent change in licking (Figure 8C). Because sucrose is hedonically positive under
wild type conditions, the lick score is calculated as licks to the stimulus minus licks to

licking at higher concentrations. Neither of the KO mice had a comparable concentration

water, therefore a score close to zero resembles water. Neither the PLC β 3-KO nor

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IP₃R3-KO mice showed a concentration-dependent effect on licking to sucrose until 288 they were presented with the highest concentration. In contrast, both sets of wild type 289 290 mice showed strong concentration dependent increases in licking to sucrose (Figure 8D). For the salt stimulus (NaCl), there were no significant differences in the licking 291 behavior for either of the KO mice compared to wild type mice, indicating that the ability 292 to respond to salt is not affected by loss of either PLCβ3 or IP₃R3 (Figure 8E). Thus 293 loss of either IP₃R3 or PLCβ3 caused comparable, and almost complete, loss of 294 preference for the bitter, sweet and umami stimuli tested but not salt taste. 295 Response profiles for the dual-responsive taste cells 296

Finally, we analyzed how DR cells respond to taste stimuli. We evaluated a 297 298 subset of DR cells that were stimulated with at least one bitter, one sweet and one 299 umami stimulus and responded to at least one of these stimuli. If the DR cells were not stimulated with all three types of taste qualities, they were excluded from this analysis. 300 We created response profiles for bitter, sweet and umami in wild type and IP₃R3-KO 301 mice (Figure S11A-C) and found that the DR taste cells were most likely to be broadly 302 tuned (respond to all three types of stimuli). Few, if any, of the DR cells responded to a 303 single taste quality (in addition to 50mM KCI). We found that the loss of IP₃R3 did not 304 influence these response profiles. 305

306 Three populations of taste cells respond to bitter, sweet and umami stimuli

We repeated the same analysis on taste cells from wild type and IP₃R3-KO mice that were stimulated with bitter, sweet, umami and 50mM KCl but *did not respond to* 50mM KCl (Figure S11D-F). Therefore, wild type taste cells analyzed included all the Type II taste cells as well as any other taste cells that responded to bitter, sweet and/or

umami stimuli but lack VGCCs. We only analyzed taste cells that responded to at least
one of these stimuli and compiled a response profile for these taste cells from wild type
and IP₃R3-KO mice. These taste cells were much less likely to be broadly tuned in both
the WT and KO taste cell populations.

While the taste-only cells from wild type mice express IP₃R3 and therefore should 315 316 have Type II cells contributing to these response profiles, the analysis of the IP₃R3-KO mice was focused on taste cells that were not functional Type II taste cells (due to the 317 absence of IP₃R3). These cells also did not respond to 50mM KCl with a Ca²⁺ signal 318 319 and are not Type III cells. Therefore, this taste cell population lacks functional Type II cells and VGCCs but responds to bitter, sweet and/or umami stimuli. These functional 320 data suggest that there are at least three populations of taste cells that can respond to 321 bitter, sweet and umami stimuli: 1) Type II cells (identified as cells using IP₃R3), 2) a 322 subset of Type III cells (which respond to 50mM KCl with a Ca²⁺ signal) and 3) another 323 population of taste cells that are not Type II or Type III cells (neither reliant on IP₃R3 nor 324 responsive to 50mM KCI). Future studies are needed to identify and characterize this 325 taste cell population. 326

327 **DISCUSSION**

Our data establish that PLC β 3 signaling in taste receptor cells is important in the detection of bitter, sweet and umami taste stimuli. The presence of a second signaling pathway that detects these stimuli resolves a long standing inconsistency in the taste field. One study concluded that the PLC β 2 signaling pathway in Type II cells is entirely responsible for the detection of bitter, sweet and umami stimuli (5). A second study focused on understanding the role of Skn-1a in taste cell formation reported that the

334 loss of Skn-1a caused a significant remodeling of the taste bud. Using in situ hybridization, the authors did not detect the canonical signaling molecules normally 335 found in Type II cells but did identify an increase in the Type III cell marker expression. 336 They also reported significant behavioral and gustatory nerve defects for bitter, sweet 337 and umami stimuli in the absence of Skn-1a, indicating that Type II cells are required to 338 339 detect these stimuli (29). However, other studies reported that knocking out any component of the GPCR/PLCβ2/IP₃R3 pathway resulted in reduced, but not abolished, 340 signaling for these stimuli (24-28, 46-49). There are deficits in taste-evoked signaling 341 when any part of the GPCR/PLC_B2/IP₃R3 pathway is nonfunctional, but they are 342 variable and there was never complete abolishment of the responses. Nevertheless, 343 the current dogma within the taste field is that only this signaling pathway is responsible 344 345 for all bitter, sweet and umami transduction (5). While this is puzzling, it is likely due to the lack of another identified signaling pathway that transmits taste signals. We have 346 347 now identified that a PLC β 3 signaling pathway also detects bitter, sweet and umami stimuli. 348

Indeed, a closer inspection of the reported findings from these earlier studies 349 350 reveals differential deficits in taste responses when the PLC β 2/IP₃R3 signaling pathway 351 is nonfunctional. Evaluations of IP₃R3 loss in mice found impaired behavior and nerve responses to different taste stimuli, but not abolishment of these signals (25). They 352 reported reduced behavioral sensitivity for bitter and sweet stimuli at some 353 concentrations but no differences from controls at other concentrations. The cranial 354 nerve recordings revealed deficits in signals that varied with the cranial nerve and taste 355 356 stimulus being tested (25). These findings confirm that IP₃R3 is required for normal

taste signaling but also suggest that it is not the only pathway involved. Similarly, work 357 focused on the loss of PLC β 2 reported reduced, but not abolished, avoidance of bitter 358 compounds by PLC β 2-KO mice (28). In this study, the PLC β 2-KO mice did not avoid 359 360 denatonium as well as wild type mice but still reduced licking as denatonium concentration increased. The PLC_β2-KO mice did not lick normally for sucrose but did 361 lick normally for salt and sour stimuli. The authors concluded that a PLC β 2-362 independent pathway also contributes to taste transduction (28) which agrees with the 363 IP₃R3-KO study (25). Other studies found that PLC β 2 and IP₃R3 co-localize (4, 12, 33) 364 in Type II taste cells, so it is not surprising that loss of either protein causes similar 365 deficits in taste-driven behaviors. 366

