Scribble is a critical cell polarity regulator that has been shown to work as either an oncogene or tumor suppressor in a context dependent manner, and also impacts cell migration, tissue architecture and immunity. Mutations in Scribble lead to neural tube defects in mice and humans, which has been attributed to a loss of interaction with the planar cell polarity regulator Vangl2. We show that the Scribble PDZ domains 1, 2 and 3 are able to interact with the C-terminal PDZ binding motif of Vangl2 and have now determined crystal structures of these Scribble PDZ domains bound to the Vangl2 peptide. Mapping of mammalian neural tube defect mutations reveal that mutations located distal to the canonical PDZ domain ligand binding groove can not only ablate binding to Vangl2 but also disrupt binding to multiple other signaling regulators. Our findings suggest that PDZ-associated neural tube defect mutations in Scribble may not simply act in a Vangl2 dependent manner but as broad-spectrum loss of function mutants by disrupting the global Scribble-mediated interaction network.

Introduction

The establishment of cell polarity, defined as the asymmetric distribution of proteins, carbohydrates and lipids within the cell, is a crucial process for the organization and development of all animal tissues [1]. In multicellular organisms, four major different types of polarity can be distinguished: apico-basal cell polarity, asymmetric cell division, front-rear cell polarity and planar cell polarity [2]. Planar cell polarity is a critical feature of the extension and axial elongation of tissues, establishing uniformly directional information through coordinating polarity across cells within the tissue plane [3]. In mammals, planar cell polarity is essential during embryonic developmental stages, where its dysregulation during this time results in the neural tube failing to close and subsequent congenital malformations, including spina bifida and craniorachischisis, a severe neural tube defect in which the midbrain, hindbrain and the entire spinal region remain open [4]. Planar cell polarity deregulation is further implicated in defects in wound closure, a process that requires coordination of multiple cells across a tissue to migrate and proliferate together, as well as the early stages of cancer [5].

The Scribble protein is a highly conserved cell polarity regulator comprising 16 Leucine-Rich Repeats and four PSD-95/Discs-large/ZO-1 (PDZ) domains and belongs to the LAP family of proteins [6,7]. Together with Discs Large (Dlg) and Lethal Giant Larvae (Lgl), Scribble acts to establish and maintain tissue homeostasis, as well as directed cell migration and tissue growth [8]. Whilst Scribble was originally characterized as a cell polarity regulator and tumor suppressor in the vinegar fly Drosophila melanogaster [9], its role in planar cell polarity was first identified in the mouse, where mutation to Scribble result in a severe, embryonic lethal form of neural tube defect known as craniorachischisis [10,11] (Reviewed in [12]). Importantly, Scribble mutations leading to neural tube defects have also been identified in humans, further highlighting its importance as a candidate gene to study planar cell polarity regulation and developmental defects [13,14].
Scribble modulates numerous cellular processes through its many protein–protein interactions, which are largely mediated by Scribble’s four PDZ domains [8]. These PDZ domains bind C-terminally located PDZ-binding motifs (PBMs) on specific interactors in a selective manner, whilst individual Scribble PDZ domains display overlapping specificities for particular ligands [8].

One such Scribble-interactor is Vangl2 (mammalian homolog of Drosophila Van Gogh), a transmembrane protein identified as a critical component of the planar cell polarity pathway [15]. Similarly to Scribble, Vangl2 is essential in early embryonic development where its loss leads to neural tube defects as well as disruption in the cochleae inner ear hair bundle orientation [10,16]. A number of mutations in human Vangl2 have also been shown to be associated to neural tube closure defects [17] with recent studies implicating disruption of planar cell polarity as the causative factor for these mutations [18]. Importantly, Vangl2 genetically interacts with Scribble to regulate planar cell polarity processes in the mouse [10,11,19,20]. Furthermore, Vangl2 has been shown to bind to Scribble PDZ domains both in vivo and in vitro [10,21,22] and its loss has been shown to lead to Vangl2 mislocalisation in mouse and zebrafish models [19,23]. Specifically, Vangl2 was shown to engage Scribble PDZ2 and 3 domains via a C-terminal PBM [21,22].

