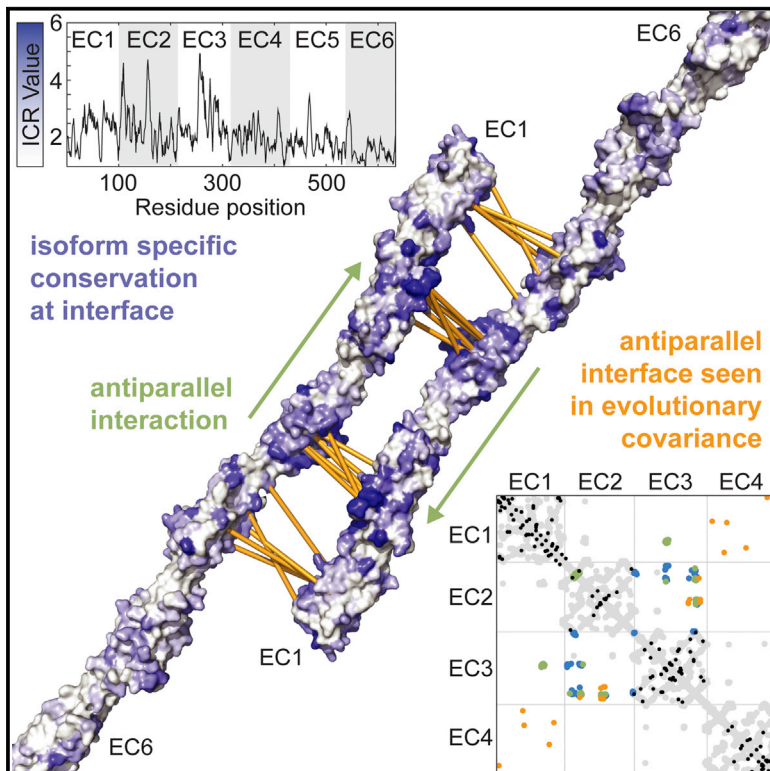


Structure

Structure and Sequence Analyses of Clustered Protocadherins Reveal Antiparallel Interactions that Mediate Homophilic Specificity

Graphical Abstract



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In Brief

Nicoludis et al. determined the structures of two clustered protocadherin fragments and found they formed antiparallel complexes. The authors provide a scaffold for isoform-specific interfaces and propose a full-length antiparallel complex that is responsible for homophilic *trans* interactions that mediate self-avoidance during synaptogenesis.

Highlights

- Clustered protocadherin EC1–3 fragments form antiparallel complexes in crystals
- A full extracellular cadherin repeat region model yields an extended molecule
- Isoform-specific conservation of interfaces explains strict homophilic specificity
- Evolutionary correlations provide support for antiparallel arrangement

Accession Numbers

4ZI9
4ZI8

Structure and Sequence Analyses of Clustered Protocadherins Reveal Antiparallel Interactions that Mediate Homophilic Specificity

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<http://dx.doi.org/10.1016/j.str.2015.09.005>

SUMMARY

Clustered protocadherin (Pcdh) proteins mediate dendritic self-avoidance in neurons via specific homophilic interactions in their extracellular cadherin (EC) domains. We determined crystal structures of EC1–EC3, containing the homophilic specificity-determining region, of two mouse clustered Pcdh isoforms (Pcdh γ A1 and Pcdh γ C3) to investigate the nature of the homophilic interaction. Within the crystal lattices, we observe antiparallel interfaces consistent with a role in *trans* cell-cell contact. Antiparallel dimerization is supported by evolutionary correlations. Two interfaces, located primarily on EC2–EC3, involve distinctive clustered Pcdh structure and sequence motifs, lack predicted glycosylation sites, and contain residues highly conserved in orthologs but not paralogs, pointing toward their biological significance as homophilic interaction interfaces. These two interfaces are similar yet distinct, reflecting a possible difference in interaction architecture between clustered Pcdh subfamilies. These structures initiate a molecular understanding of clustered Pcdh assemblies that are required to produce functional neuronal networks.

INTRODUCTION

Clustered protocadherins (Pcdhs) have an important role in cell-cell interactions in neurons (Weiner and Jontes, 2013; Yagi, 2012). The Pcdh gene cluster comprises three groups, *Pcdh α* , *Pcdh β* , and *Pcdh γ* , producing 53 variable isoforms in humans. Various studies have associated clustered Pcdhs with synapse maintenance and formation (Fernández-Monreal et al., 2009; Li et al., 2012; Weiner et al., 2005), promotion of synapse development by astrocytes (Garrett and Weiner, 2009), connectivity between terminals of type Ia afferent neurons and ventral inter-

neurons (Prasad and Weiner, 2011), and arborization of cortical pyramidal neurons (Garrett et al., 2012). Loss of the Pcdh γ cluster has been linked to apoptosis, neurodegeneration, and synapse loss in different neuron populations (Chen et al., 2012; Emond and Jontes, 2008; Lefebvre et al., 2008; Prasad et al., 2008; Su et al., 2010; Wang et al., 2002), indicating that the role of clustered Pcdhs is complex and multifaceted.

The clustered Pcdhs have also been associated with self-avoidance in Purkinje and starburst amacrine cells, which results in dendritic self/non-self discrimination during synaptogenesis (Kostadinov and Sanes, 2015; Lefebvre et al., 2012). Each neuron stochastically expresses 5–10 isoforms (Esumi et al., 2005; Kaneko et al., 2006; Yagi, 2012; Yokota et al., 2011). In addition, the five C-type isoforms are constitutively expressed (Kaneko et al., 2006). Part of the cadherin superfamily of Ca²⁺-dependent adhesion proteins, clustered Pcdhs, are thought to use specific homophilic interactions to signal self-avoidance (Lefebvre et al., 2012). In insects, the *Dscam1* gene plays a role analogous to that of the clustered Pcdhs (Zipursky and Sanes, 2010). Unlike *Dscam1*, where the homophilic interfaces are well characterized (Meijers et al., 2007; Sawaya et al., 2008), the structural determinants of clustered Pcdh interactions are largely unknown.

Each clustered Pcdh isoform contains six extracellular cadherin (EC) repeats followed by a transmembrane helix and a C-terminal intracellular domain. Although there are many known classical cadherin structures, few structures of the protocadherin subfamily are available, with the only known interface being the tip link Pcdh15–Cdh23 complex involved in mammalian hearing (Sotomayor et al., 2012). Understanding the putative homophilic interactions of clustered protocadherins requires structures of these complexes.

Specific homophilic interactions are common in the cadherin superfamily. In classical cadherins, the first repeat, EC1, mediates homophilic interactions, either through an “X-dimer” complex or a tryptophan-containing strand-swapping mechanism (Brasch et al., 2012; Sotomayor et al., 2014). In contrast, while the clustered Pcdh EC1 is required for complex formation, EC2 and EC3 determine interaction specificity (Schreiner and Weiner, 2010; Thu et al., 2014). Thus the nature of clustered Pcdh

