A Comprehensive Mutagenesis Screen of the Adhesion GPCR Latrophilin-1/ADGRL1

**HIGHLIGHTS**

- Latrophilin-1/3 decreases cAMP levels and increases SRE levels in mammalian cells
- Mutagenesis of the TM region reveals residues key for adhesion GPCR signaling
- A cancer-associated mutation exhibits increased basal activity
- The same cancer mutation abolishes latrophilin function in *C. elegans* development

**Graphical Abstract**

- Lphn decreases cAMP and increases SRE
- Lphn activation by Stachel peptide
- Basal activity decreased in mutagenesis
- Basal activity increased in mutagenesis
- Synthetic no peptide response
- Cancer mutation fails to rescue lethality phenotype

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**Article**

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A Comprehensive Mutagenesis Screen of the Adhesion GPCR Latrophilin-1/ADGRL1

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SUMMARY

Adhesion G-protein-coupled receptors (aGPCRs) play critical roles in diverse cellular processes in neurobiology, development, immunity, and numerous diseases. The lack of molecular understanding of their activation mechanisms, especially with regard to the transmembrane domains, hampers further studies to facilitate aGPCR-targeted drug development. Latrophilin-1/ADGRL1 is a model aGPCR that regulates synapse formation and embryogenesis, and its mutations are associated with cancer and attention-deficit/hyperactivity disorder. Here, we established functional assays to monitor latrophilin-1 function and showed the activation of latrophilin-1 function and showed the activation of latrophilin-1 by its endogenous agonist peptide. Via a comprehensive mutagenesis screen, we identified transmembrane domain residues essential for latrophilin-1 basal activity and for agonist peptide response. Strikingly, a cancer-associated mutation exhibited increased basal activity and failed to rescue the embryonic developmental phenotype in transgenic worms. These results provide a mechanistic foundation for future aGPCR-targeted drug design.

INTRODUCTION

Adhesion G protein-coupled receptors (aGPCRs) are cell-surface molecules that mediate intercellular communication via cell-cell and cell-matrix interactions (Hamann et al., 2015; Prömel et al., 2013). With 33 members in humans, they make up the second largest GPCR family, but are the least studied and least understood (Fredriksson et al., 2003). Genetic studies suggest critical roles for aGPCRs in development and immunity and especially in neurobiology (such as brain development [Bae et al., 2014; Piao et al., 2004], synapse maturation and elimination [Bolliger et al., 2011], myelination of neurons [Monk et al., 2009], central nervous system [CNS] angiogenesis [Kuhnert et al., 2010], and neural tube development [Chae et al., 1999; Langenhan et al., 2009; Shima et al., 2004; Usui et al., 1999]), and link them to numerous diseases including neurodevelopmental disorders, deafness, male infertility, schizophrenia, and immune disorders (Langenhan et al., 2013; Promel et al., 2013). In addition, many aGPCRs are found to be over- or underexpressed in various cancers (Kan et al., 2010; Shashidhar et al., 2005; Xu et al., 2006), and a recent study reports that aGPCRs are some of the frequently mutated genes in cancerous tumors (O’Hayre et al., 2013). Considering that many drugs target the transmembrane (TM) helices of GPCRs to regulate receptor activity, thereby eliciting the desirable therapeutic effects, aGPCRs may be promising targets for drugs to treat numerous diseases including cancer. Currently, there is no high-resolution structure for the TM domain of an aGPCR. GPCRs from the secretin family have the highest TM domain similarity to aGPCRs, suggesting that aGPCR TM domains might be activated via similar mechanisms (Fredriksson et al., 2003; Hollenstein et al., 2013; Rasmussen et al., 2011; Rosenbaum et al., 2007; Siu et al., 2013). In addition, the signaling pathways of aGPCRs are largely unknown, making any functional studies difficult to perform. Altogether, further studies of aGPCRs progress slowly owing to these obstacles, and the molecular mechanisms underlying aGPCR signal transduction remain unknown.

Unlike other GPCRs, aGPCRs have large extracellular regions (ECRs) that are autoproteolytically cleaved from their seven-pass TM regions within a conserved GPCR-autoproteolysis-inducing (GAIN) domain (Arac et al., 2012; Chang et al., 2003; Ichtchenko et al., 1999; Krasnoperov et al., 1997; Lin et al., 2004) (Figure 1A). Autoproteolysis occurs between the last two β strands of the GAIN domain, cleaving the receptor into two fragments (Arac et al., 2012): an N-terminal extracellular fragment comprising various adhesion domains and the GAIN domain (lacking its last β strand) and a membrane-anchored C-terminal fragment.
comprising the TM domain (tethered to the last β strand of the GAIN domain) and the cytoplasmic tail (Krasnoperov et al., 1997; Lelianova et al., 1997). Autoproteolysis occurs in the endoplasmic reticulum, and the cleaved fragments of an aGPCR stay associated as a heterodimer even after cleavage (Arac et al., 2012). On one of the hypothetical models for the activation of aGPCRs is the tethered agonist model, which suggests that aGPCRs are activated by a short peptide that corresponds to the last β strand of the GAIN domain (called the Stachel peptide, Figure 1A) (Liebscher et al., 2014; Stoveken et al., 2015). Ligand binding to the ECR is believed to lead to shedding of the ECR, which results in exposure of the previously hidden Stachel peptide to the TM domain, leading to receptor activation.

Both the secretin family and the aGPCR family receptors are activated by the N termini of short agonistic peptides (Hoare, 2005; Liebscher et al., 2014; Stoveken et al., 2015). However, Stachel peptides of aGPCRs are hydrophobic and prone to β strand formation (Arac et al., 2012), whereas the agonistic peptides of secretin-like receptors fold into α helices (Watkins et al., 2012), suggesting differences in the molecular details of the peptide-TM interaction. Although recent structures of secretin-like receptors in inactive and peptide-bound active conformations revealed the molecular mechanism of peptide recognition by the TM, little is known about the molecular details for the activation of aGPCRs by the Stachel peptide (Hollenstein et al., 2013; Liang et al., 2017; Rasmussen et al., 2011; Rosenbaum et al., 2007; Siu et al., 2013; Zhang et al., 2017). These secretin family structures are extremely valuable for aGPCR studies because more accurate models of aGPCR TMs can be constructed, facilitating studies like this one.