However, a more detailed molecular understanding of the relationship between Scribble and Vangl2 has been missing. We now report a systematic evaluation of Vangl2-PDZ interactions with Scribble PDZ domains, together with structural analysis of all identified Scribble–Vangl2 interactions. Furthermore, using mutagenesis we biochemically interrogate the consequences of reported Scribble mutations associated with neural tube defects, thus providing a mechanistic basis to understand the interplay between Scribble and Vangl2 during the correct establishment of planar cell polarity as well as in a disease state such as during neural tube closure defects.

**Materials and methods**

**Protein expression and purification**

Recombinant human Scribble (Uniport accession number: Q14160) domains spanning PDZ1 (728–815); PDZ2 (860–950); PDZ3 (1002–1094); PDZ4 (1099–1203) as well as Scribble mutants were expressed using *Escherichia coli* BL21 (DE3) pLysS cells (BIOLINE) as Glutathione S-transferase or Maltose Binding Protein fusions [24] and purified as previously described [25]. Scribble PDZ domain mutants PDZ1 H793A, PDZ1 E814G, PDZ2 R896A, PDZ2 H928A, PDZ3 P1043L and PDZ3 R1044Q were synthesized as codon-optimized synthetic cDNA and cloned in the pGEX-6P3 vector (Genscript). Mutant Scribble PDZ domains were expressed and purified as previously described [25].

**Isothermal titration calorimetry**

Purified wild type and mutant human Scribble PDZ domains were used in isothermal titration experiments against 8-mer peptides spanning the C terminus of human Vangl2 (Uniport accession number: Q9ULK5; RLQSETSV) or a non-binding mutant of Vangl2 (RLQSEASA) to determine the affinity for Scribble PDZ domains. Additional peptides used span the C terminus of beta-PIX [25] and APC [26] and were described previously. A synthetic pan-PDZ binding peptide referred to as superpeptide (RSWFETWV) was used as a positive control [27]. Titrations were performed at 25°C with a stirring speed of 750 rpm using the MicroCal iTC200 System (GE Healthcare). A total of 20 injections with 2 μM against peptide concentration of 0.9 mM were used. Peptides were purchased from Genscript (San Francisco, CA, USA). Raw thermograms were processed with MicroCal Origin® version 7.0 software (OriginLabTM Corporation) to obtain the binding parameters of each interaction. A 1:1 binding model was used for all interactions.

**Protein crystallisation, data collection and refinement**

PDZ1, 2 and 3 domain proteins were mixed with wild-type Vangl2 peptide at a 1 : 20 molar excess to reconstitute the relevant complexes. Samples were subjected to crystallization screening in-house using a Gryphon nanodispenser (Art Robbins Instruments). PDZ1:Vangl2 crystals were obtained at 25 mg/ml in 0.1 M Phosphate/citrate pH 4.2 and 40% (v/v) PEG 300. Both PDZ2 and PDZ2–Vangl2 complex crystals were obtained at 15 mg/ml in 0.2 M Potassium thiocyanate and 20% (w/v) PEG 3350. PDZ3–Vangl2 crystals were obtained at 10 mg/ml in 0.1 M HEPES pH 7.8 and 66% (v/v) MPD.
Diffraction data collection and structure determination

All crystals were flash cooled in mother liquor, the former supplemented with 20% (v/v) ethylene glycol (PDZ2–Vangl2 and PDZ1–Vangl2 crystals). All diffraction data were collected on the MX1 beamline at the Australian Synchrotron equipped with an ADSC Quantum 210r CCD detector (Area Detector Systems Corporation, Poway, California, USA) or the MX2 beamline equipped with the EIGER 16M detector. Diffraction data were integrated using XDS [28], followed by AIMLESS [29] for merging and scaling. The structure of the PDZ1–Vangl2 complex was solved by molecular replacement using Phaser [30] using the structure of Scribble PDZ1 (PDB code 6MTV [31]) as a search model. The structure of the PDZ3–Vangl2 complex was solved by molecular replacement using Phaser [30] using the structure of Scribble PDZ3 (PDB: 4WYT) as a search model. The structure of the PDZ2 domain on its own was also solved using the final model of the PDZ3–Vangl2 structure. The PDZ2–Vangl2 complex was then subsequently solved using the final model of the PDZ2 domain structure. The solutions produced by Phaser were manually rebuilt over multiple cycles using Coot [32] and refined using PHENIX [33]. All images were generated using the PyMOL Molecular Graphics System, Version 1.8 Schrödinger, LLC. All software was accessed using the SBGrid suite [34].