Table 1. Statistics for A1 and C3 Structures

	Pcdh γ A1 EC1–3	Pcdh γ C3 EC1–3
PDB ID	4ZI9	4ZI8
SBGrid DataBank ID	173	174
Data Collection		
Beam source	APS 24-ID-E	APS 24-ID-C
Wavelength (Å)	0.97918	0.97918
Space group	P2 ₁	P2 ₁ 2 ₁ 2 ₁
Unit cell		
a, b, c (Å)	67.751, 63.687, 107.486	44.123, 82.286, 237.69
α , β , γ (°)	90, 99.693, 90	90, 90, 90
Resolution (Å)	50–1.77 (1.833–1.77)	50–1.70 (1.73–1.70)
Total reflections	269,651 (13,875)	281,547 (8,193)
Unique reflections	77,361 (5,120)	84,132 (3,718)
Multiplicity	3.5 (2.7)	3.3 (2.2)
Completeness (%)	87 (58)	86 (39)
Mean I/ σ (I)	5.70 (0.46)	5.38 (0.53)
Resolution shell at I/ σ (I) = 2 (Å)	2.14–2.07	2.07–2.02
Wilson B factor	22.96	25
R _{merge}	0.125 (1.499)	0.112 (1.079)
R _{meas}	0.1469 (1.804)	0.1316 (1.342)
CC _{1/2}	0.995 (0.513)	0.995 (0.292)
CC*	0.999 (0.824)	0.999 (0.672)
Refinement		
Refinement resolution range	33.51–1.77 (1.833–1.77)	48.17–1.698 (1.759–1.698)
Reflections used in refinement	76,754 (4,773)	83,396 (3,379)
Reflections used for R _{free}	2,031 (130)	1,983 (79)
R _{work}	0.2274 (0.4453)	0.2043 (0.3800)
R _{free}	0.2649 (0.4659)	0.2394 (0.4085)
CC _{work}	0.964 (0.647)	0.961 (0.529)
CC _{free}	0.946 (0.550)	0.948 (0.640)
No. of non-hydrogen atoms		
Total	5,424	5,655
Macromolecules	4,935	5,001
Ligands	12	48
Waters	477	606
Protein residues	618	636
Rmsd		
Bonds (Å)	0.008	0.011
Angles (°)	1.10	1.23
Clashscore	2.03	2.4
B Factor		
Average	46.60	36.15
Macromolecules	46.78	35.03
Ligands	31.22	52.37
Solvent	45.16	44.12

Table 1. Continued

	Pcdh γ A1 EC1–3	Pcdh γ C3 EC1–3
Ramachandran Plot Regions		
Favored (%)	96	100
Allowed (%)	3.5	0.31
Outliers (%)	0	0
Rotamer outliers (%)	0	0.18 (H256)

homophilic interactions is distinct from those of classical cadherins, prompting investigation into the difference in homophilic interaction specificity between these two cadherin subfamilies.

In cell-aggregation assays, chimeric Pcdh γ isoforms were used to demonstrate that isoform concordance in both EC2 and EC3 is necessary for homophilic interactions (Schreiner and Weiner, 2010), implying the formation of parallel complexes, in which EC2 interacts with EC2 and EC3 with EC3. Concomitantly, immunoprecipitation experiments suggested that Pcdh γ isoforms form promiscuous *cis* complexes within a cell (Biswas et al., 2012; Han et al., 2010; Schalm et al., 2010; Schreiner and Weiner, 2010). These data led to a proposed model of a *cis* bundle promoting parallel homophilic *trans* interactions, i.e. across two cells. *Cis* interactions have a combinatorial behavior such that expression of combinations of isoforms leads to novel specificity (Thu et al., 2014), equivalent to an “AND” logic gate. This is strong evidence that combinatorial interactions provide sufficient diversification potential to support the clustered Pcdhs as cell-identity markers in neurons. Cell-aggregation assays cannot easily distinguish homophilic interactions that occur in *trans* from promiscuous *cis* interactions, especially if they are interdependent, indicating that we need other methods to determine the architecture of the specificity complex.

We know little about the structures of either the *cis* or *trans* interactions in clustered Pcdhs that lead to cellular self/non-self discrimination. We thus sought to identify possible interfaces present in these complexes as a way to explore the overall architecture. Using a combination of X-ray crystallography and bioinformatics, we identified and analyzed possible interfaces. Here, we describe a set of antiparallel interfaces that are similar to the *trans* Pcdh15-Cdh23 interactions (Sotomayor et al., 2012). These structures and analyses initiate a molecular understanding of the complicated interactions in the clustered Pcdhs.

RESULTS

Structures of Protocadherin γ A1 EC1–3 and γ C3 EC1–3

We obtained structures of mouse protocadherin γ A1 EC1–3 (A1) and γ C3 EC1–3 (C3) (Table 1). The A1 and C3 protomers form elongated structures containing three Greek-key β -sandwich motif EC domains (Figure 1A), consistent with other available cadherin structures and nuclear magnetic resonance structures of EC1 from several other clustered Pcdh isoforms (Figure S1) (Morishita et al., 2006). The three repeats are arranged in tandem with two inter-repeat linker regions each containing three Ca²⁺-binding sites comprising canonical Ca²⁺-binding residues (Figure 1B).

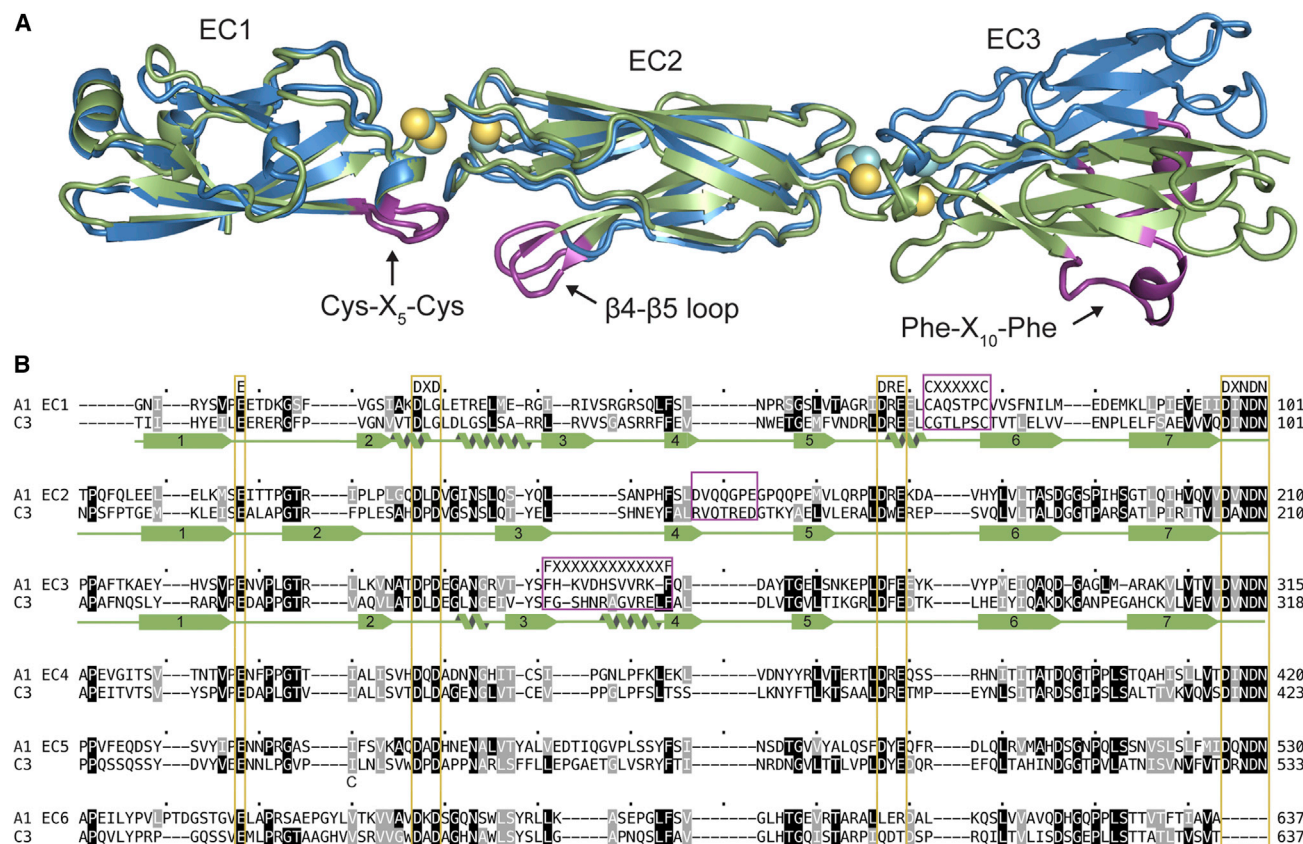


Figure 1. Sequence and Structural Properties of Clustered Pcdhs

(A) Superposition of A1 (blue with cyan Ca²⁺) and C3 (green with yellow Ca²⁺) structures showing the Cys-X₅-Cys, Phe-X₁₀-Phe and EC2 β4-β5 loops in purple. (B) Alignment of the six EC repeats of mouse PcdhγA1 and PcdhγC3. Ca²⁺-binding motifs are highlighted in yellow, the Cys-X₅-Cys, Phe-X₁₀-Phe, and EC2 β4-β5 loops in purple, and the A1 EC1-3 secondary structure is shown below the sequence.

