Supplementary Information for
A new family of neural wiring receptors in bilaterians defined by phylogenetic, biochemical and structural evidence

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Supplementary Methods

Phylogenetics

Putative Dpr and DIP homologs were identified using BLAST with *D. melanogaster* proteins as queries (1). To exclude distant IgSF homologs, the BLAST hits were used as queries to search over the *D. melanogaster* proteome (reciprocal BLAST), and only those with a Dpr or DIP as the top hit were retained. For identifying IgLON homologs in protostomes, human IgLONs were used as queries and reciprocal BLAST was performed on the human proteome. Amalgam, CG34353, CG7166, DIP, Dpr, Klingon, Lachesin, and Wrapper were identified as IgLON homologs in *D. melanogaster*. Only some of the 21 Dprs appeared as IgLON homologs, presumably due to their fast rate of evolution. Amalgam was excluded from analysis because it could not be reliably aligned to other proteins. Nectin, Necl, Kirrel, and Nephrin were the only other IgSF subfamilies that could be reliably aligned to the Wirin family.

Sequences for insect, bony fish, and tetrapod proteins were obtained from the NCBI protein database. To acquire sequences from other organisms, whose proteomes were generally poorly represented in the NCBI protein database, transcripts were assembled from RNA-seq reads in the NCBI SRA database (2). To selectively assemble specific transcripts rather than the entire transcriptome, RNA-seq reads with similarity to reference proteins were extracted using TBLASTN. Transcripts were then de novo assembled using Velvet/Oases (3, 4), and coding sequences were identified using TransDecoder (http://transdecoder.github.io) (5).

Only the Ig domains were included in the alignment because the rest of the proteins could not be aligned across paralogs. The number of Ig domains varied across proteins: two for Dpr, three for DIP, IgLON, Klingon, Lachesin, Nectin, and Necl, five for Kirrel, and eight to ten for Nephrin. The first one and a half Ig domains of Dpr aligned with the corresponding part of the 3-Ig proteins. The rest of Dpr aligned with the second half of the third Ig domain of the 3-Ig proteins. The first two Ig domains of the 3-Ig proteins aligned with the corresponding part of Kirrel and Nephrin. The alignment of the third Ig domain to the rest of Kirrel and Nephrin was ambiguous. However, the phylogeny was robust to the uncertainty in alignment. When alignments were generated using different gap penalties, the resulting phylogenies were topologically identical to that in Fig. 1. Furthermore, using only the first two Ig domains resulted in a phylogeny that is also topologically identical to that in Fig. 1.

Based on preliminary alignments and phylogenies, proteins with exceptionally long branches or nonsensical species placements were discarded. The remaining proteins were classified into paralog groups. Proteins in each paralog group were aligned separately, removing paralog-specific insertions and alignment-ambiguous regions. The full alignment was assembled from paralog alignments through sequential profile alignment. All alignments were generated using MUSCLE with default settings (6). Preliminary phylogenies were inferred using FastTree2 (7). The ML phylogenies were inferred using RAxML v8.2.12 (8). The best-fit model of evolution for the full phylogeny was WAG + G + I + X (X: ML estimation of equilibrium amino acid frequencies). The best-fit model for DIP, IgLON, and Klingon individually, however, were LG + G + I + X. Therefore, paralog relationships within each family were inferred in separate analyses. Approximate likelihood ratio statistics were calculated using PhyML v.3.0 under the topology and equilibrium amino acid frequencies inferred using RAxML (9).

The unreduced phylogenies are attached to the manuscript as datasets. The provided datasets are for the following phylogenies:

| Dataset | Supporting Figure | Phylogeny File |
|---------|-------------------|----------------|
| 1       | 1                  | Wirin family   |
| 2       | 2A                 | IgLON subfamily|
The Extracellular Interactome Assay (ECIA)

Interactions between ectodomains of Dprs, DIPs and homologs were tested using ECIA (10) with minor modifications: The promoters in the bait and prey expression vectors have been replaced with the constitutively active Actin 5C promoter from *D. melanogaster* in lieu of the inducible metallothionein promoter. The transfection agent was also changed to TransIT-Insect (Mirus), which was used according to the manufacturer’s recommended protocol. Mouse IgLON cDNAs were used in the binding experiments.

Throughout the manuscript, the outcome of the assay is reported in absorbance values at 650 nm, as the AP substrate KPL BluePhos (Seracare, catalog no. 5120-0061) is turned over to a blue product.