The knowledge of the signaling pathway of a receptor and an in vitro assay to monitor downstream signaling has been invaluable in characterizing and drugging the canonical GPCRs. The lack of a robust in vitro functional assay for aGPCRs has obstructed both structure/function studies, including the functional characterization of disease-causing mutations, and agonist/antagonist ligand discovery and characterization. In this study, we used latrophilin-1 (Lphn1)/ADGRL1, a key molecule in synapse formation and brain development, as a model aGPCR to study aGPCR function. We established two robust in vitro assays to monitor receptor signaling and showed that the endogenous agonist of Lphns, a 14-amino-acid peptide, binds to and activates the receptor. We studied a large set of bioinformatics-based point mutations and disease mutations on the TM region of Lphns and identified mutants that are constitutively active, constitutively inactive, or nonresponsive to the agonist peptide. Intriguingly, we identified a cancer-associated mutation that exhibited high basal activity and abolished the rescue of the embryonic development phenotype in transgenic worms. These results provide the basic groundwork for future drug design against aGPCRs.
RESULTS

Establishment of In Vitro Functional Assays to Monitor Latrophilin Activity

Latrophilins (Lphn1–3) regulate synapse formation and brain development (O’Sullivan et al., 2012). Lphn1 was first identified as the calcium-independent receptor for α-latrotoxin, a black widow spider toxin component that triggers massive neurotransmitter release from nerve terminals (Deak et al., 2009; Krassover et al., 1997; Lelianova et al., 1997; Sudhof, 2001; Sugita et al., 1999). Mutations of Lphns are associated with attention-deficit/hyperactivity disorder (ADHD) as well as numerous cancers in humans (Arcos-Burgos et al., 2010; Kan et al., 2010; O’Hayre et al., 2013). Lphns are one of the two aGPCRs that are conserved between vertebrates and invertebrates. In Caenorhabditis elegans, the Lphn homolog lat-1 is required for the alignment of cell division planes to the anterior-posterior axis during development (Langehan et al., 2009; Muller et al., 2015). In Drosophila, Lphn/dCIRL sensitizes the chordotonal neurons to perceive mechanical signals (Scholz et al., 2015). In vertebrates, Lphns mediate excitatory synapse formation (Lu et al., 2015; O’Sullivan et al., 2012). The intracellular signaling of Lphns via G proteins was reported by several studies, including the activation of Gs by rat Lphn1 as well as LAT-1, one of the C. elegans homologs of Lphn (Muller et al., 2015); the decrease of cyclic adenosine monophosphate (cAMP) by Lphn/dCIRL in Drosophila (Scholz et al., 2017); and the increase of intracellular Ca2+ levels by rat Lphn1 (Silva et al., 2011); however, the assays used in these studies were not suitable for large-scale mutagenesis screens.

To measure Lphn activity in a signaling assay, we screened several assays that monitor downstream G protein signaling using the basal activity of full-length rat Lphn1 and full-length human Lphn3 (these constructs are used throughout this article unless stated otherwise). The first assay in which we robustly observed basal activity of Lphn was the GloSensor assay (Promega), which reports cAMP levels in mammalian expression systems. A comparison of the background cAMP level in control HEK293 cells (transfected with empty vector) with cells transfected with Lphn1 provided the first indication that Lphn1 may couple to Gi as Lphn1 overexpression decreased cAMP levels (Figure S1A).

We modified the GloSensor assay in two ways to reliably detect a decrease in cAMP levels: (1) we elevated cAMP level in cells by co-transfecting the cells with the Gs-coupled β2-adrenergic receptor (β2AR) and activating it with its agonist, isoproterenol (ISO) (Figures 1B and S1B) or (2) we elevated cAMP level in cells by adding forskolin to activate the endogenous adenylate cyclase (Figures 1C and S1C). Relative to empty vector, overexpression of Lphn1 or Lphn3 decreased the level of cAMP in cells, suggesting the possible activation of Gi. Lphn1 DNA concentration response curves for cell-surface expression of Lphn1 and for the decrease of cAMP levels in cells are as expected (Figures S1D and S1E). In addition, we performed direct G protein coupling experiments using human Lphn3 GAIN + TM domains embedded in insect cell membranes. Comparison of empty insect cell membranes with those embedded with Lphn3 showed an increase in binding to purified Gi proteins, suggesting that human Lphn3 couples to Gi (Figure S1F). When membranes were treated with urea to induce dissociation of the ECR and Stachel exposure, we observed increased coupling to Gi, suggesting that Lphns are sensitive to Stachel-mediated activation in this assay (Figure S1F).

The second assay in which we robustly observed basal activity of Lphn was a serum response element (SRE)-luciferase assay that reports, among others, activation of G12/13, which is upstream of SRE and RhoA (Luo et al., 2014; Stachel et al., 2016). In transfection systems. A comparison of the background cAMP level in control HEK293 cells (transfected with empty vector) with cells transfected with Lphn1 provided the first indication that Lphn1 may couple to Gi as Lphn1 overexpression decreased cAMP levels (Figure S1A).

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Latrophilins Are Activated by Their Stachel Peptide in the Established Assays

Several aGPCRs besides rat LPHN1 and its homologs in C. elegans and Drosophila, including GPR126, GPR133, GPR64, GPR114, GPR56, GPR110, and GPR116, have been reported to be activated by their Stachel peptide, which functions as a tethered agonist (Demberg et al., 2015, 2017; Kishore et al., 2016; Liebscher et al., 2014; Muller et al., 2015; Scholz et al., 2017; Stoveken et al., 2015) (Figures 2A and 2B). Recent studies showed that addition of the synthesized Stachel peptide on full-length receptors in trans activated the receptors (Liebscher et al., 2014; Stoveken et al., 2015). Similarly, truncation constructs starting from the first residue after the autoproteolysis site, thus revealing the Stachel peptide free, also displayed dramatically increased signaling (Kishore et al., 2016; Paavola et al., 2011). To test whether these results can be generalized to other mammalian Lphns and to optimize our signaling assays for detecting Stachel peptide-dependent activation, we repeated similar experiments for Lphn1.
First, we added a synthesized 14-residue peptide (termed P14, TNFAVLMAHREIYQ) corresponding to the Stachel peptide of rat Lphn1 onto full-length rat Lphn1 and monitored receptor function in our cAMP assays. Addition of P14 onto Lphn1 decreased the cAMP level in both β2AR and forskolin assays (Figures 2C and 2D). Further addition of pertussis toxin (PTx), a specific inhibitor of Gi (Burns, 1988), abolished the effect of P14 and brought cAMP levels back to basal activity level, suggesting that Stachel peptide-dependent activation of Lphn1 is mediated by Gi (Figures 2C and 2D). However, PTx had no effect on the basal activity of Lphn1, suggesting that other signaling pathways mediate the decrease in cAMP levels upon Lphn1 over-expression.

To confirm direct interaction between the Stachel peptide and the TM domain of Lphns rather than another domain of Lphn, we tested whether a construct encoding the TM domain of Lphn3 binds to a synthesized Stachel peptide. Purified and detergent-solubilized Lphn3 TM domain was mixed with synthesized Stachel peptide in the presence or absence of a cross-linking reagent, and peptide binding was detected by western blot analysis (Figures S2A and S2B). Taken together, these results suggest that Stachel peptide acts as an agonist for the TM domain of Lphns when added in trans.

