Structural basis for the differential interaction of Scribble PDZ domains with the guanine nucleotide exchange factor \(\beta\)-PIX

Received for publication, May 28, 2017, and in revised form, October 11, 2017 Published, Papers in Press, October 23, 2017, DOI 10.1074/jbc.M117.799452

Krystle Y. B. Lim\(^{1,5}\), Nathan J. Gödde\(^{1,5}\), Patrick O. Humbert\(^{1,5,11,1}\), and Marc Kvansakul\(^{1,2}\)

From the \(^{1}\)Department of Biochemistry and Genetics, La Trobe Institute for Molecular Science, La Trobe University, Melbourne, Victoria 3086, the \(^{5}\)Cell Cycle and Cancer Genetics Laboratory, Research Division, Peter MacCallum Cancer Centre, Melbourne, Victoria 3002, the \(^{\ast}\)Sir Peter MacCallum Department of Oncology, University of Melbourne, Melbourne, Victoria 3002, and the Departments of \(^{11}\)Biochemistry and Molecular Biology and \(^{11,1}\)Pathology, University of Melbourne, Melbourne, Victoria 3010, Australia

Edited by Eric R. Fearon

Scribble is a highly conserved protein regulator of cell polarity that has been demonstrated to function as a tumor suppressor or, conversely, as an oncogene in a context-dependent manner, and it also controls many physiological processes ranging from immunity to memory. Scribble consists of a leucine-rich repeat domain and four PDZ domains, with the latter being responsible for most of Scribble’s complex formation with other proteins. Given the similarities of the Scribble PDZ domain sequences in their binding grooves, it is common for these domains to show overlapping preferences for the same ligand. Yet, Scribble PDZ domains can still exhibit unique binding profiles toward other ligands. This raises the fundamental question as to how these PDZ domains can still exhibit unique binding profiles toward other ligands. Given the similarities of the Scribble PDZ domain sequences in their binding grooves, it is common for these domains to show overlapping preferences for the same ligand. Yet, Scribble PDZ domains can still exhibit unique binding profiles toward other ligands. This raises the fundamental question as to how these PDZ domains discriminate ligands and exert specificities in Scribble complex formation. To better understand how Scribble PDZ domains direct cell polarity signaling, we investigated here their interactions with the well-characterized Scribble binding partner \(\beta\)-PIX, a guanine nucleotide exchange factor. We report the interaction profiles of all isolated Scribble PDZ domains with a \(\beta\)-PIX peptide. We show that Scribble PDZ1 and PDZ3 are the major interactors with \(\beta\)-PIX and reveal a distinct binding hierarchy in the interactions between the individual Scribble PDZ domains and \(\beta\)-PIX. Furthermore, using crystal structures of PDZ1 and PDZ3 bound to \(\beta\)-PIX, we define the structural basis for Scribble’s ability to specifically engage \(\beta\)-PIX via its PDZ domains and provide a mechanistic platform for understanding Scribble-\(\beta\)-PIX–coordinated cellular functions such as directional cell migration.

At the molecular level, cell polarity results in the asymmetric distribution of proteins, lipids, and carbohydrates into distinct cellular domains (1). This uneven distribution of cellular contents in epithelial cells gives rise to apical–basal cell polarity, which is important for the regulation of crucial cellular signaling pathways, including those associated with apoptosis, vesicle trafficking, cell proliferation, and migration (2). Conversely, loss of cell polarity is an important hallmark of cancer development (3). Scribble, a key regulator of cell polarity, is part of the Scribble module comprising Scribble, Dlg, and Lgl, which in conjunction with the Par and Crumbs complex controls apical–basal polarity (4–6). Scribble was initially discovered in Drosophila melanogaster as a tumor suppressor where Scribble loss disrupts epithelial organization and causes aberrant growth in the imaginal discs of the larvae (7). This tumor-suppressing activity is conserved across species, and in mouse models, loss of Scribble promotes tumor initiation and, in combination with oncogenic drivers such as RAS, tumor progression in multiple epithelial tissue types, including mammary, prostate, skin, and the lung (8–12). As Scribble localization is spatially restricted in most cell types, Scribble has been postulated to act as a site-specific adaptor protein that mediates molecular interactions and in particular functions to direct distinct signaling complexes to subcellular sites (4). In disease states such as prostate cancer, mislocalization of Scribble correlates with poor patient prognosis (10), whereas in breast cancer mislocalized Scribble can act as an oncogene to drive tumorigenesis via activation of the Akt/mTOR/S6 kinase pathway (13). Thus, although mislocalization of Scribble may not alter its function per se, it is likely to alter the pool of accessible ligands, resulting in aberrant interactions leading to oncogenic signaling (3, 4). In addition to its tumor modulatory activities, Scribble regulates polarity and signaling in a wide number of cell types and organisms, playing a crucial role in organ development and physiology. This includes the development of the cardiovascular system, skeletal muscle stem cells, myelination, skin barrier formation, as well as regulation of physiological responses ranging from mammalian immunity to memory loss in Drosophila (4, 9, 14–17). Thus, understanding how Scribble functions has broad implications for understanding a wide number of biological systems.

This work was supported in whole or part by the National Health and Medical Research Council Australia Project Grant APP1103871 (to M. K. and P. O. H.), Senior Research Fellowship APP1079133 (to P. O. H.), Australian Research Council Fellowship FT130101349 (to M. K.), and La Trobe University Scholarship (to K. Y. B. L.). The authors declare that they have no conflicts of interest with the contents of this article.

The atomic coordinates and structure factors (codes SVWC, SVWK, and SVWI) have been deposited in the Protein Data Bank (http://wwpdb.org/).

