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HAL Id: inserm-00707643
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Submitted on 13 Jun 2012

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Plant Insecticide \(\text{L}-\text{Canavanine}\) Repels \(\text{Drosophila}\) via the Insect Orphan GPCR \(\text{DmX}\)

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Abstract

For all animals, the taste sense is crucial to detect and avoid ingesting toxic molecules. Many toxins are synthesized by plants as a defense mechanism against insect predation. One example of such a natural toxic molecule is \(\text{L}-\text{canavanine}\), a nonprotein amino acid found in the seeds of many legumes. Whether and how insects are informed that some plants contain \(\text{L}-\text{canavanine}\) remains to be elucidated. In insects, the taste sense relies on gustatory receptors forming the gustatory receptor (Gr) family. Gr proteins display highly divergent sequences, suggesting that they could cover the entire range of tastants. However, one cannot exclude the possibility of evolutionarily independent taste receptors. Here, we show that \(\text{L}-\text{canavanine}\) is not only toxic, but is also a repellent for \(\text{Drosophila}\). Using a pharmacogenetic approach, we find that flies sense food containing this poison by the \(\text{DmX}\) receptor. \(\text{DmXR}\) is an insect orphan G-protein–coupled receptor that has partially diverged in its ligand binding pocket from the metabotropic glutamate receptor family. Blockade of \(\text{DmXR}\) function with an antagonist lowers the repellent effect of \(\text{L}-\text{canavanine}\). In addition, disruption of the \(\text{DmXR}\) encoding gene, called \(\text{mangetout}\), suppresses the \(\text{L}-\text{canavanine}\) repellent effect. To avoid the ingestion of \(\text{L}-\text{canavanine}\), \(\text{DmXR}\) expression is required in bitter-sensitive gustatory receptor neurons, where it triggers the premature retraction of the proboscis, thus leading to the end of food searching. These findings show that the \(\text{DmX}\) receptor, which does not belong to the Gr family, fulfills a gustatory function necessary to avoid eating a natural toxin.

Introduction

Taste is essential to distinguish between nutritious and toxic substances. To avoid eating toxins, animals are able to detect them by using a repertoire of taste receptors [1]. Although it is recognized that a bitter taste sensation is critical to avoid toxic substances [2,3], the cellular and molecular mechanisms that have been established during evolution to detect a toxin are not well understood. In particular, how a receptor becomes tuned to a toxin is not well documented, mainly because the structure of its ligand binding pocket (LBP) and the evolutionary relationship with the ancestor receptor are not known.

In \(\text{Drosophila}\), the family of gustatory receptors (Grs) is predicted to consist of 68 genes [4,5]. This family of receptors, which consist of seven transmembrane domain proteins, is characterized by a very high level of amino acid divergence, showing as little as 8%–12% amino acid identity [5]. Such diversity suggests that the Gr family could cover the entire range of taste-receptive capability of the fly. Nevertheless, the extreme divergence within this family does not exclude the possibility of evolutionarily independent insect taste receptors not belonging to the Gr family. To date, only few receptors of the Gr family have been associated with a specific taste molecule: for example, the receptor for the sugar trehalose, called Gr5a [6], and the bitter compound caffeine coreceptors, called Gr66a and Gr93a [7,8].

Plants synthesize many toxic molecules as defense mechanisms against predation [9,10]. A number of such toxic compounds are nonprotein amino acids [11,12]. The best-characterized example of nonprotein amino acid that plays a defensive role is \(\text{L}-\text{canavanine}\) (2-amino-4-guanidinooxybutyric acid) [13–15], which is massively accumulated in the seeds of many legumes (up to 143 mM in \textit{Medicago sativa} [16]). \(\text{L}-\text{Canavanine}\) is a natural insecticide because it is structurally similar enough to \(\text{L}-\text{arginine}\) (Figure S1) to interfere with \(\text{L}-\text{arginine}\) metabolism and to be incorporated by arginyl-tRNA synthase in de novo proteins resulting in dysfunctional proteins [17–19]. Thus, these properties


Academic Editor: Hugo J. Bellen, Baylor College of Medicine, United States of America

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Funding: This work was supported by fellowships from the Ministère de la Recherche and the Fondation pour la Recherche Médicale to CM, a Centre National de la Recherche Scientifique (CNRS) postdoctoral fellowship to LS, by the ANR grant DROSDYN to MLP and by grants from the CNRS and Institut National de la Santé et de la Recherche Médicale (INSERM) to the Institut de Génomique Fonctionnelle. The small animal imaging core facility (http://ipam.igf.cnrs.fr/) was supported by grants from the National Biophotonics and Imaging Platform (Ireland), Réseau National des Génapoles, Institut Féderatif de Recherche 3 (IFR3), and Région Languedoc Roussillon. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

Abbreviations: GPCR, G-protein–coupled receptor; Gr, gustatory receptor; GRN, gustatory receptor neuron; LBP, ligand binding pocket; LSO, labral sense organ; mGlR, metabotropic glutamate receptor; PER, proboscis extension reflex; PI, preference index; PR, proboscis retraction; QRT-PCR, quantitative real-time reverse-transcriptase polymerase chain reaction; RNAi, RNA interference; VCSO, ventral cibarial sense organ; WT, wild-type.

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Author Summary

Plants evolve to fend off the insects that attack them, often by synthesizing compounds toxic to insects. In turn, insects develop strategies to avoid these plants or resist their toxins. Some plant toxins are nonprotein amino acids. For example, seeds from numerous legumes contain high amounts of l-canavanine, a nonprotein amino acid that is structurally related to l-arginine and is highly toxic to most insects. How insects can detect l-canavanine remains to be elucidated. Using pharmacology, genetics, and behavioral approaches, we show that flies sense l-canavanine using the receptor DmX, an orphan G-protein–coupled receptor that has diverged in its ligand binding pocket from metabotropic glutamate receptors. Disruption of the DmXR gene, called mangetout (mtt), suppresses the l-canavanine repellent effect. DmXR is expressed and required in aversive gustatory receptor neurons, where it triggers the premature retraction of the proboscis, thus leading to the end of food searching. Our results indicate a mechanism by which some insects may detect and avoid a plant toxin.

of l-canavanine render it a highly toxic secondary plant constituent [15]. To deal with this natural poison, some insects have generated several adaptive strategies. Indeed, the tobacco budworm Heliothis virescens uses detoxification [20] and the beetle Caryaedis brasiliensis feeds exclusively on l-canavanine–containing seeds but catalyzes l-canavanine to l-canaline and urea [21]. However, these two mechanisms to circumvent the toxic properties of l-canavanine are specific to few insect species. Thus, the evidence for a protective function against predation for such nonprotein amino acids, i.e., whether and how insects are informed that plants contain l-canavanine, remains to be shown [15].

Amino acids are known to be the ligands of G-protein–coupled receptors (GPCRs) belonging to the family C [22,23]. All members of this family display a common structural architecture characterized by a long N-terminal extracellular domain containing a bilobular LBP [24], a seven transmembrane domain, and an intracellular C-terminus. This family includes metabotropic glutamate receptors (mGluRs). In mammals and in insects, mGluRs, which are activated by the neurotransmitter glutamate, play different roles in the central nervous system [25,26]. We have previously shown that one mGluR has diverged through evolution to give rise to the mX receptor, called DmXR in Drosophila [27]. Orthologs of DmXR are so far only found in insects [27]. DmXR differs from mGluRs in the distal part of the LBP, so that this receptor is an orphan receptor, which is not activated by glutamate [27]. However, we previously showed that the DmXR and mGluR LBPs share the crucial residues necessary to bind a ligand with amino acid structural properties [27]. To deorphanize the DmX receptor, we previously tested various molecules having such properties, including all the classical amino acids, and did not find any ligand [27].