Homology Modeling

Dpr and DIP orthologs were modeled based on the Dpr6-DIP-α structure (PDB: 5EO9) using MODELLER (11). For the NEGR1-NTM IG1-IG1 complex, Dpr6 was used to model NEGR1, DIP-α and was used to model NTM. Further side chain rotamer optimization was performed using SCWRL4 (12) and manual inspection of alternate rotamers in PyMOL (13).

Expression, Purification and Crystallization of RIG-5 IG1 and ZIG-8 IG1

The N-terminal domain of *C. elegans* RIG-5 was cloned into pAcGP67A with a C-terminal hexahistidine tag, and co-transfected into Sf9 cells with linearized baculoviral DNA (Expression Systems) using the TransIT-Insect transfection reagent (Mirus). Amplified virus was used to infect High Five cells. Media was collected 60 hours post-infection. RIG-5 was first purified using Ni-NTA agarose resin, followed by size-exclusion chromatography on a Superdex 75 10/300 column (GE Healthcare) in HBS (10 mM HEPES, pH 7.2 and 150 mM NaCl). RIG-5 was crystallized with the sitting-drop vapor diffusion method, using a Mosquito crystallization robot (TTP Labtech) with 100 nl protein + 100 nl crystallant drops against a 50-µl crystallant reservoir.

The N-terminal domain of *C. elegans* ZIG-8 was expressed and purified as above. Successful expression required the extension of the IG1 construct to include all of the N-terminal sequence between the domain and the signal peptide. We now recognize that this part constitute the additional N-terminal helix, which packs onto and is disulfide linked to the IG1. ZIG-8-RIG-5 IG1-IG1 complex was purified on a Superdex75 10/300 column.

RIG-5 IG1 sample was concentrated to 17 mg/ml, and crystallized by the sitting-drop vapor diffusion method in 0.1 M HEPES pH 7.0 and 1 M sodium citrate. Crystals were cryoprotected in 0.1 M HEPES pH 7.0, 1.2 M sodium citrate, 10% glycerol and vitrified in liquid nitrogen. Diffraction data were collected at SSRL beamline 9-2.

ZIG-8-RIG-5 was crystallized at 17.5 mg/ml in two different conditions. Crystal form 1 (tetragonal) was crystallized in 0.2 M disodium hydrogen phosphate, 20% PEG 3350, and cryoprotected with 0.2 M NaCl, 0.1 M HEPES pH 7.2, 22%PEG 3350, 24% Glycerol. Crystal form II (monoclinic) was crystallized in 0.2 M lithium sulfate, 0.1 M sodium cacodylate, pH 6.5, 30% PEG 400. This condition did not require cryoprotection. Diffraction data were collected at APS beamline 24-ID-C.
RIG-5 homodimer structure was solved by molecular replacement (MR) using a DIP-η IG1 monomer (PDB ID: 6NRX) (14) as the model with PHASER (15). ZIG-8-RIG-5 complex was solved by MR using the RIG-5 IG1 homodimer structure with PHASER. The models were refined with phenix.refine (16) and real-space model building was performed in Coot (17). Model validation was performed using Molprobity (18) within the PHENIX suite (19).

**Surface Plasmon Resonance (SPR)**

ZIG-8 and RIG-5 full-length ectodomain constructs were expressed and purified as above. 2430 RUs of ZIG-8 was coupled on a Biacore CM5 chip (GE Healthcare) using NHS/EDC chemistry and random amine coupling. RIG-5 ectodomain was run over the chip as analyte in HBS with 0.05% Tween-20 as surfactant and 0.1% BSA to remove non-specific binding.

**Analytical Ultracentrifugation**

Full ectodomain ZIG-8 and RIG-5 were used in sedimentation velocity experiments in a Beckman analytical ultracentrifuge with an An50-Ti rotor at 20°C. Protein samples were placed in charcoal-filled Epon centerpieces sandwiched between sapphire windows. Centrifugation was done at 50,000 rpm.

The AUC data were analyzed using the $c(s)$ methodology in SEDFIT (20, 21). Partial-specific volume, density, and viscosity were calculated using SEDNTERP (22). Partial specific volumes used for RIG-5 and ZIG-8 were 0.7192 and 0.7227 cm$^3$/g, respectively. Figures were rendered in GUSSI (23). For $K_D$ analyses, GUSSI was used to integrate the $c(s)$ distributions, which were assembled into isotherm files that were imported into SEDPHAT (24), where a monomer-dimer model was imposed, with a fixed $s$-value for the dimer (4.2 S for RIG-5, 3.65 S for ZIG-8).