See also Figure S1.

Comparing A1 and C3, each of the three EC repeat structures are very similar (root-mean-square deviation [rmsd] of 1.05, 0.824, and 0.593 Å for EC1, EC2, and EC3, respectively). However, superposition of A1 and C3 shows that A1 EC3 and C3 EC3 vary in tilt by 15° relative to EC2 (Figure 1A). There is some flexibility in fully Ca²⁺-bound linkers between cadherin repeats (Sotomayor and Schulten, 2008), which can explain the difference in relative EC3 orientation in A1 and C3. Since the overall A1 and C3 structures are similar, for simplicity figures refer to the A1 structure and sequence unless otherwise indicated.

The clustered Pcdh structures reveal distinctive features compared with classical cadherins (Figure 1). The most distinguishing feature is a conserved disulfide-clamped Cys-X₅-Cys loop in EC1. This loop, elongated by the X₅ sequence compared with classical cadherins, is required for cell-surface expression of Pcdh isoforms (Schreiner and Weiner, 2010), but its role in Pcdh interactions is unknown. Another feature of clustered Pcdhs is the Phe-X₁₀-Phe loop-helix motif in EC3. Lastly, the loop between β4 and β5 in EC2 is elongated when compared with classical cadherins. All three of these regions have low conservation between human paralogs (15%–25% average overall identity), suggesting that they may be sources of isoform diversity.

Crystallographic Interfaces Suggest Possible Biological Interfaces

The purified A1 construct behaved as a monomer (37.7 kDa) in multiangle laser light scattering (MALLS) measurements at concentrations of up to 137 μM (Figure S2). Furthermore, C3 and A1 behaved similarly in size-exclusion chromatography, indicating that C3 also behaves as a monomer in solution. This suggests that interactions between EC1-3 in solution are weak, precluding the use of solution-based experiments to investigate dimerization interfaces with these constructs. In the absence of in-solution interactions, we analyzed the physicochemical properties of the crystallographic interfaces to determine their biological relevance.

To select potential interfaces, we calculated the difference in accessible surface area (ΔASA) for all observed crystal packing interfaces. We further examined all interfaces of at least 600 Å², except one C3 interface with ΔASA = 650 Å² because it was highly solvated and the affinity tag contributed more than 80 Å². We thus selected four potential biological interfaces to evaluate (Figure 2). Interestingly, all four are antiparallel dimerization interfaces (EC1-3 directions in the protomers are antiparallel to each other) with two-fold rotational symmetry approximately perpendicular to the long axis of the protomers.

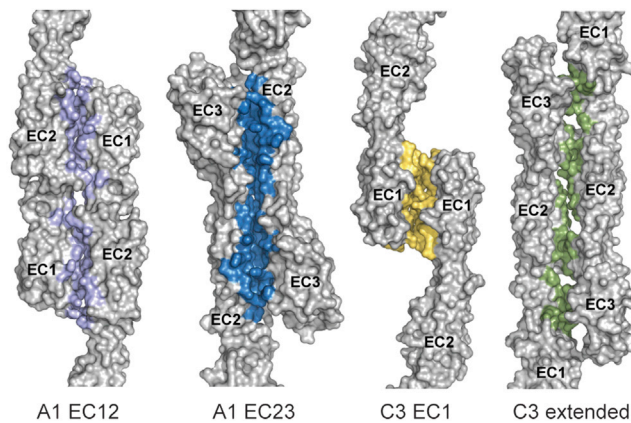


Figure 2. Observed Antiparallel Dimeric Interfaces in the A1 and C3 Crystal Lattices

Side view of the four interfaces with ΔASA greater than 600 \AA^2 : A1 EC12, A1 EC23, C3 EC1, and C3 extended. Residues with $\text{BSA} > 0 \text{ \AA}^2$ are colored. See also Figure S2.

We found two mutually compatible A1 interfaces: one where EC2 interacts with EC3 (hereafter called the A1 EC23 interface; $\Delta\text{ASA} = 1229 \text{ \AA}^2$) and the other where EC1 and EC2 interact (the A1 EC12 interface; $\Delta\text{ASA} = 1,381 \text{ \AA}^2$) (Figure 2). The combination of these two A1 interfaces yields crystal packing along one axis. We found two more potential interfaces in the C3 structure: an extended interface over the length of EC1–3 (C3 extended; $\Delta\text{ASA} = 1,156 \text{ \AA}^2$), and an EC1–EC1 interface (C3 EC1 interface; $\Delta\text{ASA} = 614 \text{ \AA}^2$) (Figure 2). These four interfaces may represent some of the interactions present in the *cis* or *trans* complexes of clustered Pcdhs.

The largest determinant, by far, of biological significance of a protein complex interface is the ΔASA (Janin and Rodier, 1995). In a survey of PDB structures, an 856-\AA^2 cut-off distinguished a biological interface from a crystal contact with 85% accuracy (Ponstingl et al., 2000). All but the C3 EC1 interface are significantly above the 856-\AA^2 threshold (Table 2), indicating that they are likely biological in nature.

The number of hydrogen bonds is indicative of specific protein interactions versus crystallization artifacts. Crystal contacts tend to have few hydrogen bonds (~ 5) while homodimers and protein complexes have $\sim 10\text{--}20$ (Bahadur et al., 2004). The interfaces have a relatively large number of hydrogen bonds (Table 2), sug-

gesting they are not crystallization artifacts, and that they are, if valid, non-obligate in nature, as obligate interactions rely more heavily on non-polar interactions (Jones and Thornton, 1996).

Strong and specific interactions typically have highly complementary surfaces. In our interfaces, the A1 EC12 interface has the highest shape correlation ($S_c = 0.70$), while the C3 extended interface has the lowest ($S_c = 0.65$) (Table 2). These are all near the range reported for antibody/antigen complexes ($S_c = 0.64\text{--}0.68$), suggesting that the interfaces are specific, yet non-obligate (Lawrence and Colman, 1993), which would agree with the expected properties of the clustered Pcdh interactions.

NOXclass is a protein-protein interaction classifier that distinguishes crystal packing from biological interactions and predicts whether an interface is likely obligate or non-obligate (Zhu et al., 2006). This prediction is useful for the non-obligate clustered Pcdhs interactions. According to NOXclass (Table 2), the C3 EC1 interface has a zero probability of being biological. A1 EC12 and A1 EC23 are both highly likely biological with 98.7% and 90% probability, respectively, while the C3 extended interface is 68% probable. All three are predicted to be obligate, but the A1 EC12 interface the least so with a 67% probability. This provides further support for the biological significance of all but the C3 EC1 interface.

The A1 EC23 and C3 extended interfaces include some of the structural features particular to the clustered Pcdhs described above (Figures 1 and 3). Polar interactions distribute toward the edge of the interface, while hydrophobic interactions are found toward the center of the interface, a property of biological interfaces (Chakrabarti and Janin, 2002). In A1 EC23 (Figure 3A), the Phe- X_{10} -Phe loop of EC3 interacts with EC2 (Figure 3B). In the C3 extended interface (Figure 3C), the elongated $\beta 4$ - $\beta 5$ loop in EC2 interacts with EC3 (Figure 3D). There is also extensive hydrogen bonding between the Cys- X_5 -Cys loop of EC1 and the Phe- X_{10} -Phe loop of EC3 (Figure 3E). In this network T70 forms hydrogen bonds with S255 and H256 of the other protomer, which holds H256 as a rotamer outlier in both protomers (Figure S3), suggesting a strong and specific interaction.

The A1 EC23 and C3 extended interfaces overlap, as they both include the $\beta 1$ - $\beta 2$ loop and $\beta 2$ of EC2, and the $\beta 3$ - $\beta 4$ and $\beta 6$ - $\beta 7$ loops of EC3. The two interfaces are related by a slide of the proteins by 22 \AA and a slight rotation about their long axis (Figure 3F). The highly dissimilar sequences in this region may explain the difference in geometry between the A1 EC23 and C3 extended interfaces.

