**Highlights**

- The LPHN olfactomedin-like domain forms a five-bladed β propeller
- A conserved calcium-binding site is located at the center of the protein
- Latrophilin-FLRT binding depends on a conserved binding site
- Mutations in the binding site inhibit Latrophilin-FLRT signaling

**In Brief**

Jackson et al. describe a crystal structure of mLPHN3 lectin and olfactomedin-like (Olf) domains, revealing the Olf β-propeller fold and calcium-binding site. Assays using HeLa cells and cortical neurons reveal a bi-functional role for Olf and its ligand FLRT, leading to HeLa cell adhesion and neuron repulsion.

**Accession Numbers**

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Structural Basis of Latrophilin-FLRT Interaction

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SUMMARY

Latrophilins, receptors for spider venom α-latrotoxin, are adhesion type G-protein-coupled receptors with emerging functions in synapse development. The N-terminal region binds the endogenous cell adhesion molecule FLRT, a major regulator of cortical and synapse development. We present crystallographic data for the mouse Latrophilin3 lectin and olfactomedin-like (Olf) domains, thereby revealing the Olf β-propeller fold and conserved calcium-binding site. We locate the FLRT-Latrophilin binding surfaces by a combination of sequence conservation analysis, point mutagenesis, and surface plasmon resonance experiments. In stripe assays, we show that wild-type Latrophilin3 and its high-affinity interactor FLRT2, but not the binding-impaired mutants we generated, promote HeLa cell adhesion. In contrast, cortical neurons expressing endogenous FLRTs are repelled by wild-type Latrophilin3 and not by the binding-impaired mutant. Taken together, we present molecular level insights into Latrophilin structure, its FLRT-binding mechanism, and a role for Latrophilin and FLRT that goes beyond a simply adhesive interaction.

INTRODUCTION

Latrophilins (LPHN1–3) are a family of adhesion type G-protein-coupled receptors (GPCRs) first characterized as calcium-independent receptors for the black widow spider toxin α-latrotoxin, which causes massive exocytosis of calcium at nerve terminals (Davletov et al., 1996; Krasnoperov et al., 1996; Lelianova et al., 1997; Sugita et al., 1998). More recently, LPHNs were shown to function as heterophilic cell adhesion molecules in processes such as synapse formation or maintenance. All LPHNs have a large ectodomain of approximately 1,000 amino acids, with an N-terminal lectin domain, followed by an olfactomedin-like (Olf) domain, a serine/threonine-rich region, HormR domain, and a GPCR-autoproteolysis inducing (GAIN) domain containing a GPCR proteolysis site (GPS). The structure of the N-terminal lectin domain was solved a few years ago using nuclear magnetic resonance (NMR) (Vakonakis et al., 2008). It folds into a β sandwich comprising two antiparallel sheets decorated with a 10-residue long α helix and two extended loops. One of the loops harbors a low-affinity rhamnose-binding site (Vakonakis et al., 2008). More recently, the crystal structure of the HormR and GAIN domains was solved by X-ray crystallography (Arac¸ et al., 2012). The structure revealed the basis for the autoproteolytic cleavage of LPHNs at the GPS via an evolutionarily ancient mechanism (Arac¸ et al., 2012). Within the LPHN ectodomain, the Olf domain is the only predicted globular domain that remains structurally uncharacterized.

LPHN1 and 3 are expressed predominantly in the brain. In humans, mutations in LPHN3 are associated with the largely hereditary attention-deficit hyperactivity disorder (ADHD) (Arcos-Burgos et al., 2010; Domène et al., 2011). In zebrafish, decreased activity of the LPHN3 homolog elicits ADHD-like behavior (Lange et al., 2012). The endogenous functions of LPHNs are linked to cell adhesion and synapse formation or maintenance, but the molecular mechanisms underlying these functions are only beginning to emerge. Known adhesive ligands of LPHNs include members of the teneurin family (Silva et al., 2011), neurexins (Boucard et al., 2012), and the fibronectin leucine-rich repeat transmembrane proteins (FLRT1–3) (O’Sullivan et al., 2012). FLRTs have recently emerged as powerful guidance factors in vascular, neural, and early embryonic development (Egea et al., 2008; Leyva-Díaz et al., 2014; Maretto et al., 2008; Müller et al., 2011; O’Sullivan et al., 2012; Seiradake et al., 2014; Yamagishi et al., 2011). They promote both cell adhesion (through homophilic interactions) and cell repulsion (through interaction with Uncoordinated-5 [Unc5] receptors). Due to their dual functionality, FLRTs are also referred to as repulsive cell adhesion molecules (ReCAMs) (Seiradake et al., 2014). LPHN3 and FLRT3 were reported to interact in trans through their ectodomains to mediate cell adhesion, an interaction that promotes the development of glutamatergic synapses (O’Sullivan et al., 2012). The LPHN Olf domain is required for the synapse-promoting function and also for FLRT binding (O’Sullivan et al., 2014).