We measured the neural activity in the NTS using c-Fos labeling as a marker for 367 activated neurons to determine how the loss of PLCB3 affects central processing of 368 taste information from the periphery (Figure 7). We used oral infusions of taste stimuli in 369 awake and behaving animals because it removes any potential confounds due to 370 anesthetization that is used in other approaches. Since the NTS receives input from the 371 gustatory neurons and is the first synaptic relay from the peripheral taste system, these 372 373 data confirm that loss of either IP₃R3 or PLCβ3 in the taste cells abolishes the taste signal that is sent to the brain. These data from the NTS agrees entirely with our 374 375 behavioral data (Figure 8) which together support the idea that both IP₃R3 and PLC β 3 have important roles in taste transduction. 376

Loss of PLCβ3 caused significant behavioral impairments for the bitter, sweet and umami stimuli that were comparable to the behavioral deficits when IP₃R3 was not present in Type II cells. The loss of either protein was specific to these taste stimuli and

did not affect salt taste (Figure 8E). These data confirm that the lack of either IP₃R3 380 signaling in Type II or PLC β 3 signaling in taste cells had a comparable impact on taste 381 behaviors (Figure 8). Since PLC β 3 and IP $_3$ R3 are not expressed in the NTS (Figure 382 383 S10) or the geniculate ganglia (50), these data strongly support the idea that input from both of these signaling molecules in taste cells is required for normal taste behavior. 384 Therefore, if signaling from either population of taste cells (those using IP₃R3 or those 385 using PLC β 3) is impaired, taste information is not properly conveyed to the brain. This 386 387 conclusion agrees with other studies which suggest that Type II cells are not the only cells that transduce bitter, sweet and umami stimuli (24, 25, 27, 28). 388 Our data support the idea that multiple taste receptor mechanisms, outside of 389 those canonically identified, are involved in the detection of different taste stimuli. 390 Multiple laboratories have independently concluded that the identified taste receptors 391 are not solely responsible for transducing all bitter, sweet and umami stimuli. For 392 instance, the detection of some carbohydrates is not impaired by the loss of T1R2 or 393 T1R3, the identified sweet receptors (47-49, 51), and umami stimuli appear to be 394 transduced by receptors in addition to the T1R1+T1R3 heterodimer (31, 52-58). 395 Moreover, because the identified T1Rs and T2Rs appear to have restricted expression 396 397 in Type II taste cells (5, 59), the presence of the PLC β 3 signaling pathway in non-Type Il cells bolsters the hypothesis that additional taste receptor mechanisms are likely 398 involved in transduction. 399

Our data clearly demonstrate that a subset of Type III cells (DR) are sensitive to
 bitter, sweet and/or umami stimuli in contrast to the idea that Type III cells only detect
 sour and salty stimuli (19, 38). Furthermore, it appears likely that PLCβ3 is expressed

in Type III cells and is responsible for the bitter, sweet and umami responses recorded
in a subset of Type III cells. We previously reported that a PLCβ3/IP₃R1 signaling
pathway is expressed in a subset of taste cells that are not Type II cells (32) which our
current data suggest is a subset of Type III cells. These physiological findings are
supported by recent RNA-sequencing analysis of individual Type II and Type III cells
(37) which revealed PLCβ3 expression in a subset of Type III cells.

We found that many of the DR Type III cells were broadly tuned and responded 409 to two or more taste qualities, including taste stimuli of different modalities (Figure S6. 410 411 S11). Importantly, our studies used isolated taste cells that were not in contact with neighboring cells. Earlier studies using intact buds or taste epithelium reported that a 412 subset of taste cells responded to multiple stimuli with some of these studies concluding 413 414 these cells likely received input from neighboring taste cells (60-62). While we do not rule out the possibility that taste cells receive input from other taste cells, our data show 415 416 that DR Type III cells do not require neighboring input to respond to bitter, sweet or umami stimuli. 417

Our study has demonstrated that 1) Type II cells as defined by the expression of 418 the IP₃R3 are not the only taste cells in the bud that detect bitter, sweet and umami 419 stimuli; 2) taste cells with a PLC β 3 pathway also detect bitter, sweet and umami stimuli; 420 3) PLCβ3 is not expressed in Type II cells but is present in a subset of Type III cells and 421 422 another population of taste cells that lack VGCCs; and 4) loss of either IP₃R3 or PLC_B3 in taste cells causes similar deficits in taste-dependent c-Fos activity in the NTS as well 423 as deficits in taste driven behaviors. We conclude that peripheral taste transduction is 424 more complex than is currently appreciated. 425

426 **EXPERIMENTAL PROCEDURES**

427 **Mice**

Animals were cared for in compliance with the University at Buffalo Institutional 428 Animal Care and Use Committee. Multiple mouse lines were used in this study. Both 429 sexes were used and mice ranged in age from 1 to 6 months. Taste cells for gPCR 430 431 analyses were collected from C57BL/6 mice. The IP₃R3-KO mouse was generated in a C57BL/6 background and was obtained from the Mutant Mouse Resources and 432 Research Center (MMRRC:032884-JAX). This mouse has a targeted mutation in which 433 exon 1 is replaced with a MAPT/GFP fusion (63) which results in the expression of GFP 434 in place of a functional IP₃R3 receptor. Thus, these mice lack a functional IP₃R3 435 receptor and express GFP in cells that would otherwise have expressed IP₃R3. 436 Heterozygous mice express both GFP and the IP₃R3 receptor. The PLCβ3-KO mouse 437 was generously provided by Dr. Sang-Kyou Han (64). The mutation in these mice 438 disrupts the catalytic domain of phospholipase C and was generated in a 129SV agouti 439 mouse strain that was crossed with CD1 mice (65). These mice were maintained in this 440 441 mixed background. Immunohistochemical analyses were also performed in the TRPM5-GFP mouse strain which expresses GFP in all taste cells that express TRPM5. 442 These mice have been backcrossed into the C57BL/6 background and were used as 443 another marker of Type II taste cells (16) to evaluate the expression patterns of PLC β 3. 444 These mice were generously provided by Dr. Robert Margolskee. 445

446 Taste Receptor Cell Isolation

Taste receptor cells were harvested from CV, Fol and Fun papillae of adult mice as previously described (32, 66-71). Briefly, mice were sacrificed using carbon dioxide

and cervical dislocation. Tongues were removed and an enzyme solution containing 449 0.7mg/mL Collagenase B (Roche, Basel, Switzerland), 3mg/mL Dispase II (Roche), and 450 1mg/mL Trypsin Inhibitor (Sigma-Aldrich, St. Louis, MO) was injected beneath the 451 lingual epithelium. After the tongues were incubated in oxygenated Tyrode's solution 452 for approximately 17 min, the epithelium was peeled from the underlying muscle and 453 pinned serosal side up before it was incubated in Ca²⁺-free Tyrode's for approximately 454 25 min. Cells were removed from taste papillae using capillary pipettes with gentle 455 suction and placed onto coverslips coated with Cell-Tak (Corning, Corning, NY). 456

