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The *Drosophila melanogaster* Phospholipid Flippase dATP8B Is Required for Odorant Receptor Function

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Abstract

The olfactory systems of insects are fundamental to all aspects of their behaviour, and insect olfactory receptor neurons (ORNs) exhibit exquisite specificity and sensitivity to a wide range of environmental cues. In *Drosophila melanogaster*, ORN responses are determined by three different receptor families, the odorant (Or), ionotropic-like (IR) and gustatory (Gr) receptors. However, the precise mechanisms of signalling by these different receptor families are not fully understood. Here we report the unexpected finding that the type 4 P-type ATPase phospholipid transporter dATP8B, the homologue of a protein associated with intrahepatic cholestasis and hearing loss in humans, is crucial for *Drosophila* olfactory responses. Mutations in dATP8B severely attenuate sensitivity of odorant detection specifically in Or-expressing ORNs, but do not affect responses mediated by IR or Gr receptors. Accordingly, we find dATP8B to be expressed in ORNs and localised to the dendritic membrane of the olfactory neurons where signal transduction occurs. Localisation of Or proteins to the dendrites is unaffected in dATP8B mutants, as is dendrite morphology, suggesting instead that dATP8B is critical for Or signalling. As dATP8B is a member of the phospholipid flippase family of ATPases, which function to determine asymmetry in phospholipid composition between the outer and inner leaflets of plasma membranes, our findings suggest a requirement for phospholipid asymmetry in the signalling of a specific family of chemoreceptor proteins.

Introduction

In insects such as *Drosophila melanogaster* the detection of environmental odours is achieved by arrays of olfactory receptor neurons (ORNs) housed in different types of chemosensory hairs (sensilla) on two olfactory organs, the antenna and the maxillary palp. Each class of ORN is tuned to specific chemical signals by expression of different olfactory receptor genes. The responses of most insect ORNs are reliant on members of two large and divergent families of olfactory receptor proteins, the odorant receptor (Or) and Ionotopic glutamate-like receptor (IR) families. Or proteins are seven trans-membrane domain proteins that are topologically inverted in comparison to G-protein-coupled receptors [1,2], raising the question as to whether they do interact with G proteins. Indeed, several studies have concluded that Or-signalling is rather ionotropic, and that the functional receptor is a ligand-gated cation channel composed of a variable Or odorant-binding subunit and a co-receptor subunit called Orco [1–4]. Orco is required for Or proteins to be transported to the dendrites [5], and heterologous expression studies suggest Orco is also part of the functional receptor and is essential for the initial fast inward current upon ligand binding [6,7]. However, there is also genetic and pharmacological evidence for a slower metabotropic transduction cascade (for an excellent recent review see [8]). For example, heterologous expression experiments demonstrated a slower metabotropic current after Or stimulation, as well as an increase in cAMP [4]. Orco has been suggested to be activated by cAMP [4], and in addition it has been suggested that phosphorylation of Orco by protein kinase C (PKC) is required for its activation [9]. However, several studies of loss of function of Gr-encoding genes in *Drosophila* have yielded conflicting results [10–12]. A recent study suggests that metabotropic regulation of Orco regulates Or sensitivity [13]. Overall, the mechanism of Or signalling appears to be complex, with both ionotropic and metabotropic pathway involvement, and despite much investigation is not fully understood.

IR genes encode a very different family of receptors, they are three trans-membrane domain ligand-gated ion channels that are related to ionotropic glutamate receptors [14]. IR proteins form heteromers but are not reliant on Orco [15,16]. A third family of receptors, the gustatory receptors (Grs), are also seven trans-membrane proteins and are evolutionarily related to the Ors [17,18]. Where the Or and IR families both detect a range of structurally diverse odorants and function in many ORNs [19,20], only one functional class of ORN, specialized for detection of carbon dioxide (CO₂), expresses *Gr* genes [21,22]. Most other *Gr*
The olfactory systems of insects are fundamental to critical behaviours such as finding mates, food and host plants. Insects can detect a wide range of environmental cues using three different families of olfactory receptor proteins. Why insects have three different families of receptor genes, and how they function together, is not fully understood. Here we identified a new gene, \textit{dATP8B}, which is critically and specifically required for the function of only one of these receptor families in \textit{Drosophila}. \textit{dATP8B} is a member of the P4-type ATPases, or phospholipid flippases; these enzymes function in establishing a difference or asymmetry in lipid composition between the outer and inner leaflets of plasma membranes. This is thought to be important for many cellular membrane processes; however, specific functions of individual flippase proteins are not well described. We find that \textit{dATP8B} is required for the function of the odorant receptor family, but not the ionotropic-like and gustatory receptor families. This further highlights the functional differences between these receptor families and suggests a role for phospholipids in the signalling of a specific family of receptor proteins.