1 To whom correspondence may be addressed: Dept. of Biochemistry and Genetics, La Trobe University, Melbourne, Victoria 3086, Australia. Tel.: 61-3-9479-5155; Fax: 61-3-9479-2467; E-mail: p.humbert@latrobe.edu.au.

2 To whom correspondence may be addressed: Dept. of Biochemistry and Genetics, La Trobe University, Melbourne, Victoria 3086, Australia. Tel.: 61-3-9479-2263; Fax: 61-3-9479-2467; E-mail: m.kvansakul@latrobe.edu.au.
Scribble includes 16 leucine-rich repeats (LRR)\(^3\) and four PSD-95/Disc-large/ZO-1 (PDZ) domains and belongs to the LAP family of proteins (Fig. 1A). These domains enable Scribble interactions with a wide range of ligands that are involved in multiple discrete signaling pathways. For example, the Scribble LRR domain can associate with the Lgl2 protein that is also involved in the regulation of cell polarity (18). During directed cell migration, vimentin interacts with Scribble PDZ domains to regulate its proteasome degradation, allowing Scribble to induce polarization of the Golgi apparatus (19). Scribble is also involved in receptor recycling by binding to thyroid-stimulating hormone receptor and activating ARF6 through the recruitment of a \(\beta\)-PIX–GIT1 complex upon thyroid-stimulating hormone stimulation (20). Again, the majority of these interactions occur through Scribble PDZ domains. All four Scribble PDZ domains share high levels of sequence identity (Fig. 1B), raising the question of how specificity for distinct interactions with Scribble ligands is achieved. Notably, the Scribble PDZ domains appear to harbor overlapping preferences for certain ligands, with each PDZ domain capable of engaging multiple binding partners. To understand how Scribble PDZ domains distinguish between ligands and achieve selectivity and specificity, we examined Scribble interactions with a well-established and characterized binding partner, \(\beta\)-PIX. \(\beta\)-PIX is a guanine nucleotide exchange factor for small GTPases, where its membrane-associated localization is dependent on Scribble interaction (21). Scribble and \(\beta\)-PIX complexes are found in various cell types ranging from epithelial and neuronal cells to endothelial cells where they regulate cellular processes such as vesicle trafficking (22), receptor recycling (20), cytoskeletal organization, and cell migration (23–25). Previous studies have shown that all four of Scribble PDZ domains can bind to \(\beta\)-PIX (22, 26). Here, we directly examine the redundancy of Scribble PDZ domain affinity toward \(\beta\)-PIX and investigate the differences in Scribble PDZ domain affinity for \(\beta\)-PIX. To this end, we purified Scribble PDZ domains as four individual domains, and we characterized the interaction of each of the PDZ

\(^3\)The abbreviations used are: LRR, leucine-rich repeat; PBM, PDZ-binding motif; PDB, Protein Data Bank; MBP, maltose-binding protein; BisTris, 2-(bis[2-hydroxyethyl]amino)-2-(hydroxymethyl)propane-1,3-diol; BisTris propane, 1,3-bis[tris(hydroxymethyl)methylamino]propane; r.m.s.d., root mean square deviation.
domains with a peptide corresponding to the β-PIX C-terminal PDZ-binding motif (PBM). Using isothermal titration calorimetry and pulldown assays, we quantitate and reveal the binding hierarchy for Scribble PDZ domain affinity for β-PIX. Finally, we determined crystal structures of Scribble PDZ1 alone, as well as PDZ1 and PDZ3 in complex with β-PIX peptide. Through these studies, we have identified the specific residues involved in the interaction and propose a model by which Scribble–β-PIX interaction specificity is achieved.

**Results**

**Isolated Scribble PDZ domains specifically interact with the β-PIX PBM**

Scribble has previously been shown to directly interact with β-PIX C-terminal PBM using all four of its PDZ domains, with the last three C-terminal amino acids of β-PIX being essential for the interaction (22, 26). However, no comprehensive quantitative affinity measurements for all Scribble PDZ domains were available. To understand how individual Scribble PDZ domains contribute to the interaction with β-PIX, and to determine whether there are functionally relevant differences for interactions between the individual Scribble PDZ domains, we recombinantly expressed and purified all four Scribble PDZ domains and examined their affinity for an 8-mer peptide corresponding to the β-PIX PBM (Fig. 2A).

Unexpectedly, we observed marked differences in the interactions between each of the Scribble PDZ domains and β-PIX. Although PDZ domains 1–3 bound to β-PIX with micromolar affinities, we were unable to observe any interaction of PDZ4 with β-PIX. Furthermore, although PDZ1–3 interacted with
β-PIX, substantial differences were detected in the affinities. PDZ1 was identified as the strongest binder at 3.3 μM followed by PDZ3 with 14.5 μM, whereas the interaction with PDZ2 was significantly weaker at 67.8 μM. As expected, PDZ1–4 did not interact with a mutant form of β-PIX where the C-terminal TNL residues were changed to ANA (Fig. 2B). To confirm that our recombinant PDZ domains were active and indeed capable of interactions with a given PBM (Fig. 3B), we also determined the affinities of PDZ1–4 for a synthetic superpeptide that displays high affinity to a vast pool of PDZ domains and is considered a pan-PDZ-binding peptide (Fig. 3A) (27). As expected, all four PDZ domains showed micromolar affinities for the superpeptide, confirming that the lack of interaction of PDZ4 for β-PIX is significant. Together our data demonstrate that PDZ1 has the highest affinity, followed by PDZ3 and PDZ2, thus establishing a binding hierarchy for Scribble PDZ domains and β-PIX C-terminal PBM interaction.