Drosophilidae are saprophagic animals, and members of dipteran families such as Tephritidae or Scatophagidae are seed predators [28], so we asked whether l-canavanine could activate DmXR. Here, we show that l-canavanine is a ligand of DmXR in vitro. We then wondered whether insects could be informed that plants contain l-canavanine via the mX receptor. We have addressed this question by using Drosophila as an insect model. First, we confirmed that l-canavanine is highly toxic when ingested. We then tested whether Drosophila avoid eating food containing l-canavanine. We found that l-canavanine is recognized by flies and mediates a behavioral avoidance response via a chemosensory mechanism. Hence, l-canavanine is a repellent. We then analyzed the molecular and cellular bases of l-canavanine–induced repulsive behavior, using gustatory behavior, pharmacology, and genetic approaches. We found that l-canavanine is detected in vivo by the DmX receptor. To control the l-canavanine avoidance behavior, the DmX receptor is expressed and required in bitter-sensitive gustatory receptor neurons (GRNs). These findings show that the gustatory detection of a natural toxin relies on DmXR, a divergent mGluR not belonging to the Gr family.

Results

The Orphan Insect Receptor DmX Is Activated In Vitro by l-Canavanine

To test whether l-canavanine could activate DmXR, we transiently expressed this receptor in human embryonic kidney (HEK) cells and assayed for l-canavanine–induced DmXR activation. We found that l-canavanine activated HEK cells expressing DmXR (Figure 1A and Figure 1B). In contrast, the close structural homolog l-arginine showed no agonist or antagonist effect on DmXR (Figure 1C). l-Canavanine did not activate HEK cells expressing the unique fly mGlu receptor, DmGluA (Figure 1A). We searched for an antagonist and found that N-methyl-l-arginine (NMA) inhibited DmXR activation by l-canavanine (Figure 1C and 1D). Previous sequence analysis, mutagenesis, and 3-D modeling studies had shown that the LBP of the DmXR is very homologous to the LBP of mGluRs [27]. The residues contacting the amino acid moiety of glutamate (the \( \text{COO}^- \) and NH3\(^+\) groups) are conserved in DmXR (e.g., Thr-176), whereas the residues interacting with the \( \gamma \)-carboxylic group are not [27]. The Thr-176 residue is conserved in all mGluRs, and its mutation strongly decreases the affinity of these receptors for glutamate [24]. Similarly, l-canavanine did not activate Thr-176–mutated DmX receptor (DmXRT176A) transfected in HEK cells (Figure 1A), although this mutated receptor is actually localized at the plasma membrane (as shown in [27]). This indicates that the plant amino acid binds into the LBP. Altogether our data show that DmXR is a l-canavanine receptor because of a partial modification of its LBP from the LBP of mGluRs. These results suggest that Drosophila may be able to detect l-canavanine in vivo through the DmX receptor.

L-Canavanine Is Toxic for Drosophila

Since l-canavanine is described as a natural insecticide [15], we first examined whether ingested l-canavanine is also toxic for Drosophila melanogaster. We maintained 50 young wild-type (WT) flies on Drosophila medium containing 10 mM l-canavanine and compared their viability and their fecundity to flies maintained on medium without l-canavanine (\( n = 8 \)). When flies fed on 10 mM l-canavanine, we did not observe massive mortality or dramatic decrease of the lifespan. However, all the offspring of flies constrained to eat 10 mM l-canavanine died during larval stages (number of offspring in control medium \( >1,000 \), number of offspring in 10 mM l-canavanine = 0). These results indicate that Drosophila is a l-canavanine–susceptible insect.

L-Canavanine Mediates a Behavioral Avoidance Response

Because of its toxicity, we hypothesized that Drosophila may avoid eating l-canavanine if they have the choice. To test this, we performed a two-choice feeding preference test. This behavioral assay measures the consumption of a sucrose solution (5 mM)
colored by two food dyes (blue or red) offered simultaneously to adult fly populations [29]. After 2 h in the dark, flies are counted on the color dye witnessed in their abdomen. In the control situation, WT flies preferred the blue solution, the preference index (PI) being 0.82 \pm 0.04 (Figure 2). We then added increasing concentrations of L-canavanine to the blue solution (1 mM to 40 mM). We found that L-canavanine inhibited the intake of the blue solution (unpublished data). The L-canavanine effect increased with concentration, reaching a plateau at 30–40 mM. At these concentrations, the PI for the blue solution was 0.13 \pm 0.06 (Figure 2). A similar repulsive effect of L-canavanine was also visible when the drug was added to the red solution (unpublished data). To determine whether this repulsive effect was mediated by a chemosensory mechanism, we used flies carrying an adult-viable mutant allele of the pox-neuro (pox) gene. In homozygous pox flies, external chemosensilla are deleted or transformed into mechanosensilla [30,31]. pox flies fed equally on

**Figure 1.** L-Canavanine activates DmXR and N-methyl-L-arginine (NMA) inhibits DmXR. (A) L-Canavanine (Cana) is an agonist of DmXR. L-Canavanine at a concentration of 10 mM (black) activated DmXR in HEK cells, but had no effect on the unique fly metabotropic glutamate receptor DmGluAR, which was activated by 1 mM glutamate (checkered). Note that 10 mM L-canavanine did not activate the DmX ligand binding pocket mutant receptor (DmXRT176A). (B) Dose-response curve of L-canavanine on DmXR. IP production by DmXR receptors activated with increasing concentrations of L-canavanine (half-maximal effective concentration [EC50] of L-canavanine = 0.5 \pm 0.2 mM). (C) N-methyl-L-arginine (NMA) inhibits DmXR. The activation of DmXR by 1 mM L-canavanine (grey) was completely inhibited by 1 mM NMA (diagonally hatched on grey background). Thus NMA, which has no effect by itself (diagonally hatched on white background), is a potent antagonist of DmXR. L-arginine (1 mM) did not activate DmXR (horizontally hatched on grey background) nor antagonize 1 mM L-canavanine effect on DmXR (horizontally hatched on white background). (D) Dose-response curve of NMA antagonistic effect on DmXR. IP production by DmXR receptors activated with 2 mM L-canavanine (L-cana) in the presence of increasing concentrations of the antagonist NMA (half-maximal inhibitory concentration [IC50] of NMA = 0.2 \pm 0.2 mM). For (A–D), data are expressed as the IP (inositol triphosphate) production in HEK cells coexpressing a chimeric G-protein \( \alpha \)-subunit Gqi9 and the indicated receptor in presence of drugs relatively to IP production in the basal conditions. The vertical bars represent the standard error of the mean (SEM of triplicate determinations from typical experiments). Asterisks indicate significant differences by t-test (\( p < 0.001 \)).

doi:10.1371/journal.pbio.1000147.g001
the two colored 5 mM sucrose solutions (PI for blue = 0.49 ± 0.09) (Figure 2). When 40 mM L-canavanine was added to the blue solution, their feeding behavior did not differ (PI for blue = 0.51 ± 0.09, Figure 2). Thus, these results show that L-canavanine is a repellent molecule and that Drosophila uses a chemosensory mechanism to detect this plant insecticide.

