Development

**Multiplane Calcium Imaging Reveals Disrupted Development of Network Topology in Zebrafish**

*pcdh19* Mutants

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**Abstract**

Functional brain networks self-assemble during development, although the molecular basis of network assembly is poorly understood. Protocadherin-19 (pcdh19) is a homophilic cell adhesion molecule that is linked to neurodevelopmental disorders, and influences multiple cellular and developmental events in zebrafish. Although loss of PCDH19 in humans and model organisms leads to functional deficits, the underlying network defects remain unknown. Here, we employ multiplane, resonant-scanning in vivo two-photon calcium imaging of developing zebrafish, and use graph theory to characterize the development of resting state functional networks in both wild-type and pcdh19 mutant larvae. We find that the brain networks of pcdh19 mutants display enhanced clustering and an altered developmental trajectory of network assembly. Our results show that functional imaging and network analysis in zebrafish larvae is an effective approach for characterizing the developmental impact of lesions in genes of clinical interest.

**Key words:** calcium imaging; functional connectomics; pcdh19; zebrafish

**Significance Statement**

Non-clustered protocadherins are linked to neurodevelopmental disorders that include microcephaly, intellectual disability, autism spectrum disorders and epilepsy. In humans, mutations in protocadherin-19 (PCDH19) cause a female limited form of infantile epileptic encephalopathy and are associated with an increased incidence of schizophrenia and autism. In this study, we use large-scale calcium imaging to reveal that mutations in zebrafish pcdh19 alter the development of brain network topology. This work is the first to use functional imaging to explore the effects of a clinically relevant mutation on brain-wide network assembly in vivo. We show that graph analysis of spontaneous network activity is a sensitive method for revealing subtle changes to network architecture in response to genetic perturbations.

**Introduction**

Brain function and behavior are determined by the organization of the underlying neuronal networks, which are assembled through evolutionarily conserved developmental genetic programs. Thus, understanding the time course of network development can provide essential information on the underlyi

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insights into both normal function and the effects of pathogenic mutations. The importance of proper neural development is clearest in some of the most prominent brain disorders, such as schizophrenia and autism spectrum disorder, which arise during the course of development and have been referred to as “connectopathies” (Friston and Frith, 1995; Geschwind and Levitt, 2007; Collin et al., 2016; Yamasaki et al., 2017). Many of the genes linked to these disorders are important for core neurodevelopmental processes, such as axon guidance, dendrite growth and synaptogenesis (Hussman et al., 2011; Lal et al., 2015). To better understand how structural and functional brain networks are constructed during development, it is necessary to visualize them during the course of their assembly. Not only will this inform our understanding of how network organization relates to brain function, but it can also provide essential insight into how genetic changes perturb brain function by altering developmental trajectories.

The zebrafish is an important vertebrate model system with several advantages for studying neural development. In addition to being well suited to both forward and reverse genetics, the small size and transparency of their embryos make them ideal for in vivo imaging of neural development (Jontes et al., 2000; Niell et al., 2004; Ahrens et al., 2013; Liu et al., 2018). With advances in imaging technology (Ahrens et al., 2013; Yang et al., 2016), data processing (Mukamel et al., 2009; Freeman et al., 2014; Pnevmatikakis et al., 2016) and the engineering of genetically-encoded calcium indicators (Chen et al., 2013), it is now possible to simultaneously image the activity of large numbers of neurons in vivo (Keller and Ahrens, 2015; Jercog et al., 2016). These advantages make it possible to follow both the structural and functional development of neural networks in vivo both during normal development and in lines harboring deleterious mutations.

The 6-protocadherins (6-pcdhs) comprise a family of homophilic cell adhesion molecules that are differentially expressed in the developing nervous system (Kim et al., 2007; Blevins et al., 2011), and it has been proposed that this differential expression contributes to an adhesive code governing neural circuit organization (Wolvert and Lalande, 2001; Vanhalst et al., 2005; Light and Jontes, 2017). Several 6-pcdhs have been linked to epilepsy (Lal et al., 2015), autism (Marshall et al., 2008; Morrow et al., 2008; Butler et al., 2015), microcephaly (Aran et al., 2016), and intellectual disability (Kasnaukiene et al., 2012), suggesting that this family plays essential roles in vertebrate brain development. In humans, mutations in protocadherin-19 (PCDH19) cause a female-limited form of infantile epileptic encephalopathy (Dibbens et al., 2008; Depienne et al., 2009), with further work suggesting that PCDH19 is also linked to a broader array of neural disorders, including autism (Piton et al., 2011). While loss of pcdh19 affects visually-guided behaviors in zebrafish (Cooper et al., 2015) and heterozygous female mice show behavioral defects (Hayashi et al., 2017; Pederick et al., 2018), mutants in both models are homozygous viable and do not exhibit overt structural or anatomic phenotypes. Given the brain-wide expression of pcdh19, the viability of mutants, and the subtle behavioral defects, we wanted to investigate the influence of pcdh19 on the overall development of network organization and activity.