Interaction of isolated PDZ domains with full-length β-PIX

To examine the significance of our observation of a differential β-PIX binding profile among the Scribble PDZ domains in a cellular context, we performed pulldown experiments of individual recombinant GST-PDZ fusion domains with lysates from HEK293T cells that harbor endogenous full-length β-PIX. Consistent with our initial findings, PDZ1 and PDZ3 interacted with endogenous β-PIX (Fig. 4), whereas PDZ4 did not. However, pulldown experiments with PDZ2 did not show detectable amounts of endogenous β-PIX (Fig. 4),
suggesting that the $K_D$ of 67.8 $\mu$m is too weak to be detected in this setting.

**Crystals structures of PDZ1–β-PIX peptide and PDZ3–β-PIX peptide**

To understand the structural basis of the differential interactions between Scribble PDZ1–3 domains with the β-PIX peptide, we determined crystal structures of PDZ1 as well as complexes of PDZ1–β-PIX and PDZ3–β-PIX (Figs. 5 and 6). PDZ1 adopts a compact globular fold comprising six β-strands and two α-helices that form a β-sandwich structure, with our determined structure being essentially identical to the previously solved structure (PDB code 2W4F) as indicated by an r.m.s.d. of 0.356 Å over 77 Ca atoms. PDZ1 engages β-PIX using the canonical ligand-binding groove found in PDZ domains that is formed by the β2 strand and helix α2 (Fig. 5A). β-PIX is oriented in an anti-parallel manner to the β2 strand, with the C terminus of β-PIX packed against the β1–2 loop. Leu-646β-PIX is located in a hydrophobic pocket formed by PDZ1 Leu-738, Ile-740, Ile-742, Val-797, and Leu-800, whereas Trp-641β-PIX packs against PDZ1 Gly-744 and Tyr-751. In addition, three hydrogen bonds are formed by Asn-645β-PIX–Ser-741PDZ1, Glu-643β-PIX–Ser-761PDZ1, and Thr-644β-PIX–His-793PDZ1 (Fig. 5E). Binding of the PBM of β-PIX to PDZ1 does not result in significant changes in the overall PDZ1 structure, with the complex of PDZ1–β-PIX superimposing onto PDZ1 alone with an r.m.s.d. of 0.9 Å over 96 Ca (Fig. 5D). The PDZ1–β-PIX complex features a number of crystal contacts that are mediated by a 14-residue segment N-terminal of the PDZ1 domain, and it packs against the loop connecting helix α2 and strand β6 from a neighboring PDZ1 molecule. The β-PIX-binding groove of PDZ1 is oriented toward a solvent channel.

The overall topology of the PDZ3–β-PIX complex is similar to PDZ1–β-PIX (Fig. 5F), and the two complexes superimpose with an r.m.s.d. of 1.5 Å over 89 Ca residues (Fig. 5G). In the PDZ3–β-PIX structure, the pocket accommodating Leu-646β-PIX is formed by PDZ3 Leu-1014, Leu-1016, Ile-1018, Val-1075, and Leu-1079, and hydrogen bonds are contributed by Asn-645β-PIX–Ser-1017PDZ3, Glu-643β-PIX–Ser-1039PDZ3, and Thr-644β-PIX–His-1071PDZ3 (Fig. 5F). Notably, Trp-641β-PIX adopts a different conformation in the PDZ3–β-PIX complex compared with the PDZ1 complex and now contributes a hydrogen bond via Ser-1026PDZ3. The PDZ3–β-PIX complex features a number of crystal contacts, including an interface where the two N-terminal Pro and Ala residues of the β-PIX peptide pack against the β1 strand of a neighboring PDZ3 domain. A second crystal interface is formed by the loop connecting strands β2 and β3, which packs against the β3–α1 loop from a neighboring PDZ3 domain.

Because the major structural differences in β-PIX conformation between the complexes with Scribble PDZ1 and PDZ3 domains are centered on the orientation of Trp-641, we generated a panel of mutations to further examine the role of Trp-641 for the interactions with Scribble PDZ1 and -3 domains (Fig. 7). Mutation of PDZ1 Y751S was predicted to interfere with the packing of β-PIX Trp-641 due to the loss of the aromatic ring, and it resulted in a $K_D$ of 28.2 $\mu$m. In contrast, mutation of PDZ3 S1026A led to a modest increase in β-PIX binding ($K_D = 9.6 \mu$m), suggesting that this hydrogen bond is not a key contributor to the binding of β-PIX.

Analysis of the configuration of the β2–3 loops that harbor key interaction residues for β-PIX binding in Scribble PDZ1 and -3 revealed that in PDZ1 this loop is shorter and more rigid. Indeed, chimeric PDZ1 with a β2–3 loop from PDZ3 (PDZ1PDZ3LoopRegion) bound β-PIX with a $K_D$ of 44.4 $\mu$m, whereas chimeric PDZ3 with a β2–3 loop from PDZ1 (PDZ3PDZ1LoopRegion) bound β-PIX with a $K_D$ of 1 $\mu$m, suggesting that the β2–3 loop is a critical regulator of β-PIX binding. To further understand the role of the loop in PDZ-binding specificities, we asked if the length of the loop is a key determining factor for PDZ1 and -3 binding to β-PIX. We generated additional chimeric proteins where we removed residues from PDZ3 corresponding the equivalent shorter area in PDZ1 to generate PDZ3PDZ1Loop–G, and conversely, the corresponding three amino acids of PDZ3 residues were inserted into the PDZ1 loop to generate PDZ1PDZ3LoopDHSS (Fig. 7). PDZ1PDZ3LoopDHSS showed a 9-fold decrease in its affinity toward WT β-PIX peptide (32.5 $\mu$m). In contrast, PDZ3PDZ1Loop–G only resulted in a slight increase in affinity (9.8 $\mu$m). These data suggest that although the length of the loop is important for specificity, the residues within the loop are crucial for determining PDZ binding specificity and affinity.