\textbf{Results} 

To identify new genes involved in olfactory neuron function we screened 482 lines carrying homoygous viable EMS-induced mutations on chromosome III (from the Zuker Collection \cite{Zuker}) for defects in olfactory responses. Electroantennograms (EAGs) were employed to measure voltage changes across the antennal epithelium in response to a set of odorants known to excite a variety of different ORN classes. We identified one line (\textit{i22}) in which homozygous mutant flies showed reduced EAG responses to most tested odorants compared to heterozygous controls (p<0.05, Fig. 1A). Neurons on the second olfactory organ, the maxillary palp, showed an equivalent reduction in response to odorants in the mutant line (p<0.05, Fig. 1B). In contrast to the general odorants, the EAG response to carbon dioxide, which is generated by one specific antennal ORN class (ab1C) and mediated by two gustatory receptor genes, was unaffected in the mutant (Fig. 1A).

We next used genomic deficiencies to map the mutation. We found that flies trans-heterozygous for the \textit{i2} chromosome and either \textit{Df(3R)Exel7312} or \textit{Df(3R)Exel8155} showed the same mutant phenotype as homozygous \textit{i2} flies (Fig. 1C). This localised the mutation to an 86.3kb genomic region that contains 14 annotated genes (Fig. 1D). Whole genome sequencing experiments identified a nonsense mutation in the \textit{CG14741} gene within this region (genomic location 7,902,447, mutation CG7902447T). \textit{CG14741} has seven predicted isoforms; the nonsense mutation is in the first coding exon common to all predicted isoforms and thus is predicted to severely truncate all proteins encoded by the locus. The \textit{CG14741} gene has not previously been functionally characterised. However, sequence comparisons reveal it belongs to the type 4 P-type ATPase family of integral membrane transporter proteins \cite{Gr}. This group of proteins includes four human members named ATPB1-4 and \textit{CG14741} as the sole \textit{Drosophila} representative. Henceforth we refer to the gene \textit{CG14741} as \textit{dATP8B}.

To confirm that the nonsense mutation in \textit{dATP8B} was the cause of the olfactory defects observed in the \textit{i2} line we examined an independent mutant allele, a line containing a \textit{piggyBac} element inserted in the coding region and predicted to affect all the isoforms (\textit{dATP8B}\textit{P02502}, Fig. 1D). Flies homozygous for the \textit{dATP8B}\textit{P02502} allele showed the same EAG defect as homozygotes for the original EMS allele (p<0.05, Fig. 1C). This phenotype was reverted when the \textit{piggyBac} insertion was precisely excised (Fig. 1E), confirming that the olfactory defect is due to the insertion in \textit{dATP8B}. We also showed that the EMS and \textit{piggyBac} insertion alleles failed to complement; trans-heterozygotes for the two mutant alleles also exhibited the olfactory defect (p<0.05, Fig. 1C).

Together, these data confirm that mutations in \textit{dATP8B} cause a severe reduction in ORN responses.

In our initial EAG recordings we noted that the \textit{dATP8B}\textit{P02} mutant flies had normal responses to carbon dioxide (CO$_2$). Unlike the other odorants tested, which are detected by ORNs that express members of the \textit{Or} receptor family, CO$_2$ is detected by the \textit{ab1C} ORN which expresses two members of the \textit{Gr} receptor family, \textit{Gr21a} and \textit{Gr63a} \cite{Gr21a,Gr63a}. These data suggested that the mutation might be specifically affecting \textit{Or}-expressing ORNs rather than all ORNs. To determine if this was the case we characterised mutant responses from selected ORN classes on the antenna whose responses are determined by members of all three different receptor gene families. In each neuron type we examined that expresses \textit{Or} genes we found that the responses to ligands were substantially reduced over a range of concentrations (Fig. 2). This was the case regardless of the morphological type of sensillum, as we found greatly reduced responses to 2-heptanone from the ab3B neuron in basiconic sensilla (Fig. 2A), to cis-vaccenyl acetate from the ab3B neuron in coeloconic sensilla (Fig. 2C). These experiments also confirmed a finding we had noted from our EAG recordings, namely that the responses of \textit{dATP8B} mutant flies, while greatly reduced, are not completely abolished. We found that at high odorant concentrations mutant neurons were still able to fire at relatively high rates (an average of 100 spikes per second; Figs. 2A–C). We also noted that the effect of the mutation can be different for different odorants activating the same receptor. For the ab3A neuron, which expresses \textit{Or22a}, we found that the sensitivity to its high affinity ligand ethyl hexanoate was reduced by three log steps (Fig. 2D). However, when we recorded responses from ab3A neurons for a lower affinity ligand, ethyl butanoate, we found that the curve is shifted much less (Fig. 2E). This suggests that sensitivity to different ligands of the \textit{Or22a} receptor is affected by the \textit{dATP8B} mutation to differing extents.