DmXR, Encoded by the mtt Gene, Is Required In Vivo to Detect L-Canavanine

To study whether DmXR was involved in L-canavanine detection in vivo, we first used a pharmacological approach. We hindered DmXR function by using the NMA antagonist in the two-choice feeding test. When present in the medium, this drug should diminish the repulsive action of L-canavanine. We first tested whether NMA alone could influence the fly feeding behavior: WT flies were allowed to choose between a blue solution containing 30 mM NMA versus a red control solution, both containing 5 mM sucrose. We found that flies were insensitive to 30 mM NMA (PI for blue = 0.80 ± 0.04 and 0.83 ± 0.04 in control and NMA-fed flies, respectively, Figure 3A).

Figure 2. L-Canavanine triggers a chemosensory repulsive effect. Two-choice feeding tests allowed the measure of the preference between the blue and the red solutions of wild-type (WT) and chemosensory defective pox-neuro (poxn) mutant flies. A preference index (PI) value of 1 or 0 indicates a complete preference or aversion for the blue solution, respectively. A value of 0.5 indicates no preference. See Materials and Methods for the calculation of the PI. In our conditions, WT flies prefer the blue solution (white). When L-canavanine (Cana, black) was added at the indicated concentration (1 to 40 mM), it inhibited the intake of the blue solution in a concentration-dependent manner, being very aversive at 30–40 mM. In contrast, poxn flies fed equally on the blue solution with (black) or without (dotted) 40 mM L-canavanine, indicating a chemosensory action for the plant amino acid. Error bars indicate SEM. The single and double asterisks indicate significant differences by t-test (p < 0.05 and p < 0.001, respectively) between WT flies that fed on the blue solution without (white) or with (black) L-canavanine.

doi:10.1371/journal.pbio.1000147.g002

Figure 3. The L-canavanine-induced repulsive behavior requires the DmXR function. (A) Pharmacological inhibition of DmXR by the NMA antagonist reduces L-canavanine repulsive effect. Histograms show the preference index (PI) for the blue solution from two-choice feeding assays in which drugs have been added to the blue solution. There was no significant difference in the behavior of WT flies (WT) that fed similarly on the blue control solution (white) and on the blue solution containing either 30 mM NMA (diagonally hatched) or 30 mM L-arginine (Arg) (horizontally hatched). Significantly more flies fed on the blue solution containing 20 mM L-canavanine (Cana) + 30 mM NMA (dotted) than on the blue solution containing 20 mM L-canavanine (grey). L-arginine (30 mM) had no effect on 20 mM L-canavanine repulsive effect (checkered). Error bars indicate SEM. Asterisks indicate significant differences by t-test (p < 0.001). (B) mtt mutant flies are insensitive to L-canavanine. Histograms show the preference index (PI) for the blue solution from two-choice feeding assays by using the blue solution without (white) or with (black) 30 mM L-canavanine. WT and the genetic background control f01266/f01266 flies are strongly repulsed by L-canavanine and avoid feeding it. In contrast, mtt flox/flox and mtt flox/DfEx7096 mutant flies are insensitive to L-canavanine, as they fed similarly on the blue solution.
with or without L-canavanine. Error bars indicate SEM. Asterisks indicate significant differences between the intakes of the blue solution without or with L-canavanine by t-test (p<0.0001).

doi:10.1371/journal.pbio.1000147.g003

However, significantly more flies fed on 20 mM L-canavanine in presence of 30 mM NMA (PI for blue = 0.68 ± 0.07) than on 20 mM L-canavanine alone (PI for blue = 0.36 ± 0.07) (Figure 3A). L-Arginine (30 mM), which is inactive on DmXR, had no effect on L-canavanine-induced repulsive behavior (PI for blue = 0.41 ± 0.03, Figure 3A). Thus, blockade of DmXR with the antagonist NMA lowered the repellent effect of L-canavanine.

We then used a genetic approach, taking advantage of two fly lines in which the DmXR encoding gene that we called mttf06268 has been inserted into the mtt gene as determined by Exelixis sequence analysis [32], and the Df(2R)Exel7096 line completely removes the mtt locus [33] (Figure 4A). Both mutants are viable in homozygous conditions. We expected that the insertion of a transposon 35 bp downstream from the third exon of mtt would disrupt the transcription of the gene. Indeed, f06268 homozygous flies are mtt loss-of-function mutants since no RNA was detected by quantitative real-time reverse-transcriptase polymerase chain reaction (QRT-PCR) in adults (Figure 4B). In homozygous Df(2R)Exel7096 flies, mtt RNA was also not detectable by QRT-PCR (Figure 4B). As a control for genetic background, we used flies homozygous for the f01266 piggyBac transposon line [32] (Figure 4A), which express normal levels of mtt RNA (Figure 4B).

In control two-choice feeding tests without L-canavanine, homozygous mttf06268, hemizygous mttf06268/Df(2R)Exel7096, and homozygous f01266 flies behaved as WT (Figure 3B). When 30 mM L-canavanine was added to the blue solution, mttf06268 and mttf06268/Df(2R)Exel7096 flies fed on this blue solution (PI for blue = 0.71 ± 0.06 and 0.73 ± 0.09, respectively), whereas WT and f01266 flies were repulsed (PI for blue = 0.16 ± 0.06 and 0.23 ± 0.08, respectively) (Figure 3B). Thus, mtt mutant flies are insensitive to 30 mM L-canavanine.

Both pharmacological and genetic data lead to the conclusion that DmXR may be the L-canavanine-sensitive receptor in chemosensory organs, because us localization hinders the feeding. Since DmXR may be the L-canavanine–sensitive receptor in chemosensory organs, we wondered whether this receptor was actually localized in these organs.

mtt Is Expressed in Gustatory Sensilla

Drosophila, like other insects, base their feeding decisions on the presence or absence of specific volatile and nonvolatile chemicals present in the food. Volatile chemicals are in general detected by olfactory neurons, located mainly on the antenna, whereas nonvolatile chemicals like amino acids are detected by gustatory receptor neurons (GRNs). GRNs are present in taste sensilla localized in the legs, the labial palps (or labellum) found on the tip of the proboscis, and within the pharynx (called internal taste organs) [34]. As flies walk on their food sources, tarsal gustatory sensilla evaluate their chemical contents. If phagostimulants are present, the fly extends its proboscis, enabling labellum sensilla to have contact with the food. In the labellum, gustatory chemosensilla house two to four GRNs as well as a single mechanosensory neuron [35–37]. In each sensillum, the different subsets of specialized taste neurons are activated by specific classes of tastants, allowing Drosophila to detect sugars, bitter compounds, and water [4,38].