Olf domains are present in at least 13 different proteins in mammals including the LPHNs (1–3), noelin (1–3; also called olfactomedins 1–3), olfactomedin 4, olfactomedin-like (1–3; with olfactomedin-like 2 separated into 2A and 2B), myocilins, and gliomedins (Tomarev and Nakaya, 2009); however, the lack of structural information has hampered their functional analysis. Here we present the crystal structure of an N-terminal fragment of murine LPHN3 comprising the lectin and Olf domains...
We expressed secreted murine LPHN3 Lec-Olf in human embryonic kidney (HEK) 293T cells using established protocols to circumvent the formation of heterogeneous glycans (Aricescu et al., 2006), and determined the crystal structure to a resolution of 2.16 Å (Table S1). The structure reveals two globular domains separated by a 17-residue linker (residues V192–K208) (Figure 1A). The structure of the LPHN3 lectin domain agrees closely with the lectin domain of LPHN1 solved using solution NMR (Vakonakis et al., 2008) (root-mean-square deviation [rmsd] is 0.86 Å for 90 aligned Cα atoms). The Olf domain folds into a five-bladed β propeller with metal ions buried in the center of the molecule. We show that, by introducing specific mutations at the surface of the Olf domain, we can reduce or impair binding to FLRT. We also show that LPHN3 Lec-Olf is sufficient to attract FLRT-expressing cells through its adhesive FLRT-binding properties. Interestingly, the FLRT-binding site on LPHN3 is required for repulsive activity of LPHN3 toward cortex-derived neurons. The structural data presented here suggest that LPHN3 is a bi-functional protein with adhesive and repulsive properties, possibly mediated by similar surfaces of the Olf domain.

RESULTS

Crystal Structure of the Murine LPHN3Lec-Olf

We expressed secreted murine LPHN3Lec-Olf in human embryonic kidney (HEK) 293T cells using established protocols to circumvent the formation of heterogeneous glycans (Aricescu et al., 2006), and determined the crystal structure to a resolution of 2.16 Å (Table S1). The structure reveals two globular domains separated by a 17-residue linker (residues V192–K208) (Figure 1A). The structure of the LPHN3 lectin domain agrees closely with the lectin domain of LPHN1 solved using solution NMR (Vakonakis et al., 2008) (root-mean-square deviation [rmsd] is 0.86 Å for 90 aligned Cα atoms). The Olf domain folds into a five-bladed β propeller, with each blade consisting of a four-stranded β sheet (Figure 1B). The five blades (I–V) are arranged successively in an anticlockwise direction around the central channel, giving rise to the disk-like architecture of the propeller. The β strands within each blade exhibit an up-and-down connectivity. The first strand packs against the first strand of the other blades in a parallel orientation to form a central channel. The fourth strands from blades I–IV largely form the periphery of the disk together with the Olf domain N terminus (β0), which contributes as the fourth strand to blade V. A disulfide bridge (C203–C385) connects the region upstream of β0 with blade IV and appears to stabilize β0 in its position. Notably for an extracellular protein, LPHN3 Lec-Olf contains a reduced cysteine (C227) buried in the core of the Olf domain. This cysteine is conserved in the three murine LPHNs, but not in other Olf domains (Figure S1A). The loops between the β strands within a sheet are generally short on the lectin-exposed side, with the exception of the β3–β4 loop which folds into an 8-residue helix sandwiched by blades I and V. The loops on the other side of the propeller are of varying length, the longest of which connects β8 and β9 and occludes the entrance to the central ion-containing channel.