Circular dichroism spectroscopy

Proteins were buffer exchanged into 15 mM Na2HPO4/NaH2PO4 buffer. Spectra were acquired at protein concentrations ranging from 0.080–0.150 mg/ml on an AVIV 420 CD spectrometer at 25°C. Wavelength scans were performed from 190 to 260 nm at 1 nm intervals and an averaging time of 4 s. Data were processed using the AVIV Biomedical software and displayed using Excel.

Cell culture

MCF10A cells were cultured in Dulbecco's Modified Eagle Medium:F12 (DMEM:F12) supplemented with 5% donor horse serum (Gibco), 10 μg/ml insulin (Actrapid, Novo Nordisk Pharmaceuticals Ltd.), 0.5 μg/ml hydrocortisone (Merck), 20 ng/ml epidermal growth factor (Peprotech), 100 ng/ml cholera toxin (List Biological Laboratories Inc.), 100 units/ml penicillin, and 100 μg/ml streptomycin and maintained at 37°C in 5% CO2.

Plasmid construct and viral transduction

Wild-type Vangl2 cDNA and Vangl2 cDNA missing the last four amino acids (ETSV) were cloned into MSCV vector using Gibson Assembly to generate N-terminally fused GFP-Vangl2 and GFP-Vangl2ΔPBM expression constructs. Retroviral particles were produced by co-transfection of Hek293T cells with retroviral packaging plasmids and MSCV-based retroviral vectors expressing GFP-Vangl2, GFP-Vangl2ΔPBM and the MSCV vector only and subsequently infecting low-passage MCF10A cells for stable expression. Infected cells were selected with 2 μg/ml puromycin for 1 week.

GST pulldowns

GST-fusions of wild type and mutant Scribble PDZ domains were expressed and purified as described above. MCF10A cells were lysed in NETN lysis buffer (20 mM Tris-Cl, pH8.0; 100 mM NaCl; 1 mM EDTA; 0.5% Nonidet P-40) supplemented with a phosphatase inhibitor and protease inhibitor cocktail (Roche Diagnostics) for 10 min. An amount of 250 μg of MCF10A cell lysate and 5 μg of individual GST-tagged recombinant proteins were mixed and incubated with glutathione Sepharose 4b beads (GE Healthcare) at 4°C overnight. Beads were washed three times with NETN buffer prior to being eluted in loading buffer.

Western blot

Eluted samples were resolved on standard sodium dodecyl sulphate-polyacrylamide gels and transferred to polyvinylidene difluoride membrane (Millipore). Membranes were blocked in 3% BSA in 0.2% Tween-20 in PBS for 1 h, then incubated with appropriate primary and secondary antibodies in blocking solution. Membranes were visualized using an Odyssey infrared imager (LiCor Biosciences). Antibodies used were rabbit polyclonal anti-GFP antibody (A-6455; Invitrogen); rabbit polyclonal anti-β-PIX antibody (4515S; Cell Signaling Technology) and rabbit polyclonal anti-GST antibody (71-7500; Invitrogen).
Results
Scribbel PDZ domains show interactions with Vangl2

To understand the molecular basis for Scribble:Vangl2 interactions and its consequences on correct tissue development, in particular neural tube closure, we systematically examined interactions between individual Scribble PDZ domains with the C-terminal PBM of Vangl2 (RLQSETVS) using isothermal titration calorimetry. We unexpectedly found that Scribble PDZ1, 2 and 3 domains are able to interact with the PBM of Vangl2, with affinities ranging from 24–45 μM (Figure 1), whereas Scribble PDZ4 did not show any detectable affinity. As expected, a mutant Vangl2 peptide (RLQSEASA) did not show any binding to Scribble PDZ domains (Table 1). These findings are in contrast to previous data indicating that Scribble engages Vangl2 only via its PDZ2 and 3 domains [21,22]. Consequently, we performed pull-down assays using recombinant GST-fusions of Scribble PDZ1 to PDZ domain 4 domains to examine binding of GFP-Vangl2 as well as β-PIX as a control. Whilst β-PIX bound to Scribble PDZ1 and PDZ3 as previously reported [25], we could also only detect binding of GFP-Vangl2 to PDZ2 and PDZ3 and this was dependent on the Vangl2 PBM (Figure 1).