To investigate whether mtt is expressed in gustatory sensilla, we first performed QRT-PCR experiments on dissected labellum and tarsi/tibiae of WT flies. As shown in Figure S2, mtt RNA is expressed in both organs bearing gustatory sensilla. To assess whether mtt would be present in GRNs, we compared the expression level of mtt RNA in WT to that found in poxn mutants, where chemosensory neurons are transformed into mechanosensory neurons. A significant decrease of the amount of mtt RNA was detected in the labellum and the tarsi/tibiae of poxn mutant compared to WT (Figure S2), suggesting that mtt is expressed in some GRNs. To visualize whether mtt is indeed expressed in gustatory chemosensilla, we then performed in situ hybridization experiments on labellum of WT flies. We found that mtt riboprobe hybridized to a single neuron-like cell within some chemosensory sensilla (Figure 5A–5D). These sensilla, which house five neurons, are clearly gustatory sensilla, because mechanosensory sensilla contain a single, mechanosensory neuron [39]. In situ labeling

Figure 4. The f06268 and the Df(2R)Exel7096 lines are mtt null mutants. (A) Schematic diagram showing the structure of the mtt gene and the insertion sites of the transposon lines used in this study. Black boxes show coding regions of mtt and CG12780 CDSs. Black arrow shows the most distal transcription start site (+1) of mtt (predicted by Flybase). Arrowhead indicates the orientation of transcription of the CG12780 gene. The f06268 line carries a piggyBac transposon inserted 35 bp downstream the third exon of mtt exon. The f01266 piggyBac transposon and the NP4288-GAL4 enhancer trap lines are inserted 1.7 Kb downstream and 3.1 Kb upstream, respectively, from the transcription start site of mtt. The Df(2R)Exel7096 line corresponds to a small deficiency that completely removes the mtt locus and some adjacent genes (CG8697 to CG2397). (B) QRT-PCR analysis on mtt mutants. The expression levels of mtt RNA on homozygous wild-type (WT), f06268, Df(2R)Exel7096, and f01266 adults flies were measured by quantitative real-time reverse-transcriptase polymerase chain reaction (QRT-PCR). WT and f01266 homozygous flies show comparable levels of mtt RNA, whereas no amplification is obtained from the f06268 and the Df(2R)Exel7096 homozygous lines. Error bar indicates SEM.

doi:10.1371/journal.pbio.1000147.g004

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appeared to be specific for mtt because chemosensory sensilla were not labeled in labellum from Df(2R)Exel7096 mtt mutant flies (Figure 5E). Altogether, QRT-PCR and in situ hybridization data indicate that mtt is expressed in only one GRN per labellar chemosensillum, consistent with a role of DmXR as a taste receptor.

Due to its very low level of expression, we were unable to use the in situ hybridization technique combined with immunocytochemistry to further analyze the mtt expression pattern. Hence, in order to investigate the nature of the GRNs expressing mtt, we took advantage of a GAL4 enhancer trap line, NP4288-GAL4, inserted 3.1 Kb upstream from the transcription start site of mtt (Figure 4A). A similar strategy had been undertaken for many other Grs, also reported to have a low level of expression [40]. The expression patterns of these receptors were analyzed using GAL4 transgenes containing taste receptor promoters [29,40–42] or enhancer traps such as NP1017 [43]. These studies have shown that Gr66a-GAL4, Gi5a-GAL4, and NP1017-GAL4 lines drive specific expression in bitter-, sugar-, and water-sensitive GRNs, respectively [29,42,43]. When NP4288-GAL4 was crossed with a green fluorescent protein (GFP) reporter line, we observed GFP-positive neurons in taste organs. In the labellum, chemosensory sensilla contained one GFP-positive neuron (Figure 5F–5I), in accordance with the in situ hybridization data. We observed around 28 GFP-positive neurons per labial palp, and two in each tarsus of the legs (Figure 5J and 5K, and Table 1). In addition, four GFP-positive neurons were present in the labral sense organs (LSO) and in the ventral cibarial sense organs (VCSO) (unpublished data), which are bilaterally symmetrical internal taste organs located in the pharynx [44]. Interestingly, Gr66a-GAL4, but not Gr5a-GAL4 or NP1017-GAL4, drives also expression in the LSO and VCSO (Table 1), suggesting that NP4288 is expressed in bitter-sensitive GRNs. To determine whether NP4288-GAL4 and Gr66a-GAL4 are indeed coexpressed, we analyzed transgenic fly lines expressing UAS-nlsGFP under the control of both NP4288 and Gr66a GAL4 drivers, and then counted and compared the number of GFP-positive neurons to that of flies

Figure 5. mtt is present in chemosensory sensilla. (A–D) and (E) show mtt in situ hybridization on WT and Df(2R)Exel7096 mtt mutant labellum, respectively. (A) In WT labellum, single cells (arrows indicate two single cells in focus) are labeled by the mtt riboprobe. (A) is a composite image of one image focusing on the labeled cells and one image focusing on the chemosensory bristles. (B) and (C) are high-magnification views of one sensilla visible in (A) (upper arrow). (B) focuses on the labeled cell. (C) is a composite image of (B) and a different focal plane in which the nearby chemosensory bristle is visible. Note that one single neuron-like cell (indicated by an arrow) is labeled near the chemosensory bristle visible in (C). This cell is likely a neuron, as an axon is weakly visible (indicated by an arrowhead). (D) shows another sensilla with a different Nomarski setting illustrating that the single labeled cell (indicated by a black arrow) is present in a chemosensilla housing at least five cells. (E) No signal is detected on Df(2R)Exel7096 mutant labellum. (F) is a composite image of one image focusing within the labellum and one image focusing on the chemosensory bristles. (F–I) Labellum of NP4288-GAL4/UAS-nlsGFP showing the GFP-expressing taste neurons. (F) In one labial palp, around 28 GRNs are present (see also Figure S3). (I) is a composite image of the fluorescent and bright-field images shown in (G) and (H), respectively. Arrowhead in (G and I) indicates the dendrite innervating the taste bristle sensilla. (J and K) NP4288-GAL4/UAS-mCD8GFP double homozygous foreleg showing that taste neurons are labeled in the tarsi. Arrowhead in (J) indicates the dendrite of the taste neuron. Note that only one of the two labeled neurons is visible at this focal plane. (K) is a composite image of the fluorescent and bright-field images.

doi:10.1371/journal.pbio.1000147.g005
containing each driver alone. In flies that express either the NP4288-GAL4 or Gr66a-GAL4 driver, an average of 28.4 and 26.6 neurons were observed per labial palp, respectively (Figure S3). In flies that express both drivers, an average of 28.8 neurons were detected per palp (Figure S3). Furthermore, we also observed coexpression of both drivers in the LSO and in the foreleg tarsi (table in Figure S3). This indicates that most, if not all, GRNs that express Gr66a also express NP4288, suggesting that this transgene reflects a role for DmXR in bitter-sensitive GRNs. Altogether, these results show that mtt is expressed in one GRN per sensilla that likely correspond to the bitter-sensitive GRNs.

### Leg GRNs Are Sensitive to l-Canavanine via DmXR

To investigate whether GRNs are sensitive to l-canavanine, we examined a direct behavioral measure of leg GRN stimulation by the proboscis extension reflex (PER) paradigm: when the leg tarsi encounter an attractive sugar solution, the proboscis often extends [42,43,45]. If a toxic or bitter compound is added to the sugar solution, the PER is inhibited [42,43,45]. This assay enables the application of the drugs only on the legs, which carry solely taste sensilla. In addition, we took care that the proboscis never touched the drugs when it extended, so that we were sure that there was no ingestion of drugs (and consequently, no central effect of these drugs). Using the classical PER paradigm (5-s stimulation by touching the leg tarsi either with a 100 mM sucrose solution or with a 100 mM sucrose+40 mM l-canavanine solution), we found that the occurrence of PER was not affected by l-canavanine in WT or mtt mutant flies (Figure 6). However, after the PER, WT flies usually sustain their proboscis extension to search for food when their legs are maintained in contact with the sugar solution as shown in Video S1. This sustained phase was strongly affected by l-canavanine, since significantly more flies prematurely retracted their proboscis (70±5% of proboscis retraction [PR]) compared to 25±4% of PR for the sucrose solution, Figure 6). This PR phenotype occurred generally just after the proboscis extension (Video S1). We then tested whether the l-canavanine–induced PR phenotype requires DmXR and found that this phenotype disappeared in the mtt loss-of-function mutants (around 25% of PR, Figure 6). Altogether, these data indicate that DmXR is required in leg GRNs for the l-canavanine detection.