457 Ca²⁺ Imaging

458 All measurements of intracellular calcium (Ca²⁺) were performed in isolated taste 459 receptor cells that were no longer in contact with other taste cells. Cells were loaded for 460 20 minutes at room temperature (RT) with 2 μ M Fura2-AM (Molecular Probes,

Invitrogen, Carlsbad, CA) containing Pluronic F-127 (Molecular Probes). Loaded cells 461 were then washed in Tyrode's solution under constant perfusion for 20min. Multiple 462 taste stimuli and high potassium (50mM KCI) solutions were individually applied and 463 Ca²⁺ responses were recorded. Cells were visualized using an Olympus IX73 464 465 microscope with a 40x oil immersion lens and images were captured with a Hamamatsu ORCA-03G camera (Hamamatsu Photonics K.K., SZK Japan). Excitation wavelengths 466 467 of 340nm and 380nm were used with an emission wavelength of 510nm. Cells were 468 kept under constant perfusion using a gravity flow perfusion system (Automate Scientific, San Francisco, CA). Images were collected every 4s using Imaging 469 470 Workbench 6.0 (Indec Biosystems, Santa Clara, CA). Experiments were graphed and 471 analyzed using Origin 9.2 software (OriginLab, Northhampton, MA).

- 472 Intracellular Ca²⁺ levels were measured as a ratio of fluorescence intensities.
- Fluorescence values were calibrated using the Fura-2 Ca²⁺ Imaging Calibration kit
- 474 (Invitrogen). The effective dissociation constant Kd was calculated to be 180nM, which
- 475 was used in the following equation to calculate Ca^{2+} concentration:
- 476 $[Ca^{2+}] = K_d [(R-R_{min})/(R_{max}-R)](S_{f2}/S_{b2})$
- 477 $[Ca^{2+}] = 0.180 [(R-0.338)/(26.81-R)](22.19)$

R is the ratio value of fluorescence obtained after exciting cells at 340 and 380nm. Data

from cells were analyzed if the cell had a stable Ca^{2+} baseline within the range of 65nM

- and 200nM. An evoked response was defined as measurable if the increase in
- 481 fluorescence was at least two standard deviations above baseline.

482 Immunohistochemistry

Mice were deeply anesthetized by intraperitoneal injections of sodium 483 pentobarbitol, 40 mg/kg (Patterson Veterinary, Mason, MI). Mice were then injected 484 intracardially with heparin (Sigma) and 1% sodium nitrate (Sigma) followed by perfusion 485 with approximately 30mL of 4% paraformaldehyde (Electron Microscopy Sciences, Ft. 486 Washington, PA) in 0.1M phosphate buffer (PB), pH 7.2. After perfusion, the tongues 487 were removed and placed in 4% paraformaldehyde/0.1M PB for 1-2h followed by a 4°C 488 overnight incubation in 20% sucrose/0.1M PB, pH 7.2. For some experiments, tongues 489 490 were immersion fixed overnight in 4% paraformaldehyde/0.1M PB, pH 7.2 at 4°C with 20% sucrose. Regardless of the fixation method, the next day, 40µm sections were cut 491 and washed in PBS 3X10 min at RT. For some experiments, antigen retrieval was 492 493 performed by placing sections in 10 mM sodium citrate, pH 8.5 at 80°C for 5 min. This

494 was done to disrupt the cross-bridges formed by fixation and expose antigen binding495 sites.

Sections were incubated in blocking solution (0.3% Triton X-100, 1% normal goat 496 serum and 1% bovine serum albumin in 0.1M PB) for 1-2h at RT. Primary antibody was 497 added to the sections in blocking solution and incubated for 2 hours at RT followed by 498 499 overnight exposure to primary antibody at 4°C. Controls with no primary antibody were included in each experiment. All primary antibodies were diluted in blocking solution. 500 Mouse anti-IP₃R3 (Transduction Labs, Lexington, KY) was used at 1:50 following 501 502 antigen retrieval. Rabbit anti-PLCB3 (Abcam) was used at 1:200, rabbit anti-PLCB2 (Santa Cruz Laboratories, Santa Cruz, CA) was used at 1:1000, rabbit anti-gustducin 503 (Santa Cruz) was used at 1:200 and anti-NTPDase2 (1:100(72). Mouse anti-SNAP-25 504 (Genway Biotech, San Diego, CA) was used at 1:200 following antigen retrieval. 505 Following overnight incubation in primary antibody, sections were washed in PBS 3X10 506 min at RT and then incubated with the appropriate secondary antibody (Cy5, 1:500; 507 Rhod, 1:250; Jackson ImmunoResearch Laboratories, West Grove, PA) at RT for 2h in 508 the dark. Controls were performed for double labeling experiments to ensure secondary 509 510 antibodies were not binding to primary antibodies raised in different organisms. After secondary antibody incubation, sections were washed in PBS (3x10 min) and mounted 511 on Superfrost Plus slides (VWR, Radnor, PA) using Fluoromount G (Southern 512 513 Biotechnology Associates, Birmingham, AL) and coverslipped. All images were obtained using a Zeiss LSM 710 Confocal Microscope (Zeiss, Oberkochen, Germany). 514 515 Stacks were collected using Zen software (Zeiss) and images were processed using 516 Adobe Photoshop CS5 software adjusting only brightness and contrast.

517 Real-Time PCR of Isolated Taste Cells

518	Taste receptor cells from CV, Fol or Fun papillae were isolated from the papillae
519	as described above and then centrifuged for 20min at 13,000 RPM. RNA was purified
520	using the NucleoSpin RNA XS Kit (Macherey-Nagel, Düren, Germany) according to kit
521	instructions. PCR analysis was performed for GAPDH to ensure sample quality and
522	check for genomic contamination. Contaminated samples were discarded and new
523	samples were collected. Real-Time PCR was performed using a BioRad MiniOpticon
524	system (Bio-Rad Laboratories, Hercules, CA), with BioRad SYBR Green reagents (Bio-
525	Rad Laboratories). Primers used for these experiments were: PLC _{β2} , Fwd:
526	CAATTGAGGGGCAGCTGAGA <i>Rev</i> : TTCTAGGCTGCATCTGGGC; PLCβ3 , <i>Fwd</i> :
527	TCCTGGTGGTCAGGGAT Rev: CTGCCTGTCTCTGCTATCCG; GAPDH Fwd:
528	ACAGTCAGCCGCATCTTCTT, Rev: ACGACCAAATCCGTTGACTC.
529	For real-time PCR analyses, each sample was run in triplicate. If there was more
530	than 5% difference between the replicates, the data were discarded. Multiple biological
531	repeats were used for each papillae type (CV, n=6; Fol, n=6; Fun, n=5). Data was
532	normalized to GAPDH expression for each sample to correct for any loading differences
533	and reported as fold differences.