Crystal structures of PDZ2:Vangl2 and PDZ3–Vangl2 complexes

To understand the structural basis and the mode of interaction for Scribble PDZ domains with Vangl2, we determined crystal structures of ScribblePDZ1:Vangl2, PDZ2–Vangl2 and PDZ3–Vangl2 as well as Scribble PDZ2 on its own (Figure 2, Supplemental Table EVS1, Supplemental Figure EVS2). As expected, all Scribble PDZ domains adopt the typical PDZ domain fold comprising a six-stranded β-sheet with two α-helices. All three Scribble PDZ domains comprise a compact globular fold featuring six β-strands and two α-helices that adopt a β-sandwich structure. The overall mode of Vangl2 peptide engagement is conserved across all three PDZ domain complexes, with PDZ1, PDZ2 and PDZ3 engaging Vangl2 in a manner where the peptide is bound in an anti-parallel orientation compared to the second β-strand whilst being sandwiched by the α2 helix in the canonical binding pocket of the PDZ domains. A comparison of the PDZ1 and PDZ3 domain structures on their own with their Vangl2 complex counterparts revealed no significant movement of secondary structure elements upon Vangl2 binding, as previously observed in complexes of PDZ1 and PDZ3 with β-PIX [25] and PDZ1 complexes with APC [26] and MCC [31]. Similarly, superimposition of the Scribble PDZ2 domain structure with PDZ2:Vangl2 yielded an r.m.s.d of 0.453 Å over 73 Ca atoms. A detailed examination of the Scribble PDZ1:Vangl2 complex (Figure 2A) reveals that Vangl2 forms a number of direct contacts with the PDZ1 domain. The C-terminal carboxyl group of Vangl2 makes contacts with the main chains of L738PDZ1, G739PDZ1 and I740PDZ1, whilst the V521Vangl2 side chain is accommodated in a hydrophobic pocket formed by L738PDZ1, I740PDZ1, I742PDZ1, V797PDZ1 and L800PDZ1. Furthermore, hydrogen bonds between T519Vangl2: I742PDZ1, T519Vangl2:H793PDZ1, Q516Vangl2:T749PDZ1, E518Vangl2:S761PDZ1, E518Vangl2:R762PDZ1 are observed.

In the Scribble PDZ2:Vangl2 complex (Figure 2B), key features of the PDZ1:Vangl2 complex are conserved, with V521Vangl2 being located in a pocket formed by L872PDZ2, F874PDZ2, I876PDZ2, V932PDZ2 and L935PDZ2, whilst the C-terminal carboxyl group forms hydrogen bonds with the main chain of L872PDZ2, G873PDZ2 and F874PDZ2. Additional interactions are formed by S520Vangl2:S875PDZ2, T519Vangl2:H928PDZ2, as well as a salt bridge between E518Vangl2:S895PDZ2 in the PDZ3:Vangl2 complex (Figure 2C), V521Vangl2 is engaged by a hydrophobic pocket formed by L1014PDZ3, L1016PDZ3, I1018PDZ3, V1075PDZ3 and L1079PDZ3 with the V521Vangl2 carboxyl group forming hydrogens with the main chains of L1014PDZ3, L1016PDZ3, G1015PDZ3 and L1016PDZ3. Other contacts are formed by T519Vangl2: I1018PDZ3, S520Vangl2:S1017PDZ3, T519Vangl2:H1071PDZ3, and E518Vangl2:S1039PDZ3.

Whilst previous studies verified the mode of binding to Scribble PDZ1 and PDZ3 via mutagenesis [25,27], this has not been performed for Scribble PDZ2. Consequently, we generated a PDZ2 mutant where H928 was substituted by Ala (PDZ2H928A), which showed complete loss of binding to the Vangl2 PBM peptide (Figure 3). To validate our affinity measurements, we performed pull-down assays using recombinant GST-fusions of Scribble PDZ2 and GFP-Vangl2 and confirmed that PDZ2H928A could no longer bind to Vangl2 (Figure 3).