Table 2. Physicochemical Properties of Clustered Pcdh Interactions

Interface	$\Delta\text{ASA} (\text{\AA}^2)$	#HB ^a	#SB ^b	S_c ^c	NOXclass				
					Interface Area Ratio	Interface/Surface Correlation	GAP ^d	Biological (%)	Obligate (%)
A1 EC12	1,381	26	6	0.701	0.0845	0.5773	8.9	98.73	66.89
A1 EC23	1,229	11	4	0.662	0.0718	0.6007	12.2	90.03	85.64
C3 extended	1,156	17	2	0.648	0.0672	0.6941	15.1	68.14	91.73
C3 EC1	614	16	13	0.666	0.0356	0.7195	21.7	0	91.81

^aNumber of hydrogen bonds at interface.

^bNumber of salt bridges at interfaces.

^cShape correlation statistic.

^dGap volume index.

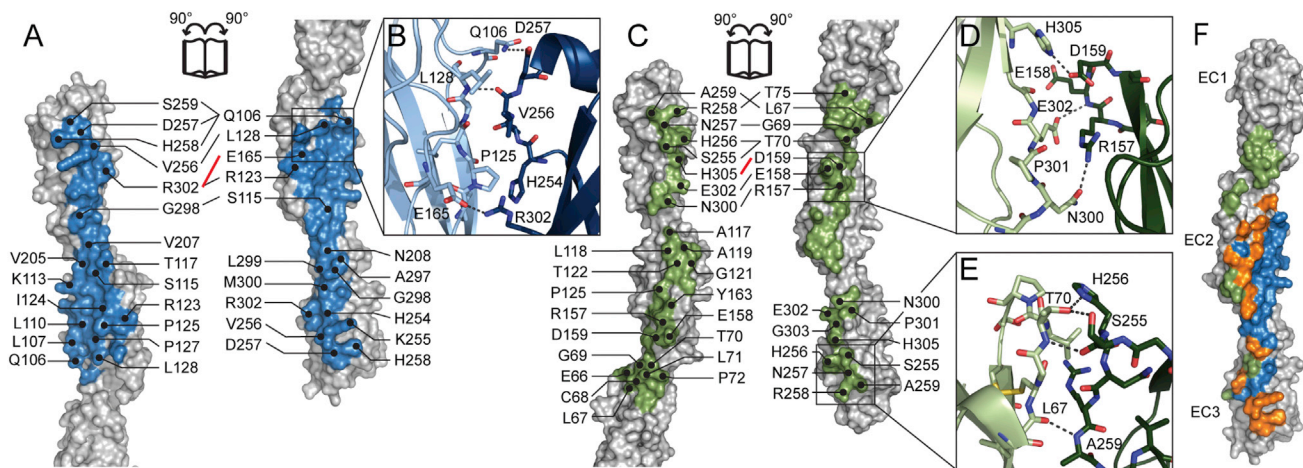


Figure 3. Structural Details of the A1 EC23 and C3 Extended Interfaces

(A–E) A1 EC23 (A) and C3 extended (C) interfaces are shown in an “open-book” orientation. Hydrogen-bonding (black) and salt-bridge (red) interactions are labeled on the top half of the interfaces, and residues with BSA >20 Å² on the bottom half. (B) In A1 EC23, the Phe-X₁₀-Phe loop of EC3 (dark blue) makes specific interactions with β1 and β2 of EC2 (light blue). In the C3 extended interface, (D) the β4–β5 loop in EC2 of one protomer (dark green) interacts with β7 of EC3 in the other protomer (light green), and (E) the Cys-X₅-Cys loop of EC1 (light green) forms a specific hydrogen-bonding network with Phe-X₁₀-Phe loop of EC3 (dark green).

(F) The A1 EC23 (blue) and C3 extended (green) interfaces overlap extensively (orange).

See also Figure S3.

Homology Modeling of EC1–6 Clustered Protocadherin Predicts a Linear Structure

To gain further insights into the complete structure of a clustered protocadherin extracellular domain, we used the A1 and C3 EC1–3 structures along with sequence and structural analysis of other cadherin structures to construct a homology model of EC1–6.

The Ca²⁺-binding sites rigidify the linkers between EC repeats (Cailliez and Lavery, 2005; Sotomayor and Schulten, 2008). We predict that all clustered Pcdh EC linkers are fully occupied with three Ca²⁺ ions because all Ca²⁺-binding motifs are strictly conserved in each of the five EC linker regions (96% conservation of canonical Ca²⁺-binding residues in Figure 1B). To determine how to orient EC4–6 relative to each other and EC3, we surveyed cadherin structures with at least two ECs and canonical three-Ca²⁺ linker regions (Table S1). We defined the tilt angle as the angle between the long axes of two adjacent ovoid EC repeats (Figure 4A). Rotation of each EC around this primary axis determines the azimuthal rotation of adjacent ECs, which reflects the orientation of the cadherin fold for consecutive repeats. The average tilt and azimuthal angles for 17 canonical classical cadherin EC repeat pairs are 153° ± 6° and 239° ± 9°, respectively, versus 165° ± 6° and 217° ± 7° for the 13 non-classical cadherins and protocadherins (protocadherin-15, cadherin-23, A1 and C3). Thus adjacent EC pairs of non-classical and protocadherins are more linear than classical cadherins. The origin of this difference is unknown, but could denote an important structural difference between cadherin subfamilies.

We used two EC pairs as a starting model to position clustered Pcdh EC4–6: one model using the C3 EC2–3 pair, closest to the non-classical/protocadherin average (tilt = 164°, azimuthal = 212°), and one using mouse cadherin-6 EC1–2 pair (3Ind_a), closest to the classical cadherin average (tilt = 154°, azimuthal =

239°). We then used these models to construct homology models of the full mouse PcdhA1 extracellular domain (Figure 4C). The two overall linear structures vary slightly in their overall curvature, but neither is incompatible with any of the potential interfaces in the A1 or C3 structures (Figure 4D).

The resulting clustered Pcdh model differs significantly from the five EC classical cadherin structures, due to the relative orientations of the repeats (Figure 4C). Non-canonical Ca²⁺-binding sites in classical cadherins cause a significantly smaller EC3–EC4 tilt angle (~135°), yielding a nearly perpendicular orientation of EC1 and EC5. The clustered Pcdh models are more linear, reflecting the fact that all canonical Ca²⁺-binding motifs are completely conserved across all linker regions in clustered Pcdhs.

The A1 EC23 and C3 Extended Interfaces Lack Potential Glycosylation Sites

Post-translationally-modified N- and O-glycosylation sites in classical cadherins expressed in *Xenopus laevis* or in HEK293 cells are distributed away from the homophilic binding interface (Boggon et al., 2002; Harrison et al., 2011), indicating that glycosylation is disfavored at cadherin interfaces. Since our recombinantly expressed Pcdh constructs are not glycosylated, we mapped predicted glycosylation sites on the Pcdh structures. We used the NetNGlyc v1.0 and NetOGlyc v4.0 servers to predict N- and O-glycosylation sites, respectively, in all human clustered Pcdhs (Blom et al., 2004; Steentoft et al., 2013). Fourteen predicted N-glycosylation sites distributed across EC2–6 are well conserved across similar isoforms, and none overlap with the interfaces (Figure 5). Surprisingly, only two O-glycosylation sites were predicted: T325 in PcdhA1 and position 357 (mouse PcdhA1 numbering) in Pcdhα2, Pcdhα9, and Pcdhα10 (Figure 5). This contrasts with the high