534 **Oral Infusions**

535 Wild type, IP₃R3-KO and PLC β 3-KO mice (n=3 for each) were tested to measure 536 the effects of quinine stimulation on the c-Fos immunoreactivity in the nucleus of the 537 solitary tract (NTS). Water infusions (n=3) were performed to measure background 538 responses. Surgical procedures were conducted using sterile technique while mice 539 were anesthetized via 2% - 4% isoflurane anesthesia. Guided by a 21G needle, a small

length of PE-90 tubing was passed through the animal's cheek, emerging into the oral
cavity by the rear molars. The tubing was threaded through Teflon washers and heat
flared to secure on both the external and oral sides. Mice were given an analgesic
(0.5mg/ml of carprofen) and allowed to recover. After recovery, they were infused with
either quinine (5mM) or water into the oral catheter for 30 minutes (0.2ml/min, 30 min
infusion). Following the infusion, animals were returned to their cages and left
undisturbed for 45 minutes.

547 Brain Histology and Analysis

After the 45 minute post-infusion period, mice were perfused as described above 548 and the hindbrains were removed. The following day, hindbrains were sectioned into 549 40µm coronal sections which were washed in TBS (pH 7.5), 3x10 min each. Sections 550 were incubated in 1:100 H_2O_2 in TBS for 15 min, followed by 3X10 min TBS washes. 551 Sections were then incubated in anti-c-Fos (ab190289; Abcam, 1:1000) diluted in 552 blocking buffer (3% Normal Donkey Serum in TBS-TritonX) for 1 hr at RT. This was 553 followed by an overnight incubation at 4°C. The next day, sections were washed 3x10 554 min in TBS and then incubated for 2 hrs in anti-rabbit secondary antibody (711-065-152, 555 556 Jackson Immunoresearch, 1:1000) in blocking buffer. After incubation, sections were kept in an avidin-biotin mixture (Elite kit; Vector Laboratories) in TBS-TritonX for 1 hr. 557 558 Tissue sections were washed (3x10 min in TBS) and then stained with DAB (3,3'-559 diaminobenzidine-HCL; Vector Laboratories) for 5 min. Stained tissue sections were washed (3x5 min in TBS) and mounted onto slides using Fluoromount-G (Southern 560 561 Biotechnology Associates). Sections were examined using a light microscope (10-100 562 X) equipped with a digital camera. Analyses of digital images were performed on the

intermediate rostral nucleus tractus solitarius (iRNTS; ~500 µm caudal to the dorsal 563 cochlear nucleus) which receives afferents from both the glossopharyngeal and chorda 564 tympani nerves and displays dense c-Fos-like immunoreactivity in response to intraoral 565 delivery of quinine (42, 73). A template tracing of the nucleus was made by an 566 experimenter blind to genotype and stimulus condition and this template was applied to 567 568 all sections. The number of FLI-positive neurons within the template was counted by hand by an experimenter blind to genotype. Data were analyzed with a one way 569 570 ANOVA with follow up Student's t tests to identify individual differences. Significance 571 level was set at p<0.05.

572 Analysis of Licking Behavior

Unconditioned licking responses to varying concentrations of taste stimuli were 573 recorded in a test chamber designed to measure brief-access licking (Davis MS80 Rig; 574 Dilog Instruments and Systems, Tallahassee, FL). This apparatus consisted of a 575 Plexiglas cage with a wire mesh floor. An opening at the front of the cage allowed 576 access to one of sixteen spill-proof glass drinking tubes that reside on a sliding platform. 577 A mechanical shutter opened and closed to allow the mouse access to one of the tubes 578 579 for a user-specified length of time. A computer controlled the movement of the platform, order of tube presentation, opening and closing of the shutter, duration of tube access 580 581 and interval between tube presentations. Each individual lick was detected by a contact 582 lickometer and recorded on a computer via DavisPro collection software (Dilog Instruments and Systems). 583

584 Mice were adapted to the test chamber and trained to drink from the sipper tubes 585 for 5 consecutive days as previously described (74, 75). During training, mice were 20-

h water deprived. On the first day of training, the mouse was presented with a single 586 stationary bottle of water for 30 min. On the second day, a tube containing water was 587 588 presented but this time the mouse was given 180s to initiate licking and once licking was recorded the mouse was given 30s access to the tube. At the conclusion of either 589 the 30s access or the 180s limit, the shutter was closed again for 10s. Each of the 8 590 591 tubes, all containing water, was presented 3 times. During the remaining three days of training, the mouse was given 30 min to initiate licking to one of eight tubes of water. 592 593 Once the mouse began licking, it was given 10s to lick before the shutter closed for 10s, 594 after which a new tube was presented.

During testing, animals were allowed to take as many trials as possible in 30 min. 595 Mice were tested on varying concentrations of sucrose (0,3,10,30,60,100,300,1000 596 mM), acesulfame K (0,0.5,2,6,8,16,20,32 mM), MSG with 10 µM amiloride 597 (0,25,50,75,100,400,600,800 mM), denatonium benzoate (0,0.1,0.3,1,3,5,10,20 mM), 598 599 and NaCl (0,3,10,30,60,100,300,1000 mM), in that order. Each stimulus was presented in randomized blocks on Monday, Wednesday and Friday in a single week. Animals 600 were 22-h water deprived for all testing except sucrose, when animals were tested 601 602 water replete. Once the animal began licking the tube, they were allowed 10 seconds of access before the shutter closed. 603

For stimuli tested in the water deprived condition (acesulfame K, MSG + amiloride, denatonium benzoate, and NaCl), lick ratios were calculated by dividing the average number of licks at each concentration by the average number of licks to water. For stimuli tested while the animals were water replete (sucrose) licks scores were calculated by subtracting the average number of licks at each concentration by the

average number of licks to water. These corrections are used to standardize for
individual differences in lick rate and are based on water need. Lick scores and licks
relative to water are compared by repeated measures ANOVA with genotype as
between factors variable and concentration as a repeated measures within factors
variable. Significant interaction terms were followed by Tukey's Honestly Significant
Difference tests. Statistical analyses were performed in Statistica.