In view of the reported physical and genetic interaction of Scribble and Vangl2 and their proposed role in coordinating neural tube closure in mice and humans, we then examined the ability of reported neural tube closure point-mutants in Scribble PDZ domains on their ability to bind the PBM of Vangl2 using ITC (Figure 3). The mutations examined identified in human neural tube closure defects were PDZ1Q808H [35],
Figure 1. Interactions of Scribble PDZ domains with Vangl2.

(A) Binding profiles of isolated Scribble PDZ domains interaction with Vangl2 peptides are displayed. Each profile is represented by a raw thermogram (top panel) and a binding isotherm fitted with a one-site binding model (bottom panels). $K_D$: 24.5 $\mu$M ± 3.9 for SCRIB PDZ1:Vangl2, 40.2 $\mu$M ± 3.9 for SCRIB PDZ3:Vangl2, 45.1 $\mu$M ± 4.9 for SCRIB PDZ2:Vangl2, and NB for SCRIB PDZ4:Vangl2.
PDZ3P1043L [14], and PDZ3R1044Q [36]. We also engineered the equivalent mutation to the mouse *scrib1* ENU mutant, *crn2* [37] into human PDZ1, as PDZ1E814G, and examined engineered mutants PDZ1H793A, PDZ2R896A and PDZ2H928A. Scribble mutants PDZ1E814G and PDZ3P1043L revealed a loss of binding to Vangl2, whereas PDZ1Q808H maintained an affinity comparable to the wild-type interaction. Furthermore, despite its close proximity to PDZ3P1043L, PDZ3R1044Q maintained an affinity comparable to the wild-type interaction indicating the specificity of the PDZ3P1043L mutation on ligand binding. To confirm that the loss of binding of certain Scribble mutants was not due to an unfolding defect, we confirmed that all Scribble mutants were folded using circular dichroism spectroscopy (Supplemental Figure EVS3). Furthermore, all mutants maintained the ability to bind a synthetic pan PDZ domain binding peptide referred to as superpeptide [27] that binds PDZ domains including those of Scribble with high affinity, suggesting that all mutants maintained a functional canonical ligand-binding groove that is capable of interacting with PDZ binding motif peptides. To determine whether or not the mutations in Scribble PDZ domain mutants were specific for Vangl2 or acted more broadly we examined the ability of PDZ1E814G, PDZ1Q808H, PDZ3P1043L and PDZ3R1044Q as well as our engineered mutants for other well-known Scribble interactors, such as β-Pix [25] and APC [26]. ITC measurements revealed that PDZ1E814G and PDZ3P1043L lost binding to both β-Pix and APC in addition to Vangl2. In contrast, PDZ1Q808H and PDZ3R1044Q bound interactors with affinities comparable to wild-type PDZ [25] and APC [26], with PDZ1Q808H binding β-Pix with 10.4 μM, APC with 6.0 μM and Vangl2 with 10.4 μM, whereas PDZ3R1044Q bound β-Pix with 14.1 μM, APC with 27.4 μM and Vangl2 with 20.9 μM. PDZ1H793A showed only a significant decrease in binding to β-Pix (62.77 μM), whereas PDZ2R896A showed decreased affinity to β-Pix and APC compared to wild-type Scribble PDZ2 binding (189.24 μM and 30.59 μM, respectively). PDZ2H928A lost binding to all three interactors. To validate our affinity measurements we performed pull-down assays using recombinant GST-fusions of Scribble PDZ1 to

Table 1 Summary of affinities of Vangl2, β-Pix, APC and superpeptide [27] peptides for WT and mutant Scrib PDZ domains measured at pH 7.5 and 25°C

