Subsets of clustered protocadherin isoforms (α-., β-., and γ-Pcdhs) are stochastically expressed in individual vertebrate neurons. These cell-surface proteins provide a basis for neuronal self-recognition and non-self discrimination, which underpin neuronal self-avoidance. Using cell aggregation assays, X-ray crystallography, cryo-electron microscopy, biophysical measurements, and computational modeling, we have determined much of the molecular logic by which Pcdhs mediate neuronal self-vs-non-self-discrimination. Pcdh isoforms mediate cell-cell recognition through strictly homophilic trans-interactions involving extracellular cadherin domains 1–4 (EC1–4) [1]. Crystal structures of multiple α-, β-, and γ-isoforms revealed the molecular basis of their homophilic specificity [2]. Pcdh isoforms also associate promiscuously in cis via their membrane-proximal EC5–6 domains, generating cis-dimeric recognition units [1,3]. Coupling of Pcdh cis and trans interactions results in the formation of a zipper-like assembly between contacting cell surfaces [3]. Computational experiments showed that the size of this assembly is very sensitive to the presence of mismatched isoforms between contacting cell surfaces, suggesting a mechanism for self-vs-non-self-discrimination among vertebrate neurons [1].


**Keywords:** Cell-cell recognition, Neuronal self-avoidance, Protein interaction specificity