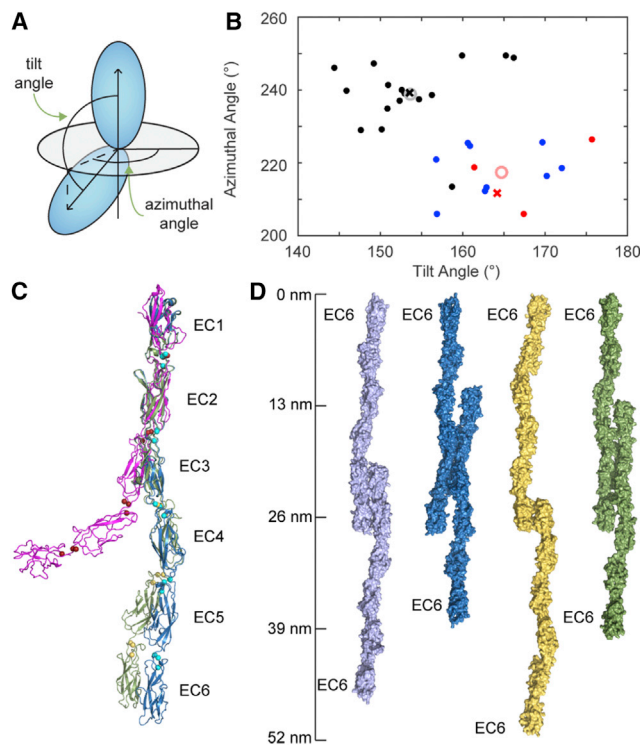


Figure 4. Clustered Pcdh EC1-6 Homology Models as Elongated Molecules

(A) The orientation of adjacent EC repeats can be defined by the tilt and azimuthal rotation of the EC domain principal axes.

(B) Distribution of tilt and azimuthal angles from adjacent EC repeat pairs of classical (black), Pcdh15 and Cdh23 (blue), and clustered Pcdh (red). Gray and pink open circles represent averages of classical and non-classical/clustered Pcdh tilt and azimuthal angles; these pairs were used to generate the homology models. Crosses show the EC repeat pairs closest to the average values; these pairs were used to generate the homology models.

(C) Homology models of A1 using the C3 EC2-3 (blue) and cadherin-6 EC1-2 (PDB: 3lnd_a; green) model pairs. Classical EP-cadherin (PDB: 1l3w; magenta; five EC repeats) is shown for comparison.

(D) Models of *trans*-interacting antiparallel full-length dimers corresponding to the A1 EC12 (lilac), A1 EC23 (blue), C3 EC1 (yellow), and C3 extended (green) interfaces based on the full-length A1 homology model. See also Table S1.

O-glycosylation seen in C-, N-, and E-cadherin structures (Boggon et al., 2002; Harrison et al., 2011). The NetOGlyc server predicts no O-glycosylation sites for mouse C-, N-, or E-cadherin, indicating that it is not well trained for predicting O-glycosylation sites in cadherins.

O-Mannosylation sites in clustered Pcdhs were identified by the SimpleCell strategy (Vester-Christensen et al., 2013). Many O-mannosylation sites were found in EC2, EC3, EC5, and EC6 of various isoforms. Several O-mannosylated residues overlap the A1 EC12 interface (T185, S195, and T197) (Figure 5). A site at position 300, found in Pcdh γ B1, overlaps the A1 EC23 and C3 extended interfaces, but a methionine in γ A1 (M300) or arginine in γ C3 indicates this glycosylation is absent in these isoforms. While glycosylation site prediction could not provide striking support for a particular interface, it does suggest that glycosylation may prevent the A1 EC12 interaction.

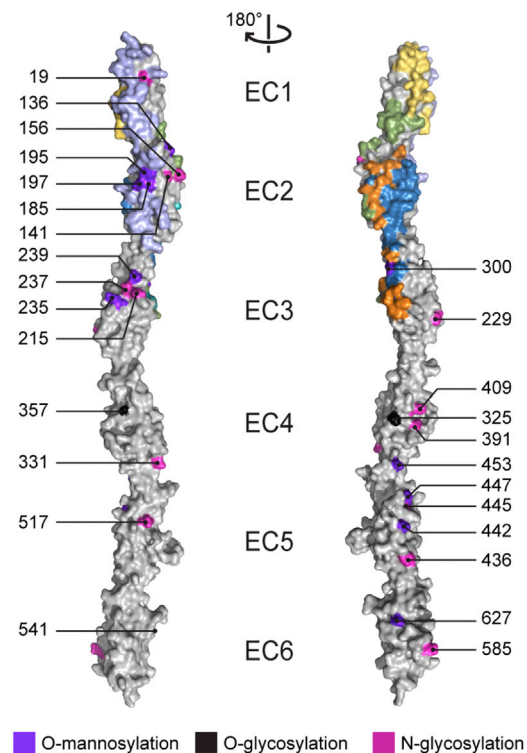


Figure 5. Predicted Clustered Pcdh Glycosylation Sites Largely Do Not Overlap with the Identified Antiparallel Interfaces

N-Glycosylation (magenta) or O-glycosylation (black) sites predicted in at least five human clustered Pcdh isoforms or a single C-type isoform, and O-mannosylation sites (purple) from the SimpleCell strategy (Vester-Christensen et al., 2013) in clustered Pcdh EC repeats are overlaid with the observed interfaces (A1 EC12, lilac; A1 EC23, blue; C3 EC1, yellow; C3 extended, green; and A1 EC23 and C3 extended overlapping, orange). The left and right panels are related by a 180° rotation; numbering corresponds to mouse Pcdh γ A1.

A1 EC23 and C3 Extended Interfaces Correlate with Isoform-Specific Conservation

Sequence conservation analysis confirmed the homophilic interaction epitopes in the *Drosophila Dscam1* variable exons (Meijers et al., 2007). We searched for regions of high isoform-specific conservation, which may signify regions that impart homophilic specificity, by comparing the conservation of Pcdh paralogs in humans with orthologs of each isoform in mammalian species. Conservation in mammalian orthologs is higher (average overall identity for all orthologs is 78% \pm 6%) than in human paralogs (50% overall identity). Thus, we defined an isoform conservation ratio (ICR), the ratio of ortholog conservation of an isoform to paralog conservation of all isoforms in a species (in our case, humans), as an index for residue-by-residue sequence diversification. We then calculated residue-by-residue average ICR for all 60 mammalian isoforms (Figure 6A).

Average ICR values for EC1-3 are significantly higher than in EC4-6, and highest for EC2 and EC3 (Figure 6A). This preponderance of isoform-specific positions corroborates clustered Pcdh-driven cell-aggregation assays that demonstrated the importance of EC2-3 in homophilic specificity (Schreiner and Weiner, 2010). These data also correlate with the presence of positively selected sites in EC2 and EC3, where the rate of

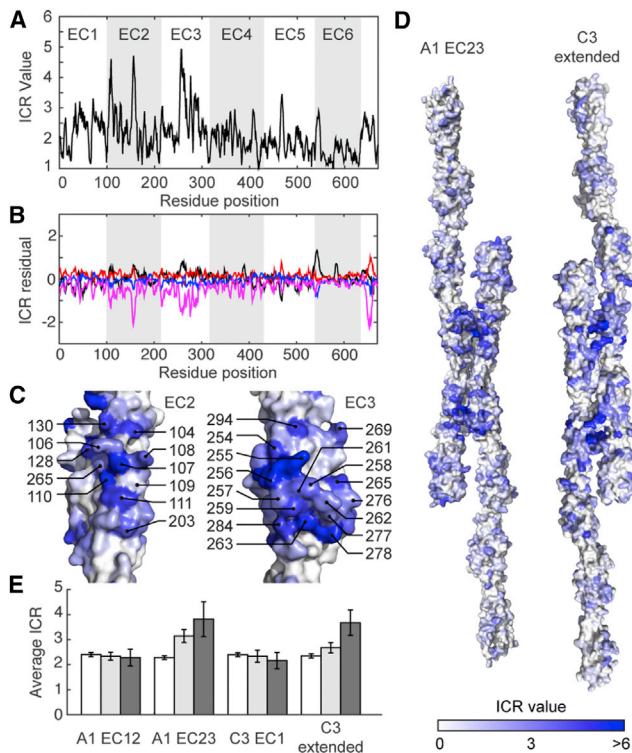


Figure 6. Isoform-Specific Conservation in Clustered Pcdhs Localizes to EC2 and EC3 Surfaces