In contrast, we found that mutations in \textit{dATP8B} have no effect on responses of ORNs that express \textit{IR} or \textit{Gr} genes. This was the case for two ORN types that express \textit{IR} genes; the responses to
Drosophila Flippase and Odorant Receptors

A

EAG Amplitude (mV)

Odorants

B

EPG Amplitude (mV)

Odorants

C

EAG Amplitude (mV)

Odorants

D

3R

7940k 7920k 7900k 7880k 7860k 7840k 7820k 7800k

DF(3R)Exel8155

DF(3R)Exel7312

dATP8B (CG14741)

dATP8B gene

transcripts

RG

RH

RC

RF

RI

RB

II2 / f05203

E

EAG Amplitude (mV)

Odorants

+ / + f05203 / + II2 / f05203

+ / + f05203ex
1,4-diaminobutane of the ac2A neuron (Fig. 2F) and to propionic acid of the ac3A neuron (Fig. 2G) were unaffected in the mutant. Consistent with the initial EAG data, in dATP8B mutants the CO2 response from the ab1C neuron, which is determined by Gr genes, is not significantly reduced for any of the concentrations that evoke a wide range of excitation levels (Fig. 2H). We also performed recordings from gustatory neurons on the labellum expressing members of the Gr family, and showed that in dATP8B mutants the response to both sucrose and caffeine, mediated by different Gr receptors [23], was unaffected (Fig. 2I). Taken together, these data strongly suggest that dATP8B is specifically required for the function of Or-expressing neurons, and not for chemosensory neurons in general.

dATP8B was identified in a previous study as being present in the antennal proteome [27]. To confirm its expression in antennae and to determine the cell type in which dATP8B is expressed we performed immunohistochemistry using an anti-dATP8B antibody. Strong staining was seen within the shafts of the sensilla, the location of the outer dendrites of the ORNs (Fig. 3A). Staining was observed in both basiconic and trichoid sensilla, which both house neurons expressing Or genes. We could not easily visualise the sensilla shafts of the coeloconic sensilla. Staining was absent from the outer dendrites in the sensilla shafts in mutants for dATP8B (Fig. 3B). Signal was also observed in the inner dendrites and the cell bodies, however this signal was also observed in mutants for dATP8B, albeit more weakly, (compare Figs. 3A and B), and thus may be largely background staining.

Given dATP8B function seems to only be required in the Or-expressing neurons we next asked if it is specifically expressed in Or-expressing, but not IR or Gr-expressing, neurons. We confirmed that dATP8B is expressed in the Or-expressing neurons by showing that it co-localises with Orco (Fig. 3C). We then asked if dATP8B is expressed in the ab1C neurons that express Gr21a. These neurons are housed in ab1 basiconic sensilla together with three other neurons, which all express Or genes. We visualised the ab1C neurons by using Gr21a-Gal4 to drive the membrane-localised mCD8:GFP and staining with anti-GFP. Strong GFP signal was observed in the outer and inner dendrites and cell bodies (Fig. 3D). Double staining with anti-GFP and anti-dATP8B showed that dATP8B is absent from the outer dendrites of Gr21a-expressing (ab1C) neurons (Fig. 3D). We saw many examples where the GFP-positive dendrites of the ab1C neurons ran parallel to but did not overlap dATP8B-positive dendrites within the sensilla shafts. We conclude that dATP8B localises to the outer dendrites in Or-expressing neurons, and likely does not in Gr-expressing neurons.