**Discussion**

Scribble is a critical cell polarity regulator that integrates a number of signaling pathways via interactions using its four PDZ domains. Our data suggest that Scribble PDZ domains 1–4 harbor differential affinities for the PBM of β-PIX, an important guanine nucleotide exchange factor involved in cell migration and intracellular trafficking. Previous studies (22, 26) showed that Scribble PDZ1–4 domains were all able to engage β-PIX, raising the question of how specificity is achieved in this interaction and whether or not Scribble PDZ domains are able to discriminate for β-PIX binding or alternatively act in an indiscriminate manner. We now quantitatively show that Scribble PDZ1 and -3 domains are the highest affinity β-PIX binders, whereas PDZ2 displays 20-fold weaker affinity, and
PDZ4 is unable to interact with β-PIX (Fig. 2). Importantly, in a cellular context only, PDZ1 and -3 are able to pull down endogenous β-PIX. Overall, these findings identify Scribble PDZ1 and -3 as the primary interaction sites for β-PIX and suggest that the engagement of Scribble PDZ domains with β-PIX occurs in a highly specific manner.

At the structural level, binding of β-PIX to Scribble PDZ1 and -3 is consistent with the binding mode observed for the canonical class I PDZ domains, where side chains of Leu-0, Thr-2, Glu-3, and Trp-5 are involved in key interactions with the receiving PDZ domain. A number of the interactions with β-PIX are conserved between the PDZ1 and PDZ3 domain, including hydrogen bonds between β-PIX Asn-645 and a PDZ domain Ser, β-PIX Thr-644 and a PDZ domain His, as well as β-PIX Glu-643 and PDZ domain Ser (Fig. 5, E–G). The major difference between Scribble PDZ1 and PDZ3 binding to β-PIX centers on β-PIX Trp-641, which in PDZ1 forms hydrophobic interactions by packing against PDZ1 Gly-744 and Tyr-751, whereas in PDZ3 Trp-641 adopts a different rotamer and instead only contributes a hydrogen bond with Ser-1026, a difference that is also reflected in the ~4-fold lower affinity observed with PDZ3 and β-PIX over the PDZ1 interaction (Fig. 5G).

Interestingly, the β-PIX Trp at position −5 closely mimics the behavior of a key Trp at position −4 in the high affinity nanomolar Erbin complex with a synthetic high-affinity peptide (28). This Trp in position −4 has been shown to be critical for the high-affinity interaction by engaging the β2–3 loop (Fig.
5H). Furthermore, in the high-affinity complex of ZO-1 with a different synthetic peptide (29), a Trp in the -6 position plays a similarly critical role to the Trp in the Erbin complex at position -4 by also engaging the β2–3 loop. These data suggest that the ability to form hydrophobic interactions with the β2–3 loop is an important determinant for high affinity interactions of the PDZ domain-binding ligands. Indeed, these interactions can be achieved by suitable hydrophobic residues such as Trp in the N-terminal region of the PDZ2-binding motif and are not limited to a particular position, because Trp residues in the -4, -5, and -6 positions are able to engage the β2–3 loop. However, despite utilizing a Trp to contact the β2–3 loop, the affinity of the Scribble PDZ1–β-PIX interaction is an order of magnitude lower than those by Erbin and ZO-1 with synthetic peptides. In part, this can be attributed to the observation that β-PIX does not form an ionic interaction via Glu-643 with PDZ1 due to a lack of a suitable Arg residue, as observed in both Erbin and ZO-1 complexes. However, loss of the β-PIX Trp-641 hydrophobic interaction with Scribble PDZ1 Tyr-751 substantially impacts binding, with a PDZ1 Y751S mutant showing an ~8-fold lower $K_D$. Furthermore, the β2–3 loop also plays a critical role in controlling β-PIX binding to Scribble PDZ1 and -3 domains, because swapping of this loop between Scribble PDZ1 and -3 impairs binding of β-PIX to the resultant PDZ1PDZ3LoopRegion and PDZ3PDZ1LoopDH5 chimeras, but it increases binding to the PDZ3PDZ1LoopRegion and PDZ3PDZ1LoopG chimeras (Fig. 7). This opens the possibility of selectively engineering the length and/or identity of the residues within the PDZ β2–3 loop as shown through PDZ1Y751S to modulate binding to β-PIX.

A sequence alignment of all four Scribble PDZ domains reveals that key residues involved in interactions from PDZ1 to -3 with β-PIX are not fully conserved in PDZ2 and -4. In particular, the residues involved in interactions with β-PIX Trp-641 such as Tyr-751 and Ser-1026 are not found in PDZ4. Our mutagenesis data support the notion that Tyr-751 is important for binding of β-PIX, and the substitution of this residue is one of the underlying reasons for the lack of an interaction between PDZ4 and β-PIX, and possibly for the substantially lower affinity observed for PDZ2–β-PIX.

Previous studies of the shank PDZ domain interaction with β-PIX revealed that only a single β-PIX chain within the trimeric β-PIX coiled-coil engages with a shank PDZ domain at a given time (30). If a similar mechanism is involved in the PDZ domain-mediated interaction of Scribble with β-PIX, we speculate that PDZ1 is the primary interaction site for β-PIX; however, such a model awaits experimental verification.