Here, we used multiplane in vivo two-photon microscopy of zebrafish larvae expressing GCaMP6s (Ahrens et al., 2013; Chen et al., 2013) to visualize the development of network activity at the level of single neurons in wild-type larvae and in mutants lacking pcdh19. Using graph theory to analyze resting state functional networks (Bullmore and Sporns, 2009; Rubinov and Sporns, 2010), we show that pcdh19 mutants display significant differences in several network measures as early as 3 dpf (post-fertilization), as well as altered developmental trajectories of network properties. These data show that in vivo functional imaging in larvae harboring clinically relevant mutations can reveal quantitative changes during the development of functional networks.

Materials and Methods

Fish maintenance and transgenic lines

Adult zebrafish (Danio rerio) were maintained at 28.5°C and staged according to Westerfield (1995). All animal procedures were performed in accordance with the Ohio State University animal care committee’s regulations. Embryos were raised in E3 embryo medium (Westerfield, 1995) with 0.003% phenylthiourea (Sigma-Aldrich) to inhibit pigment formation.

The transgenic line Tg(elavl3:GCaMP6s) was established using the plasmid Tol2-elavl3-GCaMP6s, kindly provided by M. Ahrens (Janelia Farms, Addgene, plasmid #59531). This transgenic line was crossed with the pcdh19 mutant line, pcdh19mutant (Cooper et al., 2015). All experiments were performed in pcdh19mutant larvae.

Imaging

Unanaesthetized larvae were embedded dorsal side up in 2% low melting point agarose, made up in E3 embryo medium. Imaging was performed on a custom-built resonant-scanning two-photon microscope. Briefly, excitation was provided by a Chameleon-XR Ti:Sapphire laser (Coherent, Inc.) tuned to 900 nm. We used a Nikon Apochromat 25×/1.1NA water-immersion objective for imaging. The resonant scanhead and controller, 3DMS robotic stage, GaAsP photomultiplier tubes and power supply were obtained from Sutter Instruments. A piezo-electric objective positioner (nPFocus250) was obtained from nPoint. Laser power was controlled by a Pocket’s cell (Conoptics Inc.). The microscope was run with ScanImage 5.2 (Vidrio Technologies). All other parts were obtained from Thorlabs. Image stacks of 12 optical sections (512 × 512) were collected at 1-s intervals for 15 min, with a pixel size of 1.3 μm. In older fish, there was a transient increase in fluorescence that accompanied the start of scanning (a stimulus artifact), so the first minute of imaging was removed before analysis for all movies.

For each group, we included data from eight to 11 larvae, which is in line with the number of fish used in comparable studies (Romano et al., 2015; Avitan et al., 2017). These were derived from at least two, separate crosses. The data presented here do not represent a
longitudinal study, i.e., we did not image the same fish on successive days. Thus, the number of independent larvae imaged was 38 and 35 for wild-type and pcdh19 mutants, respectively. Some movies showed drift in the z-direction over the first few minutes of imaging; these were excluded from the analysis presented here.

Image segmentation and extraction of calcium signals

To extract ΔF/F traces from our calcium imaging movies, we used the constrained non-negative matrix factorization method (CNMF), as described in Pnevmatikakis et al. (2016), after the volumetric data were separated into timeseries for each imaged plane. Individual planes were cropped to exclude the tectal synaptic neuropil and autofluorescent skin, then processed using CNMF. For the CNMF pipeline, we used the following parameters: \( \tau = 5 \), \( K = 300 \), merge_thr = 0.95. We overestimated the number of cells, as the program would discard cells during the refinement process. The processing pipeline was used as described at https://github.com/flatironinstitute/CaImAn-MATLAB. The data from each plane were then combined to produce a single dataset for an imaged larva.

Network analysis

We used the Brain Connectivity Toolbox (https://sites.google.com/site/bctnet/) to calculate all network measures (Rubinov and Sporns, 2010). Correlation matrices were generated for each multiplane dataset, with entries being the pairwise Pearson’s correlation coefficient for the ΔF/F traces of neurons i and j. As the number of detected components varied among our datasets, the sizes of our corresponding networks also varied. To compare the distributions of edge weights, we generated a cumulative probability distribution for each dataset (50 bins between -1.0 and 1.0), which could then be averaged with other distributions of edge weights, we generated a cumulative probability distribution for each dataset (50 bins between -1.0 and 1.0), which could then be averaged with other distributions of edge weights. To calculate network measures, we binarized our networks and calculated network measures across a series of global thresholds (0–0.7). The change in the distributions of edge weights between wild-type and pcdh19 mutant networks meant that network densities differed for a given threshold. To calculate network measures, we used the undirected binary versions of each function. To normalize complex network measures, such as clustering coefficient, path length, small-worldness, and transitivity, we generated time-series randomized surrogate networks (Louie and Wilson, 2001; Zalesky et al., 2012) for each experimental network, then calculated an average clustering coefficient and path length for each ensemble of randomized networks at each threshold. The thresholds of the randomized networks were adjusted to match the corresponding network densities of the experimental networks. The normalized clustering coefficient and path length were calculated as \( C_{norm} = C / C_{rand} \) and \( L_{norm} = L / L_{rand} \) respectively. Small-worldness was calculated as \( C_{norm} / L_{norm} \). Network transitivity is related to the clustering coefficient and is defined as the ratio of observed triangles in the network to all possible triangles:

\[
T = \frac{\sum_{i \in N} 2t_i}{\sum_{i \in N} k_i(k_i - 1)}
\]