We also showed that dATP8B function is required in the ORNs, rather than another cell type, by using RNA interference (RNAi) to knock down dATP8B in the Or-expressing ORNs using an Orco driver. In Orco-Gal4:UAS-dATP8B RNAi flies the EAG response was significantly reduced for some of the odorants for which we also saw reduced responses in the two dATP8B mutant alleles, for example methyl salicylate and ethyl hexanoate (Fig. 3E). Although this defect is much less severe than seen in the loss of function mutants (Fig. 1A), taken together with the immunohistochemistry data these results suggest that dATP8B has a functional role in ORNs, rather than in another cell type such as support cells. As Orco has a relatively late onset of expression in pupal development [5], this result also suggests that dATP8B plays a role in these neurons after the initial development of olfactory sensilla.

The phenotype of dATP8B mutants bears a strong resemblance to that of mutations in the Orco gene, which is required for the localisation and function of the Or receptors, but not for the IR or Gr receptors [5,15,16]. We thus asked if loss of dATP8B caused Orco itself and/or the other Or proteins to be incorrectly localized. We tested this by using antibodies to examine the localization of both Orco and Or22a. In wild type flies anti-Orco staining is seen in both the outer dendrites and the cell bodies, for anti-Or22a there is strong staining in the outer dendrites only (Fig. 4). No difference in localization of either Orco or Or22a was observed in dATP8B mutants (Fig. 4). In addition, no noticeable difference was observed in the length or shape of the dendrites in the mutant. This confirmed an initial finding that the overall appearance of the olfactory sensilla in the mutant is normal (Fig. S1).

Discussion

dATP8B is a member of the type 4 subfamily (P4-ATPases) of P-type ATPases. Unlike the other subfamilies, most of which encode ion transporters, the P4-ATPases are believed to share a distinct function as phospholipid translocases or “flippases” [26]. Eukaroytic plasma membranes have an asymmetrical distribution of phospholipids across the bilayer, with sphingolipids and phosphatidylcholine enriched in the exoplasmic leaflet, and more polar lipids such as phosphatidylserine and phosphatidyethanolamine enriched in the cytoplasmic leaflet. Phospholipid flippases contribute to asymmetry by “flipping” phospholipids from the exoplasmic to the cytoplasmic leaflet [28]. The physiological significance of this asymmetry is not well understood but it seems to be important for critical membrane processes such as vesicle trafficking and intracellular signalling [26,29]. Disruption of the asymmetry may affect the conformation, membrane insertion, or trafficking, of integral membrane proteins. Alternatively it could affect lipid-signalling molecules, or membrane fluidity or bending. There are six members of the phospholipid flippase family in Drosophila and C. elegans and 14 in humans [26]. Genetic studies in C. elegans have suggested that the different members have different functions [30]. One of the six Drosophila flippases (CG33298) has
been implicated in secretory vesicle formation and cholesterol homeostasis [31], the others have been uncharacterized to date. dATP8B is the *Drosophila* homologue of four mammalian ATP8B proteins [26]. Of these only ATP8B1 has been substantially studied. Mutations in human ATP8B1 are associated with intrahepatic cholestasis [32] and also cause hearing loss [33]. The protein localizes to the apical plasma membrane of hepatocytes [32,34], where it is thought to play a role in protection from the detergent effects of bile salts [35]. In hair cells, the sensory cells of the inner ear, ATP8B1 has been shown to localize to the stereocilia that transduce the mechanical vibrations in the cochlea [33]. ATP8B1 deficient mice exhibit a progressive degeneration of cochlea hair cells, possibly due to changes in the mechanical properties of stereocilia or the disruption of a Ca2+ transporter crucial for sensory transduction [33].

Here we have found that dATP8B is essential specifically for the responses of *Or*-expressing neurons, and not for *IR* or *Gr*-expressing neurons in general. Our expression studies suggest that