Second, to test the role of the Stachel peptide as an agonist tethered to the Lphn TM domain, i.e., in cis, we designed an active-like construct encoding the TM domain with the tethered Stachel peptide of Lphn1 (termed P + TM) and an inactive-like construct encoding only the TM domain of Lphn1 (termed TM) (Figure 2A). To avoid expression problems and potential effects on signaling, both constructs were FLAG-tagged at the C terminus, rather than the N terminus. Overexpression of the P + TM construct displayed increased basal activity compared with full-length Lphn1, whereas the TM construct was less active, although not completely inactive. All mutants responded to P14 peptide addition in trans, and peptide response was sensitive to PTx exposure (Figures 2C and 2D). The cell-surface expression levels of TM
and P + TM constructs were quantified by biotinylation experiments (Figures S2C and S2D). The cell-surface expression level of TM and P + TM were at similar levels, although both showed decreased expression when compared with the full-length Lphn1. These results suggest that the Stachel peptide acts as an agonist when presented to the receptor in a tethered conformation in cis.

**Mutagenesis of the TM Domain Affects Basal Activity and Peptide Response**

Ligand-induced conformational changes of GPCRs transduce extracellular signals to the intracellular part of the receptor and facilitate G protein interaction. Conserved motifs in the rhodopsin family of GPCRs (such as the CWxP motif in TM6, NPxxY motif in TM7, and DRY motif in TM3) are critical for receptor activation upon ligand binding (Audet and Bouvier, 2012). To identify essential residues for basal activity and for agonist peptide response in adhesion GPCRs, we performed a mutagenesis screen of the TM domain within the full-length Lphn1 construct using our β2AR cAMP-based signaling assay (Figure S1C). A homology model of Lphn1 TM domain was generated based on the crystal structure of the corticotropin-releasing factor receptor (CRFR), a member of the secretin family with 29% sequence identity to Lphn1 TM domain (Ebejer et al., 2013; Hollenstein et al., 2013). Over 60 point mutations were designed based on various criteria, including homology to functionally important conserved motifs in the rhodopsin family (Audet and Bouvier, 2012), homology to the secretin family in general (Krishnan et al., 2012; Schioth et al., 2011), homology to CRFR residues reported to interact with its agonist peptide (Coin et al., 2013), conservation within aGPCRs (including the following motifs: GWGxP [helix IV], K/RKLH [intracellular loop I], and discontinuous RHE motif [helix II]) (Peeters et al., 2016), and visual analysis of extracellular loops. In addition, previously reported mutations associated with cancer and ADHD were generated (Kan et al., 2010; O’Hayre et al., 2013) (Figure S3 and Table S1).

The effect of mutations on the function of the full-length Lphn1 was monitored by performing the β2AR cAMP-based signaling assay (Figures 3A, 3D, and 3G). Cell-surface expression level for all mutants was quantified using flow cytometry by detecting N-terminally FLAG-tagged Lphn1 on non-permeabilized cells (Figures 3C, 3F, and 3I). Approximately half of the mutations had an effect on Lphn1 signaling in the cAMP assay. These mutants were further tested in the SRE-luciferase assay (Figures 3B, 3E, and 3H). Our screen revealed mutants that either increased basal activity (colored red throughout Figures 4 and 6 and Table S1), decreased basal activity (colored green), or affected response of the receptor to the agonist peptide (colored magenta for confirmed mutants and orange for likely mutants). Some mutants had an effect on both the basal activity and response to the peptide (colored blue). All cAMP signaling, SRE signaling, and cell-surface expression data for each mutant as well as the information about the location and conservation of the mutated residues can be found in Figures 3 and 5, Table S1, and Figures S5 and S6. Please note that color coding of residues in figures and tables oversimplifies the results, and we recommend referring to the raw data presented in Figures 3 and 5 and Table S1.

**Homologous Yet Different Molecular Features within the TM Mediate Basal Activity**

Although aGPCRs do not have the motifs conserved in the rhodopsin family, mutagenesis of homologous residues resulted in altered basal activity in Lphn1, suggesting that aGPCRs are activated by molecular mechanisms that are mediated by homologous motifs (Figures 3, 4, and S4). For instance, the NPxxY motif that is conserved in the rhodopsin family is replaced by “IFVFH” in Lphn1 (Figures 3A–3C). Mutations in this region affected basal activity severely (Figures 3A and 4C). V1094I and nearby V1090A increased basal activity (Figures 3A–3C). Furthermore, I1092A, V1094I, and nearby C1097Y abolished response to the peptide. Among all mutants, the most active mutant, V1094I, is one that was associated with upper aerodigestive tract cancer, previously (O’Hayre et al., 2013) (Figures 3G–3I and 4E). C1097Y is also associated with endometrium cancer. These results suggest that a motif homologous to the NPxxY motif is critical for aGPCR activation.

Another example of a unique aGPCR motif that might act similarly to a homologous one in the rhodopsin family is the DRY motif, which is replaced by a more hydrophobic “HLY” motif in Lphn1 (Figures 3A–3C and 4D). In conventional GPCRs, the DRY motif on TM3 and a glutamate residue from TM6 (replaced by D1043 in Lphn1) contribute to an ionic lock that traps the receptor in an inactive conformation (Ballesteros et al., 2001; Greasley et al., 2002). In Lphn1, both L947A and Y948A mutations within the “HLY” motif lead to increased basal activity (Figures 3A–3C). Furthermore, mutation of a hydrophobic residue on TM6, I1045N, a liver carcinoma mutation, abolished response of the receptor to the agonist peptide (Figures 3G–3I and 4E).
Figure 3. TM Mutations That Affect Basal Activity of Lphn1

(A–I) TM mutations that have an effect on the basal activity of full-length Lphn1 are shown for the cAMP signaling assay (A, D, and G), SRE signaling assay (B, E, and H), and cell-surface expression quantification (C, F, and I). The mutations are categorized as: (A–C) mutations that map to the indicated conserved motifs from rhodopsin, secretin, and adhesion families; (D–F) mutations that lead to constitutive activity; and (G–I) previously reported cancer-associated mutations that affect Lphn1 signaling. Mutations that abolish receptor response to the agonist peptide are shown in Figure 5. One mutation may belong to more than one category. See Table S1 for raw data and for other mutants that had no effect. See Figure 4 for structural visualization of basal activity mutants. See Figures S4 and S5 for raw cell-surface expression data.
The altered activity upon mutagenesis of these hydrophobic residues in Lphn1 suggests a possible hydrophobic lock in aGPCRs instead of the classical ionic lock.

Overall analysis of the dataset showed that the majority of basal activity-altering mutations resulted in increased basal activity, suggesting that mutations disturb the inactive conformation and promote an active conformation (Figures 3 and 4B). For instance, only W1063A, G1066A, and H1096A decreased basal activity, whereas V1010A, W939A, V1090A, M1077A, Y970A, and Y948A are among the many mutations that increased basal activity. Mapping these basal activity-altering mutations on the Lphn1 model revealed that the mutations were distributed throughout the TM domain, including distant positions, suggesting long-distance allosteric pathways within the TM fold (Figure 4B). Our results largely agree with the results of a recent study that mutagenized the active-like truncated construct (P + TM) for another aGPCR, GPR112, with some differences that are likely due to the use of full-length versus truncated constructs in these studies (Peeters et al., 2016).