(A) ICR average of all clustered Pcdh isoforms compared with human clustered Pcdhs, as a sliding average with a five-residue window size. (B) Residual of the ICR value between the overall and subfamily averages (α = black, β = red, γ = blue, and C-type = magenta). (C) ICR values mapped on the EC2 and EC3 surfaces with a high density of high ICR values, numbered according to mouse Pcdh γ A1. (D) ICR values mapped on full-length homology models of the A1 EC23 and C3 extended interfaces. (E) Comparison of the average ICR values (\pm SEM) of non-interface (white) and interface (gray) residues. Interface residues are defined as those having a BSA of at least 0 \AA^2 (light gray) or 55 \AA^2 (dark gray). See also [Files S3](#) and [S4](#).

non-synonymous substitution at some positions is higher than the rate of synonymous mutations, implying that isoforms are under selective diversification (Wu, 2005). The average ICR of these previously identified positively selected sites (4.5 ± 0.5) is significantly higher than the overall average ICR (2.09 ± 0.05), supporting the ICR as a metric for selective diversification. We also compared the ICR values of clustered Pcdh subfamilies (α , β , γ , C-type) with that of entire cluster by calculating the residual of the subfamily ICR from the entire cluster ICR (Figure 6B). No region has significantly different ICR values in the α , β , and γ subfamilies, supporting the notion that the homophilic specificity regions are common to all isoforms. In contrast, the C-type isoforms contain some regions with significantly decreased ICR values, specifically in EC2 and EC3. These regions could suggest a difference in C-type subfamily homophilic interactions and/or could reflect the fact that C types may be under stronger conservation selection due to their essentiality in neuronal development (Chen et al., 2012).

The highest concentration of large ICRs is on a small portion of EC2 and a face on EC3 (Figure 6C), two regions proximal to both the A1 EC23 and C3 extended interfaces (Figure 6D). For example, the previously noted hydrogen-bond network in the C3 extended interface (Figure 3D) includes residues with high ICR values (T70 = 5.48, S255 = 6.18, and H256 = 5.49), indicating that this interaction is highly specific to the γ C3 isoform.

To quantify the correlation between the observed interfaces and the ICR values, we calculated the average ICR value for non-interface and interface residues (judged as residues with a buried surface area [BSA] of more than 0 \AA^2). The average ICR of non-interface residues is 2.36 ± 0.08 , while the average ICR of interface residues of the A1 EC23 and C3 extended complexes are 3.1 ± 0.3 and 2.7 ± 0.2 , respectively (Figure 6E). In contrast, the average interfaces ICR for the A1 EC12 and C3 EC1 complexes are both 2.3 ± 0.2 . To determine whether the correlation between ICR and our interfaces was amplified in residues that contribute most significantly to the interface, we reduced our selection of interface residues to those with a BSA of at least 55 \AA^2 per residue (Figure 6E). At this BSA cut-off, the average ICR of interface residues increases to 3.8 ± 0.7 for A1 EC23 and 3.7 ± 0.5 for C3 extended, whereas the A1 EC12 and C3 EC1 interface ICR values at this BSA cut-off are indistinguishable from non-interface ICR values. This result indicates that residues in the A1 EC23 and C3 extended complexes contributing appreciably to the interface surface area are more likely to have a high ICR value, which supports the functional significance of these interfaces.

Evolutionary Correlations Corroborate Antiparallel Complex Architecture

Pairs of residues near each other in a structure can be identified via evolutionary covariance caused by compensatory mutations. These evolutionary correlations can indicate structural contacts within a given protein (Aurell and Ekeberg, 2012; Hopf et al., 2012; Kamisetty et al., 2013; Marks et al., 2011) or between proteins that interact (Hopf et al., 2014; Ovchinnikov et al., 2014). We used EVcouplings (adapted from (Hopf et al., 2014; Marks et al., 2011)) to search for evolutionary correlations arising from homotypic interactions from an alignment generated with mouse Pcdh γ C3 as a query. The top 320 (~alignment length/2) evolutionary correlations (Figure S4A) predominantly reflect structural features of the cadherin fold (Figure 7A). Eleven pairs in EC1–4 are not close to each other in the monomer, but nevertheless score higher than many true positives, suggesting they represent oligomeric contacts (Figure 7A). These pairs—between EC2 and EC3, or EC1 and EC4—support an antiparallel complex. Our structures corroborate the EC2–EC3 pairs, and although we did not crystallize EC4 our full-length homology model suggests that EC1 and EC4 interact (Figure 4D).

The 11 non-structural correlated pairs in EC1–4 come in close proximity in the A1 EC23 and C3 extended interfaces (Figures 7B and 7C), corroborating our posited interfaces. A plot of coupling strength versus pair distance reveals that these pairs are outliers compared with the rest of the observed monomeric pairs (Figure S4B). Using the A1 EC23 and C3 extended full-length complex models to define the pair distance of these 11 correlated pairs as intermolecular contacts, we find that the pair distance-coupling strength distribution is better preserved, supporting

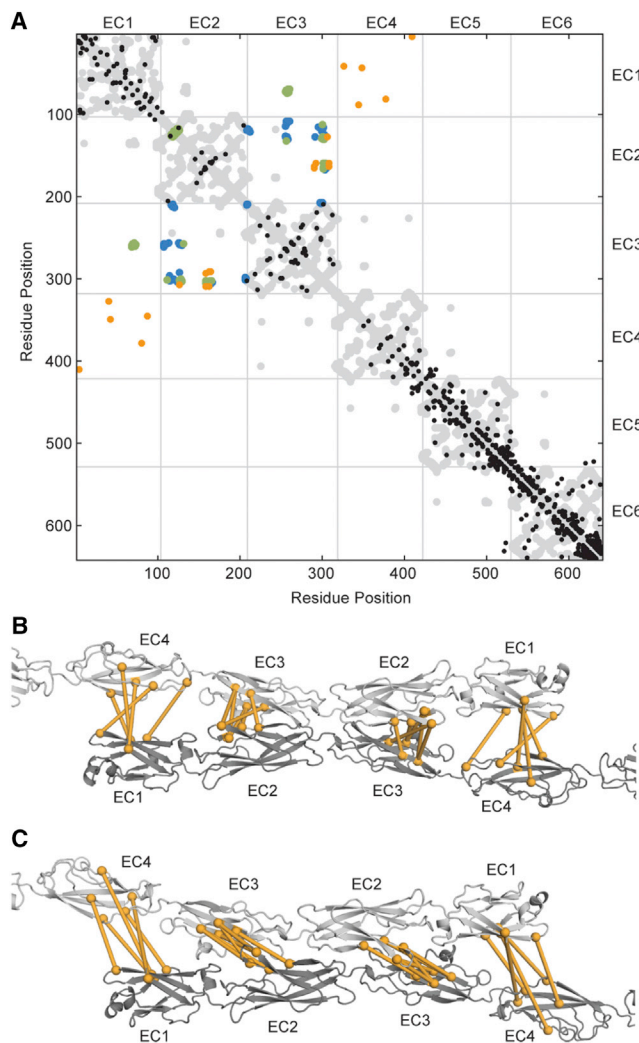


Figure 7. Evolutionary Covariance Supports Antiparallel Oligomeric Contacts

(A) The top 320 correlated pairs (sequence length/2) are shown with the contact map of the EC1–6 clustered Pcdh homology model (gray) for reference. Correlated pairs matching structural constraints are shown in black while predicted oligomeric contact pairs are in orange. The observed interface contact residues are also mapped (A1 EC23, blue; C3 extended, green). (B and C) The predicted oligomeric contact correlated pairs are mapped (orange spheres) on the A1 EC23 (B) and C3 extended (C) complex models with a line between coupled residues. See also [Figure S4](#); [Files S5](#) and [S6](#).

their significance as oligomeric contacts instead of intramolecular structural ones. Interestingly, the A1 EC23 model yields shorter distances overall for the 11 correlated pairs, which is evident from the sliding of A1 EC23 and C3 extended interfaces relative to each other along the long cadherin axis ([Figures 7B and 7C](#)). This could indicate that A1 EC23 better represents homodimeric clustered Pcdh interfaces. However, because the clustered Pcdh locus has relatively few C-type isoforms, our sequence alignment contains many more non-C-type isoform sequences. This may manifest as a bias of observed correlated pairs toward non-C-type versus C-type interfaces if, as biolog-

ical data suggest ([Chen et al., 2012](#)), C-type and non-C-type have different modes of interaction.