### DmXR Is Required in Bitter-Sensitive GRNs for l-Canavanine Sensitivity

To determine in which GRNs mtt is required, we established transgenic flies carrying a mtt RNA interference (RNAi) construct under the control of UAS sequence [46]. We first expressed mtt RNAi with NP4288-GAL4 in heterozygous mtt^F0268^ flies and tested the effect of l-canavanine by the PER/PR behavioral assay. The occurrence of PER was not affected by l-canavanine in controls (Cana) were put in contact with the leg tarsi during the 5-s assay. l-Canavanine (Cana) does not affect the percentage of proboscis extension reflex (PER) in WT and mtt mutant (mtt^F0268^ and mtt^F0268^/DfEx7096) flies. After the PER response, l-canavanine triggers the proboscis retraction (PR) in WT, but not in mtt mutant flies. Error bars indicate SEM. Asterisks indicate significant differences by t-test (p<0.001).

### Table 1. Expression patterns of the GAL4 lines described in this study.

<table>
<thead>
<tr>
<th>Labelum</th>
<th>Internal Taste Sensilla</th>
<th>Legs</th>
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<tbody>
<tr>
<td></td>
<td>LSO</td>
<td>VCSO</td>
</tr>
<tr>
<td>gr5a</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>gr66a</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>NP1017</td>
<td>Yes</td>
<td>No*</td>
</tr>
<tr>
<td>NP4288</td>
<td>Yes*</td>
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To analyze the expression of NP1017 and NP4288 driver lines, we crossed each of them with UAS-nlsGFP flies to visualize the presence of GFP-positive neurons. Also included are observations made by the groups that originally characterized the expression pattern of Gr5a, Gr66a, and NP1017 drivers [29,40,43,60].

This clearly indicates a requirement of mtt for l-canavanine detection. Thus, we next used the Gr66a-GAL4 driver to specifically express the mtt RNAi in Gr66a-GRNs of heterozygous mtt^F0268^ flies. As already observed with NP4288-GAL4–driven knockdown of mtt, the occurrence of PER was unmodified in this genetic condition (Figure S4A). Importantly, Gr66a-GAL4–induced knockdown of mtt significantly reduced the occurrence of l-canavanine–induced premature PR (Figure 7A). This clearly indicates a requirement of mtt in Gr66a-GRNs for l-canavanine sensitivity.

To clearly demonstrate that l-canavanine detection required the presence of DmXR only in Gr66a GRNs, we performed rescue experiments by targeting mtt expression in distinct types of GRNs of mtt homozygous mutants by using the GAL4/UAS system [46]. Indeed, several types of GRNs are present in the tarsi, such as sucrose-, bitter-, and water-sensitive GRNs (Figure S5).

![Figure 6. l-Canavanine does not affect the occurrence of PER, but induces a DmXR-dependent retraction of the proboscis after the reflex.](image-url)
Figure 7. mtt is required in Gr66a-GRNs. (A) Knockdown of mtt expression by RNAi in NP4288-GRNs and Gr66a-GRNs suppresses the L-canavanine (Cana)-induced PR gustatory phenotype. Histograms show the percentage of PR of controls (NP4288/+; UAS-mtt RNAi/+), UAS-mtt RNAi/+ and Gr66a:mttf06268/+/mtt f06268 and mtt heterozygous flies expressing the mtt RNAi construct in NP4288-positive GRNs (NP4288/mtt f06268;UAS-mtt RNAi/+). Compared to controls, the down-regulation of mtt in NP4288-GRNs or in Gr66a-GRNs suppresses the L-canavanine–induced PR. For all genotypes, L-canavanine does not significantly affect the percentage of PER (Figure S4). Behavioral analyses were performed as described in Figure 6. Error bars indicate SEM. Double asterisks indicate significant differences by t-test (p<0.001). (B) Expression of mtt in bitter-sensitive Gr66a-GRNs rescues the PR mutant gustatory phenotype of mtt mutant flies. Histograms show the percentage of PR of control flies (mtt f06268/+/UAS-mtt/+), mtt mutant flies carrying one copy of each GRN GAL4 (Gr66a:mtt f06268;mtt f06268, Gr5a:mtt f06268;mtt f06268, and NP1017:/mtt f06268) and mtt mutant flies expressing mtt in bitter-, sugar-, and water sensitive GRNs (Gr66a:mtt f06268;UAS-mtt+). Note that only mtt mutant flies expressing mtt in Gr66a-GRNs show a rescue of the L-canavanine–induced PR phenotype. doi:10.1371/journal.pbio.1000147.g007

Expression of mtt in the different subsets of GRNs did not affect the percentage of PER (Figure S4). We then analyzed the PR and found that expression of mtt in sugar or water GRNs did not rescue the mutant phenotype (Figure 7B). In contrast, expression of mtt in Gr66a-GRNs rescued L-canavanine sensitivity (Figure 7B). Thus, it is the stimulation of DmXR, in Gr66a-GRNs, which is responsible for L-canavanine–induced PR. To further verify that only Gr66a-GRNs are necessary for L-canavanine sensitivity, we expressed the hid and epr proapoptotic genes [47] in these neurons, or inhibited their neurotransmitter release with the tetanus toxin transgene [48]. In both cases, the PER was not affected (Figure S6), but the L-canavanine–induced PR was lost (Figure S6). Altogether, these results demonstrate that DmXR is expressed and required only in Gr66a-GRNs for L-canavanine detection.

DmXR Is a Gustatory Receptor

The sites of taste reception are localized to the dendrites of GRNs [36], which are bipolar neurons containing a single dendrite and a single axon. To confirm that DmXR was actually the L-canavanine taste receptor and not a regulatory receptor modulating Gr66a-GRN synaptic transmission, we performed two kinds of experiments. First, we expressed a HA-tagged receptor in Gr66a-GRNs to determine its subcellular localization in the labellum. As shown in Figure 8A, the receptor was highly concentrated at the dendrite and not detected in the GRN axon in accordance with a gustatory function. Second, we tested whether DmXR could modify Gr66a activation by other repellents than L-canavanine. It has been shown that Gr66a-GRNs are required for caffeine-aversive behavior, Gr66a being a gustatory receptor for caffeine [8]. As already published, we could find that caffeine inhibited sucrose-induced PER (Figure 8B). However, when the PER occurred, we noticed that there was a high rate of premature PR. The caffeine-induced PR phenotype occurred generally just after the PER, similar to what was observed with L-canavanine (Video S2). We then tested the caffeine-induced phenotypes in mtt mutants and did not find changes in caffeine-induced PER inhibition and caffeine-induced PR (Figure 8B). These data clearly demonstrate that DmXR acts as a L-canavanine gustatory receptor in Gr66a-GRNs.

Discussion

The ability to avoid ingestion of toxic plants compounds is crucial for insect survival. However, before the current study, only two receptors, Gr66a and Gr93a, which are essential for the caffeine response, were associated with a specific bitter tastant [7,8]. The nonprotein amino acid L-canavanine is known to be toxic to insects, when ingested ([18] and this study). Here, our results show that L-canavanine is detected as a repulsive molecule. With a pharmacogenetic approach, we have shown that Drosophila uses a taste detection mechanism mediated by the orphan GPCR, DmXR, which is activated by L-canavanine to trigger this avoidance behavior. This process occurs in bitter-sensitive GRNs where this receptor is expressed.