Transitivity is a scalar value generated by the function transitivity_bin for binary undirected networks. We normalized transitivity to the mean calculated from an ensemble of 100 time-series randomized networks, as was done for CI and L. Assortativity is a correlation coefficient that measures the tendency of nodes to link to other nodes of similar degree:

\[
r = \frac{\left[ \sum_{u \in L} \frac{1}{2}(k_i + k_j) \right]^2 - \left[ \sum_{u \in L} \sum_{j \in L} \frac{1}{2}(k_i + k_j) \right]^2}{\left[ \sum_{u \in L} \frac{1}{2}(k_i^2 + k_j^2) \right] - \left[ \sum_{u \in L} \sum_{j \in L} \frac{1}{2}(k_i + k_j) \right]^2}
\]

Assortativity was calculated with the function assortativity_bin. In all cases \( (C_{norm}, L_{norm}, T, T) \), values for networks at each threshold were averaged within a group, and pairwise comparisons were made at each threshold between wild-type and pcdh19 mutant groups for a given developmental time. Data were not analyzed blind, as data processing by CNMF was largely automated, and analysis was performed identically across all datasets with minimal user intervention. To determine the statistical significance of differences between wild-type and mutant networks, we used the unpaired, two-tailed Student’s t test for the normally distributed graph metrics, calculated at each threshold. For the comparisons of network measures across developmental time in Figure 3, we used ANOVA with Tukey HSD. The Kolmogorov–Smirnov test was used to determine the significance of changes in the
distributions of correlation coefficients. Statistics were calculated either with RStudio or with Igor Pro (WaveMetrics).

Results

Multiplane calcium imaging in developing zebrafish larvae

To investigate the development of functional brain networks in the zebrafish, we employed multiplane two-photon calcium imaging in transgenic zebrafish that expressed the genetically-encoded calcium indicator GCaMP6s under the control of the elav3l promoter, Tg(elav3l:GCaMP6s). This promoter drives expression in nearly all neurons (Park et al., 2000; Ahrens et al., 2013). To provide a baseline for comparison to pcdh19 mutants, we first visualized spontaneous neural activity in unanaesthetized wild-type larvae at 3, 4, 5, and 6 dpf that were immobilized in agarose. To sample neural activity patterns throughout the midbrain and hindbrain, we collected 12 optical sections spaced at 10 μm at 1-s intervals for 15 min (Fig. 1A; Movie 1). Constrained non-negative matrix factorization (Pnevmatikakis et al., 2016) was used to segment movies of each imaged plane (Fig. 1B,C) and to extract ΔF/F traces from identified spatial components [regions of interest (ROIs); Fig. 1D]. For each imaged larva, fluorescence traces from all imaged planes were combined into a single dataset. We analyzed 8–11 larvae per timepoint for a total of 38 wild-type datasets.

Development of zebrafish network organization

Individual datasets consisted of an array of fluorescence traces that represented a broad sampling of neural activity in the midbrain and hindbrain of zebrafish larvae (Fig. 2A). A prominent feature of these data was the presence of synchronous bursts of activity, which we observe as early as 3 dpf. This synchronous activity has
been previously reported in zebrafish (Dunn et al., 2016; Avitan et al., 2017), and is similar to correlated activity that has been observed in other developing systems (Golshani et al., 2009). Using these datasets, we constructed correlation-based functional networks, with identified ROIs serving as nodes and pairwise correlation coefficients serving as edges (Fig. 2B). To begin to understand how these functional networks mature during development, we compared the distributions of correlation coefficients. As the number of detected cells and the sizes of corresponding networks varied among individual datasets, we generated cumulative probability distributions of edge weights for each network, allowing us to average them within an experimental group (Fig. 2C,D). The distribution of correlation coefficients was similar for 3 and 4 dpf and for 5 and 6 dpf, but showed a prominent shift between 4 and 5 dpf.

To explore the topology of network development, we applied graph theory to our datasets, as it provides a robust mathematical framework for analyzing functional brain networks and allows quantitative comparisons between experimental groups (Bullmore and Sporns, 2009; Rubinov and Sporns, 2010). A previous study characterized the development of neuronal cultures using multi-electrode array recordings and graph theory, revealing the emergence of rich-club topology, the tendency of highly connected nodes to be linked to other highly connected nodes (Schroeter et al., 2015). More recently, graph measures were used to assess the influence of visual experience on the development of the zebrafish optic tectum (Avitan et al., 2017). To calculate network measures, we binarized each network across a range of thresholds (van Wijk et al., 2010). The network density is the ratio of connections in a network to all possible connections. At a given threshold, densities varied as a function of developmental age (Fig. 2D; Table 1), in accord with the time course of changes in the distribution of correlation coefficients (Fig. 2C). The number of ROIs detected increased between 3 and 4 dpf, but was stable between 4 and 6 dpf (Fig. 2E).