Numerous Residues within the TM Are Involved in the Response to Agonist Peptide

To identify the residues that are involved in the response of the receptor to the agonist peptide, we looked for mutations that have similar basal activity in the presence or absence of the exogenously added peptide on the full-length receptor (Figure 5). These residues were mapped onto the modeled structure of Lphn1 that was generated based on the agonist peptide-bound active structure of the Glucagon-like peptide (GLP1) receptor (Zhang et al., 2017) (Figure 6). Visual analysis revealed that residues that affect peptide response map onto the extracellular face, the middle layers, or the cytoplasmic face of the receptor (Figures 6A and 6B). We expect that lack of receptor response to the agonist peptide can be caused by the following different mechanisms. Residues that are on the extracellular face (such as Y1001F, F1069A, A1078G, and A926G) are likely involved in direct peptide binding, which their mutagenesis may abolish (Figures 6B–6D). Noticeably, Y1001 is a lung cancer-associated mutation (Kan et al., 2010). Residues that map to the core of the receptor (such as G1060A, G1008A, G1089A, G969A, H946A, and V1094I) likely mediate transmission of the signal from the extracellular face to the intracellular face (Figure 6B). These glycine residues likely enable the flexibility of the receptor, which allows the signal to be transduced down the protein. Finally, residues that map to the cytoplasmic face of the receptor (such as R891A, H1096A, H895A, R885A, H946A, Y964A, Y965A, I1092A, and C1097Y) likely mediate receptor activation and G protein binding (Figures 6C and 6E). Several mutants including G1060A and V1094I were constitutively active and did not respond to exogenous peptide (Figure 5). These results suggest that a large set of residues are involved in the response of the receptor to the agonist peptide.

The peptide hormone-bound structures of GLP1 receptor and calcitonin receptor revealed that the peptide agonist engages the receptor by binding to a hydrophobic pocket that is generated by the large outward movement of the extracellular ends of TM helices 6 and 7 (Liang et al., 2017). This conformation is accompanied by a 60° kink in helix 6. A large outward movement of the intracellular end of this helix opens the TM to accommodate interactions with the a5 helix of Gas (Liang et al., 2017; Zhang et al., 2017). In Lphn1, A926 in TM3, Y1001 in TM5, F1069 in TM6, and A1078 in TM7 are residues on the extracellular face of the receptor (Figure 6D). Superimposition of the modeled Lphn1 structure with the peptide-bound GLP1 receptor structure suggests that these residues might be involved in direct peptide binding (Figures 6C and 6D). F1069 and A1078 are located at the tips of TM6 and TM7, respectively, where the helices bend out in the calcitonin receptor (Liang et al., 2017). G1060, on the other hand, sits in the middle of the kink in TM6, which enables opening of the TM bundle (Liang et al., 2017). Although Stachel peptides of aGPCRs are more hydrophobic and more prone to β strand formation than agonist peptides of secretin receptors, our results suggest that they may share similar molecular signatures for binding to their agonist peptides.

A Cancer-Associated Mutation with Increased Basal Activity Abolishes Lphn Function in C. elegans Development

To get insights in the physiological impact of point mutations that alter Lphn1 signaling on an entire organism, we selected the mutations that exhibited the most effect in our in vitro assay and mutated them in the
Lphn homolog lat-1 of the nematode C. elegans: (1) V1094I (L790A in C. elegans), which severely increased basal activity and abolished the response of the receptor to the agonist peptide; importantly, this is a mutation associated with upper aerodigestive tract cancer and maps to TM7 at the NPxxY homology region (O’Hayre et al., 2013). (2) H1096A (H792A in C. elegans), which is one of the few mutations that decreased basal activity, although not completely; this mutation abolished agonist peptide response and maps to TM7 at the NPxxY homology region.

Figure 4. Basal Activity Mutations Mapped on the Modeled TM Structure of Lphn1
(A) Snake plot for visualization of basal activity mutations on the transmembrane helices.
(B) All basal activity mutations mapped on the Lphn1 TM domain, which is modeled based on the inactive CRFR structure (PDB ID: 4K5Y). Mutations that increase basal activity are colored red, mutations that decrease basal activity are colored green, and mutations that increase basal activity and also decrease response to the agonist peptide are colored blue.
(C) A cytoplasmic view of the modeled Lphn1 structure showing the basal activity mutants and the mutants that map to conserved region homologous to the NPxxY motif.
(D) A view of the modeled Lphn1 structure showing the basal activity mutants that map to conserved regions homologous to the DRY motif and the ionic lock. I1045N mutation (colored magenta) affects peptide response.
(E) Cancer-associated mutations on Lphn1 TM domain that affect signaling. V1094I (V1095I in human Lphn1 in upper aerodigestive tract carcinoma), R885L (R886L in human Lphn1 in endometrioid carcinoma), C1098Y (C1098Y in human Lphn1 in endometrioid carcinoma), Y1001F (Y1019F in human Lphn3 in lung adenocarcinoma), and I1045N (I1146N in human Lphn1 in hepatocellular carcinoma) were reported previously (Kan et al., 2010; O’Hayre et al., 2013) (colored yellow).

Mutations that are introduced into transgenic worm are labeled with a cyan star.
TM7 at the NPxxY homology region. (3) F1069A (F763A in *C. elegans*), which maps to the extracellular end of TM6 and results in abolished agonist peptide response (these three residues are colored cyan or indicated by a cyan asterisk throughout Figures 3, 4, 5, and 6). All three mutations yield receptors with expression levels and localization similar to wild-type *lat-1* in *C. elegans* (Figure 7A).

LAT-1 plays essential roles in fertility as well as in oriented cell division in embryogenesis (Langenhan et al., 2009; Prömel et al., 2012). During embryonic development, LAT-1 transduces a signal cell autonomously via the TM and the cytoplasmic tail mediating a G protein-dependent signal (Muller et al., 2015; Prömel et al., 2012), whereas its role in fertility requires only the ECR (Prömel et al., 2012). Consistently, a *lat-1* null mutant, *lat-1*(ok1565) (hereafter referred to as *lat-1* mutant), shows severe defects in fertility, characterized by a reduced brood size, and in cell polarity during embryogenesis, reflected in a high lethality rate (Langenhan et al., 2009; Prömel et al., 2012) (Figures 7B and 7C). Expression of transgenic *lat-1* constructs ameliorates these phenotypes and leads to a rescue. This rescue can be read out through the total brood size and the level of developmental lethality in the progeny of transformants. We used these assays to assess the functionality of the three generated LAT-1 variants F763A, L790A, and H792A in vivo. All three mutations rescue the fertility defects of *lat-1* mutants and thus are able to fulfill LAT-1 function in this context (Figure 7B). Variants carrying F763A and H792A are fully functional receptors as they also rescue the lethality phenotype (Figure 7C). Strikingly, the variant carrying L790A failed to rescue the lethality phenotype. Thus, the point mutation L790A leads to a receptor that is not capable of transducing signals into the cell, suggesting that L790A mutation, which corresponds to the V1094I cancer-associated mutation in rat and exhibits high basal activity and no response to the agonist, is critical for proper LAT-1 function during *C. elegans* development.