DISCUSSION

There is strong evidence that EC1–3 mediate clustered Pcdh specificity of cell-surface signaling from cell-aggregation assays and sequence analysis ([Schreiner and Weiner, 2010](#); [Wu, 2005](#)). Our structures of mouse Pcdh γ A1 and Pcdh γ C3 EC1–3 and bioinformatics analyses provide further evidence for the importance of EC2 and EC3 and offer the first hypotheses for these interactions. We propose two biologically relevant interactions: A1 EC23 and C3 extended. The physicochemical properties of these interfaces are consistent with canonical biological interactions, and interface residues are not predicted glycosylation sites and are conserved in isoform orthologs. The antiparallel arrangement is consistent with evolutionary correlations observed between EC2 and EC3, and EC1 and EC4. Lastly, isoform-specific conservation suggests that all subfamilies have similar modes of homophilic recognition, with the possible exception of the C-type isoforms.

The combination of an antiparallel arrangement of the specificity-determining homodimer and the linearity of our EC1–6 homology model limits the geometry of a *trans* complex. To satisfy both constraints, the overall length of A1 EC23 or C3 extended complexes is approximately 40 nm ([Figure 4D](#)). This distance is consistent with intercellular distances of chemical synapses (20–40 nm), suggesting that this *trans* complex architecture would be effective in determining synaptic specificity at nascent synapses ([Phillips et al., 2003](#)).

The two interfaces we identified comprise similar sets of residues with high ICR values, but only partially overlap ([Figures 3F and 5](#)). At present, we cannot offer any explanation for these similar yet distinct interfaces. However, the biological behaviors of the two isoforms we crystallized are different: C-type Pcdh isoforms are necessary for neonatal neuronal survival in mice, while A-type Pcdh isoforms show redundancy in behavior ([Chen et al., 2012](#)). Thus, A1 EC23 and C3 extended interfaces may be representative of γ -type and C-type Pcdh isoforms, respectively, and reflect structural adaptations to use overlapping surfaces to encode a large diversity of preferentially homotypic dimerization interfaces.

In our data we find some indirect support for two potential *cis* architectures in relation to the identified antiparallel *trans* interactions ([Figure 8](#)). The lack of predicted glycosylation sites on a face of EC4–6 opposite the homophilic EC2–EC3 interface ([Figure 5](#)) and evolutionary correlations in EC5 and EC6 that are not part of intramolecular contacts ([Figure 7A](#)) suggest that EC5–6 could be part of promiscuous *cis* interactions ([Figure 8A](#)). Supporting this model is the finding that α isoforms require EC6 of a carrier isoform (β or γ) to be surface expressed ([Thu et al., 2014](#)), suggesting *cis* EC6 interactions across paralogs. Furthermore, deletion of EC1–3 does not impair *cis* multimers ([Schreiner and Weiner, 2010](#)), consistent with a model in which the C-terminal EC domains provide further multimerization interfaces. An alternative model derives from the A1 EC12 interface, which is statistically assessed as potentially biological ([Table 2](#)). The A1 EC12 interface is on the opposite side of the A1 EC23 interface, allowing a zipper of Pcdhs at an intercellular junction to form

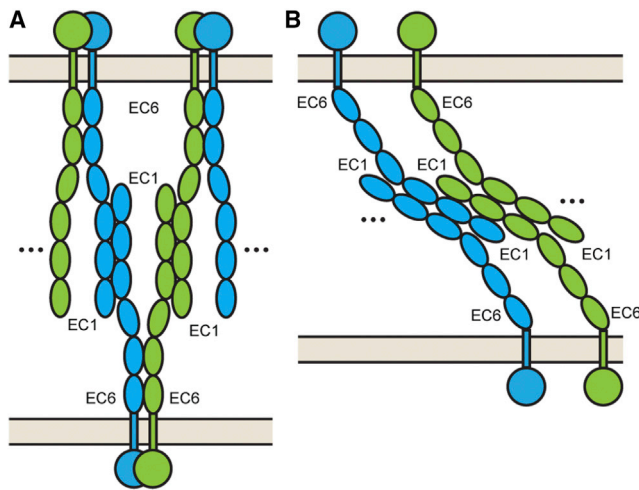


Figure 8. Model of Clustered Protocadherin Antiparallel *trans* and Possible *cis* Interactions

(A) Lack of predicted glycosylation sites on one face of EC4-6 allows a model where EC4-6 engage in promiscuous *cis* interactions to promote oligomerization.

(B) The A1 EC12 interaction, which does not show preponderance for high ICR value residues, could support a model for *trans*-mediated *cis* oligomerization through EC1-2. Isoform types are shown as either blue or green to indicate homophilic (same color) or promiscuous (different color) interactions.

an oligomer (Figure 8B), implying the presence of *trans*-mediated promiscuous *cis* interactions. This model is supported by a relatively conserved interface across isoforms, but is challenged by predicted glycosylation sites on the A1 EC12 interface. While neither model fully clarifies the mechanism used to enforce combinatorial homophilic interactions or specifies the possible multimeric state of these complexes, they can serve as testable hypotheses. Future experiments will require the development of biochemical and biophysical methods for investigating weak adhesion at a molecular level.

The interactions we identified are novel for the cadherin superfamily. Unlike classical cadherins that interact through EC1 in an X-dimer or strand-swapping mechanism (Brasch et al., 2012; Sotomayor et al., 2014), these interactions are mediated by EC2 and EC3. The *cis* interactions of classical cadherins are also distinct, as they occur through EC1 and EC2 and are parallel in nature (Boggon et al., 2002; Nagar et al., 1996). The most similar interface described thus far is the antiparallel “handshake” Pcdh15-Cdh23 interaction in the tip link of hair cells (Sotomayor et al., 2012). The elongated antiparallel interaction spanning multiple EC repeats of the Pcdh15-Cdh23 complex and in the clustered Pcdh interfaces described here may be a general adhesion strategy for other cadherin superfamily interactions.

We have described homophilic interactions of two clustered Pcdh isoforms and have presented physicochemical and bioinformatics evidence for their biological significance. Overall, we showed that the A1 EC23 and C3 extended interfaces are the most likely to be relevant *in vivo*. It is clear from these interfaces and the presence of both *cis* and *trans* interactions that the architecture of the entire clustered Pcdh signaling complex is multifaceted. In addition, clustered Pcdhs play many roles in neuronal

development, suggesting that different complex architectures and interactions with other proteins provide further layers of complexity, which we need to better define before we reach a full structural understanding of these proteins. While cell-based aggregation assays have been indispensable for studying clustered Pcdh interactions, these assays have not yet been used to distinguish between *cis* and *trans* interactions (Schreiner and Weiner, 2010; Thu et al., 2014). Thus, better biochemical methods are needed to verify these and other interactions. The structures, homology models, and interactions we describe provide a glimpse of the details of this complex system and constraints to better study clustered Pcdh signaling assemblies.