By using the two-choice feeding test, the repulsive effect of L-canavanine was clearly demonstrated. Contrary to the known repellents (caffeine, quinine [8,49]), L-canavanine does not affect...
the PER. However, l-canavanine triggers the retraction of the proboscis following its initial extension impairing the food intake. Indeed, after the PER, WT flies usually sustain their proboscis extension to search for food when their legs are maintained in contact with the sugar solution. This sustained phase is strongly affected by l-canavanine, since significantly more flies prematurely retracted their proboscis at this stage. This inhibition of sustained proboscis extension is not specific for l-canavanine, but is also observed, with the same rate, in the presence of caffeine. Hence, caffeine induces a fast response, which is the PER inhibition, and a slow response, which is the PR, whereas l-canavanine only induces the slow response. One open question is to understand the molecular mechanisms responsible for such a difference between caffeine and l-canavanine in the response dynamics, knowing that both drugs act on the same cell type (Gr66a-GRNs). GPCR-induced signal transduction pathways rely on the activation of intracellular heterotrimeric G-proteins [50–52]. To date, two G-proteins have been implicated in the taste pathway, and both are required for sugar perception [53,54]. However, a direct evidence for a coupling between Grs and G-proteins has not been demonstrated [38]. As we have shown in our cell transfection assay ([27] and this study), DmXR is a genuine GPCR. It is thought that the Gr66a receptor, like other Grs, is a putative GPCR [8,38]. However, we do not know to which type of G-protein these two receptors are coupled in vivo. A first explanation about the different dynamics induced by l-canavanine and caffeine could be that that DmXR and Gr66a are coupled to distinct G-proteins or that both receptors are coupled to the same G-protein, but with different efficiencies. A more speculative explanation may be due to the different structural features of the two receptors, taking into account that Grs are structurally related to the olfactory receptors in Drosophila [40]. As was recently shown, Drosophila olfactory receptors may act as ligand-gated channels instead of being coupled to a G-protein [55,56]. Thus, Gr66a receptor may also be a ligand-gated channel. Because of the absence of any intermediate, changes in membrane excitability would be more rapid in presence of caffeine compared to l-canavanine. This may explain the difference in the response dynamics between these two drugs.

This study shows that DmXR is expressed and required in bitter-sensitive leg GRNs. However, DmXR is also known to be expressed in the adult brain, in agreement with our observations that NP4288-GAL4 is expressed in this tissue (unpublished data). This suggests the existence of an unknown endogenous ligand, different from l-canavanine, triggering DmXR activation in the brain. To exclude any action of l-canavanine in the brain, we took care that flies avoided ingesting the drug solutions by applying them only on legs during the PER analysis. In addition, we used GRN-restricted drivers, allowing us to specifically analyze the peripheral function of DmXR. Finally, the absence of any defects in the caffeine-induced response of mtt mutants flies excludes a role of DmXR in second and higher order neurons involved in the resistance.

Figure 8. DmXR is localized to GRN dendrites and is dispensable for caffeine sensitivity. (A) Subcellular localization of the DmX receptor. Anti-HA immunostaining on labellum of Gr66a-GAL4/+;UAS-HAmtt/+ flies. HAmtt is a HA-tagged version of the DmX receptor. When DmXR is expressed in Gr66a-GRNs, it is principally targeted to the dendrite. Note the strong localization of the receptor at the tip of the dendrite (arrowhead). (B) The aversive effect of caffeine is not affected in mtt mutant flies. Behavioral analyses were performed as described in Figure 6 except that 100 mM sucrose (white) and 100 mM sucrose+50 mM caffeine (grey) solutions were used. Histograms show the percentage of PER and PR of control (WT) and mtt mutant (mtf06268/mtf06268) flies. For both control and mtt mutant flies, the caffeine inhibits the sucrose-induced PER. This effect is not complete as some flies still performed a PER in the presence of caffeine. Most of these flies retract their proboscis in response to caffeine, similar to what is observed with l-canavanine. Error bars indicate SEM. Double asterisks indicate significant differences by t-test (p<0.001).

doi:10.1371/journal.pbio.1000147.g008
control of the studied gustatory behavior. As we observed mtt expression in the labellum and in internal taste organs (LSO and VCSO), it is very likely that these taste sensilla also play a role in the L-canavanine–induced aversive behavior. In agreement with this, we observed that flies did not drink a L-canavanine/sucrose-containing solution when directly applied on the labellum (unpublished data), confirming the presence of L-canavanine-sensitive GRNs. So, we assume that DmXR is a L-canavanine-tuned gustatory receptor in all these taste organs.

Surprisingly, it is not a Gr member that has been selected to detect L-canavanine, despite the very high sequence diversity of this family. Indeed, DmXR belongs to the mGluR GPCR subfamily because of its close sequence relationship [27]. DmXR and mGluR LBP sequences and 3-D model comparisons have shown that DmXR has diverged only in the LBP part interacting with the γ-carboxylic group of glutamate [27]. These modifications have targeted and changed two residues that are conserved in all mGluRs and are crucial for glutamate-induced activation [24,27]. Our study shows that these structural changes are correlated with the ligand selectivity of the receptor. Indeed, DmXR has a divergent LBP so that glutamate is no more an agonist but L-canavanine is. Conversely, the Drosophila mGlu ortholog receptor, DmGluA, is not activated by L-canavanine. This suggests that the original conformation of the mGluR LBP was more adapted to diverge and to recognize L-canavanine than the one of the Grs. In addition, to give rise to this new type of gustatory receptor, appropriate GRN expression has also been added during the evolution of the DmXR function since mGluR expression is mainly found in the central nervous system [25]. Thus, our study suggests that other GPCRs, different from Grs, may have evolved in insects to recognize specific tastants.

Finally, the insect-borne diseases are largely increasing, partly due to the development of insecticide resistance. Thus, it becomes urgent to identify insect-specific targets for the design of new drugs against insects. Our work illustrates that the pharmacological and functional characterization of the insect-specific GPCRs, which may be promising targets. These findings may help to develop new strategies of chemical communication in the insect kingdom.

Materials and Methods

Chemicals and Pharmacology
1-glutamate, t-canavanine, t-arginine, and γ-N-methyl-t-arginine (NMA) were from Sigma. Human embryonic kidney (HEK 293) cells were cultured and transiently transfected by electroporation as previously described [27]. Carrier plasmid DNA (pRK5) (14 µg), plasmid DNA containing HA-DmXR WT, HA-DmXRT176A WT, DmGluRA (2 µg), and plasmid DNA containing Gzqi9 (2 µg) (to enable the artificial coupling of DmXR and DmGluRA to phospholipase C, [57]) were used for the transfection of 10^5 cells. Determination of inositol phosphate (IP) accumulation in transfected cells was performed as previously described [27].