We next calculated more complex network measures at each developmental time, across a range of thresholds. For example, the normalized clustering coefficient measures the proportion of closed triangles between a node and its neighbors (Watts and Strogatz, 1998). As correlation-based networks exhibit enhanced clustering due to the transitive nature of correlation coefficients (Zalesky et al., 2012), we normalized the clustering coefficient ($C_{norm}$) for each network against the mean calculated from 100 corresponding, time-series randomized networks (Louie and Wilson, 2001; Zalesky et al., 2012), adjusting the thresholds of the randomized networks to match the network density of the experimental network (Fig. 3A–D). Similarly, we calculated normalized characteristic path lengths ($L_{norm}$), which represents the average shortest path linking any two nodes in the network (Fig. 3A–D). While the path length remained relatively constant for the investigated time-points, the clustering coefficient increased between 3 and 4 dpf (Fig. 3A,B), but remained stable thereafter (Fig. 3B–E).

Nervous systems ranging from C.elegans to the human brain exhibit small-world properties, characterized by high clustering of connections and short path lengths (Sporns, 2010; Fornito et al., 2016). To explore this property in developing zebrafish brain networks, we calculated small-worldness ($\omega$) as the ratio of $C_{norm}/L_{norm}$, with values above 1 taken as evidence for small-world organization (Watts and Strogatz, 1998). Zebrafish larvae showed evidence of small-worldness across a range of thresholds at all ages (Fig. 3F), with an evident enhancement between 3 and 4 dpf, driven primarily by changes in the clustering coefficient at high thresholds (Fig. 3E).

Transitivity is a network measure similar to the clustering coefficient, measuring the probability that if a node is connected to two other nodes, those nodes will also be connected. The normalized transitivity exhibited a similar developmental time course (Fig. 3G) to what was found for clustering (Fig. 3E), increasing between 3 and 4 dpf and remaining stable thereafter. We additionally calculated the degree assortativity, which is similar to rich-club organization and measures the tendency of nodes to connect to other nodes of similar degree: highly connected nodes will tend to be connected to other highly connected nodes and sparsely connected nodes will tend to be connected to other sparsely connected nodes. We observed positive assortativity across a range of thresholds at all ages, but found little evidence for developmental changes (Fig. 3H). Overall, our data show that the complex network measures are relatively stable between 4 and 6 dpf, suggesting that the basic scaffold of the brain is established early during development.

**Altered network development in pcdh19 mutants**

To determine how loss of *pcdh19* affects brain development in zebrafish larvae, we crossed the Tg(elav3l:GCaMP6s) fish with a previously published pcdh19
mutant line, *pcdh19* and performed calcium imaging, as described above (Movie 2). In all, we imaged 8–10 larvae at each developmental time, for a total of 35 larvae imaged. Calcium imaging datasets from *pcdh19* mutants were qualitatively similar to those obtained from wild type: similar numbers of cells were identified, and these cells had comparable levels of activity (Fig. 2E). Like wild-type larvae, the *pcdh19* mutants exhibited a developmental increase in correlated activity (Fig. 4A). However, mutant networks show reduced correlation at 3 dpf, but enhanced correlation at 6 dpf (Fig. 4A,D), revealing that the developmental trajectory of functional connectivity is altered in *pcdh19* mutants. The change in the distribution of correlation coefficients was also reflected by changes in network density across a range of thresholds (Fig. 4E–H).

To determine the effects of *pcdh19* loss on network assembly, we compared complex network measures between mutant and wild-type larvae (Fig. 5). At 3 dpf, the *pcdh19* mutants displayed both higher clustering and a longer path length over a range of thresholds (Fig. 5A), The slight enhancement of clustering in the mutants was maintained at 4 dpf (Fig. 5B) and 5 dpf (Fig. 5C), but became more pronounced at 6 dpf at the higher thresholds (Fig. 5D). Like wild-type networks, the mutant networks had small-world properties across a range of thresholds at all ages (Fig. 5E–H). At 3 dpf (Fig. 5E) and 4...