615 Solutions

All chemicals were purchased from Sigma Chemical (Sigma-Aldrich, St. Louis, 616 MO) unless otherwise noted. Tyrode's solution contained 140mM NaCl, 5mM KCl, 3mM 617 CaCl₂, 1mM MgCl₂, 10mM HEPES, 10mM glucose, and 1mM pyruvate, pH 7.4. Ca²⁺⁻ 618 free Tyrode's contained 140mM NaCl, 5mM KCl, 2.7mM BAPTA, 2mM EGTA, 10mM 619 HEPES, 10mM glucose, 1mM pyruvate, pH 7.4. Nominal Ca²⁺ free Tyrode's contained 620 140mM NaCl, 5mM KCl, 10mM HEPES, 10mM glucose, 1mM pyruvate, pH 7.4. Hi KCl 621 622 solution contained 50mM KCI, 90mM NaCI, 3mM CaCl₂, 1mM MgCl₂, 10mM HEPES, 10mM glucose, 1mM pyruvate, pH 7.4. All taste-solutions, 2µM U73122 (Tocris, Bristol, 623 United Kingdom), and 2µM thapsigargin (Tocris) were prepared in normal Tyrode's. 624 Multiple taste stimuli were analyzed for bitter (5mM denatonium benzoate (Den), sweet 625 (20mM sucralose, 2mM saccharin, 50mM sucrose, 20mM Acesulfame K (Ace K)), and 626 umami (10mM monopotassium glutamate (MPG)). 50mM sucrose contained 90mM 627 628 NaCl instead of 140mM. Salt (250mM sodium chloride, NaCl) and sour (50mM citric acid, CA, pH4) stimuli were tested using the same protocol as Lewandowski et al (19). 629 **Statistics** 630

631	Analysis of the taste responsiveness was performed using an interactive Chi-
632	square analysis with Yate's correction for continuity (76). Significant differences were
633	reported if p<0.05. For real-time PCR analyses, a one-way ANOVA with p<0.05 set as
634	the limit of significance was used to identify any significant differences between the
635	relative expression of PLC β 2 and PLC β 3 in the different papillae types. Comparisons
636	between two experimental conditions were made using a two-tailed Student's T test with
637	p<0.05 set as the limit of significance.
638	Colocalization analysis
639	Image stacks (n=5, 1 μm each) were acquired from labelled sections obtained
640	from three different animals and regions of interest (ROI) were drawn to include the
641	area inside clearly identifiable taste buds. Colocalization analysis was performed on the

- ROIs using ImageJ Fiji software (NIH). Colocalization was determined based on
- 643 Pearson's coefficient using the Colocalization_Finder plugin. If the Pearson's coefficient
- value was greater than 0.9, we considered it to be 100% colocalization due to the
- variability in immunofluorescence intensity. If the Pearson's coefficient was less than
- 646 0.05, we considered this to be no colocalization. A Pearson's coefficient higher than
- 647 0.05 but lower than 0.9 was considered as partial colocalization.

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654 AUTHOR CONTRIBUTIONS

DDB and EDB did live cell imaging, immunohistochemistry, data analysis and
 contributed to writing the manuscript. DDB also did infusion experiments. LEM
 performed brief-access lick behavior experiments, infusion experiments and data
 analysis. KEK performed infusion experiments and data analysis. GCL conceived and

- analyzed infusion experiments. ARN did immunohistochemistry, qPCR and data
- analysis. ZCA and BRK did live cell imaging and data analysis. BTK did
- 661 immunohistochemistry. AV did experiments and analyzed data. AMT supervised brief-
- access lick behavior, infusion experiments, analyzed and interpreted the data, and
- edited the manuscript. KFM conceived and supervised the project, did live cell imaging,
- 664 data analysis and wrote the manuscript.
- 665
- 666 The authors declare no competing interests.
- 667

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879 FIGURE LEGENDS

880	Figure 1. Current taste cell models and experimental setup. A) Current models of
881	the signal transduction pathways in Type II cells (left) that respond to bitter, sweet, and
882	umami stimuli as well as Type III cells (right) that respond to sour and salty stimuli. B)
883	Example of an isolated taste cell that was stimulated with MPG (10mM) and KCI
884	(50mM). The DIC image is shown first, followed by image of Fura 2-AM loading. The
885	next three images show the ratio (340/380) at different time points as the cell is
886	stimulated. The corresponding calcium (Ca ²⁺) changes are shown in 1C.
887	
888	Figure 2. Loss of IP ₃ R3 causes a reduction, but not abolishment, of taste cell
889	sensitivity to different taste stimuli. Chi square analysis with Yate's correction for
890	continuity was used to compare the frequency of evoked Ca ²⁺ responses to different
891	taste stimuli between wild type (WT) and IP $_3$ R3-KO (KO) mice. A) Table of the mouse
892	line, number of mice tested, number of cells tested and number of responses recorded.
893	The P value for each comparison between WT and IP $_3$ R3-GFP-KO is shown as well as
894	the percent response value. These data include all cells from IP_3R3 -GFP-KO mice,
895	both GFP+ and GFP B) Graph shows the percentage of overall responsiveness to
896	stimuli for taste cells from the CV, Fol and Fun papillae in WT and IP $_3$ R3-KO. The
897	responsiveness to multiple stimuli was evaluated and is shown for CV (C), FOL (D) and
898	FUN (E), Specific numbers and P values for each are shown in Figure S3. (*, p<0.05; **,
899	p<0.01; ***, p<0.001).

900

901 Figure 3. Taste-evoked Ca²⁺ release in IP₃R3-KO mice is dependent upon PLC

activity and Ca²⁺ release from internal stores. Open columns represent the time that 902 the taste stimulus is presented. The application of Ca²⁺ free Tyrode's is indicated by the 903 dashed lines. The stimulus presented during this time is also in Ca²⁺ free Tyrode's. 904 The gray hatched columns represent the application of either thapsigargin or U73122, 905 906 both of which are irreversible inhibitors. A) Bitter-evoked taste responses (5mM Den) persist in the absence of extracellular calcium (Ca²⁺-free) and are abolished by the 907 SERCA pump inhibitor thapsigargin (B) as well as the PLC blocker U73122 (C). D) 908 Responses to sweet stimuli (20mM sucrose) persist in Ca²⁺-free and are abolished by 909 thapsigargin (E) and U73122 (F). G) Umami stimuli (10mM MPG) persist in Ca²⁺-free 910 and were abolished by thapsigargin (H) and U73122 (I). Individual data points are 911 shown in Figure S4. 912

913

Figure 4. Expression of PLC\beta3 in taste cells. Laser scanning confocal micrographs (LSCMs, stack of 5 slices, 1 μ m each) of PLC β 3 immunostaining in the IP₃R3-KO-GFP mice reveals that PLC β 3 is expressed in a separate population from the GFP positive taste cells in the CV (A, n=5), Fol (B, n=3) and Fun papillae (C, n=4). Scale bar=20 μ M. D) Co-localization analysis is shown.