In the electron density map, two potential metal atoms were visible approximately halfway through the central Olf channel. We analyzed the coordination patterns of these ions and calculated calcium bond-valence values (CBVS) to infer their chemical identities (Müller et al., 2003). The result of this analysis points to calcium (CBVS = 1.767) and sodium (CBVS = 1.560) for the octahedrally coordinated and pentacoordinated ion, respectively. The calcium ion equatorial ligands are two water molecules, the backbone carbonyl oxygen of A381, and the oxygen of the carbamamide N380. The backbone carbonyl oxygen of V435 and the carboxylic oxygen of D332 occupy the apical coordination sites (Figure 1C). The sodium ion is coordinated by the backbone carbonyl oxygens of G278 and L333 in addition to the carboxylic oxygens of D332 and D436, with the remaining coordination site filled by a water molecule (Figure 1C). Two additional water molecules are further away, shared with the calcium ion, and are not strictly speaking first-sphere coordination ligands of the sodium ion. The three amino acid side chains involved in coordinating the metal ions, D332, N380, and D436, are highly conserved (Figure 1D) among LPHNs, myocilin, olfactomedin-like 2A/B, and, to some extent, noelins, suggesting that the Olf domains from these proteins could also contain metal-binding sites.
Comparison of the LPHN3Lec-Olf structure with structures in the Dali database (Holm and Rosenström, 2010) reveals highest structural homology with a range of enzymes, especially the type I glutaminyl cyclases (QCs) from the plant pathogenic bacterium, Xanthomonas campestris (PDB accession code 3MBR) (Huang et al., 2010); the rmsd is 3.2 Å for 205 aligned Cα atoms. The catalytic site of this enzyme also contains a calcium ion in the central channel of the propeller (Figures S1B and S1C).

Mapping and Mutagenesis of the LPHN-FLRT-Binding Site

Given the importance of the FLRT-LPHN interaction in brain wiring and synaptic development (O’Sullivan et al., 2012, 2014), and the fact that FLRT and LPHN interactions were observed among multiple homologs (O’Sullivan et al., 2012), we hypothesized that the binding surfaces on FLRT and LPHN would be conserved across species. We generated sequence conservation scores (Glaser et al., 2003) using alignments of FLRTs and Latrophilins from mouse, chicken, frog, and fish, and mapped these onto the structures of LPHN3Lec-Olf and the previously solved FLRT2 leucine-rich repeat (LRR) domain (Seiradake et al., 2014) (Figures 2A and 2B). LPHN3Lec-Olf exhibits a sequence-conserved patch on the Olf domain, stretching across blades II and III of the propeller. Surfaces of blades I, IV, and V are relatively less conserved (Figure 2A). On FLRT2, a conserved surface patch extends from the concave to one lateral side of the LRR domain (Figure 2B). We previously showed that the conserved lateral side harbors the Unc5-binding side while the conserved concave side promotes FLRT-FLRT interaction (Seiradake et al., 2014). To probe these conserved surface regions, we generated a series of mutant proteins in which N-linked glycosylation sites were introduced at various positions. Introduction of N-linked glycans in protein-protein binding sites is an established way of disrupting protein binding (Seiradake et al., 2010, 2011, 2013, 2014). For LPHN3 these mutations were R292N+R294T (LPHN3L5), R324N+G326T (LPHN3LF2δ), T267N+K269T (LPHN3LF3δ), and P244N+R246T (LPHN3LF4δ) (Figures 2A and 2C). For FLRT2 we used previously described mutants FLRT2FF and FLRT2UF (Seiradake et al., 2014).

Surface plasmon resonance (SPR) experiments revealed high-affinity binding between wild-type FLRT2 and LPHN3 proteins, with Kd values in the range 18–40 nM depending on the experimental setup (Figures 3A–3E). In contrast, the LPHN3Lec-Olf mutants LPHN3L and LPHN3LF2δ do not have measurable affinity to the FLRT2 ectodomain. LPHN3LF3δ and LPHN3LF4δ bind FLRT2, but with greatly reduced affinity (Figures 3C–3E). We conclude that the FLRT-binding site involves conserved surface regions on blades II and III of the LPHN Olf domain. We also tested the binding of previously described FLRT2 mutants to the LPHN3 Olf domain and found that the non-Unc5 binding FLRT mutant (UF) binds with high affinity. In contrast, the non-FLRT-binding FLRT mutant (FF) lost measurable affinity for LPHN3. Thus the same FLRT2 surface that promotes adhesive homophilic interaction also promotes FLRT2-LPHN3 interaction. For subsequent functional analysis we used wild-type and LF mutant LPHN3Lec-Olf proteins. Consistent with the SPR data, wild-type LPHN3Lec-Olf, but not the mutant, bound to full-length FLRT2 expressed at the surface of cells (Figures 3F and 3G).