Genetics, Fly Lines, and Constructs
Drosophila stocks were raised on standard fly food medium at 25°C on a 12-h light/dark cycle. WT Canton S flies were used as control flies in all behavioral assays. For experiments using mttneo [poxn] adult mutant flies, homozygous flies carrying the poxn−/− mutant allele were used [30]. mtt[br] and f01266 lines carry PBac(WH) transposon and are described in [32]. The Df(2R)E0179066 line carries a small deficiency that completely removes the mtt locus and some adjacent genes (CG8897 to CG2397) [33]. The p(UAS-mtt) transgene construct was generated by cloning the hemagglutinin N-terminally tagged full coding sequence of DmXR (HA-mtt) [27] into the pUAST transformation vector and injected into w^1118 embryos. Several insertion lines were obtained. After QRT-PCR analysis (unpublished data), the w^1118,UAS-mttRNAi1 line was chosen for this study. The mtt RNAi line was obtained after amplifying DmXR cDNA sequence with the sense primer 5’-ACT CCT GAT GAC GAG GAA A-3’ and the antisense primer 5’-CCG GGC TCT AGA ACA ACA G-3’. This sequence was digested with the XbaI restriction enzyme and subcloned into AvrII-digested pWIZ. This new construct was then digested with the NheI restriction enzyme and ligated with the same XbaI-digested PCR product. A clone with the second insert oriented opposite to the first was then selected and used for injection of w^1118 embryos. Several insertion lines were obtained. After QRT-PCR analysis (unpublished data), the w^1118,UAS-mtt RNAi1 line was chosen for this study. The Gr66a-GAL4 (II chromosome) and Gr5a-GAL4 (II chromosome) promoter GAL4 lines are generous gifts from H. Amrein (Duke University, United States). The NPl1017-GAL4 (X chromosome) enhancer trap line was kindly provided by T. Tanimura (Kyushu University, Japan). The NPl288 enhancer trap was obtained from the GETDB Stock Center (Kyoto, Japan) [58]. The UAS-hid,UAS-rpr RNAi line was a gift from J. R. Martin (Paris Sud University, France). The UAS-Te-F line was kindly provided by C. J. O’Kane (Cambridge University, England). The w;UAS-mCD8-GFP and the w;UAS-nlsGFP were obtained from the Bloomington Stock Center.

Behavioral Assays
Two-choice feeding preference test. The two-choice feeding preference test was performed as described elsewhere, with minor modifications [29]. To enhance the feeding activity during the tests, we used a 5 mM sucrose solution instead of 1 mM. At this concentration of sucrose, WT flies showed a preference for the blue solution as shown in Figure 1. For each trial, 3- to 5-d-old adult flies (30 males and 30 females) were starved on water-saturated cotton for 24 h. Flies were then placed on a 96-well microplate at 23°C to 25°C in the dark. Wells contained 5 mM sucrose test solutions in 0.3% agarose with 5 mg/ml erioglaucine dye (blue) or 20 mg/ml sulforhodamine B dye (red) in the alternating wells, both at pH 7.5. Sucrose, erioglaucine, and sulforhodamine B were from Sigma. After 2 h on the plates, the numbers of flies that were blue (N^b), red (N^r), or purple (N^o) were determined on the basis of the colors of the abdomens. The preference index (PI) values for the blue solution were calculated according to the following equation: (N^b+0.5N^o)/(N^b+N^r+N^o). In all the tests shown, the drugs (pH 7.5) were added to the blue solution. Similar results were obtained when L-canavanine was added to the red solution (unpublished data). At least eight independent trials were carried out for each point. Only trials in which at least 35% of the flies had fed were included for statistical analysis. The percentage of flies with purple abdomens, having ingested both blue and red food, was approximately invariant in our experiments (15%–25%). All error bars represent the standard error of the mean (SEM). Unpaired Student t-tests were used to check for significant differences between the indicated pairs of data.

Proboscis extension reflex (PER) and proboscis retraction (PR) tests. The PER was examined principally as described in [42,43]. The 3- to 6-d-old adult flies were maintained on fresh medium for 1 d. Flies were then starved on water-saturated cotton for 20 h. Before the assay, the tarsi of the flies were touched with a
water-saturated 3MM Whatman paper. If the drop of water induced PER, the fly was allowed to intake sufficient water. Each fly was tested during 5 s by touching only the leg tarsi with either a 100 mM sucrose solution or 100 mM sucrose+drug solution. Six to eight batches of 40–60 flies were tested for each solution and each genotype. The occurrence of premature PR was also determined during the assay. The percentage of PER represents the number of flies from a given genotype that showed the PER phenotype divided by the total number of flies. The percentage of PR represents the number of flies from a given genotype that showed the PER phenotype divided by the number of flies that have shown a PER. Unpaired Student t-tests were used to check for significant differences between the indicated pairs of data.

Quantitative RT-PCR (QRT-PCR) Analyses

Total RNAs were extracted from whole adult flies (for the analysis of mtt mutants) or dissected labella, tarsi, and tibiae (for the analysis of mtt expression) by using Trizol (Sigma). cDNAs were generated from 1 μg of total RNAs treated with DNase I (Ambion) by using random decamers (Ambion) and Moloney murine leukemia virus reverse transcriptase (Invitrogen). Real-time PCR was done using Applied Biosystems SYBR Green PCR mix according to the manufacturer’s instructions. PCR was done as follows: 10 min at 95°C followed by 40 cycles: 15 s at 95°C, 60 s at 60°C. Housekeeping genes used to normalize DmXR expression levels were RpL13, Tbp, and Pkh. Sequences of the primers are RpL13 5'-AGGAGGCGCAAGACAATCC and 5'-CTTGCCTGCAGTCTCCTGAG, Tbp 5'-CGTCGCTCCGGCAATTCC and 5'-TTTTCCTGCTGACCTTCCCA, Pkh 5'-TCCTGAAAGGTCTCAACAAAAATG and 5'-TCCACACGGTTTCGACGATCT, and DmXR 5'-CGAATGCAACTG-3' and Tbp 5'-TGAAGAAATCTCCTCGAAC-3'.

In Situ Hybridization

Labella were dissected from flies and collected in 4% paraformaldehyde in PBS with 0.05% Triton X-100. Fixation overnight at 4°C, samples were washed 6×10 min in PTX (PBS, 2% Triton X-100) at room temperature. Prehybridization was then done for 2 h at 55°C in hybridization buffer (HB) (50% formamide, 5× SSC, 0.5 mg/ml yeast RNA, 0.1 mg/ml Salmon Sperm DNA, 0.05 mg/ml heparin, 0.3% Triton X-100). Hybridization was performed overnight at 55°C with digoxigenin-labeled antisense mtt riboprobe derived from mtt cDNA and prepared according to the manufacturer’s instructions (Roche). Washes were performed at 58°C in HB followed by washes in HB/PTX mix (3/1, 1/1, and 1/3, respectively). After blocking in 0.5% Blocking Reagent (Roche) in PTX, samples were then incubated with anti-Dig-AP (Roche) overnight at 4°C. Samples were then washed 6×10 min in PTX, NBT/BCIP mix (Roche) was used to visualize the digoxigenin-labeled probe. Samples were mounted in 90% glycerol.

Immunohistochemistry

To visualize HA-DmXR protein expression, we dissected the labella from Gr66a-GAL4/UAS-HA+mtt/+ flies from adult head, fixed them overnight in 4% paraformaldehyde in 1× phosphate-buffered saline (PBS), 0.3% Triton X-100. Immunostaining was performed in 1× PBS 3% Triton X-100 and 0.5% Blocking Reagent (Roche). The following antibodies were used: monoclonal rat anti-HA (Roche; 1:200) and Cy3-conjugated donkey anti-rat (Jackson Immunoresearch; 1:300). Samples were mounted in Vectashield. Images were acquired using a Leica microscope and CoolSNAP camera.