**DISCUSSION**

GPCRs are the largest family of cell-surface receptors and are the targets for approximately 30% of currently marketed drugs that treat a very broad spectrum of diseases including neuropsychiatric, cardiovascular, pulmonary, and metabolic disorders; cancer; and acquired immune deficiency syndrome (AIDS). Adhesion GPCRs recently emerged as a novel subfamily of GPCRs with important roles in diverse cellular...
processes, and they carry the potential to be the targets for the next generation of drugs. However, a molecular understanding of their activation mechanisms is a first step for moving forward. In this study, we focused on a model aGPCR subfamily, Lphns, and completed a thorough study that links in vitro molecular understanding of the receptor to in vivo function in whole animals.

Figure 6. Peptide Response Mutations Mapped on the Modeled TM Structure of Lphn1
(A) Snake plot for visualization of peptide response mutations on the transmembrane helices.
(B) All mutations that affect response of the receptor to the agonist peptide mapped on the Lphn1 TM domain, which is modeled based on the peptide-bound active GLP receptor structure (PDB ID: 5VAI). Mutations that affect peptide response are colored magenta, mutations that affect both peptide response and basal activity are colored blue, and mutations that likely affect peptide response are colored orange.
(C) Superimposition of the model of Lphn1 TM domain (gray) with the GLP1 peptide (red)-bound GLP1 receptor/G protein (green/tan) complex structure (PDB ID: 5VAI). Lphn1 residues are colored as in B.
(D) Close-up extracellular view of (C) showing the modeled Lphn1 TM domain and the GLP1 peptide. Peptide response mutants that map to the extracellular side of the receptor as well as GLP peptide residues important for its interaction with the GLP receptor are shown by sticks.
(E) Close-up cytoplasmic view of (C) showing the modeled Lphn1 TM domain and the G protein. Peptide response mutants that map to the cytoplasmic side of the receptor are shown by sticks. Mutations that are introduced into transgenic worms are labeled with cyan asterisk.
Our data show an activation of a Gi signaling cascade by Lphn1, which can be stimulated by a 14-amino-acid endogenous agonist peptide. This peptide compares well with a similar 13-amino-acid-long peptide, which has already been successfully used in previous studies on Lphn1 (Muller et al., 2015). The work by Muller et al. has shown that the receptor activates a Gs cascade leading to the formation of cAMP. These results might be explained by the different assays used in these studies. Owing to the different cell systems, expression levels can vary, and thus the possibility of coupling to other G protein families was not fully excluded in the study. We successfully monitored Lphn downstream signaling by intracellular cAMP and SRE-luciferase assays (Figure 1). A carefully designed mutagenesis screen performed on both assays revealed that approximately half of the mutated residues have an effect on the activity of the receptor (Figures 3 and 5). Importantly, the aGPCR residues that are homologous to the conserved motifs in the rhodopsin family that have diverged significantly, such as the DRY and the NPxxY motifs, are indeed critical for aGPCR function (Figures 4C and 4D). For instance, the ionic lock that the DRY motif seems to be replaced by is a possible hydrophobic lock that is still critical for receptor function. Analysis of the entire dataset suggests that although the TM fold of the aGPCRs carries a similar skeleton and works in a similar fashion to other GPCR families, there are major differences in the fine details within the fold.

Figure 7. An Overactive Cancer-Associated Mutant Fails to Rescue the Developmental Phenotype of LAT-1 Knockdown in C. elegans

(A) Expression and protein localization of all three point mutation-containing variants is indistinguishable from wild-type lat-1:gfp. Fluorescence images show presence of LAT-1 and the variants at the plasma membrane of pharyngeal muscle cells, neurons in the nerve ring, and the pharyngeal nervous system. Scale bars represent 10 μm.

(B and C) The point mutations within the TM of LAT-1 lead to different abilities to rescue fertility (brood size, B) and lethality (individuals reaching adulthood, C) of lat-1 mutants. All three variants ameliorate the fertility defects observed in lat-1 mutants similar to a wild-type lat-1 transgene or a construct comprising the extracellular region tethered to the membrane via the first TM (lat-1 [aa1–581]) (B). Only LAT-1(L790A) does not rescue lethality, while LAT-1(F763A) and LAT-1(H792A) display the same functionality as a wild-type lat-1 transgene. Data are shown as percentage of the original brood sizes (C). The wild-type lat-1 transgene, which rescues fertility and lethality, and a lat-1[aa1–581] construct, which only rescues fertility, served as controls. Data are shown as means ± SEM, n ≥ 20, n.s., not significant; **p < 0.01; ***p < 0.001. SEM, standard error of the mean.
To study the mechanism of aGPCR activation by the agonist peptide, we first identified the agonist peptide and showed that the peptide can bind and activate Lphn1 both in trans (i.e., as added synthetic peptide) and in cis (i.e., as tethered to the receptor) (Figure 2). The residues that are involved in the response of the receptor localize to the entire receptor, as they are likely involved in direct peptide binding, signal transduction, or interaction with the G protein (Figure 6). This large dataset is consistent with the numerous allosteric pathways within the TM fold (DeVree et al., 2016). Comparison of the GLP receptor bound to GLP with a model of Lphn1 suggests that the mutations on the extracellular face (A926, Y1001, F1069, and A1078) are likely involved in direct peptide binding (Figure 6D). However, the structure that the agonist peptide adopts when bound to an aGPCR might be different from that of secretin receptor peptides.

Aberrant expression and altered signaling profile of mutated GPCRs contribute to cancer progression and metastasis (O’Hayre et al., 2013). Mutant GPCRs are often overexpressed and constitutively active in various cancers (Dorsam and Gutkind, 2007; Kan et al., 2010; O’Hayre et al., 2013). Disease-associated mutations were previously reported to change other adhesion GPCR signaling (Kishore and Hall, 2017; Purcell et al., 2017). We tested cancer-associated mutations of Lphns reported in different types of carcinomas (Kan et al., 2010; O’Hayre et al., 2013) and found that the cancer-associated mutations affected basal activity and/or peptide response (Figure 4E). Remarkably, V1094I (O’Hayre et al., 2013) mutant was constitutively active in both the cAMP and SRE assays, suggesting a possible role for high activity of Lphns in cancer progression.

Introduction of this highly active cancer-associated mutation to transgenic worms (corresponding to L790A mutation in C. elegans Lat-1) led to a LAT-1 variant devoid of its function in cell polarity during embryonic development but not in fertility (Figures 7B and 7C). As it has been shown that the role of LAT-1 in fertility is solely dependent on its ECR but independent of TM or cytoplasmic tail (Promel et al., 2012), it is conceivable that any mutation within the TM does not impair LAT-1 function in this context. However, cell polarity during embryogenesis is controlled by LAT-1 via a signal requiring TM and cytoplasmic tail (Muller et al., 2015; Promel et al., 2012). L790/V1094 is located in TM7 at a position homologous to the NPxxY motif essential for transducing the signal into the cell. An amino acid change at this position likely alters LAT-1 signaling capabilities in a way such that it cannot regulate cell polarity.