Figure 2. Surface Sequence Conservation Analysis and Mutagenesis

(A) The structure of LPHN3Lec-Olf is shown as surface views. A sequence alignment of mouse LPHN3 and 1, chicken LPHN2, and LPHN from fish (Danio rerio) and frog (Xenopus tropicalis) was used to produce surface conservation scores for LPHN3Lec-Olf with Consurf (Ashkenazy et al., 2010). Black indicates highest conservation scores and white the lowest conservation scores. Residue positions that were mutated to asparagine to introduce an artificial N-linked glycosylation site are labeled and colored red or cyan. (B) The structure of the FLRT2 LRR domain (FLRT2LRR) (Seiradake et al., 2014) is shown as surface views. A sequence alignment of FLRT2 and FLRT3 from mouse, frog, and fish was used to produce surface conservation scores. The conserved Unc5-binding site on FLRT2 is marked with a purple arrowhead. Mutants carrying an artificial N-linked glycosylation site at R186 (red) were previously shown to reduce adhesive properties of FLRTs (Seiradake et al., 2014).

(C and D) Cartoon views of LPHN3Lec-Olf and FLRT2LRR. Colors are according to the rainbow, as in Figure 1. Mutated residues are colored separately and depicted as spheres.

FLRT2-LPHN3 Interaction Controls Cell Adhesion and Repulsion

To assess the adhesive function previously postulated for the LPHN-FLRT interaction, we performed stripe assays in which HeLa cells transiently transfected with FLRT2-ires-GFP or the non-LPHN3-LPHN3 binding FLRT2 mutant (FLRT2FF-ires-GFP) chose between growth on alternating stripes of LPHN3Lec-Olf containing wild-type and mutant proteins with Cy3-conjugated anti-hFc antibody. The remainder of the dish was coated with control Fc protein and anti-hFc, resulting in the non-LPHN3-Lec-Olf-containing areas appearing black. Expression of FLRT2 in HeLa cells led to a significant increase in the preference for growth on the LPHN3Lec-Olf stripes (Figures 4B, 4E, and 4F).
We also performed stripe assays in which FLRT2-transfected cells were asked to choose between growth on the non-FLRT-binding LPHN3 mutant protein, LPHN3 LF, or control protein (Figures 4 C, 4E, and 4F). In these experiments, the strong adhesive effect was abolished. Similarly, overexpression in HeLa cells of the non-LPHN3-binding FLRT2 mutant (FLRT2 FF) led only to a mild preference for LPHN3 Lec-Olf stripes (Figures 4 D and 4E).

HeLa cells express endogenous FLRT1 and FLRT3 (data not shown), which may explain the small adhesive effect of LPHN3 Lec-Olf observed also in the presence of the mutant FLRT2. Thus our stripe assay data with transfected HeLa cells indicate that interaction between the LPHN3 Olf domain and the FLRTs promotes adhesion, consistent with the proposed role of LPHN3 as a positive regulator of synapse development.

Figure 3. Surface Plasmon Resonance Data Reveal FLRT2-LPHN3 Binding Surfaces
(A) Full ectodomains of wild-type murine FLRT2, the non-Unc5-binding mutant FLRT2 UF or the non-dimerizing mutant FLRT2 FF were immobilized on a CM5 Biacore chip and the purified olfactomedin domain of murine LPHN3 was injected as analyte at different concentrations. Response units are shown for each injection.
(B) Binding curves were fitted using maximum response units for each injection and a 1:1 binding model.
(C) Biotinylated wild-type or mutant LPHN3 sec-Olf was immobilized on streptavidin-coated Biacore chips and the ectodomain of FLRT2 was injected at different concentrations. Response units are shown as in (A).
(D) Binding curves from (C) were fitted using maximum response units for each injection and a 1:1 binding model.
(E) Calculated $K_d$ values are shown for binding of FLRT2 constructs to the LPHN3 olfactomedin domain (top three rows) and for LPHN3 sec-Olf constructs binding to FLRT2 ectodomain.
(F and G) We tested the binding of wild-type or LF mutant LPHN3 sec-Olf to HeLa cells transfected with a FLRT2-ires-GFP vector (Seiradake et al., 2014). Bound LPHN3 sec-Olf proteins were visualized using an antibody against the polyhistidine tag (red). Scale bar, 10 μm.