**Figure 2.** Development of zebrafish brain networks. A, ΔF/F fluorescence traces for 2241 ROIs identified in a movie collected from a 4 dpf zebrafish larva. B, A correlation matrix showing the pairwise correlation for all components in the network. C, The average cumulative probability histograms for the correlation coefficients determined at each developmental time for wild-type larvae. Plots represent mean ± SEM. D, The average network density as a function of threshold for edge weights. Plots represent mean ± SEM. Statistical significance was determined by ANOVA and a Tukey test. Exact *p* values are provided in the statistical table. E, Shown are the numbers of segmented spatial components (ROIs) for wild-type and mutant larvae at each developmental time point. The numbers *n* are the number of larvae analyzed. The error bars represent SEM. *p* < 0.05.
### Table 1. Statistical Table

| Figure | Test | n | Statistical significance |
|--------|------|---|--------------------------|
| 2C     | Kolmogorov–Smirnov | 3 dpf = 8,536,216; 5 dpf = 21,599,014 | 2.20E-16 |
|        |      | 3 dpf = 8,536,216; 6 dpf = 26,218,604 | 2.20E-16 |
|        |      | 4 dpf = 27,511,930; 5 dpf = 21,599,014 | 2.20E-16 |
|        |      | 4 dpf = 27,511,930; 6 dpf = 26,218,604 | 2.20E-16 |
| 2D     | One-way ANOVA + Tukey | 3 dpf = 10; 5 dpf = 11 | 0.00883753 |
|        |      | 4 dpf = 8; 5 dpf = 11 | 0.00883753 |
|        |      | 0.1 3 dpf = 10; 5 dpf = 11 | 0.0257874 |
|        |      | 4 dpf = 8; 5 dpf = 11 | 0.0236042 |
|        |      | 0.2 3 dpf = 10; 5 dpf = 11 | 0.0145531 |
|        |      | 4 dpf = 8; 5 dpf = 11 | 0.0308327 |
|        |      | 0.3 3 dpf = 10; 5 dpf = 11 | 0.00743578 |
|        |      | 4 dpf = 8; 5 dpf = 11 | 0.0172431 |
|        |      | 0.4 3 dpf = 10; 5 dpf = 11 | 0.0230188 |
|        |      | 4 dpf = 8; 5 dpf = 11 | 0.0497095 |
| 3E     | One-way ANOVA + Tukey | 3 dpf = 10; 4 dpf = 8 | 0.230188 |
|        |      | 3 dpf = 10; 5 dpf = 11 | 0.0479055 |
| 3F     | One-way ANOVA + Tukey | 3 dpf = 10; 4 dpf = 8 | 0.0100918 |
|        |      | 3 dpf = 10; 5 dpf = 11 | 0.0460378 |
|        |      | 0.3 3 dpf = 10; 5 dpf = 11 | 0.0475936 |
|        |      | 0.4 3 dpf = 10; 5 dpf = 11 | 0.0187628 |
|        |      | 0.5 3 dpf = 10; 5 dpf = 11 | 0.0226596 |
|        |      | 0.6 3 dpf = 10; 5 dpf = 11 | 0.0383578 |
| 4A     | Kolmogorov–Smirnov | WT = 8,536,216; mut = 5,925,875 | 2.20E-16 |
| 4D     | t test, unpaired, two-tailed | WT = 26,218,604; mut = 20,481,645 | 2.20E-16 |
| 5A     | WT = 10; mut = 8 | WT = 9; mut = 8 | 0.0288994 |
| 5D     | WT = 10; mut = 8 | WT = 9; mut = 8 | 0.0485647 |
|        | WT = 9; mut = 8 | WT = 9; mut = 8 | 0.0395246 |
|        | WT = 9; mut = 8 | WT = 9; mut = 8 | 0.0264769 |
|        | WT = 8; mut = 8 | WT = 9; mut = 8 | 0.0255265 |
|        | WT = 8; mut = 8 | WT = 9; mut = 8 | 0.0249533 |
| 5E     | WT = 10; mut = 8 | WT = 9; mut = 8 | 0.025265 |
| 5F     | WT = 10; mut = 8 | WT = 9; mut = 8 | 0.024533 |
|        | WT = 9; mut = 8 | WT = 9; mut = 8 | 0.0249533 |
|        | WT = 9; mut = 8 | WT = 9; mut = 8 | 0.0118865 |
| 6A     | WT = 10; mut = 8 | WT = 9; mut = 8 | 0.0264769 |
| 6B     | WT = 10; mut = 8 | WT = 9; mut = 8 | 0.025265 |

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dpf (Fig. 5F), there was no difference in small-worldness between wild-type and pcdh19 mutant larvae. However, in contrast to wild type, pcdh19 mutants showed a steady increase in small-worldness at 5 dpf (Fig. 5G) and 6 dpf (Fig. 5H). While normalized transitivity was largely stable in wild-type mutants between 4 and 6 dpf, pcdh19 mutants showed an enhancement relative to wild type across all developmental times, as well as a steady increase between 3 and 6 dpf that was not observed in wild type (Fig. 6A–D). The increased transitivity at 6 dpf is consistent with the observed increase in clustering and small-worldness at 6 dpf. Degree assortativity was increased in pcdh19 mutants at 3 dpf, compared to wild-type larvae (Fig. 7A). Subsequently, this difference was lost (Fig. 7A–D); degree assortativity was similar between wild-type and mutant larvae at 5 dpf (Fig. 7C) and 6 dpf (Fig. 7D).