In summary, we show that Scribble PDZ domains display differential binding affinities and capacities for binding to β-PIX and that Trp-641 of the β-PIX PBM is a key determinant for the observed differences in affinities. These findings provide a clear structural basis for Scribble–β-PIX interactions and will form the platform for detailed structure-guided investigations to understand how the differences in ability of individual PDZ domains to engage β-PIX impact the control of cell polarity and directed migration.

**Experimental procedures**

**DNA constructs**

Expression constructs encoding PDZ1, PDZ2, and PDZ4 residues 2108–2455, 2501–2896, and 3302–3616, respectively, are based on the hScrib transcript variant 1 (NM_182706). PDZ1t (residues 2180–2449) and PDZ3 (residues 3011–3283) and all PDZ1 and PDZ3 mutants were codon-optimized for protein expression in *Escherichia coli* BL21 (GenScript). For bacterial expression, all constructs were cloned into pGil-MBP (31) and pGex-6P-3 (GE Healthcare), except for PDZ1t, which was cloned into pGex-6P-1 (GE Healthcare).
Crystal structures of Scribble PDZ1 and PDZ3 with β-PIX

A

| PROTEIN | SEQUENCE | K_D |
|---------|----------|-----|
| PDZ1    | GLGISIAGGK--GSTPYKDDEGI | 758 3.3 ± 0.3 μM |
| PDZ1<sub>PZD3LoopRegion</sub> | GLGISIAGGS DHS HPF GVQ EP GI | 758 44.4 ± 8.1 μM |
| PDZ1<sub>PZD3LoopDHS</sub> | GLGISIAGGGSS HPF GVQ EP GI | 758 32.5 ± 2.2 μM |
| PDZ1<sub>Y751S</sub> | GLGISIAGGK--GSTPS KDDEGI | 758 28.2 ± 6.4 μM |
| PDZ3    | PLGLS IVGSDSHS HPF GVQ EP GV | 1036 14.5 ± 2.1 μM |
| PDZ3<sub>PZD1LoopRegion</sub> | PLGLS IVGSGK--GSTPYKDDEGV | 1036 1.0 ± 0.2 μM |
| PDZ3<sub>PZD1Loop--G</sub> | PLGLS IVGGS--GHPF GVQ EP GV | 1036 9.8 ± 0.5 μM |
| PDZ3<sub>S1026A</sub> | PLGLS IVGSDHSHPF GVQ EP GV | 1036 9.6 ± 1.3 μM |
Protein expression and purification

Protein overexpression was performed using *E. coli* BL21 (DE3) pLYS S cells (BIOLINE) in super broth supplemented with 200 μg/ml ampicillin (AMRESCO) using auto-induction media (10 mM Tris-Cl, pH 7.6, 100 mM NaCl, 1 mM MgSO4, 0.2% (w/v) d-lactose, 0.05% (w/v) glucose, 0.5% (v/v) glycerol) (32) at 37 °C until the absorbance at 600 nm (A600) reached 1.0 before transferring cultures to 16 or 25 °C for 24 h for protein expression. Proteins were also expressed via manual induction using 1 mM isopropyl 1-thio-D-galactopyranoside bacterial cultures at A600 0.6 at 37 °C for 4 h or at 20 °C for 24 h.

Bacterial cells were harvested by centrifugation and lysed in the presence of deoxyribonuclease I (Sigma) from bovine pancreas using TS Series 0.75 kw model cabinet (Constant Systems Ltd.) at 25,000 p.s.i., Qsonica Q700 sonicator at amplitude 50 for 4 min process time on ice (5 s pulse-on time and 30 s pulse-off time) or a Fastprep®-24 (MP Biomedicals) using lysing matrix B for 20 s. Lysates were clarified by centrifugation at 20,000 x g for 20 min using an Avanti® J-E (Beckman Coulter).

The resulting supernatant was filtered using Millex-GP syringe filter unit 0.22 μm (Merck Millipore) prior to loading onto equilibrated columns for affinity purification.

Glutathione S-transferase (GST)-tagged recombinant proteins were captured using glutathione-Sepharose 4B (GE Healthcare) in buffer A (50 mM Tris-Cl, pH 8.0, 150 mM NaCl, and 1 mM EDTA) and eluted with buffer A supplemented with 20 mM reduced l-glutathione. Recombinant proteins were subsequently dialyzed into buffer A to remove glutathione for the pulldown assay. GST-tagged PDZ1 was cleaved on-column with HRV 3C protease to remove the GST tag.

Hexahistidine and maltose-binding protein (His-MBP)-tagged recombinant proteins were purified using HisTrap HP columns (GE Healthcare) in buffer B (50 mM Tris-Cl, pH 8.5, 300 mM NaCl, 5 mM β-mercaptoethanol) and washed with buffer B supplemented with 20 mM imidazole before eluting in buffer B supplemented with 250 mM imidazole. His-MBP-tagged recombinant proteins were cleaved with tobacco etch virus protease in buffer B supplemented with 0.5 mM EDTA and 1 mM DTT before being subjected to a second round of affinity chromatography to remove the cleaved His-MBP tag and uncleaved fusion protein. All cleaved target proteins were subjected to size-exclusion chromatography using the HiLoad 16/600 Superdex 75 (GE Healthcare) equilibrated in 25 mM HEPES, pH 7.0, 150 mM NaCl, and eluted as a single peak.

Cell culture

Human embryonic kidneys transformed with SV40 large T antigen cell lines (HEK293T) were cultured and maintained in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 20 mM HEPES, 10% fetal bovine serum (Gibco), 2 mM GlutaMAX (Gibco), 100 units/ml penicillin (Gibco), and 100 units/ml streptomycin (Gibco) at 37 °C, 5% CO2.