919

920 Figure 5. Some Type III cells respond to bitter, sweet and umami stimuli.

A-C) Representative traces of taste cells that responded to bitter, sweet and/or umami stimuli and Hi KCI (50mM) depolarization with Ca²⁺ signals. Some cells responded to 1 stimulus while others responded to multiple stimuli. D) Representative trace of taste

cells from IP₃R3-KO mice that responded to 50mM citric acid (CA), 50mM KCI, bitter 924 (5mM denatonium), sweet (20mM sucralose) and umami (10mM monopotassium 925 glutamate, MPG) stimuli. E) Representative trace of a separate subset of cells that 926 responded to 250mM NaCl, 50mM citric acid (CA) and 50mM KCl but were not sensitive 927 to the bitter, sweet, umami stimuli tested. F) Summary of taste cells from IP₃R3-KO 928 mice that responded to 50mM KCl with a Ca^{2+} signal (n=20), 25% only responded to 929 CA, 35% responded to CA and NaCl, and 40% responded to CA and bitter, sweet 930 and/or umami stimuli (B,U,S) while no cells responded to all the taste stimuli. G) The 931 932 overall responsiveness of taste cells from the CV, Fol and Fun papillae of the IP₃R3-KO mice and WT mice was measured. This is defined as the percentage of stimulated cells 933 that responded to at least one stimulus. The percentage of DR cells in both lines was 934 also measured using the criteria that the cell responded to at least one stimulus AND 935 50mM KCI with Ca²⁺ signals. The overall responsiveness to taste stimuli was 936 significantly reduced in the IP₃R3-KO mice compared to controls (**,p<0.01) while the 937 number of responsive DR cells did not change in the absence of IP₃R3. H) The 938 percentage of responsiveness in the DR taste cells to different taste stimuli in the WT 939 and IP₃R3-KO mice. Chi-square analysis found no significant differences. 940

941

Figure 6. PLC β 3 is required for the detection of bitter, sweet and umami stimuli.

Chi square analysis with Yate's correction for continuity was used to compare the
frequency of evoked Ca²⁺ responses to different taste stimuli between wild type (WT)
and PLCβ3-KO (KO) mice. A) Table of the mouse line, number of mice tested, number
of cells tested and number of responses recorded. The P value for each comparison

947	between WT and PLC β 3-KO is shown as well as the percent response value. B) The
948	overall responsiveness and the percentage of DR taste cells from the CV, Fol and Fun
949	papillae of the PLC β 3-KO mice and WT mice was measured. Both the overall
950	responsiveness to taste stimuli and the number of DR cells were significantly reduced in
951	the PLC β 3-KO mice compared to controls. The responsiveness to multiple stimuli was
952	evaluated and is shown for CV (C), FOL (D) and FUN (E). Specific numbers and P
953	values for each are shown in Figure S8. (*, p<0.05; **, p<0.01; ***, p<0.001).
954	
955	Figure 7. Loss of PLC β 3 or IP $_3$ R3 causes deficits in the taste-evoked neural
956	activity in the nucleus of solitary tract.
957	Oral infusion of bitter (quinine) and water elicited c-Fos immunoreactivity in the
958	intermediate rostral nucleus tractus solitarius (IRNTS). (A, B) Loss of either PLC β 3
959	(PLC β 3-KO) or IP $_3$ R3 (IP $_3$ R3-KO) caused a significant reduction in the c-Fos
960	immunoreactivity in the IRNTS compared to WT mice. The residual c-Fos labeling in
961	each of the KO mice was comparable to the c-Fos labelling generated by water infusion.
962	For all experiments, 3 mice of each genotype were used (***, p<0.001; **, p<0.01). The
963	red outlined area denotes the analyzed region for each. Data were analyzed with a one
964	way ANOVA with follow up Student's t tests to identify individual differences.
965	Significance level was set at p<0.05. C) A schematic diagram showing the analysis
966	region in the hindbrain that was analyzed.
967	
968	Figure 8. Loss of PLC β 3 or IP $_3$ R3 affects behavioral responses to taste stimuli.

969 Lick data (±standard deviation) from brief-access behavioral tests compare the

970	responses of IP ₃ R3-KO (red line) and PLC β 3-KO (pink line) to WT (IP ₃ R3-WT, black
971	line; PLC β 3-WT, blue line). A) Lick ratios (stimulus/water) of the WT mice for the bitter
972	stimulus denatonium (0, 0.1, 0.3, 1, 3, 5, 10, 20mM) were significantly different from the
973	IP ₃ R3-KO and PLC β 3-KO responses. B) Lick ratios of the WT mice for the umami
974	stimulus monosodium glutamate + $10\mu M$ amiloride (MSG, 0, 25, 50, 75, 100, 400, 600,
975	800mM) were significantly different from the IP ₃ R3-KO and PLC β 3-KO mice. C) Lick
976	ratios of the WT mice for the artificial sweetener acesulfame K (0, 0.5, 2, 6, 8, 16, 20,
977	32mM) were significantly different from the IP ₃ R3-KO and PLC β 3-KO mice. D) Lick
978	scores (stimulus-water) of the WT mice for sucrose (0, 3, 10, 30, 60,100, 300, 1000mM)
979	were significantly different from the IP ₃ R3-KO and PLC β 3-KO responses. E) No
980	significant differences were detected between the WT and KO mice for any
981	concentration of NaCl (0, 3, 10, 30, 60, 100, 300, 1000mM) tested. For all experiments,
982	5 mice of each genotype were used (***, p<0.001; **, p<0.01; *, p<0.05). Data were
983	compared by repeated measures ANOVA. Significant interaction terms were followed
984	by Tukey's Honestly Significant Difference tests.
005	





Papillae type	Mouse genotype	Cells tested	# of Resp	P value	% value		eness	75 -	WT (all ce IP ₃ R3-KO GFP+ in I	lls, including Typ (non green cells) P ₃ R3-KO	e II)
All cells	WT (n=105 mice)	407	259	P<0.0001	63.6		siv		_		
	KO (n=115 mice)	690	213		30.9		DOC		64	_	68
CV	WT (n=63 mice)	250	159	P=0.0003	63.6		est	50 -	**	55	5
	KO (n=70 mice)	443	172		38.8		t R	-	44	46	
FOL	WT (n=28 mice)	53	29	P=0.44	54.7		Sen	25 -			
	KO (n=37 mice)	123	52		42.3		erc	-			
FUN	WT (n=32 mice)	104	71	P=0.17	68.2		Δ.	0	0	0	
	KO (n=30 mice)	124	61		49.2				ĊV	FOL	FÚ
2 ¹⁰⁰	WT IP ₃ R3-KO	CV	C	100 Senerss 75	WT IP ₃ R3-KO	FC)L		100	WT IP ₃ R3-KO	FUN
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Comparison	Average overlap, standard deviation						
GFP vs anti- PLCβ3	0.03±0.02						