<table>
<thead>
<tr>
<th>SCRIB</th>
<th>K&lt;sub&gt;D&lt;/sub&gt; (μM)</th>
<th>Vangl2</th>
<th>mut Vangl2</th>
<th>β-Pix</th>
<th>APC</th>
<th>Superpeptide</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDZ1 WT</td>
<td>24.49 ± 3.90</td>
<td>NB</td>
<td>3.33 ± 0.30&lt;sup&gt;a&lt;/sup&gt;</td>
<td>5.97 ± 1.10&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.72 ± 0.06&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>PDZ1 H793A</td>
<td>29.94 ± 8.27</td>
<td>n.d.</td>
<td>62.77 ± 22.43</td>
<td>18.80 ± 6.16&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2.18 ± 0.44</td>
<td></td>
</tr>
<tr>
<td>PDZ1 Q808H</td>
<td>10.40 ± 0.27</td>
<td>n.d.</td>
<td>3.11 ± 0.1</td>
<td>6.06 ± 0.65</td>
<td>2.73 ± 1.02</td>
<td></td>
</tr>
<tr>
<td>PDZ1 E814G</td>
<td>NB</td>
<td>n.d.</td>
<td>NB</td>
<td>NB</td>
<td>2.36 ± 0.48</td>
<td></td>
</tr>
<tr>
<td>PDZ2 WT</td>
<td>45.13 ± 4.92</td>
<td>NB</td>
<td>67.84 ± 7.90&lt;sup&gt;a&lt;/sup&gt;</td>
<td>35.94 ± 1.10&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.42 ± 0.35&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>PDZ2 R896A</td>
<td>26.74 ± 4.01</td>
<td>n.d.</td>
<td>189.24 ± 24.89</td>
<td>30.59 ± 8.1</td>
<td>7.80 ± 4.0</td>
<td></td>
</tr>
<tr>
<td>PDZ2 H928A</td>
<td>NB</td>
<td>n.d.</td>
<td>NB</td>
<td>NB</td>
<td>27.68 ± 8.4</td>
<td></td>
</tr>
<tr>
<td>PDZ3 WT</td>
<td>40.16 ± 3.92</td>
<td>NB</td>
<td>14.47 ± 2.09&lt;sup&gt;a&lt;/sup&gt;</td>
<td>18.28 ± 3.33&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1.84 ± 0.77&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>PDZ3 P1043L</td>
<td>NB</td>
<td>n.d.</td>
<td>NB</td>
<td>NB</td>
<td>9.75 ± 3.90</td>
<td></td>
</tr>
<tr>
<td>PDZ4 WT</td>
<td>NB</td>
<td>NB</td>
<td>25.85 ± 3.98&lt;sup&gt;a&lt;/sup&gt;</td>
<td>NB</td>
<td>25.85 ± 3.98&lt;sup&gt;a&lt;/sup&gt;</td>
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NB denotes no binding, n.d. denotes not determined. Each of the values was calculated from at least three independent experiments. Values denoted by a and b were taken from [25] and [26], respectively.

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PDZ4 domains to examine binding of GFP-Vangl2 as well as β-PIX as a control (Figure 3E). We observed that PDZ3R1044Q, which displayed largely unchanged affinity to β-PIX, APC and Vangl2, was able to pull down β-PIX and Vangl2, similar to PDZ1Q808H. Other mutants including PDZ1E814G and PDZ3P1043L did not pull down either β-PIX or Vangl2. However, not all pull-down assays agreed with the measured affinities from ITC, with PDZ2R896A maintaining Vangl2 binding based on ITC but not in the pull-down assay.

Discussion

Scribble is a large multi-domain scaffold protein that has been shown to play a pivotal role in cell polarity by integrating signals from a multitude of interactors to control processes such as cell migration and wound healing. This is predominantly achieved via Scribble’s four PDZ domains, which mediate the vast majority of Scribble interactions [8]. Whilst apicobasal cell polarity control is the most prominent of Scribble’s function, Scribble also plays an important role in the establishment and control of planar cell polarity [12]. The importance of Scribble function on planar cell polarity is illustrated by the impact of specific point mutations in Scribble, which manifest themselves as major developmental defects such as neural tube closure defects and disruption in the cochleae inner ear hair bundle orientation [10,11]. Intriguingly, similar defects are observed for mutations in the transmembrane protein Vangl2, leading to the establishment of a genetic link between both proteins [11,16]. Although interactions between Scribble and Vangl2 have previously been shown to involve Scribble’s PDZ2 and 3 domains, the detailed molecular and structural basis for these interactions remains to be clarified.
Figure 3. Interaction profiles of mutant Scribble PDZ with Vangl2 peptides.