GST pulldown assay

HEK293T cells were washed with Dulbecco’s PBS prior to lysis with NETN lysis buffer (20 mM Tris-Cl, pH 8.0, 100 mM NaCl, 1 mM EDTA, 0.5% Nonidet P-40) supplemented with phosphatase and protease inhibitor mixture (Roche Diagnostics) for 10 min on ice. The lysate was centrifuged for 10 min at high speed using the benchtop centrifuge to remove cell debris from the sample. The protein concentration of the resulting supernatant was determined by Lowry assay using the DC™ protein assay kit (Bio-Rad). 250 μg of HEK293T supernatant was incubated with 5 μg of each GST-tagged recombinant proteins for an hour, rotating at 4 °C. 75 μl of 50% glutathione-Sepharose 4B slurry was added to each sample and incubated overnight at 4 °C. The glutathione-Sepharose 4B beads were washed three times with NETN lysis buffer before liberating the bound protein with sample loading buffer.

The samples were resolved in Bolt 4–12% BisTris Plus Gels (Invitrogen) in Bolt MOPS SDS Running buffer. Proteins were transferred to a polyvinylidene fluoride (PVDF) membrane (Millipore) via 30 V overnight at 4 °C using a Mini Trans-Blot Electroblot Electrophoretic Transfer Cell (Bio-Rad).

Western blotting

All Western blottings were performed in the presence of 5% skim milk in TBS with 0.2% Tween 20. The antibodies used were rabbit polyclonal anti-β-PIX (Cell Signaling Technology), rabbit polyclonal anti-GST (Invitrogen), and secondary goat anti-rabbit IgG (Bio-Rad). Membranes were initially probed with anti-β-PIX and then stripped with stripping buffer (20 mM Tris-Cl, pH 6.8, 1% SDS, and 100 mM β-mercaptoethanol) at 60 °C for an hour for subsequent anti-GST probing. Lumi-Light enhanced chemiluminescence (Roche Diagnostics) was used to reveal the membrane.

Isothermal titration calorimetry

PDZ1 concentration was quantitated at 280 nm absorbance (A280 nm) using a NanoDrop 2000/2000c UV-visible spectrophotometer (Thermo Fisher Scientific). Because the concentration of PDZ2, PDZ3, and PDZ4 could not be determined by A280 nm, the Direct Detect® infrared spectrometer (Merck Millipore) method was used. Titrations were performed at 25 °C with a stirring speed of 1000 rpm using the MicroCal™iTC200 system (GE Healthcare). A total of 20 injections with 2 μl each and a spacing of 180 s were titrated into the 270-μl protein sample (25 mM HEPES, pH 7.0, 150 mM NaCl), except for the first injection that was only 0.4 μl. A protein concentration of 20–100 μM against a peptide concentration of 0.5–1.4 mM was used according to the titration requirements. Peptides

Figure 7. Interaction profiles of Scribble mutant PDZ1 and -3 domains with β-PIX peptides. A, binding profiles of isolated mutant Scribble PDZ1 domain interactions with β-PIX peptides. Each profile is represented by a raw thermogram (top panel) and a binding isotherm fitted with a one-site binding model (bottom panel). B, binding profiles of isolated mutant Scribble PDZ3 domain interactions with β-PIX peptides. Each profile is represented by a raw thermogram (top panel) and a binding isotherm fitted with a one-site binding model (bottom panel). Kd, dissociation constant; ±, standard deviation; NB, no binding. Each of the values was calculated from at least three independent experiments. C, sequence details of mutant PDZ1 and PDZ3 domains used in this study. The original sequences of PDZ1 are shown in black, and the original sequences of PDZ3 are shown in gray. Mutated residues are shown as white font in a black box.
Crystal structures of Scribble PDZ1 and PDZ3 with β-PIX

Table 1
Data collection and refinement statistics

|                   | PDZ1–β-PIX | PDZ3–β-PIX | PDZ1 |
|-------------------|------------|------------|------|
| **Data collection** |            |            |      |
| Space group       | P 4 2 2   | P 2 1     | P 2 1 2 |
| No. of molecules in asymmetric units | 4 + 4     | 2 + 2     | 1   |
| Cell dimensions   |            |            |      |
| a, b, c (Å)       | 74.76, 74.76, 222.96 | 37.87, 44.16, 60.47 | 35.49, 41.69, 51.57 |
| α, β, γ (°)       | 90.00, 90.00, 90.00 | 90.00, 93.61, 90.00 | 90.00, 90.00, 90.00 |
| Wavelength (Å)    | 0.9537     | 0.9537     | 0.9537 |
| Resolution (Å)    | 47.77–2.35 (2.43–2.35 | 44.16–1.75 (1.78–1.75) | 41.69–1.91 (1.91–1.95) |
| Rmerge or Rmerge | 0.126 (0.668) | 0.072 (1.759) | 0.076 (0.425) |
| I/σ               | 7.3 (1.8) | 17.6 (1.0) | 16.2 (4.5) |
| CC(1/2)           | 0.996 (0.738) | 0.999 (0.307) | 0.999 (0.856) |
| Completeness (%)  | 97.5 (99.2) | 99.8 (99.9) | 99.3 (95.6) |
| Redundancy        | 4.1 (4.1) | 6.2 (6.1) | 5.7 (5.7) |
| Wilson B-factor   | 26.9       | 24.7       | 20.2  |
| **Refinement**    |            |            |      |
| Resolution (Å)    | 38.36–2.35 | 31.16–1.75 | 32.42–1.91 |
| No. of reflections | 26459     | 20310     | 6245 |
| Rmerge/Rmerge     | 0.223/0.256 | 0.191/0.237 | 0.195/0.227 |
| No. of non-hydrogen atoms | 3290 | 1467 | 1492 |
| Protein           | 207        | 207       | 12   |
| Ligand/ion        | 263        | 263       | 48   |
| Water             |            |            |      |
| B-Factors         |            |            |      |
| Protein           | 23.70      | 42.74      | 30.95 |
| Ligand/ion        | 48.90      | 48.90      | 33.31 |
| Water             | 33.80      | 33.80      | 30.07 |
| r.m.s.d.          |            |            |      |
| Bond lengths (Å)  | 0.656      | 1.169      | 0.923 |
| Bond angle (°)    | 0.004      | 0.011      | 0.008 |
| Ramachandran plot (%) | 99.1   | 97.8       | 100  |
| Favored           | 99.1       | 97.8       | 100  |
| Allowed           | 0.9        | 2.2        | 0    |
| Disallowed        | 0          | 0          | 0    |