EXPERIMENTAL PROCEDURES

Protein Expression and Purification

Mouse Pcdh γ A1 EC1-3 (residues 1-311 referred to as A1) and Pcdh γ C3 EC1-3 (residues 1-314, referred to as C3) with C- and N-terminal hexahistidine tags, respectively, were expressed in BL21 Gold (DE3) *Escherichia coli* cells in Terrific Broth, induced at OD₆₀₀ = 0.8 with 0.5 mM isopropyl β -D-1-thiogalactopyranoside at 37°C for 4 hr. Cells were lysed by sonication in 8 M guanidine hydrochloride (GuHCl), 20 mM imidazole, 50 mM HEPES (pH 7.5), 2 mM CaCl₂, 1 mM benzamide, and 1 mM phenylmethanesulfonyl fluoride. Cleared lysates were diluted to 5 M GuHCl, loaded onto Ni-Sepharose, washed with 5 M GuHCl, 250 mM NaCl, 25 mM imidazole (pH 8), and 50 mM Tris (pH 8.5), and eluted with 200 mM imidazole. A1 was refolded overnight at room temperature by drop dilution from 2 mg/ml protein in elution buffer to a final concentration of 0.2 M GuHCl with refolding buffer (100 mM Tris [pH 8.5], 10 mM CaCl₂, 1 mM EDTA, 3 mM reduced glutathione, 0.3 mM oxidized glutathione). C3 was refolded at 1 mg/ml by 12-hr dialysis steps into 3, 1.5, and 0 M GuHCl in refolding buffer with 0.2 M sucrose and 0.5 M L-arginine monohydrochloride. Concentrated refolded protein was purified by size-exclusion chromatography on a Superdex 200 column (GE Healthcare) in 20 mM Tris (pH 8.5), 200 mM NaCl, and 2 mM CaCl₂. C3 was further purified on a Resource Q column (GE Healthcare) with a gradient of 50–500 mM NaCl in 20 mM Tris (pH 8.5) and 2 mM CaCl₂. The A1 protein molecular mass was determined in runs at 5 mg/ml (137 μ M) over a Superdex S200 10/300 column with an in-line MALLS instrument (Wyatt) in 20 mM Tris (pH 8.5), 200 mM NaCl, and 2 mM CaCl₂.

Crystallization, Data Collection, and Structure Determination

A1 crystals were grown by vapor diffusion at room temperature in 100 mM Tris (pH 8.5), 7.5% polyethylene glycol 8000, and 200 mM NDSB-221 (Hampton Research). Crystals were cryoprotected in reservoir plus 20% glycerol. C3 crystals were grown by vapor diffusion at 4°C. Initial crystals, obtained with a drop ratio of 0.5 μ l 4.3 mg/ml C3 to 0.3- μ l reservoir (100 mM HEPES [pH 7], 60% 2-methyl-2,4-pentanediol [MPD], 100 mM CaCl₂), were used as seeds to grow larger crystals in 100 mM HEPES (pH 7), 20% MPD with a 0.5 μ l protein/0.2 μ l reservoir/0.1 μ l seeds ratio. Crystals were cryoprotected in reservoir plus 15% MPD. Crystals were cryocooled in liquid N₂. X-Ray data were collected according to Table 1 and processed in an HKL2000 (Otwinowski and Minor, 1997). The A1 structure was determined by molecular replacement in PHENIX (Adams et al., 2010), using cadherin-23 EC1-2 as a starting model (Sotomayor et al., 2010). The C3 structure was solved by molecular replacement using a C3 homology model based on A1. Model building was done in Coot (Emsley and Cowtan, 2004) and refinement in PHENIX (Adams et al., 2010). Both structures contain two protomers per asymmetric unit. In addition to canonical inter-EC linker Ca²⁺ ions, the C3 structure has three Ca²⁺ ions bridging an EC3-EC3, tail-end-to-tail-end, crystal packing interface between the two chains, chelated by EC3-EC4 linker residues in a non-physiological way. Validation reports for the A1 (File S1) and C3 (File S2) are provided in the Supplemental Information.

Structural Analysis of Interface Significance

The PISA webserver was used to analyze the observed Pcdh interfaces (Krisinel and Henrick, 2007). Δ ASA is (ASA_A + ASA_B – ASA_{AB})/2, where AB is a

complex of A and B. BSA is the solvent-accessible surface area buried upon complex formation for a given residue. The SC program in CCP4 was used to calculate surface complementarity according to the following parameters (Lawrence and Colman, 1993): dot density = 15 \AA^{-2} , interface separation = 8 \AA , trim width = 1.5 \AA , probe radius = 1.7 \AA , weight factor 0.5 \AA^{-2} . NOXclass was used to evaluate the observed interfaces using interface area, interface area ratio, correlation between interface and protein surface, and gap volume index parameters (Zhu et al., 2006).

Homology Model Construction

Structures of cadherin proteins containing at least two EC repeats were collected from the PDB (Table S1). EC N was aligned to EC N+1 for each pair, and singular value decomposition was used to calculate the principal components of the EC N repeat and the original and aligned positions. The angles between the respective components were calculated to determine the tilt (between the first components) and azimuthal (average of the second and third components) angles. Models were constructed in PyMOL (Schrodinger), after which Modeller (Sali and Blundell, 1993) was used to generate the EC1-6 homology model.

Sequence Analyses

For ICR analysis, Pcdh isoform sequences were collected from the UCSC Genome Browser (Kent et al., 2002) to create a multiple sequence alignment. Overall % identity of the most common residue at each position was used to calculate ICR values. The alignment (File S3) and ICR data (File S4) are provided in the Supplemental Information.

Evolutionary couplings were predicted using EVcouplings (Aurell and Ekeberg, 2012; Marks et al., 2011; Morcos et al., 2011), applying a pseudolikelihood maximization approximation (Balakrishnan et al., 2011; Ekeberg et al., 2013; Kamisetty et al., 2013). The multiple sequence alignment used was generated starting with EC1-6 of mouse Pcdh γ C3 (UniProt: Q91XX1_MOUSE, residues 30–670) as a query sequence for Jackhmmer 3.1 (Johnson et al., 2010). The alignment was created in five iterations against the UniProt Reference Cluster UniRef100 (release February 2015; Suzek et al., 2007) and then filtered to exclude sequences that are not part of the clustered protocadherin family. To build the global maximum entropy model, we clustered and weighted sequences with at least 90% identity according to their cluster size (theta 0.1). Alignment columns that contained gaps in 50% or more of the members were excluded from the model and later predictions. The effective number of sequences in the final alignment was 3,525.48 (sequences/alignment length = 5.56). Correlated pairs were normalized by the magnitude of the minimum coupling score to suppress background noise (Hopf et al., 2014). Pairs with normalized scores above 1.6 (\sim alignment length/2 = 320 couplings) are considered statistically significant ($>80\%$ predictive of contact; true positive rate [$<8 \text{ \AA}$ distance] = 66% in EC1-3, 84% in EC2-3). The alignment (File S5) and raw coupling scores (File S6) for all pairs are presented in the Supplemental Information.

ACCESSION NUMBERS

The accession numbers for the Pcdh γ A1 EC1-3 and Pcdh γ C3 EC1-3 structures reported in this paper are PDB: 4ZI9 and 4ZI8, respectively.

The unprocessed diffraction images used to determine the Pcdh γ A1 EC1-3 and Pcdh γ C3 EC1-3 structures were deposited in the SBGrid Data Bank (<https://data.sbgrid.org/>) with accession numbers SBGrid Data Bank: 173 and 174, respectively.

SUPPLEMENTAL INFORMATION

Supplemental Information includes four figures, one table, and six supplemental data files and can be found with this article online at <http://dx.doi.org/10.1016/j.str.2015.09.005>.

AUTHOR CONTRIBUTIONS

W.A.W. and R.G. initiated the project; J.M.N., W.A.W., and S.L. screened constructs for purification and crystallization; J.M.N. and W.A.W. crystallized and solved structures; S.L. performed MALLS measurements; J.M.N. constructed

homology model and completed ICR analysis; J.M.N., C.P.I.S., D.S.M., and R.G. completed evolutionary correlation analysis; R.G. directed the research; J.M.N. and R.G. wrote the manuscript with input from all authors.

ACKNOWLEDGMENTS

We thank Dr. Marcos Sotomayor and Dr. Josh Sanes for helpful discussions. We thank Sriram Srikant for help with phylogenetic analysis. Financial support (to J.M.N.) was provided by the National Defense Science and Engineering Graduate Fellowship. We thank the beamline staff of NE-CAT at the Advanced Photon Source (Argonne, IL, USA) for help with data collection. NE-CAT is funded by NIH (P41 GM103403 and S10 RR029205), and the Advanced Photon Source by the US Department of Energy (DE-AC02-06CH11357).

Received: April 29, 2015

Revised: September 14, 2015

Accepted: September 15, 2015

Published: October 15, 2015

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