Introduction of the V1094I mutation into rat LPHN1 leads to an increase in basal activity, whereas the equivalent mutation (L790A) in the homolog LAT-1 in C. elegans leads to a loss of receptor function in vivo. Several reasons can account for this fact. First, in a living organism, over-activation of the receptor might not have scorable effects. It also cannot be excluded that a highly active variant is detrimental to the organism and that the observed developmental lethality is a consequence of this rather than to a non-active LAT-1. Furthermore, it is conceivable that the motif identified in mammalian Lphns and its impact on receptor integrity and importance for signaling is not comparable to its function in C. elegans LAT-1. Both homologs could have evolved differently during evolution, and thus the variants might not exhibit the same properties.

Altogether, these results provide the groundwork for a thorough characterization of the model aGPCR, Lphn1. The data shed light on signaling pathways and Stachel peptide-mediated activation mechanism of aGPCRs, and correlate disease-associated mutations to in vivo function.

**METHODS**

All methods can be found in the accompanying Transparent Methods supplemental file.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Transparent Methods, six figures, and one table and can be found with this article online at https://doi.org/10.1016/j.isci.2018.04.019.

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AUTHOR CONTRIBUTIONS
O.N. identified the signaling pathway and performed the signaling assays. A.K. performed bioinformatics analysis, designed mutations, and performed mutagenesis (with assistance from G.S. and P.N.) J.W. and S.P. designed and performed C. elegans experiments. H.S. and G.T. designed and performed direct G protein-coupling experiments with assistance from K.L. and Y.L. K.L., K.M., and Y.L. performed the peptide-binding experiments. D.A. designed all experiments, performed data analysis, and wrote the paper.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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Supplemental Information

A Comprehensive Mutagenesis Screen
of the Adhesion GPCR
Latrophilin-1/ADGRL1

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Figure S1
Figure S1, Related to Figure 1. Development of the cAMP assay to monitor Latrophilin activity

(A) Lphn1 overexpression decreases basal cAMP level in HEK293 cells. Cells were transfected with empty vector or different Lphn1 constructs and cAMP level was measured by the Glosensor assay. (B-C) Schematic representation of the modified cAMP assays to increase cAMP level by activation of co-transfected B2-adrenergic receptor with isoproterenol (B) or by activation of endogenous adenylate cyclase with forskolin (C). (D) The cell-surface expression of Lphn1 on non-permeabilized HEK293 cells quantified by flow cytometry using an N-terminal FLAG tag as a function of DNA concentration. (E) Transiently transfected HEK293 cells display DNA amount-dependent cAMP signaling of Lphn1 determined by β2-adrenergic receptor cAMP assay.

(F) G protein coupling of Lphn3. High-Five cell membranes with Lphn3 were subject to urea treatment to induce ECR-shedding and compared to membranes with no receptor or no treatment. Membranes were reconstituted with purified heterotrimeric G proteins, including the αi subunit and the βγ subunits. The receptor-mediated G protein activation kinetics were measured using the [35S]-GTPγS binding assay for Goi. Data are presented as mean ± S.E.M.
Figure S2

A: WB: Neutavidin

B: WB: anti-His

C: rLPHN1 FL rLPHN1 TM rLPHN1 P+TM

D: Optical density, %

Figure S2
**Figure S2, Related to Figure 2, Stachel peptide binds and activates the TM domain of Latrophilin**

(A) Synthesized human Lphn3 Stachel peptide can be crosslinked to purified human Lphn3 TM in the presence of glutaraldehyde. Western blot of Lphn3 peptide, Lphn3 TM, and mixture of Lphn3 peptide and Lphn3 TM in presence or absence of 0.01% glutaraldehyde blotted with (A) labeled NeutrAvidin to detect biotinylated Stachel peptide, black arrow, or (B) anti-His antibody to detect His-tagged hLphn3 7TM, black and gray arrows indicate Lphn3 TM domain, respectively.

(C) Western blot for quantification of cell-surface expressed Lphn1 constructs. HEK293 cells were transfected with full length, TM or P+TM Lphn1 constructs. Cell-surface expressed proteins were biotynilated 48 hrs after transfection, pulled-down with streptavidin beads and eluted with urea. Eluted protein bands are highlighted with yellow dashed frames. Though, full length Lphn1 did not elute from streptavidin beads and thus cannot be quantified, it is clear that some was surface-expressed and biotinylated as it was depleted upon incubation with streptavidin beads (right). Numbers indicate repeats for the same construct.

(D) Optical density of eluted protein bands (yellow dashed frames from B). Bar heights represent the mean of two measurements. Error bars indicate ??.
**Figure S3**, Related to Figures 3 and 5. Snake plot showing all mutants that were cloned. All mutations are represented by grey color on the snake diagram.
Figure S4, Related to Figures 3 and 5. Multiple sequence alignment comparing Lphns and rhodopsin family GPCRs. PROMALS3D multiple sequence and structure alignment server was used for alignment. Four mammalian Lphn1 constructs from the Adhesion family (O88917 AGRL1_RAT_A, Q80TR1 AGRL1_MOUSE, O94910_AGRL1_HUMAN, O97831_AGRL1_BOVIN) and three human Rhodopsin family constructs, beta 1 and 2 adrenergic and rhodopsin (P08100 OPSD_HUMAN, P08588 ADRB1_HUMAN, P07550 ADRB2_HUMAN) were used for alignment.
A. Conserved motifs mutations

| Mutant | Normalized Count |
|--------|-----------------|
| W1063A | 125% 112%       |
| G1063A | 112% 65%        |
| G1056A | 157% 115%       |
| H1092A | 103%           |
| V1084A | 69% 81% 90%     |
| H1086A | 131% 100%       |
| H946A  | 73% 81% 48%     |
| L947A  | 69% 50% 78%     |
| Y948A  | 36% 78% 33%     |
| G969A  | 113% 119% 83%   |
| V1094A | 69%            |
| H946A  | 73% 81% 48%     |
| G969A  | 113% 119% 83%   |

B. DNA titration

| Mutant | Normalized Count |
|--------|-----------------|
| L947A  | 76%            |
| Y948A  | 98%            |
| Y970A  | 77%            |

Figure S5
Figure S5, Related to Figures 3 and 5. Expression data for Lphn1 mutants.

(A) Surface expression of Lphn1 conserved motif mutants presented as overlay of typical flow cytometry histograms. HEK293 cells transfected with N-terminally FLAG-tagged full-length Lphn1 constructs (wt or mutant) or empty vector were stained with anti-FLAG antibody without permeabilization. Lphn1 expression (i.e. FLAG signal) for cells transfected with each mutant is directly compared to cells transfected with wild-type Lphn1 and empty vector. (B) DNA amount experiments for L947A, Y948A and Y970A mutants. Lphn1 wt DNA amount used for transfection was decreased 2.5-fold to achieve expression level comparable with mutants.
Figure S6
Figure S6, Related to Figures 3 and 5. Expression data for Lphn1 mutants.