(A–D) Location of Scribble PDZ domain mutations used in this study. Mutated residues are colored on a gray surface representation of the relevant Scribble PDZ domain structure. Binding profiles of isolated mutant Scribble PDZ domain interactions with Vangl2 peptides. Each profile is represented by a raw thermogram (top panel) and a binding isotherm fitted with a one-site binding model (bottom panels). $K_D$: dissociation constant.
We now show that in addition to the previously identified interactions of Scribble PDZ2 and 3 domains with the C-terminal PBM of Vangl2, Scribble’s PDZ1 domain is also able to bind the Vangl2 PBM. Notably, affinities for all three interactions are comparable with a range of 24–45 μM K_D, indicating that no distinct hierarchy in affinities exists for the Scribble:Vangl2 interactions. This is unusual in comparison with other Scribble interactions with e.g. β-PIX [25] or Gukh [27]. Whilst Scribble also binds β-PIX with its PDZ1, 2 and 3 domains, PDZ1 is the highest affinity interactor with a K_D of 3.3 μM whereas PDZ2 is the lowest affinity site with a K_D of 67.8 μM [25]. In contrast, Scribble engages tightly Gukh with its PDZ1 domain (K_D = 0.66 μM) whilst PDZ3 bound Gukh with only 27.8 μM, with no binding detected with its PDZ2 and four domains [27].

To understand the structural basis of Scribble’s PDZ domain interactions with the Vangl2 PBM, we determined complexes of Scribble PDZ1,2 and 3 domains with Vangl2. A comparison of the three PDZ domain complexes with Vangl2 reveals that whilst key family defining interactions are maintained and are near identical in all three including the engagement of the C-terminal Val (0) as well as the Thr in the −2 position, other interactions vary and offer scope for distinguishing between the domains. A comparison of PDZ1:Vangl2 with other determined Scribble PDZ1 complexes indicates that specific patterns can be seen depending on the nature of the specific PBM that is bound. For example, residues at the −5 position are shown to interact with the β2–β3 loop of PDZ1, whereas such interactions could not be observed in other Scribble PDZ domains. This behavior is also observed in D. melanogaster. This may be linked to the higher affinity of PDZ1 for its interactors compared to the other domains. A notable difference between PDZ1 and PDZ2 and 3 is that the latter utilized non-aromatic residues to engage the −5 position of bound PBM sequences, which preclude π-stacking as observed in Scribble PDZ1–βPIX [25] and D. melanogaster Scribble PDZ1–GukH [27]. This supports the notion that the β2–β3 loop is a significant modulator of Scribble PDZ1 domain binding affinities, which is supported by mutagenesis data where β2–β3 loop mutations reduce binding affinity [25].

With the availability of high-resolution structures of all key Scribble interacting domains with Vangl2, we revisited the previously identified mutations in Scribble that lead to neural tube closure defects. Mapping of these point mutants [12,38,39] indicates that in addition to mutations resulting in premature termination of Scribble, several mutations mapped to individual PDZ domains. Our analysis of these mutations using ITC revealed that although PDZ1Q808H, PDZ1E814G, PDZ3P1043L and PDZ3R1044Q are located distal from the canonical PDZ domain binding site, both PDZ1E814G and PDZ3P1043L are unable to bind to Vangl2 or indeed any other interactor tested including β-PIX and APC even though both mutants are folded, whereas PDZ1Q808H and PDZ3R1044Q displayed affinity that was comparable to the wild-type interaction. In contrast, a control mutant that mutated a key H928 residue in PDZ2 to Ala showed no binding. We note that a second engineered mutant, PDZ2R896A maintained binding to the Vangl2 PBM peptide when examined by ITC, but lost binding in pull-down assays with full-length Vangl2 as well as to β-PIX. Furthermore, whilst we were able to detect binding of the Vangl2 PBM peptide to Scri PDZ1 and determine a crystal structure of the resultant complex, GST pull-down assays did not clearly detect this interaction as noted by others. This suggests that multiple factors may contribute to the interactions of Scribble PDZ domains with their interactors in addition to outright affinity, which impacts the ability to detect such interactions with a given approach. We also note that during pull-down assays, competitor interactors for Scrib present at significantly lower levels would allow such a behavior was reported for the Tiam1 PDZ domain, which undergoes significant conformational changes on its secondary structures when interacting with different binding partners [40]. Importantly, NMR and molecular dynamics experiments. (E) GST-tagged Scribble PDZ domains, wildtype and with single-point mutations incubated with MCF10A cells stably expressing GFP-Vangl2. Bound proteins were recovered with glutathione resin and revealed by western transfer using anti-GST, anti-β-PIX and anti-GFP antibodies.