Supporting Information

Figure S1. Structure of L-canavanine and L-arginine. L-canavanine (2-amino-4-guanidinoxybutyric acid) is a nonprotein amino acid synthesized by leguminous plants that are members of the Loteidae, a subfamily of the Leguminosae [13,14,16]. L-Canavanine has a structural analogy to L-arginine in that the terminal methylene group of arginine is replaced with oxygen. Found at: doi:10.1371/journal.pbio.1000128.s001 (0.56 MB PDF)

Figure S2. The expression levels of mtt RNA are strongly decreased in poxn mutant tarsi/tibiae and labellum. The relative RNA expression levels of mtt were evaluated in WT and poxn mutant flies by QRT-PCR. The RNA extraction was exclusively made from dissected tarsi/tibiae as well as labellum. Normalized gene expression of mtt was standardized to the relative quantities of three housekeeping genes (Rpl13a, Tbp, and Pkh). WT was arbitrarily assigned a value of 100%. Note the strong reduction of mtt expression in poxn mutant. Error bars indicate SEM. Double asterisks indicate significant differences by t-test (p < 0.001). Found at: doi:10.1371/journal.pbio.1000128.s002 (0.05 MB PDF)

Figure S3. NP1288-GAL4 and Gr66a-GAL4 drive GFP expression in the same GRNs. (A) Images showing the distribution of the GFP-positive cells in a labial palp of NP1288-Gal4/UAS-nlsGFP (NP1288), Gr66a-Gal4/UAS-nlsGFP (Gr66a), and Gr66a-Gal4+NP1288-Gal4/UAS-nlsGFP (NP1288+Gr66a) flies. (B) Table showing the average number of GFP-positive cells counted in taste organs. WT was previously done for the analyses of Gr expression [29,59], we compared the number of GFP-positive cells present in NP1288-Gal4/UAS-nlsGFP (NP1288), Gr66a-Gal4/UAS-nlsGFP (Gr66a), and Gr66a-Gal4+NP1288-Gal4/UAS-nlsGFP (NP1288+Gr66a) taste organs. Note that the number of GFP-positive cells observed in the foreleg tarsi of NP1288-GAL4, UAS-nlsGFP flies is higher than what is observed in NP1288-GAL4, UAS-mCD8GFP homozygous flies (see Figure 5E and 5F). This is likely due to GFP concentration in the nucleus compared to the membrane-targeted GFP. However, we can not confirm that all these cells are neuronal cells because axons and dendrites were not visible in the leg. We also observed that the number of GFP-positive cells was lower in NP293-GAL4, Gr66a-GAL4, UAS-nlsGFP (average = 2.6) than in NP1288-GAL4, UAS-nlsGFP forelegs (average number = 4.5). Although we do not have an explanation for this result, this discrepancy was already observed for Gr5a-related receptors: the number of GFP-positive cells was higher in Gr61a-GAL4, UAS-GFP (average number = 12) forelegs compared to Gr61a-GAL4, Gr5a-GAL4, UAS-GFP (average number = 9.6) and to Gr61a-GAL4, Gr64a-GAL4, UAS-GFP (average number = 9.6) [59]. Found at: doi:10.1371/journal.pbio.1000128.s003 (2.36 MB PDF)

Figure S4. The PER is not affected in rescue and loss of function experiments. (A) Knockdown of mtt expression by RNAi did not affect the PER response. Histograms show the percentage of PER of controls (NP1288+/+, mtt00528+/+; UAS-mtt RNAi/+, UAS-mtt RNAi/+, and Gr66a-mtt00528+/+) and mtt heterozygous flies expressing the mtt RNAi construct either in NP1288-positive GRNs (NP1288/mtt00528+/+; UAS-mtt RNAi/+ or in Gr66a-GRNs (Gr66a-mtt00528+/+; UAS-mtt RNAi/+). Compared to controls, the down-regulation of mtt in NP1288-GRNs or in Gr66a-GRNs did not affect the PER response. Behavioral analyses were performed as described in Figure 6. Error bars indicate SEM. (B) Expression of DmXR in Gr66a, Gr5a, and NP1017 GRNs of mtt mutant flies did not affect the PER response. Histograms show the
percentage of PER of controls (mttf06268/+;UAS-mtt/+/), mtt mutant flies carrying one copy of each GRN GAL4 (Gr66a:mttf06268/+;mttf06268, Gr5a:mttf06268/+;mttf0628, NP1017/+;mttf0628/mttf0628), and NP1017/+;mttf0628/mttf0628 and mtt mutant flies expressing mtt in bitter-, sugar-, and water-sensitive GRNs (Gr66a:mttf06268/+;mttf0628;UAS-mtt/+, Gr5a:mttf06268/+;mttf0628;UAS-mtt/+, and NP1017/+;mttf0628/mttf0628;UAS-mtt/+), respectively. For all genotypes, t-camavanine did not significantly affect the percentage of PER. Behavioral analyses were performed as described in Figure 6.

Found at: doi:10.1371/journal.pbio.1000128.s004 (0.05 MB PDF)

Figure S5. The GRN GAL4 lines drive expression of a GFP reporter in GRNs of the first leg tarsi. The expression patterns of GRN-GAL4 drivers in the first leg tarsi were visualized by GFP epifluorescence. GFP-positive neurons (arrowheads) are lateral views. Gr66a-GAL4/UAS-mCD8GFP is an anterior view. Found at: doi:10.1371/journal.pbio.1000128.s005 (0.48 MB PDF)

Figure S6. Gr66a-GRNs are necessary for L-camavanine-induced PR. By using the Gr66a-GAL4 driver, we targeted the expression of the proapoptotic genes (spr and hid) and the tetanus toxin light chain (TeT) to kill and inactivate Gr66a-GRNs, respectively. Results show the percentage of PER and PR on controls (UAS-srp;UAS-hid/+ and UAS-TeT/+), ablated (Gr66a/ UAS-srp;UAS-hid), and silenced (Gr66a/UAS-TeT) Gr66a-GRNs. Note that the PER was not affected in controls as well as when Gr66a-GRNs were ablated or silenced. Compared to controls, which show a high percentage of PR in presence of L-camavanin, the absence or the inactivation of Gr66a-GRNs abolishes the PR response. Behavioral analyses were performed as described in Figure 6. Error bars indicate SEM. Double asterisks indicate significant differences by t-test (p < 0.001).

Found at: doi:10.1371/journal.pbio.1000128.s006 (0.06 MB PDF)

Video S1. L-camavanin triggers the premature proboscis retraction after the PER. This movie illustrates the behavior of WT flies when their leg tarsi are in contact with either a 100 mM sucrose solution (sucrose) or a 100 mM t-camavanine (sucrose/camavanine) solution. Generally, caffeine inhibits the sucrose-induced PER. However, this effect is not complete as some flies extend their probosces as shown in this video. Similar to the t-camavanine-induced effect, the majority of those flies retract their probosces just after PER in the presence of caffeine.

Found at: doi:10.1371/journal.pbio.1000128.s008 (1.03 MB MOV)

Acknowledgments

We thank H. Amrein, T. Tanimura, J. R. Martin, and C. J. O’Kane for generously sharing fly stocks. We thank Jean-Philippe Pin for helpful advice in pharmacology. Many thanks to Chrystel Lafont and Muriel Gien-Assari for their assistance in video acquisition and processing. We thank Ayméric Clairier and Jean-Maurice Dura for helpful discussions. We are grateful to Christine Damibly-Chaudire for the fly mutants.

Author Contributions

The author(s) have made the following declarations about their contributions: Conceived and designed the experiments: CM LS JB MLP YG. Performed the experiments: CM LS BF MLP YG. Analyzed the data: CM LS JB MLP YG. Contributed reagents/materials/analysis tools: CM LS BF MLP YG. Wrote the paper: CM LS JB MLP YG.

References

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