We also performed stripe assays in which FLRT2-transfected cells were asked to choose between growth on the non-FLRT-binding LPHN3 mutant protein, LPHN3 sec-Olf, or control protein (Figures 4C, 4E, and 4F). In these experiments, the strong adhesive effect was abolished. Similarly, overexpression in HeLa cells of the non-LPHN3-binding FLRT2 mutant (FLRT2 FF) led only to a mild preference for LPHN3 sec-Olf stripes (Figures 4D and 4E).
O’Sullivan et al., 2012). Next, we sought to test the behavior of cultured neurons expressing FLRTs on LPHN3-Lec-Olf stripes (Figure 5A). We chose cortical neurons, which express high levels of FLRT2 and FLRT3 (Seiradake et al., 2014). In contrast to HeLa cells, the neurons were repelled by LPHN3 Lec-Olf stripes (Figures 5B–5D). The repulsive effect was abrogated when the neurons were grown on non-FLRT-binding LPHN3 LF stripes (Figures 5B–5D), indicating that the FLRT-binding site on the LPHN3 Olf domain was responsible for the repulsive activity.

DISCUSSION

Our crystallographic data reveal the structure of a mammalian olfactomedin-like domain. The β-propeller fold reveals surprising structural homology with a range of enzymes, especially type I QCs from plant and bacteria. QCs catalyze the formation of pyroglutamate from glutamine or glutamate residues at the N terminus of proteins and peptides, and contain calcium or zinc in the catalytic site (Schilling et al., 2008a). Mammals express type II QCs, which are structurally unrelated and linked to neurodegenerative diseases such as Alzheimer’s disease (Schilling et al., 2008b). Given the structural similarity with the bacterial and plant enzymes, and the approximate conservation of the metal ions, it is tempting to speculate that the LPHN3 Olf also harbors an enzymatic activity. However, the ion-coordinating pockets are not structurally conserved between LPHN3 Olf and its closest structural relative, the QC from X. campestris (Figure S1C). Furthermore, a hydrophobic pocket located at the edge of the X. campestris QC active site, important for both substrate selection and the cyclization reaction (Huang et al., 2010), is markedly different in the LPHN3 Olf structure. In X. campestris QC, a glutamate residue (E89) acts as a general acid and base to assist in the intramolecular cyclization reaction. No such residue is found in the putative active site of LPHN3, meaning it is unlikely that LPHN3Olf and bacterial QCs have exactly the same function. The “entrance” to the ion-binding pocket of LPHN3 Olf is obscured by loops. Should the LPHN3 Olf harbor an enzymatic function, the binding of substrate would likely involve structural rearrangement of these loops, perhaps providing substrate specificity. We expect that future studies will shed light on this exciting question.

Among other mammalian Olf-containing protein families, the ion-coordinating residues of LPHN are conserved in at least three: myocilin, olfactomedin-like 2, and, to a lesser degree, noelins (Figure 1D). Indeed, previous biophysical analysis suggested that myocilin binds calcium (Donegan et al., 2012). Mutation of residue D380, which is equivalent to LPHN3 D332, leads to loss of bound calcium (Donegan et al., 2012). Interestingly, the myocilin mutant D380A leads to the development of glaucoma (Donegan et al., 2012), which is characterized by the progressive degeneration of the optic nerve (Stone et al., 1997). Thus the calcium-binding site we have described here for LPHN3 is of critical functional relevance, at least in myocilin. Whereas in myocilin, olfactomedin-like 2, and LPHNs the metal-binding site appears fully conserved, noelins contain an aspartic acid in the place of mLPHN3 N380. As a result, the predicted ion-binding pocket in noelins is likely to be more acidic, although it could possibly still bind calcium. Olfactomedin 4 contains an asparagine in the place of LPHN3 D332 (or myocilin D380), thereby lacking the negative charge that appears to be essential for calcium binding at that position. Olfactomedin-like 4 contains a basic residue, specific for calcium binding, which is not present in other forms of the protein. Olfactomedin-like 1, olfactomedin-like 3, and gliomedins differ even more substantially at the putative residues.
Figure 5. The FLRT-Binding Site of LPHN3Lec-Olf Is Required for Repulsion of Cortical Neurons

(A) Cartoon outlining the experimental setup for cortical neurons grown on alternating LPHN3 and FLRT stripes.