-		-	-		_	
Papillae type	Mouse genotype	Cells tested	# of Resp.	P value	% value	
All cells	WT (n=53 mice)	577	344	P=0.002	59.6	
	KO (n=68 mice)	375	155		41.3	
CV	WT (n=40 mice)	221	137	P=0.246	61.9	
	KO (n=55 mice)	138	68		49.2	
FOL	WT (n=40 mice)	243	136	P=0.039	55.9	
	KO (n=34 mice)	120	43		35.8	
FUN	WT (n=35 mice)	113	71	P=0.035	62.8	
	KO (n=48 mice)	117	44		37.6	



В





Α





PLCβ3-KO

Water



1000



Concentration (mM)

0

1

Cell populations	Identifiers/Classification					
IP ₃ R3-KO-GFP cells	Type II cells with GFP expression replacing IP ₃ R3					
	expression.					
WT cells	All cells that were tested and/or responded to					
	stimuli, including Type II cells, Type III cells, other					
	cells.					
IP ₃ R3-KO cells	Taste cells from mice that lack functional Type II					
	cells and do not express GFP. These include Type					
	III cells and other cells.					
Dual-responsive cells	Taste cells that respond to 50mM KCI (indicating					
	the presence of VGCC) and bitter, sweet and/or					
	umami stimuli.					
Taste-only cells	Taste cells from the IP ₃ R3-KO mouse that lack GFP					
	expression (are not Type II cells) and do not					
	respond to 50mM KCI (are not Type III cells).					

Table S1. Definition of taste cell populations analyzed.



Figure S1. Characterization of the IP₃R3-KO mice. A) LSCMs (stack of 5 slices, 1µm each) from WT mice identified anti-IP₃R3 labeling in taste receptor cells from the CV (n=3). B) IP₃R3-KO mice express GFP in lieu of IP₃R3 and were not labeled by anti-IP₃R3 (n=6). C) LSCMs of the IP₃R3-het mouse identified strong co-localization between anti-IP₃R3 labeling and GFP expression. D) Anti-PLCβ2 labeling in IP₃R3-KO mice found that PLCβ2 co-localizes with GFP, indicating that IP₃R3-KO-GFP is specific to Type II cells (LSCMs: stack of 5 slices, 1µm each; n=4). E) α -gustducin is present in a subset of the IP₃R3-KO-GFP taste cells in the CV (LSCMs: stack of 10 slices, 1µm each; n=3). Asterisks identify some GFP expressing cells that do not express gustducin. Scale bars = 20 µm. F) Co-localization analysis identified the average (± standard deviation) overlapping expression for each target protein with the GFP expression, n=3 for each.



Figure S2. Taste-evoked Ca²⁺ signals are specific to the stimulus.

A concentration gradient analysis of (A) a bitter stimulus (denatonium), (B) an umami stimulus (MPG) and (C) a sweet stimulus (saccharin) demonstrates that DR cells respond to a wide range of stimulus concentrations. The GFP+ cell (lacks IP₃R3) in C did not respond to any concentration of saccharin applied. (D-E) Identical concentrations of the compound cellobiose did not generate a Ca²⁺ signal in DR cells, indicating that the measured responses are taste-activated and not due to non-specific osmotic effects.

	IP ₃ R3										
FOL											
lue		FOL	MS	Cells	# of	P val					

CV

FUN	
-----	--

CV Stimulus	MS	Cells tested	# of Resp.	P value	FOL Stimulus	MS	Cells tested	# of Resp.	P value	FUN Stimulus	MS	Cells tested	# of Resp.	P value
DEN	WT	238	102	0.06	DEN	WT	46	18	0.93	DEN	WТ	76	42	0.89
	КО	341	107			КО	100	38			ко	59	30	
SUC	WT	72	16	0.0004	SAC	WT	15	1	0.69	SUC	WT	41	10	0.89
	КО	87	1			КО	50	4			ко	19	6	
SAC	WT	106	23	0.005	SUCRAL	WT	23	6	0.08	SAC	WT	65	23	0.73
	ко	174	13			КО	30	1			ко	55	16	
SUCRAL	WT	121	47	p<0.001	MPG	WT	18	4	0.48	SUCRAL	WT	85	25	0.04
	КО	137	10			КО	73	8			ко	99	13	
MPG	WT	81	13	0.82	ні к	WT	53	14	0.93	MPG	WT	75	17	0.4
	КО	301	42			КО	113	31			ко	81	12	
НΙΚ	WT	245	75	0.68						ΗΚ	WT	104	33	0.7
	КО	384	108								КО	103	28	

Figure S3. Tables of the mouse line, number of mice tested, number of cells tested and number of responses recorded from the CV, FOL, and FUN papillae of the WT and IP₃R3-KO mice data shown in Figure 2. Chi-square analysis with Yate's correction for continuity was used to identify significant differences in the response frequency between WT and KO mice for each stimulus in each papillae.

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Figure S4. Taste-evoked Ca²⁺ release in IP₃R3-KO mice is dependent upon PLC activity and Ca²⁺ release from internal stores. Individual data points of the data presented in Figure 3. A) Bitter-evoked taste responses (5mM Den, n=6) persist in the absence of extracellular calcium (Ca²⁺-free) and are abolished by the SERCA pump inhibitor thapsigargin (B, n=7) as well as the PLC blocker U73122 (C, n=7). D) Responses to sweet stimuli (50mM sucrose, n=6) persist in Ca²⁺-free and are abolished by thapsigargin (E, n=4) and U73122 (F, n=6). G) Umami stimuli (10mM MPG) persist in Ca²⁺-free (n=7) and were abolished by thapsigargin (H, n=7) and U73122 (I, n=4). A two-tailed Student's t test was used to identify significant differences for each experimental condition.