**Figure 2.** dATP8B is required for responses of *Or* but not *IR* or *Gr*-expressing sensory neurons. (A–C) Mutations in dATP8B reduce the sensitivity of *Or*-expressing neurons to their major ligands. Dose-response curves for neurons located in three different morphological types of sensilla are shown; (A) ab3B neurons in basiconic sensilla (*Or85b*) to 2-heptanone, (B) at1A neurons in trichoid sensilla (*Or67d*) to cis-vaccenyl acetate, (C) ac3B neurons in coeloconic sensilla (*Or35a*) to Z3-hexenol. In all cases sensitivity is significantly lowered for all doses in homozygous dATP8Bf05203 flies (red) compared to controls (black, t-test, Bonferroni, n = 6–9, recorded from 4–8 flies). (D–E) Mutations in dATP8B differentially alter responses of ab3A neurons (*Or22a*) to a major ligand ethyl hexanoate (D) and a minor ligand ethyl butanoate (E). (n = 7 sensilla, recorded from 5–6 flies). (F–I) Neurons expressing *IR* or *Gr* receptors are not affected by dATP8B mutations. Four different types of neurons are shown; (F) responses of ac2A neurons in coeloconic sensilla (*IR41a* and *IR76b*) to propionic acid, (G) responses of ac3A neurons in coeloconic sensilla (*IR75abc* and *IR76b*) to Z3-hexenol, (H) responses of ac3A neurons in coeloconic sensilla (*IR75abc* and *IR76b*) to CO2, (I) responses from single neurons in labellar taste sensilla expressing *Gr* genes to 100 mM sucrose (Suc) or 10 mM caffeine (Caff) (mean ± SEM). In all cases control and mutant responses are not significantly different (n = 6–10 sensilla from 3–5 flies, t-tests, Bonferroni). Controls are either wild type or heterozygous dATP8Bf05203 mutants.

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Figure 3. dATP8B is expressed and required in Or-expressing olfactory receptor neurons. (A–A') dATP8B protein localises to the dendrites of ORNs. 14 μm thick antennal sections from wild type flies (CS-S) were stained for anti-dATP8B (magenta). Strong anti-dATP8B staining is seen in the shafts of the sensilla (the location of the outer dendrites) of basiconic (arrows) and trichoid (arrowhead) sensilla. Staining is also seen in the inner dendrites and cell bodies, however this staining is also present in dATP8B mutants. (B) The strong outer dendrite staining of anti-dATP8B is absent in dATP8B mutants (ll2/Df(3R)Exel8155), indicating it is specific for dATP8B. (C) dATP8B and Orco co-localise in the outer dendrites. 14 μm thick antennal sections from wild type flies were stained for anti-dATP8B (magenta) and Orco (green). (D) dATP8B is absent from the outer dendrites of Gr21a-expressing (ab1C) neurons. 14 μm thick antennal sections from Gr21a->mCD8:GFP flies were stained for anti-dATP8B (magenta) and anti-GFP (green). (E) In Orco-GAL4: UAS-dATP8B RNAi flies (red bars) the EAG response is significantly reduced compared to controls (black and grey bars, t-test,
dATP8B is not expressed in the latter, although further studies are needed to confirm the generality of this. Several lines of evidence suggest that mutation in dATP8B is not disrupting the development or morphology of the Or-expressing neurons and that its function is required in adult ORNs. First, in spite of the dramatic effects on sensitivity to odorants, the affected ORNs are still functional neurons, as evidenced by some response at high odorant doses, and have normal morphological appearance. Second, we found two examples where Gr or IR-expressing unaffected neurons (ac3A and ab1C) are housed in the same sensilla as affected Or-expressing neurons. This indicates that the defect is intrinsic to the Or-expressing neurons, and is not due to altered sensillum morphology or perireceptor processes.

Our data suggest that dATP8B is not required for the localization of the Or proteins to the dendrites, although we note that we have only examined Orco and one ligand-binding Or protein (Or22a) and thus cannot completely rule out effects on localization of other Or proteins. Nonetheless, it seems most likely that dATP8B is necessary instead for the function of the Or receptor complex and for primary receptor signal transduction processes. This is further supported by our finding that dATP8B mutations differentially affect sensitivity of an individual receptor to different odorants. At present there is no biochemical evidence that dATP8B functions as a phospholipid flippase, however all members of this family of P4-ATPases for which biochemical assays have been performed do function to translocate phospholipids [26,28,29]. If dATP8B functions as expected there are several ways it could affect Or signal transduction. An altered phospholipid composition of the plasma membrane in dATP8B mutants may affect Or-Orco interactions, or interfere with binding of odorants to the receptor complex. Alternatively, reduced availability or activity of membrane-localised signalling molecules may affect Or signalling. For example, phosphatidylinositol (PS) is required for activation of PKC once it translocates to the plasma membrane in response to increases in diacylglycerol [36]. In dATP8B mutants reduced availability of PS in the inner leaflet could thus lead to loss of PKC activation. PKC has functions in many signalling pathways, and has been suggested to be important for Orco activation [8]. Another possibility is that mutations in dATP8B affect the minor plasma membrane phospholipid phosphatidylinositol 4,5-bisphosphate (PIP2). PIP2 is found primarily in the cytoplasmic leaflet of the membrane and does not flip between leaflets, but levels could potentially be disrupted by flippase disfunction. PIP2 has many signalling roles. Its cleavage products inositol 1,4,5-trisphosphate and diacylglycerol are key components of G protein-activated signalling pathways. In addition, a number of families of ion channel and ion transporter proteins, for example the transient receptor potential channels, are dependent on PIP2 for their activation [37].