were purchased from GenScript (San Francisco). The raw thermograms were processed with MicroCal Origin® version 7.0 software (OriginLabTM Corp.) to obtain the binding parameters of each interaction. Peptides used are wild-type β-PIX (NP_003890.1): 639PAWDETNL 646, mutant β-PIX: PAWDEANA, superpeptide (27): RSWFETWV and mutant superpeptide: RSWFETWV.

Protein crystallization

Crystallization screens were performed at the CSIRO C3 facility (Parkville, Australia) using the Phoenix Liquid Handling System (Art Robbins Instruments) as well as in-house using the Crystal Gryphon (Art Robbins Instruments). All crystals were grown at 20 °C. For PDZ1t crystals, β-PIX peptide was added to 2.92 mg/ml PDZ1t (in buffer A) with a peptide to protein ratio of 1:1.07. The final crystals were obtained in 20% (w/v) polyethylene glycol 3350, 0.2 M magnesium chloride hexahydrate. For the protein complex crystals, a protein (25 mM HEPES, pH 7.0, 150 mM NaCl) to peptide ratio of 1:5 was used for crystallization. PDZ1–β-PIX complex was prepared at a concentration of 150 mM NaCl to peptide ratio of 1:5 was used for crystallization. PDZ1–β-PIX was also solved by molecular replacement using the structure of Scribble PDZ3 (PDB code 4WYT) as a search model (TFZ = 39.3 and LLG = 2216). The solutions produced by Phaser were manually rebuilt over multiple cycles using Coot (37) and refined using PHENIX (38). Data collection and refinement statistics details are summarized in Table 1. Coordinate files have been deposited in the Protein Data Bank under the accession codes 5VWC, 5VWK, and 5VWI. All images were generated using the PyMOL Molecular Graphics System, Version 1.8, Schrödinger, LLC. All software was accessed using the SBGrid suite (39). All raw diffraction images were deposited on the SBGrid Data Bank (40) under their PDB codes 5VWC, 5VWK, and 5VWI.

Author contributions—K. Y. B. L.: acquisition of data, analysis and interpretation of data, drafting and revising the article; N. J. G.: acquisition of data, analysis and interpretation of data; P. O. H.: conception and design, acquisition of data, analysis and interpretation of data, drafting and revising the article; M. K.: conception and design, acquisition of data, analysis and interpretation of data, drafting and revising the article.
Acknowledgments—We thank staff at the MX beamlines at the Australian Synchrotron for help with X-ray data collection; the CSIRO C3 Collaborative Crystalization Centre for assistance with crystallization; and the Comprehensive Proteomics Platform at La Trobe University for core instrument support.