Figure S5. Expression of PLCβ3 in taste papillae. A) Anti-PLCβ3 labeling in the CV of TRPM5-GFP mice determined that PLCβ3 is expressed in taste cells lacking GFP expression (LSCMs: stack of 5 slices, 1µm each; n=4). B) Co-labeling with anti-PLCβ2 and anti-PLCβ3 in the CV of C57BL/6 mice revealed that these PLCβs are expressed in separate taste cell populations (LSCMs: stack of 5 slices, 1µm each; n=3). C) Co-labeling with anti-NTDPase2 and anti-PLCβ3 in the CV of C57BL/6 mice determined that these markers are expressed in separate taste cell populations (LSCMs: stack of 5 slices, 1µm each; n=3). C) Co-labeling with anti-NTDPase2 and anti-PLCβ3 in the CV of C57BL/6 mice determined that these markers are expressed in separate taste cell populations (LSCMs: stack of 5 slices, 1µm each; n=3). Scale bar = 10µM. D) Co-localization analysis identified the average (\pm standard deviation) overlapping expression for PLCβ3 with TRPM5-GFP or anti-PLCβ2 expression, n=3 for each. mRNA was isolated from taste cells originating in the different papillae types from C57BL/6 mice. Taste cells were analyzed from at least five different mice for each. Values were normalized to GAPDH expression and are presented as a ratio to values from the CV papillae for (E) PLCβ2 and (F) PLCβ3. (***, p<0.001). A one-way ANOVA with p<0.05 set as the limit of significance was used to identify any differences in the relative mRNA levels between papillae types.



Figure S6. Dual responsive cells are present in multiple mouse lines. Control imaging experiments were performed using GAD67-GFP mice and C57BL/6 mice. GAD67-GFP is expressed in a large subset of Type III mouse taste cells (34). A-B) Representative traces of DR taste cells that responded to bitter, sweet and/or umami stimuli and Hi KCI (50mM) in GAD67-GFP mice. DR taste cells were present in both GAD67-GFP + (A) and -GFP- (B) taste cells. C) Experiments in C57BL/6 mice also identified the presence of DR taste cells.

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Figure S7. Stimulus evoked Ca²⁺ responses in taste cells in PLCβ3-KO mice. A) Representative trace of taste cells from PLCβ3-KO mice that responded to 50mM citric acid (CA) and KCI (50mM) but did not respond to bitter (5mM denatonium), sweet (20mM sucralose) or umami (10mM monopotassium glutamate, MPG) stimuli. B) Representative trace of a separate subset of cells that responded to 250mM NaCl, CA and KCl but were not sensitive to the bitter, sweet, umami stimuli tested. C) Summary of responsive taste cells from PLCβ3-KO mice (n=38): CA and KCl (n=18); CA, NaCl and KCl (n=3); CA, KCl, and bitter, sweet and/or umami stimuli (B,U,S) (n=0); and B,U,S only (n=17). Representative taste-evoked responses for bitter (D), sweet (E) and umami (F) in the PLCβ3-KO mice. None of these cells responded to CA or KCl.

PLCβ3 FOL

CV



CV Stimulus	MS	Cells tested	# of Resp.	P value	FOL Stimulus	MS	Cells tested	# of Resp.	P value		FUN Stimulus	MS	Cells tested	# of Resp.	P value
DEN	WT	221	86	0.018	DEN	WT	233	85	.144	Γ	DEN	WT	112	52	0.003
	ко	138	30			ко	113	28		Γ		КО	117	23	
SUC	WT	191	36	0.039	SUC	WT	173	21	0.97	Γ	SUC	WT	86	17	0.91
	ко	112	9			ко	65	7		Γ		КО	107	19	
SAC	WT	198	22	0.02	SAC	WT	156	27	0.04	Γ	SAC	WT	93	22	0.015
	ко	130	4			ко	86	5		Γ		КО	111	10	
ACEK	WT	181	27	0.04	ACEK	WT	163	28	0.04	Γ	ACEK	WT	84	12	0.015
	ко	111	6			ко	76	4		Γ		КО	108	3	
MPG	WT	182	32	0.031	MPG	WT	179	34	0.21	Γ	MPG	WT	81	10	0.016
	ко	116	8			ко	67	3		Γ		ко	107	2	
ні к	WT	221	82	0.06	НΙК	WT	243	68	0.08	Γ	HI K	WT	113	15	0.053
	ко	138	32			ко	120	20		Γ		КО	116	5	
DR	WT	221	59	0.0008	DR	WT	243	45	0.002	Γ	DR	WT	113	13	0.001
	КО	138	12			КО	120	5				КО	116	0	

Figure S8. Tables of the mouse line, number of mice tested, number of cells tested and number of responses recorded from the CV, FOL, and FUN papillae of the WT and PLC β 3-KO mice data shown in Figure 6. Chi-square analysis with Yate's correction for continuity was used to identify significant differences in the response frequency between WT and KO mice for each stimulus in each papillae.



Figure S9. PLC β 3 is expressed in some Type III taste cells. Immunohistochemical analyses (LSCMs: stack of 5 slices, 1µm each) using anti-PLC β 3 and anti-SNAP25 revealed some co-localization between PLC β 3 and SNAP25, confirming that PLC β 3 is expressed in some Type III cells but is not restricted in its expression to this taste cell type (A, CV papillae, n=3; B, Fun papillae, n=3) Scale bar=20µM. Labeling with anti-SNAP25 in the CV of PLC β 3-WT (**C**) and PLC β 3-KO (**D**) mice revealed no differences in the expression level of this synaptic marker (n=5 mice for each). Scale bars = 50 µm. E) Co-localization analysis identified the average (± standard deviation) overlapping expression for PLC β 3 with anti-SNAP25 expression, n=3.



Figure S10. Neither IP₃R3 nor PLC β 3 is expressed in IRNTS.

CV papillae and IRNTS sections were evaluated in parallel. A) Analyses of the brain sections from the IP₃R3-KO mice revealed no GFP labelling in the IRNTS. Immunohistochemical analysis using anti-PLC β 3 also did not detect any PLC β 3 expression. Scale bar=50µm. B) LSCM analyses (n=5 sections, 1 µM each) of the tongues from the same mice identified the expression of IP₃R3-KO-GFP and PLC β 3 in the taste receptor cells from the CV papillae (n=3). Scale bar=10µm.



Figure S11. Analyses of the different taste cell populations that responded to taste stimuli. Graphs show the percent of the responses for each taste quality tested. A-C) Response profiles for bitter, sweet and umami stimuli from the DR taste cells in WT (n=29 mice, 62 cells) and IP₃R3-KO (n=18 mice, 31 cells) mice. Taste cells that were stimulated with bitter, sweet and umami stimuli as well as 50mM KCI and responded to at least one of the taste stimuli and KCI were analyzed. The relative percentage of DR taste cells that responded to 1, 2, or 3 taste qualities is shown for each taste quality analyzed. Chi-square analysis with Yate's correction for continuity found no significant differences between the WT and IP₃R3-KO mice for bitter (A), sweet (B), or umami (C) response profiles. The same analysis was applied to taste cells that responded to at least one bitter, sweet or umami stimulus but did not respond to 50mM KCI for WT (n=37 mice, 85 cells) and IP₃R3-KO (n=33 mice, 78 cells) mice. These data are shown for bitter (D), sweet (E), and umami (F). No significant differences were found between WT and IP₃R3-KO mice.