Collectively, our data show that loss of pcdh19 dramatically alters the development of functional connectivity in zebrafish larvae. Differences in some network measures, such as transitivity and degree assortativity, are evident at 3 dpf, while differences in network clustering and small-worldness arise gradually during development. These results show that lesioning pcdh19 alters networks organization, with an increased propensity for the clustering of neuronal connections.

Spatial distribution of network changes

We previously showed that pcdh19 is expressed in distinct domains along the anterior neural tube during segmentation stages; expression is enriched in the anterior neural tube and forebrain, as well as stripes in the midbrain and hindbrain (Emond et al., 2009; Biswas et al.,

| Figure | Test | n | Statistical significance |
|--------|------|---|------------------------|
| 0.6    |      |   | WT = 8; mut = 9 | 0.0400735 |
| 0.7    |      |   | WT = 8; mut = 9 | 0.0336544 |
| 6C     |      |   | WT = 11; mut = 10 | 0.0418721 |
| 7D     |      |   | WT = 9; mut = 8 | 0.0366925 |
| 0.2    |      |   | WT = 10; mut = 8 | 0.0251808 |
| 0.3    |      |   | WT = 10; mut = 8 | 0.0019526 |
| 0.4    |      |   | WT = 10; mut = 8 | 0.00027553 |
| 0.5    |      |   | WT = 10; mut = 8 | 0.00055079 |
| 0.6    |      |   | WT = 10; mut = 8 | 0.0111991 |
| 7C     |      |   | WT = 11; mut = 10 | 0.0357318 |

Summary of the statistics calculated in each figure, including n, p value, and type of test.
By 2 dpf, *pcdh19* is strongly expressed in the forebrain and hindbrain, as well as in cells lining the hindbrain and midbrain ventricles. We also showed that, later in development, *pcdh19* is expressed in columns of neurons in the optic tectum, generating a striped appearance (Cooper et al., 2015). Here, we further characterize a transgenic line, TgBAC(pcdh19:Gal4-VP16, 5xUAS:Lifeact-GFP), that drives expression of the F-actin marker Lifeact-GFP in *pcdh19/*H11001 cells (Fig. 8). In the 5 dpf larval brain, *pcdh19* is expressed in the vasculature, as well as progenitor cells and neurons (Fig. 8). Expression is strongest in the ventral midbrain (Fig. 8A,F), the optic tectum (Fig. 8C,D), and the hindbrain (Fig. 8B,C,E). In each region, *pcdh19* is expressed in only a subpopulation of neural progenitor cells and neurons. For example, as we previously reported, *pcdh19* is present in radial stripes of tectal neurons, that are typically associated with one or more radial glia (Fig. 8D). Similarly, in the hindbrain, we observe *pcdh19* expression in subpopulations of neurons within bilateral clusters (Fig. 8B,C,E). Among these clusters, *pcdh19* is expressed in neurons that likely correspond to the anterior rhobencephalic turning region (ARTR; Dunn et al., 2016). In contrast to the broad expression of *pcdh19* in multiple brain regions, it does not appear to be expressed in the cerebellum.

To better understand the network alterations observed in *pcdh19* mutants, we investigated the spatial distributions of the normalized clustering coefficients and network degrees (Fig. 9). Both the clustering coefficient and degree are node-based metrics that can be mapped back on to the coordinates of the corresponding spatial components. To generate heat maps, we calculated the clustering coefficients and node degrees from binarized networks that were thresholded at 0.7, and averaged values for all spatial components within sliding 20 × 20 μm windows. In wild-type larvae, the normalized clustering coefficient (Fig. 9A) did not exhibit a marked spatial pattern. Although there were scattered “hotspots” in both the midbrain and hindbrain in some embryos, there was not a consistent pattern. In *pcdh19* mutants, we find here that the observed elevation in the normalized clustering coefficient is distributed across the brain and is not localized to a specific region (Fig. 9B), suggesting that the increase in clustering is distributed across the entire network. To compare the spatial distribution of degrees in networks of

![Movie 2. GCaMP6s fluorescence in a pcdh19 mutant 3 dpf zebrafish larva.](image-url)

**Figure 4.** Development of basic network properties in wild-type and *pcdh19* mutant larvae. **A–D.** The cumulative probability distributions for correlation coefficients of wild-type (black) and *pcdh19* mutant (pink) larvae at 3 dpf (A), 4 dpf (B), 5 dpf (C), and 6 dpf (D). Plots represent mean ± SEM. Statistical significance was computed using a Kolmogorov–Smirnov test. Exact p values are provided in the statistical table. **E–H.** The network density as a function of correlation coefficient threshold for wild-type (black) and *pcdh19* mutant (pink) networks at 3 dpf (E), 4 dpf (F), 5 dpf (G), and 6 dpf (H). Plots represent mean ± SEM. Significance was computed using an unpaired, two-tailed Student’s t test for each 0.1 increment. Exact p values are provided in the statistical table. *p < 0.05.*
different sizes, we normalized each map to the highest degree in the individual network (values vary between 0 and 1). In contrast to the normalized clustering coefficient, the normalized node degree was less isotropic, with high-degree nodes consistently concentrated in the hindbrain and rostral midbrain of wild-type larvae (Fig. 9C). The pcdh19 mutants exhibited a shift in the spatial distribution of high degree nodes, with a large expansion of area in the hindbrain (Fig. 9D). As pcdh19 is expressed in clusters of neurons in the hindbrain (Fig. 8), this shift could be due to increased interconnectivity of hindbrain neurons.