**Signal peptide and transmembrane helix predictions; Sequence numbering for RIG-5**

There is ambiguity with regards to the N-terminal end of RIG-5 covering the signal peptide. The C36F7.4f.1 transcript on the Wormbase database (25) only allows for a weak prediction of a signal peptide with Phobius or SignalP (26, 27), while the C36F7.4g.1 transcript yields a strongly predicted signal peptide (“MYLFALLCGVLLVFKQACSRG”) if the second methionine in the transcript is used as the start methionine and therefore sixty amino acids are removed from the transcript. We used a numbering scheme throughout the manuscript that uses the C36F7.4g.1 sequence with 60 amino acids removed from the N terminus. It should be noted that the mature proteins (i.e. after the signal peptides are processed) for both transcripts have identical sequences. In the manuscript, we do not make a call about which transcript(s) are actually expressed in worms.

**Definitions for protein families within the IgSF**

Nectins and Necls are two related families within the IgSF found in vertebrates. In humans, the family include nine members, Nectins 1 to 5 and Necls 1 to 4. It should be noted that Necl5 is more closely related to Nectins than Necls.

Here, we define Kirrels as orthologs of the *C. elegans* protein SYG-1. Kirrels are found across bilaterians, including SYG-1, Rst and Kirre (Duf) in *Drosophila*, and Kirrel1 (Neph1), Kirrel2 (Neph3) and Kirrel3 (Neph2) in vertebrates.

Nephhrins are heterophilic binding partners of Kirrels, and are orthologs of the *C. elegans* protein SYG-2. Nephhrins are found across bilaterians, including SYG-2, SNS and Hibris in *Drosophila*, and Nephrin in vertebrates.

**Constructs used in ECIA experiments for Wirins**

For ECIA experiments, expression constructs included the following residues.
ZIG-8 full-length ectodomain: Ala22 to Ser249 (Wormbase CDS Y39E4B.8).
ZIG-8 IG1: Ala22 to Pro137.
RIG-5 full-length ectodomain: Arg20 to Arg397 (Wormbase CDS C36F7.4e minus the N-terminal 60 amino acids).
RIG-5 IG1: Arg20 to Pro130.
Mouse OBCAM full-length ectodomain: Thr30 to Asn14 (NCBI Accession NP_808574.2)
Mouse NTM full-length ectodomain: Gly34 to Asn321 (NCBI Accession NP_758494.2)
Mouse NEGR1 full-length ectodomain: Val32 to Gly318 (NCBI Accession NP_001034183.1).
**Table S1.** Data and refinement statistics for x-ray crystallography of the RIG-5 homodimer and two crystal forms for ZIG-8–RIG-5 IG1-IG1 heterodimers.