In conclusion, we have identified a new olfactory gene, dATP8B, which is specifically required for odour responses of Or-expressing, but not Gr or IR-expressing, sensory neurons in Drosophila. Given the very high level of homology of dATP8B to known phospholipid flippases, our findings suggest a specific role for cell membrane phospholipids in Or receptor signalling, as well as providing further evidence for fundamental differences between the signalling mechanisms of the different families of insect olfactory receptors. Further studies of this interesting gene may provide insight into the potentially complex mechanisms of Or signalling.

Materials and Methods

Drosophila stocks

Drosophila stocks were reared on yeast and cornmeal in 30 ml vials at 22°C under a natural daylight cycle. All crosses were performed at 22°C. Flies carrying the dATP8B102 mutation were part of a collection of mutagenized stocks obtained from Charles Zuker’s laboratory [25]. The piggyBac insertion line dATP8B5201 (BL18847) and deficiency lines Df(3R)Exel7312 (BL7966) and Df(3R)Exel155 (BL7967) were obtained from Bloomington Stock Center. The RNAi line for dATP8B was obtained from the Vienna Drosophila RNAi Center (v102648). The Orco-Gal4 line was obtained from Leslie Vosshall (Rockefeller University) and the Gr21a-Gal4 line from Kristin Scott at University of California Berkeley.

Electrophysiological recordings and data analysis

Recordings from whole olfactory organs. We recorded electrical signals from whole antennae (electroantennogram, EAG) and maxillary palps (electropalpogram, EPG) as described in Tom et al. 2011 [38]. A single fly was immobilized and a reference electrode inserted in the eye. For EAGs the recording electrode was placed on the surface of the antenna and for EPGs on the palp. Changes in voltage (mV) in response to 1 s stimulation with odorants were amplified using an active probe and a serial-
amplifier (Syntech, Hilversum, the Netherlands). EAGs and EPGs represent the summed activity of a population of ORNs.

Recordings from single olfactory sensilla. Activity of individual olfactory receptor neurons was studied using the single sensillum recording (SSR) technique as described elsewhere [39]. Action potentials were recorded by a glass electrode inserted at the base of an olfactory sensillum and amplified via an IDAC-4 amplifier (Syntech). Action potential firing rates during 500 ms stimulations were analysed by subtracting firing rates during the 2 seconds before stimulation. Action potentials from the different neurons in a single sensillum were separated as in de Bruyne et al. [39]. A two-tailed Student’s t-test with a Bonferroni correction for multiple comparisons was used to compare firing rates. Odor stimulation for SSR, EAG and EPG recordings was by injecting volatiles from 5 ml disposable syringes into a airstream blown over the preparation. All odorants were at highest available purity (>98%, Sigma-Aldrich) and dissolved in paraffin oil at different dilution from 0.0001 to 10% v/v. Because of its low volatility, the Drosophila pheromone cis-vaccenyl acetate (>98%, Cayman Chemicals) was dissolved in hexane and delivered from a Pasteur pipette that was briefly heated prior to use. Male flies of age 3–7 days were used for all electrophysiological recordings, except for the RNAi experiment where newly emerged male flies were incubated at 25°C for exactly 7 days before recordings were performed.

Recordings from taste sensilla. Single sensillum tip recordings were performed from large (L-type) and intermediate (I-type) sensilla in the labellum as described earlier [20]. Male flies were aged 3–7 days and prepared for recordings by insertion of a glass micropipette reference electrode filled with Ringer’s solution. Tastants were dissolved in 30 mM tricholine citrate, which was used as the electrolyte for the recording electrode. Action potentials obtained by using a TasteProbe and IDAC-4 amplifier (Syntech), were counted during the 500-ms period after initial contact with the stimulus solution in the recording electrode, and multiplied by two to obtain firing rates in spikes per second. A two-tailed Student’s t-test was used to compare firing rates between mutant and wild-type flies. Sucrose and caffeine were purchased from Sigma Aldrich.