Surface expression of Lphn1 (A) peptide response mutations, (B) cancer-associated mutations and (C) constitutively active mutants presented as overlay of typical flow cytometry histograms. HEK293 cells transfected with N-terminally FLAG-tagged full-length Lphn1 constructs (wt or mutant) or empty vector were stained with anti-FLAG antibody without permeabilization. Lphn1 expression (i.e. FLAG signal) for cells transfected with each mutant is directly compared to cells transfected with wild-type Lphn1 and empty vector. (D) DNA amount experiments for F1069A, G1089A and F1081A mutants. Lphn1 WT DNA amount used for transfection was decreased 2.5 fold to achieve expression level comparable with F1081A mutant. F1069A and G1089A DNA amount were decreased 1.5 times to achieve expression level comparable with WT.
Table S1, Related to Figures 3 and 5. Complete signaling and expression data.

Table encodes full raw data. Annotated for each mutation is the location of the mutation, the justification for performing the mutation, the Cell-Surface Expression, Signaling (Raw and Normalized), whether there is an observed peptide response and possible references. Basal activity means cAMP level without addition of ligand peptide and it was calculated by taking the luminescence value of wildtype over the luminescence value of the mutant. A value of <1 means mutant is more active than WT, while a value of >1 means mutant is less active than WT. The slashes between values separate individual experiments. Each individual experiment is at least three repeats. Colors match to the residue colors in main figures.
**Transparent Methods**

**Lphn1 homology modeling**
The modeling was done using the MEMOIR: Membrane protein modeling pipeline. This server, maintained by the University of Oxford, combines four different protein software programs iMembrane for membrane annotation, MP-T for sequence alignment, Medeller for coordinate generation, and Completionist for loop modeling.

**Site-directed mutagenesis**
Site-directed mutagenesis was performed using QuickChange approach. Constructs for site-directed mutagenesis were generated from full length rat Lphn1 on a PCMV5 vector. The mutants were verified through sequencing at the UChicago DNA Sequencing Facility.

**Cell culture and transfection**
HEK293 and HEK293T cell lines (ATCC; generous gifts from S. Koide lab) were cultured in Dulbecco’s modified Eagle’s medium (DMEM; Gibco; 11965092) supplemented with 10% FBS (F0926; Sigma) at 37°C in 5% CO2 humidified incubator.

For both cAMP signaling assay and western blots, HEK293 cells were plated in 6-well plates and incubated until 60% confluent. For forskolin cAMP assays, HEK293 cells were co-transfected with 350 ng Lphn, 350 ng 22F Glosensor reporter plasmid (E2301, Promega; a generous gift from R. Lefkowitz lab), and 2.8 μL transfection reagent Fugene 6 (PRE2693; Promega). For β2-adrenergic receptor cAMP assays, the cells were co-transfected with the same reagents along with 9 ng β2-adrenergic receptor. For western blot and flow cytometry experiments, HEK293 cells were transfected with 2μg Lphn and 8μL Fugene 6.

For SRE signaling assays, HEK293T cells were plated in a 96-well plate and incubated until 35-50% confluent. Cells were co-transfected with 2.5 ng Lphn, 10 ng Dual-Glo lucerifase reporter plasmid, and 0.3 μl transfection reagent LipoD293 (SL100668; SignaGen Laboratories). The Dual-Glo luciferase reporter plasmid was cloned from plasmids (FJ376737 and FJ773212) purchased from Promega.

**cAMP assay**
24 hrs after transfection, HEK293 cells were detached and re-plated in a white 96-well plate (50,000 cells per well) and incubated for another 24 hrs. For pertussis toxin inhibition, during the last 5 hrs of
incubation the transfected HEK293 cells were treated with 500 ng/mL pertussis toxin (Bordetella pertussis; PHZ1174; Life Technologies) to specifically block Gi coupling. Two types of cAMP assay were performed: β2-adrenergic receptor assay and forskolin assay.

For β2-adrenergic receptor assay: After the 24 hr incubation period, the medium was changed to 100 μL CO₂-independent Opti-MEM I Reduced-Serum Medium (31985070, Life Technologies). Following a 30 min incubation period, 1 μL Glosensor substrate (Promega, PRE1290) and 11 μL FBS (Sigma-Aldrich, F0926) were added to the cells. After pre-equilibration for 10-15 min, basal cAMP signal was measured. Cells were then treated with either 100 μM agonist peptide (synthesized by GenScript) or DMSO for 5 min, then activated with 50 nM isoproterenol (I650450, Sigma).

For forskolin assay: After the 24 hr incubation period, the medium was changed to 100 μL CO₂-independent Opti-MEM I Reduced-Serum Medium (31985070, Life Technologies) supplemented with 0.3 mM phosphodiesterase inhibitor 3-isobutyl-1-methylxanthine (IBMX; I5879; Sigma). Following a 30 min incubation period, 1 μL Glosensor substrate (Promega, PRE1290) and 11 μL FBS (Sigma-Aldrich, F0926) were added to the cells. After pre-equilibration for 10-15 min, basal cAMP signal was measured. Cells were then treated with either 100 μM agonist peptide (synthesized by GenScript) or DMSO for 5 min, then activated with 0.5 μM forskolin (Coleus forskohlii, F6886, Sigma).

For both assays, measurements were taken using Synergy HTX BioTeck plate reader at 25°C, GAIN 200 in 5 min intervals. Data from the 15 min time point were analysed in Excel and presented as average values from 3 repeats with standard errors for each sample.

**SRE assay**
Assay was performed as previously described (cite Salzman, et al., Neuron 2016). Briefly, HEK293T cells were co-transfected with a Dual-Glo SRE luciferase reporter plasmid and Lphn3. Prior to lysis, cells were subject to serum starvation for 15 hours. Firefly and renilla luminescence signals were measured on Synergy HTX BioTeck plate reader. The firefly:renilla ratio for each well was calculated and normalized to empty vector. Data are presented as average values from 3 repeats with standard errors.

**Western blot**
HEK293 cells were transfected in 6 well plates with 2 μg of receptor of interest or empty vector incubated 48 hrs, and washed with ice-cold PBS (1mM MgCl₂, 0.1mM CaCl₂), and stored at -80C. Cells
were quickly thawed at 37C, resuspended on ice in PBS+0.01% BSA+protease inhibitors, and subjected to centrifugation @6500xg for 5 min. The pellet was resuspended in 500 μl solubilization buffer (20mM HEPES, pH 7.4; 150 mM NaCl; 2 mM MgCl; 0.1 mM EDTA, 2 mM CaCl2; 1% Triton X-100; protease inhibitors) and rotated at 30 min and centrifugated at 20,000xg, for 15 min. Supernatant was mixed with 6x loading buffer and subject to western blotting. Expressed proteins were detected with mouse anti-FLAG antibodies 1:2000 (F3165; Sigma).