| PDB ID       | RIG-5 IG1 homodimer | ZIG-8–RIG-5 IG1–IG1 crystal form 1 | ZIG-8–RIG-5 IG1–IG1 crystal form 2 |
|--------------|---------------------|-----------------------------------|-----------------------------------|
| **Data Collection** |                     |                                   |                                   |
| Space Group  | P622                | P422                              | C2                                |
| **Cell Dimensions** |                   |                                   |                                   |
| a, b, c (Å)  | 46.61, 46.61, 196.85| 80.08, 80.08, 166.62              | 87.46, 79.10, 89.98               |
| a, β, γ (°)  | 90, 90, 120         | 90, 90, 90                        | 90, 92.22, 90                     |
| Resolution (Å)| 50-1.42 (1.51-1.42)*| 100-2.00 (2.12-2.00)             | 100-1.70 (1.80-1.70)              |
| Rsym (%)     | 2.7 (20.2)          | 17.8 (169.3)                      | 4.7 (58.7)                        |
| <I>/<σ(I)>  | 55.5 (9.5)          | 7.94 (1.31)                       | 14.15 (1.62)                      |
| CC1/2 (%)    | 100.0 (99.6)        | 99.5 (67.2)                       | 99.9 (81.2)                       |
| Completeness (%) | 99.6 (97.7)        | 99.5 (97.6)                       | 96.8 (93.8)                       |
| Redundancy   | 17.7 (15.2)         | 13.2 (12.4)                       | 3.6 (3.4)                         |
| **Refinement** |                     |                                   |                                   |
| Resolution (Å)| 50-1.42 (1.48-1.42)*| 100-2.00 (2.05-2.00)             | 100-1.70 (1.72-1.70)              |
| Reflections  | 24787               | 37511                             | 65514                             |
| Rcryst (%)   | 19.70 (17.63)       | 17.42 (28.65)                     | 17.39 (42.52)                     |
| Rfree (%)†   | 22.61 (24.18)       | 20.76 (30.81)                     | 20.73 (43.50)                     |
| **Number of atoms** |             |                                   |                                   |
| Protein      | 841                 | 1772                              | 3562                              |
| Ligand/Glycans | 20                  | 30                                | 67                                |
| Water        | 94                  | 180                               | 404                               |
| **Average B-factors (Å²)** |             |                                   |                                   |
| All          | 34.08               | 52.05                             | 38.41                             |
| Protein      | 33.12               | 51.21                             | 37.40                             |
| Ligand/Glycans | 44.73              | 77.53                             | 58.25                             |
| Solvent      | 40.47               | 56.16                             | 43.98                             |
| **R.m.s. deviations from ideality** |             |                                   |                                   |
| Bond Lengths (Å)| 0.013              | 0.018                             | 0.006                             |
| Bond Angles (°)| 1.302              | 1.367                             | 0.821                             |
| **Ramachandran Statistics** |             |                                   |                                   |
| Favored (%)  | 96.23               | 95.50                             | 96.15                             |
| Outliers (%) | 0.0                 | 0.0                               | 0.0                               |
| Rotamer outliers (%) | 0.0         | 1.02                              | 0.25                              |
| All-atom Clashscore‡ | 3.51             | 2.82                              | 2.10                              |
| Coordinate error (Å)§ | 0.15           | 0.20                              | 0.24                              |

* The values in parentheses are for reflections in the highest resolution bin.
† 1240, 1961, and 3179 reflections (5%) were not used during refinement for cross validation.
‡ As reported by Molprobity.
§ Maximum-likelihood-based error estimate by phenix.refine version 1.15.

None of the models contained Cβ outliers.
**Fig. S1.** The ML phylogeny of the Dpr family. Approximate likelihood ratio statistics (aLRS) are shown as branch supports: ** < 9.2 (=2 ln100), * < 4.6 (=2 ln10), ~ < 2.2 (=2 ln3). Unmarked branches have aLRS > 9.2. The insets show the arthropod and mollusk phylogenies. The arthropod paralogs are labeled following the *D. melanogaster* Dpr nomenclature. The numbers next to some clades show the maximum number of paralogs in a clade when there are gene duplications in its subclades.
Fig. S2. The ML phylogeny of the DIP family shown as in Fig. S1.
Fig. S3. A. The ML phylogeny of the Klingon family shown as in Fig. S1.
B. The ML phylogeny of the Lachesin family shown as in Fig. S1.
Fig. S4. Homology modeling of the mouse NTM-NEGR1 complex.
A. Sequence alignment and secondary structural elements of Dpr6, DIP-α, and the five mouse IgLONs. Amino acids mutated in Fig. 2D are labeled with an asterisk. The sequence alignment (also in Fig 2B) visually demonstrates that IgLONs are co-orthologous to Dprs and DIPs, and are not only DIP orthologs.
B. Binding experiments for mouse IgLONs, Nectins and Nectin-like proteins using ECIA. The homodimerization of Rst, the fly ortholog of mammalian Kirrels, serves as a positive control.
C. Homology model of the NTM IG1-NEGR1 IG1 heterodimer based on the structure of the Dpr6-DIP-α complex. Side chains of amino acids mutated in Fig. 2C are shown as sticks.
Fig. S5. A. The domain structure of ZIG-8 and RIG-5. The gray arches represent predicted disulfide bonds. SP: Signal Peptide. TMH: Transmembrane helix. IG: Immunoglobulin domain.
B. Signal peptide and transmembrane helix predictions by the *Phobius* server (26).

C. Binding experiments for ZIG-8 and RIG-5, performed using ECIA. The heterophilic interaction between ZIG-8 and RIG-5 is observed between both ectodomains (C) and IG1 domains only (D). RIG-5 and ZIG-8 form weaker IG1-IG1 homodimers, similar to DIPs in *Drosophila*. Fc (bait) and AP<sub>5</sub> (prey) concentrations were normalized by dilutions.