(B) Cortex-derived cultures (E15.5) were grown on alternating stripes of LPHN3Lec-Olf and FLRT control proteins. Cortical neurons expressing high levels of endogenous FLRTs (Seiradake et al., 2014) were immunostained for the neuron-specific III-tubulin (green). The neurons are repelled from LPHN3Lec-Olf, but not LPHN3Lec-Olf mutant, stripes. Insets are higher magnification images showing the distribution of cells on the stripes. Red bars indicate the locations of the red (LPHN3-containing) stripes. Scale bar, 200 μm.

(C) Quantification of the data shown in (B), calculated for III-tubulin-stained neurons on LPHN3 stripes, essentially as done for GFP-expressing HeLa cells in Figure 4E. *p < 0.01, two-tailed Student t test. Error bars represent the SEM.

(D) Cartoon summarizing how wild-type LPHN3Lec-Olf, but not the FLRT-binding-impaired mutant, repels cortical neurons.

### EXPERIMENTAL PROCEDURES

#### Cloning and Crystallography

Murine LPHN3 (LPHN3Lec-Olf) for crystallization (residues 92–463) and LPHN3Off (residues 202–463) were cloned into the pHLSec vector and transiently expressed in kifunensine-treated HEK293T cells using previously described methods (Aricescu et al., 2006) (see Supplemental Experimental Procedures for protein expression and crystallization details). X-Ray diffraction data were collected from native and derivative crystals at the peak wavelength of the platinum L-II edge at the Diamond Light Source beamline I04, processing the data using Xia2 (Winter et al., 2013). Autosharp (Vonrhein et al., 2007), Autobuster (Blanc et al., 2004), and Coot (Emsley and Cowtan, 2004) were used to solve the structure of LPHN3Lec-Olf (see Supplemental Experimental Procedures).

For binding studies, the full-length FLRT2 ectodomain (residues 35–540) was cloned into the same vector and transiently expressed in HEK293S GlnTI- cells.

#### SPR

Equilibrium experiments were performed at 25 °C using a Biacore T200 instrument (GE Healthcare). The experiments were carried out at pH 7.5 (PBS, 0.005% (v/v) polysorbate 20). The regeneration buffer was 2 M MgCl₂. FLRT2 ectodomain proteins were coupled to CMS Biacore chips via direct amine coupling. LPHN3Lec-Olf proteins were biotinylated enzymatically at a C-terminal avidity tag (Avi-Tag) and attached via the resulting biotin label to streptavidin-coated Biacore chip surfaces to mimic the native membrane insertion topology. Data were analyzed with Scrubber2 (BioLogic).

#### Protein Cell Binding

HeLa cells in six-well plates were transfected with 2.0 μg of pCAGIG vector DNA encoding full-length FLRT2 using FuGENE 6 transfection reagent (Promega), according to the manufacturer’s instructions. Thirty-six hours after transfection, cells were incubated for 20 min at 4 °C in Hank’s balanced salt solution containing LPHN3Lec-Olf or LPHN3Off LF mutant protein pre-clustered (ratio 1:4) with His-probe rabbit polyclonal immunoglobulin G (IgG) (Santa Cruz Biotechnology). Cells were washed twice with PBS and fixed with 4% paraformaldehyde for 20 min at 4 °C. Bound LPHN3Lec-Olf proteins were then visualized with Alexa561-conjugated anti-rabbit IgG antibody (Molecular Probes, 1:400) after 30 min of blocking with 1% BSA. Nuclei were counterstained with DAPI before mounting.
Stripe Assays

Stripe assays were prepared essentially as previously described (Yamagishi et al., 2011). In brief, stripe matrices were placed on 60-mm dishes and proteins were injected. After 30 min incubation at 37°C, dishes were washed with PBS and matrices were removed. The dishes were coated with Fc protein mixed with anti-hFc for 30 min at 37°C, washed with PBS and, for culturing neurons only, coated with 20 μg/ml of laminin overnight at 37°C. HeLa cells or dissociated cortical (E15.5) neurons were cultured on the stripes for 3 hr (HeLa cells) or 16 hr (neurons), fixed, and stained. Details are provided in the Supplemental Experimental Procedures.