Genome sequencing and data analysis

Genomic DNA was extracted from adult heterozygous males using a QIAGEN Genomic-tip 20/G. A paired-end library with ~300 bp insert size was prepared and sequenced by the Australian Genome Research Facility. In total ~19 million 100 bp paired-end reads were generated using 0.5 lane on the Illumina HiSeq system. Sequencing reads were mapped to the Drosophila reference genome (Release 5 assembly) using BWA (Version 0.5.9) with default settings [40]. Integrative Genomics Viewer (Version 1.5.64) was used to visually inspect overall mapping quality of the candidate region [41]. After quality validation, consensus was generated using SAMtools (Version 0.1.13) with default settings [42]. Sequence variations were annotated using ANNOVAR’s gene-based annotation option [43] with FlyBase Release 5.36 annotation [44]. SNPs from the Drosophila melanogaster Genetic Reference Panel [45] were used to filter naturally occurring variations. All computations were performed on the Monash Sun Grid. The nonsense mutation in dATP8B was verified with Sanger Sequencing using an independently prepared genomic DNA sample. The following primers were used to amplify a 655 bp region flanking the mutation site: forward primer 5’CATACG-CATCCCTTAAGACCGC3’, reverse primer 5’ACCGAACAAT-CGGATGACCC3’.

Antibody production
cDNAs encoding six different regions of the CG14741-PB isoform (PE1, a.a. 2–236; PE2, 261–450; PE3, 527–630; PE4, 655–1113; PE5, 967–1359; PE6, 1562–1726) that are not part of predicted transmembrane domains were cloned into a pET100/D-TOPO vector (Invitrogen, Carlsbad, CA) such that the expressed peptide was tagged N-terminally with 6xHis. The 6xHis-PE1-6 peptides were expressed in E. coli and purified using a Ni-NTA column (Invitrogen). Peptides PE3, 5, and 6 were soluble and were individually injected into two guinea pigs by Cocalico Biologicals (Reamstown, PA). The obtained antisera (90 day protocol) were screened for antibodies against dATP8B using Western blots of both fly head protein extracts and the bacterially expressed peptides. The antisera obtained from injecting PE5 and PE6 showed positive signals and PE6 was used for immunostaining.

Immunohistochemistry
Antibodies and dilutions used were as follows: guinea pig anti-dATP8B (1:10,000); rabbit anti-Orco (1:5,000; Voshall lab); rabbit anti-Or22a (1:1,000; Voshall lab); rabbit anti-GFP (1:1,000; Life Technologies). Secondary antibodies raised in mouse, rabbit and guinea pig were Alexa-conjugated (Alexa Fluor 488 at 1:250–500, Alexa Fluor 568 at 1:500) (Molecular Probes). 14 μM cryo-sectioned adult heads were mounted on SuperFrost Plus slides (Thermo Scientific), dried for up to 3 hours, and then fixed in 4% paraformaldehyde/PBS for 30 mins at room temperature. Samples were washed for 10 minutes three times with PBST (PBS, 0.3% Triton-X-100), incubated in Block (5% normal goat serum in PBST) for 2 hours at room temperature and then incubated with the primary antibodies diluted in Block overnight at 4°C. After three 10 minutes washes with PBST, samples were incubated with secondary antibodies diluted in Block for 2 hours at room temperature. Sections were washed for 10 minutes three times with PBST before being mounted in Vectashield (Vector Labs). Samples were viewed and images acquired using a Nikon C1 confocal microscope.

Supporting Information

Figure S1 The olfactory sensilla of the antenna show normal morphology in dATP8B mutants. Scanning electron micrographs of third antennal segments and maxillary palps from wild type and homozygous dATP8B mutants. No obvious abnormalities are seen in the mutant flies. Scale bars are 20 μm. (TIF)

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Author Contributions
Conceived and designed the experiments: YCL TH REB AR AD MdB CGW. Performed the experiments: YCL MWP TJK SC KRS MdB. Analyzed the data: YCL MWP TJK AR MdB CGW. Contributed reagents/materials/analysis tools: MI KEZ. Wrote the paper: YCL MdB CGW.
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