D. Western blots of ZIG-8 and RIG-5 ECIA constructs used in Fig. S5C. Both bait and prey constructs were detected by an iFluor 488-coupled anti-His tag antibody (Genscript, A01800). The bands were quantified for normalization of protein amounts used in the ECIA experiment.

E. Mutations at the observed ZIG-8 and RIG-5 interface affect heterophilic binding. This panel includes a more extensive set of mutations than, and including, those in Fig. 3C. To effectively compare wild-type to mutants, protein concentrations within each mutant bait series (rows) were normalized. Each prey was tested at two concentrations (1x and 0.25 or 0.125x) to ensure that binding affinities are compared at non-saturating concentrations.

F. The structure of the ZIG-8 IG1-RIG-5 IG1 complex observed in crystal form #1 (tetragonal form) and the two NCS copies in crystal form #2 (monoclinic). The three structure models are within 0.3 Å rmsd of each other. The sticks on RIG-5 represent N-linked glycosylation.

G. The N-terminal α-helical addition to the IG1 domain in ZIG-8 (drawn in red).
Fig. S6. The heterodimeric ZIG-8-RIG-5 complex. 
A,B. ZIG-8 and RIG-5 interface amino acids L77, F85 (ZIG-8) and F70, F75 (RIG-5) are at the hydrophobic core (yellow residues) of the ZIG-8-RIG-5 interface, and fill in surface cavities in their corresponding binding partners.
C. Alignment of the entire IG domains for ZIG-8 and RIG-5 with Dpr6 and DIP-α. The amino acids mutated in Fig. 3C are labeled with an asterisk.
D. SPR data for the binding of RIG-5 ECD on a ZIG-8 ECD-coupled SPR chip. Thin black lines represent kinetic model fits to the binding data, collected at concentrations ranging from 10.8 nM to 7.9 µM. The calculated parameters are, $K_D = 7.9 \mu M$, $k_{on} = 7.59 \times 10^4 \pm 2.1 \times 10^3 M^{-1}s^{-1}$, $k_{off} = 0.60 \pm 0.02 s^{-1}$, with a mass transfer constant, $t_c = 2.9 \times 10^6 RU M^{-1}s^{-1}$, indicating that the mass transport limitation is dominant, and the kinetic parameters might be suspect.
E. Binding isotherm for equilibrium fits to the SPR data in (D). The calculated $K_D$ is $10.3 \pm 0.3 \mu M$ and maximal response is 2047 RU.
Fig. S7. The homodimeric ZIG-8 and RIG-5 complexes. A, B. $s_w$ data for RIG-5 ectodomain (A) and ZIG-8 ectodomain (B) are shown as circles, obtained by integration of the $c(s)$ distributions. The line is the fit to the data assuming a monomer-dimer model. The values between square brackets correspond to 68.3% confidence intervals. C, D. A view of the hydrophobic core of RIG-5 (C) and DIP-η (D) homodimers. Despite zero sequence identity at the core, the two structures take similar shapes and adapt nearly identical interaction geometries.
**Fig. S8.** Comparison of interfaces of IG1-IG1 complexes from Wirins and related families.

**A-F.** Amino acids at the interface are shown with side chains as sticks. In Wirins, yellow-colored amino acids are at the hydrophobic core, as previously defined in Cheng et al. (14); cyan represents periphery. Since there is no clear hydrophobic core at the interface for the Necl and SYG complexes, and all of their interface amino acids were colored light orange. The NEGR-1-NTM structure is a homology model (Fig. S4B), while all others are crystallographically determined. The structures were aligned so that the subunits depicted at the bottom are superimposed on to each other. The PDB IDs of the structures shown are 6NRW (A), 6NRX (B), 6ONB (C), 5ZO2 (E), and 4OFY (F).

**G.** Amino acid positions at the “hydrophobic core” is compared.
**Fig. S9.** Wirins and the four related IgSF protein families. Comparison of IG1-IG1 complexes from Wirins and related families. Family names are noted above the structures. The structures were aligned so that the subunits depicted on the left are superimposed on to each other. The PDB IDs of the structures shown include 6ONB, 6ON6, 6NRX, 6NRW, 4OF8, 4OFY, 3M45, and 5B21.
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