Figure 5. Altered development of small-worldness in pcdh19 mutants. A–D, Comparison of the normalized clustering coefficient (solid) and path length (dotted) between wild-type (black) and pcdh19 mutant (pink) larvae at 3 dpf (A), 4 dpf (B), 5 dpf (C), and 6 dpf (D). Plots represent mean ± SEM. Significance was computed using an unpaired, two-tailed Student’s t test. Exact p values are provided in the statistical table. E–H, Comparison of small-worldness between wild-type (black) and pcdh19 mutant (pink) larvae at 3 dpf (E), 4 dpf (F), 5 dpf (G), and 6 dpf (H). Values above 1 (dotted-line) indicate the presence of small-world organization. Plots represent mean ± SEM. Significance was computed using an unpaired, two-tailed Student’s t test. Exact p values are provided in the statistical table, although these differences were not statistically significant. *p < 0.05.

Figure 6. Developmental changes in transitivity in pcdh19 mutants. A–D, Comparison of transitivity between wild-type (black) and pcdh19 mutant (pink) larvae at 3 dpf (A), 4 dpf (B), 5 dpf (C), and 6 dpf (D). Plots represent mean ± SEM. Significance was computed using an unpaired, two-tailed Student’s t test. Exact p values are provided in the statistical table. *p < 0.05.

Figure 7. Development of assortativity in wild-type and pcdh19 mutant larvae. Comparison of assortativity between wild-type (black) and pcdh19 mutant (pink) larvae at 3 dpf (A), 4 dpf (B), 5 dpf (C), and 6 dpf (D). Plots represent mean ± SEM. Significance was computed using an unpaired, two-tailed Student’s t test. Exact p values are provided in the statistical table. *p < 0.05.
Figure 8. Expression of pcdh19 in BAC transgenic line. **A–C**, Distribution of pcdh19 expression in the midbrain and hindbrain of *TgBAC(pcdh19:Gal4-VP16, 5xUAS:Lifeact-GFP)* larvae at 4 dpf. In addition to neurons and neural progenitor cells, pcdh19 is
Non-clustered protocadherins are essential for nervous system development, as mutations in these genes have been linked to autism (Marshall et al., 2008; Morrow et al., 2008; Piton et al., 2011; Butler et al., 2015), schizophrenia (Gregório et al., 2009), intellectual disability (Kasnauskiene et al., 2012), microcephaly (Aran et al., 2016), and epilepsy (Dibbens et al., 2008; Lal et al., 2015). Our long-term objectives are to understand the cellular roles of these genes and how defects in these cellular roles lead to brain-wide defects during development. Pcdh19 is a δ2-pcdh and mutations in human PCDH19 cause a female-limited form of early onset epileptic encephalopathy. Our goal here was to explore how loss of pcdh19 influences brain network organization and development. We used in vivo multiplane two-photon calcium imaging to record neural activity from thousands of neurons in the midbrain and hindbrain regions of developing zebrafish, in both wild-type and pcdh19 mutant larvae. To

**Figure 9.** Spatial distribution of node-based measures in a 6 dpf larva. **A, B.** The normalized correlation coefficients were averaged in sliding 20 × 20 μm windows. Maps were generated from binarized networks that were thresholded at 0.7. The corresponding spatial maps show that there is little spatial organization in the distribution of clustering coefficients. Compared to wild-type larvae (A), there is a uniform elevation of the clustering coefficient throughout the brain in pcdh19 mutants (B). **C, D.** The node degree for each network was normalized to the largest degree in that network. These normalized node degrees were averaged in sliding 20 × 20 μm windows. Wild-type larvae (C) consistently showed regions of high degree nodes in the anterior midbrain and in the hindbrain. In pcdh19 mutants (D), there was a consistent spread of high degree nodes throughout the hindbrain.
gain insight into the trajectory of network development, we collected data between 3 and 6 dpf, a period of dramatic growth and network maturation; by 6 dpf, zebrafish larvae already exhibit a rich array of behaviors (Orger and de Polavieja, 2017).