ACCESSION NUMBERS

The RCSB PDB accession number for the Latrophilin 3 structure reported in this paper is 5afb.

SUPPLEMENTAL INFORMATION

Supplemental Information includes one figure, one table, Supplemental Experimental Procedures, and 3D Molecular Model and can be found with this article online at http://dx.doi.org/10.1016/j.str.2015.01.013.

AUTHOR CONTRIBUTIONS

V.A.J. performed the crystallographic analysis, protein engineering, SPR experiments, and cell stripe assays. D.dT. aided and oversaw all stripe assay experiments. M.C. purified proteins for functional analysis. P.R. analyzed the proteins and nucleic acids. Nucleic Acids Res.

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Supplemental Information

Structural Basis of Latrophilin-FLRT Interaction

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Figure S1, related to Figure 1. Sequence alignment of selected murine olfactomedin-like (Olf) domains and superposition of the mLPHN3 Olf and a glutaminyl cyclase from Xanthomonas campestris.

A Sequences from murine Latrophilins (mLPHN1-3), olfactomedin-like 2 (mOlfml2A,B), noelins (Noe1-3) and myocilin (mMyoc) were aligned and coloured according to sequence conservation using default parameters in ESPRIPT (Gouet et al., 2003). The unpaired cysteine 227 found in mLPHNs is marked with a black arrow head.

B LPHN3 Olf domain is shown in dark blue, with the sodium and calcium ions coloured in purple and skyblue, respectively. The structure of the X.campestris glutaminyl cyclase (PDB accession code 3MBR) (Huang et al., 2010) is shown in cyan, with the active site calcium ion shown in orange.

C Close-up view of the calcium-binding sites in the aligned LPHN3 Olf domain and X.campestris glutaminyl cyclase (QC) structures highlights the structural differences. The Ca\(^{2+}\)-coordinating residues E175 and E177 of X.campestris QC are sequence conserved among other bacterial and plant QCs, but not in LPHN3 Olf. Ca\(^{2+}\)-coordinating side chains are shown as sticks and aligned structures are shown in transparent cartoon representation. The colour scheme is as in panel B.
Supplemental Table:

|                  | LPHN3\textsuperscript{lec-off} (native) | LPHN3\textsuperscript{lec-off} (K\textsubscript{2}PtCl\textsubscript{6} derivative) |
|------------------|----------------------------------------|----------------------------------|
| **Data collection** |                                        |                                  |
| Wavelength (Å)   | 1.0711                                 | 1.0716                           |
| Space group      | I222                                   | I222                             |
| Cell dimensions  |                                        |                                  |
| $a, b, c$ (Å)    | 78.4, 96.6, 101.64                     | 77.09, 97.15, 101.66             |
| $\alpha, \beta, \gamma$ (°) | 90, 90, 90                         | 90, 90, 90                      |
| Crystal solvent content (%) | 46                                  | 46                               |
| Resolution (Å)*  | 62-2.16 (2.22-2.16)                    | 70-2.6 (2.66-2.6)                |
| $I / \sigma I$*  | 10.2 (1.7)                             | 14.7 (1.7)                      |
| Completeness (%)*| 99.5 (98.9)                            | 99.5 (95.0)                     |
| Redundancy*      | 3.49 (3.5)                             | 32.17 (18.36)                   |
| R-meas (%)       | 10.4 (86.1)                            | 25.3 (174.9)                    |
| **Refinement**   |                                        |                                  |
| Resolution (Å)*  | 62.1-2.16 (2.27-2.16)                  | -                               |
| No. of reflections* | 21031 (2605)                      | -                               |
| $R_{work} / R_{free}$* | 0.205 / 0.244                | -                               |
| **No. of atoms** |                                        |                                  |
| Protein          | 2814                                   | -                               |
| Heterogen        | 116                                    | -                               |
| Water            | 88                                     | -                               |
| Average $B$-factor | 42.7                                   | -                               |
| R.m.s. deviations|                                        |                                  |
| Bond lengths (Å) | 0.023                                  | -                               |
| Bond angles (°)  | 1.47                                   | -                               |
| Ramachandran plot for residues |                            |                                  |
| Favoured regions (%) | 95.01                                   | -                               |
| Allowed regions (%) | 4.99                                   | -                               |
| Disallowed regions (%) | -                                      | -                               |

Values in parentheses are for the highest-resolution shell.
For the calculation of $R_{free}$, 5% of reflections across all resolution shells were randomly assigned and used only for validation. $R_{free} = \frac{\sum_{Test} |F_{obs} - |F_{calc}|/\sum_{Test} |F_{obs}|.}$

Table S1, related to Figure 1. Crystallographic statistics.