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approaches suggested that the β1–β2 and β3–α1 regions are dynamically linked, thus providing for a mechanism where changes distal to the canonical ligand binding site of the TIAM1 PDZ domain modulate ligand binding and interactions. In addition, allosteric regulation on the surface [41] of Scribble PDZ domains may be affected by the mutations we examined. PDZ domains such as PDZ3 from PSD97/DLG4 have been shown to harbor sparse networks of coevolving amino acids, which when mutated may substantially impact ligand binding behavior [42,43]. Whilst comparable analyses to PSD97/DLG4 PDZ3 have not been performed for Scribble PDZ domains, considering that the networks of coevolving amino acids in PSD97/DLG4 are tightly linked to the structure and function of the PDZ domain it seems plausible that analogous networks may also be found in Scribble PDZ domains.

In order to gain further insight into the potential impact of specific mutations on Scribble PDZ domain structures and their interactions, we next examined the effect of these mutations using the mCSM server (Table 2) [44]. mCSM utilizes the Cutoff Scanning Matrix approach to describe protein structural signatures [45], and encodes distance patterns between atoms to represent protein residue environments. This allows the prediction of the impact of single-point mutations on protein stability and protein–protein interactions. Interestingly, an analysis of our Scribble PDZ domain structures revealed that PDZ1E814G, PDZ3P1043L and PDZ3R1044Q were all predicted to be destabilizing mutations that would result in a reduced affinity for an interactor, whereas PDZ1Q808H was predicted to be a stabilizing mutation with increased affinity for an interactor (Table 2). Nevertheless, the observed loss of binding of Scribble PDZ1E814G, PDZ3P1043L and PDZ3R1044Q were all predicted to be destabilizing mutations that would result in a reduced affinity for an interactor, whereas PDZ1Q808H was predicted to be a stabilizing mutation with increased affinity for an interactor (Table 2).

In summary, we showed that biochemically Scribble is able to bind the Vangl2 PBM using its PDZ1,2 and 3 domains. Furthermore, we determined the structural basis for Vangl2 binding for all the interacting Scribble PDZ domains, and establish that well known NTD mutants of Scribble outside the canonical ligand-binding groove affect its PDZ domain ligand binding behavior, with mutant PDZ domains unable to engage a broad set of known Scribble interactors and not only Vangl2. Our findings raise the possibility that Scribble mutations associated with diseases such as NTDs are not disease causing due to loss of a single interaction, and instead support a view where a given mutation perturbs the overall interaction network of the adaptor protein Scribble by impacting multiple interactions and the dynamic interplay of different interactor with Scribble. Overall these findings form a mechanistic platform to understand how dysregulation of the Scribble interactions impacts planar cell polarity establishment, and may cause tissue disruption that manifests itself as neural tube closure defects.

**Data Availability**

Coordinate files have been deposited in the Protein Data Bank under the accession code 6XA6, 6XA7, 6XA8 and 7JO7. Raw diffraction images were deposited on the SBGrid Data Bank [47] using their PDB accession numbers.

**Competing Interests**

The authors declare that there are no competing interests associated with the manuscript.
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CRediT Contribution
Marc Kvansakul: Conceptualization, Supervision, Funding acquisition, Investigation, Writing — original draft, Project administration, Writing — review and editing. Jing Yuan How: Investigation, Writing — original draft. Rebecca Stephens: Investigation, Writing — original draft. Krystle Y.B. Lim: Supervision, Investigation, Writing — review and editing. Patrick O. Humbert: Conceptualization, Supervision, Funding acquisition, Investigation, Writing — original draft, Project administration, Writing — review and editing.

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Abbreviations
PBMs, PDZ-binding motifs; PDZ, PSD-95/Discs-large/ZO-1.

References