Neural activity in pcdh19 mutants is qualitatively similar to that of wild-type larvae, as comparable numbers of active neurons are detected, which have similar levels of activity. Additionally, the brains of pcdh19 mutant larvae do not display any overt morphogenetic or neuroanatomical defects. Yet, despite the fact that pcdh19 is expressed in only a subpopulation of neurons, mutants exhibit several quantitative differences in network organization starting at 3 dpf, the earliest timepoint that we investigated. Both wild-type and mutant larvae show positive assortativity across all developmental times; pcdh19 mutants show a significant elevation of assortativity at 3 dpf, but is indistinguishable from wild type at later times. At 3 dpf, pcdh19 mutants have reduced levels of correlated activity and correspondingly lower network densities. This could reflect a delay or decrease in synaptogenesis at this early stage. By 6 dpf, the most prominent difference was the increase in network clustering in mutant larvae, as shown by increases in the clustering coefficient, small-worldness, and transitivity. In each case, the differences increase progressively over time, and are most pronounced at 6 dpf. These data suggest an aberrant patterning of synaptic connectivity in mutants lacking Pcdh19 function.

In functional MRI studies, resting state functional networks are routinely used to explore the topology of brain networks, relying on statistical correlations of activity patterns to identify functional connectivity between regions (Fornito et al., 2016). We have adopted this approach in the analysis of the larval zebrafish brain, as it allows the calculation of network properties for quantitative comparisons between genotypes and across development. Our calcium imaging data offered a number of obstacles to quantitative comparison between groups. While a powerful aspect of our approach is that our networks are assembled from calcium signals obtained largely from single neurons, the number of neurons detected and the size of the corresponding networks varied from one embryo to the next, restricting the network measures that could be averaged and compared across experimental groups. However, we observe robust quantitative differences in network properties between wild-type and pcdh19 mutant larvae at 6 dpf. Moreover, we also see differences in the developmental trajectories of pcdh19 mutants. Importantly, this is a relatively fast and efficient method to investigate changes in the development of network structure and dynamics, as quantitative differences can be observed even where structural defects are not apparent (no discernible change to synapse number or axon trajectories). Several groups have developed methods for registering transgenic and antibody-labeled larvae to model zebrafish brains (Randlett et al., 2015; Tabor et al., 2019). As markers for different brain regions and cell types become available, this will enable the assembly of cellular resolution atlases of the developing zebrafish brain. In future, as these atlases become better annotated, registration of our calcium imaging datasets will facilitate more detailed analyses and the construction of mesoscale functional networks.

The δ-pcdhs play diverse roles during development, having been implicated in nearly every aspect of brain assembly (Light and Jontes, 2017; Jontes, 2018). A major unresolved question for δ-pcdhs, as for other molecules, is the relationship of cellular phenotypes to changes in network-wide functional connectivity. Several studies showed that δ-pcdhs regulate cell movements and cell migration during development (Bradley et al., 1998; Aamar and Dawid, 2008; Emond et al., 2009; Biswas et al., 2010; Williams et al., 2018). Multiple lines of evidence suggest that δ-pcdhs are expressed in neural progenitor cells and regulate proliferation and neurogenesis (Zhang et al., 2014; Cooper et al., 2015; Fujitani et al., 2017). After neurogenesis, δ-pcdh family members have been implicated in axon outgrowth, guidance and arborization (Biswas et al., 2014; Hayashi et al., 2014; Leung et al., 2015). In addition, Pcdh11 has been shown to regulate dendritic branching in cultured cortical neurons (Wu et al., 2015). More broadly, in vitro studies show that δ-pcdhs can mediate cell sorting (Bisogni et al., 2018; Pederick et al., 2018). Defects in any of these developmental processes are likely to have an impact on patterns of synaptic connectivity and network topology. It has previously been shown that pcdh19 mutants exhibit increased neurogenesis in the optic tectum, as well as disorganization in neuronal columns and patterns of arborization. We hypothesize that loss of pcdh19 leads to network defects through the accumulation of several cellular perturbations. Loss of adhesion by Pcdh19 results in increased proliferation and differentiation of those neural progenitors that would normally have expressed pcdh19. Loss of contact-dependent axonal and dendritic growth could result in reduced fasciculation and inappropriate arborization, leading to promiscuous patterns of synaptogenesis. The increased number of neurons that form inappropriate synapses could lead to increased clustering in the mutant networks.

A host of genes have been linked to neurodevelopmental disorders through linkage studies and genome-wide association studies or whole genome sequencing approaches. Frequently, these genes participate in fundamental aspects of neural development (e.g., neurogenesis, neuronal migration or axon guidance), yet the mechanisms linking the genetic lesions to impaired brain function remain obscure. With the ease of making targeted genomic lesions with CRISPR/Cas9, the ability to screen large scale patterns of functional connectivity in the zebrafish brain and to follow the effects longitudinally during development offers a powerful approach to explore the biological mechanisms underlying complex brain disorders. Moreover, the zebrafish offers the possibility of combining these functional imaging approaches with simultaneous imaging and analyses of changes in axon tract development, synaptogenesis or neurogenesis. Thus, it should be possible to understand defects in functional networks in terms of the underlying changes in neural organization, and to link these changes to the cellular roles of altered genes.
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