**Biotinylation of membrane proteins**

HEK293 cells were transfected in 6 well plates with 2 μg of receptor of interest or empty vector and incubated 48 hrs, then washed twice with ice-cold PBS (1mM MgCl2, 0.1mM CaCl2) and incubated for 30 min, 4°C with Sulfo-NHS-LC-Biotin reagent (21335; Life Technologies), final concentration 0.5 mg/ml, to allow biotinylation of membrane proteins. Biotinylating reagent was then washed out with ice-cold PBS (1mM MgCl2, 0.1mM CaCl2) and cell lysate was obtained after treatment of cells with solubilization buffer (with 1% Triton X-100) for 1 hr. Biotinilated proteins were captured with streptavidin paramagnetic beads (PR Z5482; Promega), eluted from beads with 4.5 M urea and 3x SDS-PAGE loading buffer, and subject to western blotting. Expressed Lphn1 FL, TM and P+TM proteins were detected with mouse anti-FLAG antibodies 1:2000.

**Flow cytometry**

HEK293 cells were transfected with 2 μg of DNA /well in 6-well plates using Fugene6. After 48 hrs of incubation, cells were detached with citric saline solution and washed with PBS and PBS+0.1% BSA (Bovine serum albumin, A3803, Sigma). The pellet was then stained with mouse anti-FLAG M2 primary antibody, 1:1000 (F3165) in PBS+0.1% BSA for 30 min (with rotation at room temperature), washed twice with PBS+0.1% BSA and incubated with anti-mouse FITC, 1:100 (F0257) secondary antibody for another 30 min and washed twice again. Pellets were resuspended in PBS+0.1% BSA. Flow cytometry data were collected on Guava Easycyte and BD Accuri C6 flow cytometers. Data were analyzed in FlowJo.

**GTPγS assay**

Protocol was implemented as previously described (Stoveken et al., 2015), the only difference being that for the present study, insect cell membranes were pre-incubated with G proteins for 5 minutes before starting the assay.

**Snake-plot and helix box diagrams**
Snake-plot and helix box diagrams as well as coloring of mutations were done using the Protter visualization tool.

**Receptor-peptide crosslinking experiments**

Purified hLphn3 7TM (40uM) and biotinylated hLphn3 peptide (80uM) were incubated for 30 minutes at 4°C. Glutaraldehyde (Sigma-Aldrich #G5882, 0.01%) was added to the sample and incubated for 5 minutes at room temperature. The crosslinking reaction was quenched with 0.5M Tris pH 8.0. The sample was subjected to SDS-PAGE and transferred to PVDF membrane for western blotting. Peptide was detected with NeutrAvidin DyLight 650 (Invitrogen #84607, 1:2000 dilution) and hLphn3 was detected with mouse anti-His primary antibody (Qiagen #34660, 1:2000 dilution) and donkey anti-mouse Alexa Fluor 488 secondary antibody (Invitrogen #A-21202, 1:5000 dilution).

**C. elegans experiments**

**Generation of plasmids and transgenes in C. elegans**

For generation of constructs for *in vivo* analyses recombineering was employed (Dolphin and Hope, 2006; Tursun et al., 2009) and accompanying protocols were modified as previously described (Langenhan et al., 2009; Prömel et al., 2012) to construct the respective Lphn transgenes using cosmids, PCR-amplified targeting cassettes and positive antibiotic selection. To introduce the point mutations into a cosmid containing the complete *lat-1* locus, a three-step strategy was followed. Firstly, an intermediate plasmid was constructed by inserting a FRT-kanamycin resistance gene (kanR)-FRT cassette amplified from pIGCN21 (Lee et al., 2001) into the intronic region of *lat-1* between exon 6 and 7 of cosmid pTL2 (Langenhan et al., 2009) and the entire exons 5, 6-FRT-kanR-FRT-exon 7 segment was retrieved into an outward-PCR-amplified pUC18 backbone (ThermoFisher Scientific). This intermediate vector was subject to QuikChange site-directed mutagenesis (Stratagene) introducing the respective point mutation in exon 6 of *lat-1* (CTA to GCT for L790A, CAC to GCT for H792A, TTC to GCT for F763A). A 2.5 kb fragment containing the respective point mutation was amplified, gel-purified and used as a recombineering targeting cassette on cosmid pSP5 containing the entire *lat-1* locus with a *gfp* fused to the second intracellular loop of the receptor (Langenhan et al., 2009). pSP5 had been transformed into SW105 cells beforehand (Warming et al., 2005), bacteria were made recombination-competent and the targeting cassette transformed for recombineering.

**C. elegans strains**

*C. elegans* strains were cultured and manipulated according to standard protocols (Brenner, 1974). Wild-type worms were Bristol, N2. The allele *lat-1(ok1465)* was generated by the *C. elegans* Gene
Knockout Consortium and provided by the Caenorhabditis Genetics Center (CGC), which is funded by NIH Office of Research Infrastructure Programs (P40 OD010440). The extrachromosomal transgenes aprEx196[lat-1::gfp(L790A) (pFF9) rol-6(su1006) pBSK], aprEx197[lat-1::gfp(H792A) (pFF10) rol-6(su1006) pBSK] and aprEx199[lat-1::gfp(F763A) (pFF11) rol-6(su1006) pBSK] were generated for this study. The following transgenes have been previously described: aprEx77[lat-1::gfp (pSP5) rol-6(su1006) pBSK] (Langenhan et al., 2009) and aprEx47[lat-1(1-581) (pTL20) rol-6(su1006) pBSK] (Prömel et al., 2012).

Generation of transgenic lines
All transgenic strains with stably transmitting extrachromosomal arrays were generated by DNA microinjection as described (Mello and Fire, 1995; Mello et al., 1991). Cosmids were injected at a concentration of 1 ng/μl together with the coinjection marker pRF4[rol-6(su1006)+] (100 ng/μl) and pBluescript II SK+ vector DNA (Stratagene) as stuffer DNA to achieve a final concentration of 120 ng/μl. DNA was injected into the syncytical gonad of lat-1(ok1465)/mIn1[mIs14 dpy-10(e128)] hermaphrodites. Transgenic progeny were isolated and stable lines selected. Multiple independent transgenic lines were established for each transgene tested.

Microscopy
For analysis of transgene expression young adult hermaphrodites were mounted in M9 onto a 2% agarose pad. Images were acquired with an Olympus Fluoview FV1000 confocal microscope.

Lethality and fertility rescue assays
Lethality and fertility rescue assays were conducted as previously described (Langenhan et al., 2009). For the lethality rescue assay fifty L4 hermaphrodites were transferred into wells of a 72-well flat-bottom Terasaki plates (Greiner Bio-One) containing E. coli OP50 in M9 and allowed to lay eggs for 24 h at 22 °C. Five to ten eggs were transferred into fresh wells with corresponding solutions and incubated at 22 °C. The number of dead/surviving embryos was scored 24 hours later, the number of adult animals 48 hours later.

For the fertility rescue assay ten L4 hermaphrodites were allowed to lay eggs at 22 °C on separate NGM plates seeded with E. coli OP50. Every 24 h hermaphrodites were transferred onto fresh plates until egg-laying ceased and embryos were scored. Experiments were conducted at least in independent triplicates. Data were examined using ANOVA with Bonferroni post-hoc test.