Supplemental Experimental Procedures.

**Protein purification and crystallisation.** Cell culture medium containing recombinant proteins was collected and cleared by centrifugation and filtration. Secreted recombinant proteins were isolated from conditioned medium by immobilized metal affinity chromatography and purified further by size-exclusion chromatography in 10 mM Tris-HCl (pH 8.0), 150 mM NaCl. Prior to crystallization, recombinant endoglycosidase F1 (Chang et al., 2007) was added at a concentration of 0.01 mg/ml to all samples. LPHN3Lec-Off was concentrated to 9.3 mg/ml in 150 mM NaCl, 10 mM Tris-HCl pH 8.0. Crystals were grown by the vapour diffusion method at 20 °C by mixing protein and crystallisation solution (0.1 M MES buffer pH 6, 4M NaCl, 0.6 M non-detergent sulfo-betaine NDSB195) in a 1:1 (v/v) ratio. The crystals were flash-frozen in a cryoprotectant solution containing 75% crystallisation solution and 25% glycerol. To produce Pt derivative crystals, $K_2PtCl_6$ was added in powder form to the drop containing LPHN3Lec-Off crystals and the crystals were frozen in cryoprotectant solution six hours later.

**X-ray diffraction data processing.** One native and one Pt-derivative dataset were collected from LPHN3Lec-Off crystals and used as input for single isomorphous replacement phasing in Autosharp (Vonrhein et al., 2007). Autosharp identified five platinum positions and performed automatic phasing and model building. The model was refined against the native data with Autobuster (Blanc et al., 2004) and further manually built in Coot (Emsley and Cowtan, 2004). Programs from the Collaborative Computational Project 4 (CCP4) and Molprobity (Davis et al.,
2007) were used to validate the resulting structure. Crystallographic details are summarized in Table 1. Root mean square deviations (rmsd) between C-alpha atoms were calculated with Superpose (Maiti et al., 2004). The identities of the metal-ion sites were inferred using calcium bond-valence sum (CBVS), a convenient method for analysis of the geometric environment of potential sites with a view to metal-ion assignment (Müller et al., 2003). Analysis of the ion-coordination geometries in the LPHN3^Lec-Off crystal was performed by implementing formula (3) of reference (Müller et al., 2003) in a simple Unix shell script, and computing the value of CBVS from the experimentally observed ligand occupancies and ligand-ion bond distances. These values were used to infer the ions’ chemical identities by comparison with the expected CBVS values (CBVS_{Ca^{2+}}=2.0 and CBVS_{Na^+=1.57}).

**Stripe assays.** 50 μg/ml of LPHN3^Lec-Off (wild type or mutant) were mixed with Cy3-conjugated anti-hF_{C} antibody (Invitrogen) in PBS. Matrices (90 μm width) were placed on 60 mm dishes (Knöll et al., 2007) and proteins injected, resulting in red fluorescent stripes. After 30 min incubation at 37°C, dishes were washed with PBS and matrices were removed. The dishes were coated with 50 μg/ml of F_{C} protein mixed with anti-hF_{C} for 30 min at 37°C and then washed three times with PBS and, for culturing neurons only, coated with 20 μg/ml of laminin overnight at 37°C. HeLa cells (30,000 cells/experiment) or dissociated cortical (E15.5) neurons (50,000 cells/experiment) were cultured for 3 h (HeLa cells) or 16 h (neurons) on the stripes. Samples were fixed with 2% sucrose/4% PFA in PBS for 10-20 min at room temperature (RT). Cells were washed and the nuclei counterstained with DAPI before mounting. Neurons were washed and incubated with mouse monoclonal anti- beta-III tubulin antibody (Sigma) after 10 min permeabilization in 1% BSA, 0.1% Triton X-100/PBS. Cy2 or Cy3 anti-rabbit IgG secondary antibody (Jackson) was used to visualize the tubulin signal. The number of GFP+ pixels on red stripes were quantified with ImageJ
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