References

1. Nelson, W. J. (2003) Adaptation of core mechanisms to generate cell polarity. *Nature* 422, 766–774.
2. McCaffrey, L. M., and Macara, I. G. (2012) Signaling pathways in cell polarity. *Cold Spring Harb. Perspect. Biol.* 4, a009654.
3. Halauoi, R., and McCaffrey, L. (2015) Rewiring cell polarity signaling in cancer. *Oncogene* 34, 939–950.
4. Humbert, P. O., Russell, S. M., Smith, L., and Richardson, H. E. (2015) in *Cell Polarity 1: Biological Role and Basic Mechanisms* (Ebneth, K., ed) pp. 65–111, Springer International Publishing, Cham, Switzerland.
5. Ohno, S., Goulas, S., and Hori, T. (2015) in *Cell Polarity 1: Biological Role and Basic Mechanisms* (Ebneth, K., ed) pp. 3–23, Springer International Publishing, Cham, Switzerland.
6. Vaccu, B., Barthélémy-Quémin, M., Burcklé, C., Massey-Harroche, D., and Bivic, A. L. (2015) in *Cell Polarity 1: Biological Role and Basic Mechanisms* (Ebneth, K., ed) pp. 51–63, Springer International Publishing, Cham, Switzerland.
7. Bilder, D., Li, M., and Perrimon, N. (2000) Cooperative regulation of cell polarity and growth by *Drosophila* tumor suppressors. *Science* 289, 113–116.
8. Zhan, L., Rosenberg, A., Bergami, K. C., Yu, M., Xuan, Z., Jaffe, A. B., Allred, C., and Muthuswamy, S. K. (2008) Deregulement of scribble promotes mammary tumorigenesis and reveals a role for cell polarity in carcinogenesis. *Cell* 135, 865–878.
9. Pearson, H. B., McGlinn, E., Phesse, T. J., Schlüter, H., Srikumar, A., O'Donoghue, R., Ernst, M., Cullinane, C., and Humbert, P. O. (2014) Scribble modulates the MAPK/Fra1 pathway to disrupt luminal and ductal integrity and suppress tumour formation in the mammary gland. *PLoS Genet.* 10, e1004323.
10. Elsum, I. A., Yates, L. L., Pearson, H. B., Phesse, T. J., Long, F., O'Donoghue, R., Ernst, M., Cullinane, C., and Humbert, P. O. (2014) Scribble heterozygosity predisposes to lung cancer and cooperates with KRas hyperactivation to accelerate lung cancer progression in vivo. *Oncogene* 33, 5523–5533.
11. Feigin, M. E., Akshinthala, S. D., Araki, K., Rosenberg, A. Z., Muthuswamy, L. B., Martin, B., Lehmann, B. D., Berman, H. K., Pieterpol, J. A., Cardiff, R. D., and Muthuswamy, S. K. (2014) Mislocalization of the cell polarity protein scribble promotes mammary tumorigenesis and is associated with basal breast cancer. *Cancer Res.* 74, 3180–3194.
12. Ono, Y., Urata, Y., Goto, S., Nakagawa, S., Humbert, P. O., Li, T. S., and Zammit, P. S. (2015) Muscle stem cell fate is controlled by the cell-polarity protein Scrib. *Cell Rep.* 10, 1135–1148.
13. Jarjour, A. A., Boyd, A., Dow, L. E., Holloway, R. K., Goebels, S., Humbert, P. O., Williams, A., and ffrench-Constant, C. (2015) The polarity protein Scribble regulates myelination and remyelination in the central nervous system. *PLoS Biol.* 13, e1002107.
14. Zheng, W., Umitsu, M., Jagan, I., Tran, C. W., Ishiyama, N., BeGora, M., Araki, K., Ohashi, P. S., Ikura, M., and Muthuswamy, S. K. (2016) An interaction between Scribble and the NAPDH oxidase complex controls M1 macrophage polarization and function. *Nat. Cell Biol.* 18, 1244–1252.
15. Cervantes-Sandoval, L., Chakraborty, M., MacMullen, C., and Davis, R. L. (2016) Scribble scaffolds a signalosome for active forgetting. *Neuron* 93, 1200–1204.
16. Kallay, L. M., McNickle, A., Brennwald, P. J., Hubbard, A. L., and Braiterman, L. T. (2006) Scribble associates with two polarity proteins, Lgl2 and Vangl2, via distinct molecular domains. *J. Cell. Biochem.* 99, 647–664.
17. Phua, D. C., Humbert, P. O., and Hunziker, W. (2009) Vimentin regulates scribble activity by protecting it from proteasomal degradation. *Mol. Biol. Cell* 20, 2841–2855.
18. Lahou, O., Quellari, M., Achard, C., Nola, S., Méduri, G., Navarro, C., Vitale, N., Borg, J. P., and Misrahi, M. (2005) Thyrotropin receptor trafficking relies on the hScrib-βPIX-GIT1-ARF6 pathway. *EMBO J.* 24, 1364–1374.
19. Ohno, S., Goulas, S., and Hori, T. (2015) in *Cell Polarity 1: Biological Role and Basic Mechanisms* (Ebneth, K., ed) pp. 3–23, Springer International Publishing, Cham, Switzerland.
20. Bilder, D., Li, M., and Perrimon, N. (2000) Cooperative regulation of cell polarity and growth by *Drosophila* tumor suppressors. *Science* 289, 113–116.
21. Arnaud, C., Sebbagh, M., Marchetto, S., Nola, S., Nourry, C., Audebert, S., Navarro, C., Rachel, R., Montcouquiol, M., Sans, N., Etienne-Manneville, S., Borg, J. P., and Santoni, M. J. (2008) Scrib regulates PKA activity during the cell migration process. *Hum. Mol. Genet.* 17, 3552–3565.
22. Michaelis, U. R., Chavakis, E., Kruse, C., Jungblut, B., Kalyuzhka, D., Wandzioch, K., Manavsky, Y., Heide, H., Santoni, M. J., Potente, M., Eble, J. A., Borg, J. P., and Brandes, R. P. (2013) The polarity protein Scrib is essential for directed endothelial cell migration. *Circ. Res.* 112, 924–934.
23. P. O., Williams, A., and ffrench-Constant, C. (2015) The polarity protein Scrib is essential for directed endothelial cell migration. *Circ. Res.* 112, 924–934.
Crystal structures of Scribble PDZ1 and PDZ3 with β-PIX

(2011) Overview of the CCP4 suite and current developments. Acta Crystallogr. D Biol. Crystallogr. 67, 235–242
36. McCoy, A. J. (2007) Solving structures of protein complexes by molecular replacement with Phaser. Acta Crystallogr. D Biol. Crystallogr. 63, 32–41
37. Emsley, P., Lohkamp, B., Scott, W. G., and Cowtan, K. (2010) Features and development of Coot. Acta Crystallogr. D Biol. Crystallogr. 66, 486–501
38. Afonine, P. V., Grosse-Kunstleve, R. W., Echols, N., Headd, J. J., Moriarty, N. W., Mustyakimov, M., Terwilliger, T. C., Urzhumtsev, A., Zwart, P. H., and Adams, P. D. (2012) Towards automated crystallographic structure refinement with phenix.refine. Acta Crystallogr. D Biol. Crystallogr. 68, 352–367
39. Morin, A., Eisenbraun, B., Key, J., Sanschagrin, P. C., Timony, M. A., Ottaviano, M., and Sliz, P. (2013) Collaboration gets the most out of software. Elife 2, e01456
40. Meyer, P. A., Socias, S., Key, J., Ransey, E., Tjon, E. C., Buschiazzo, A., Lei, M., Botka, C., Withrow, J., Neau, D., Rajashankar, K., Anderson, K. S., Baxter, R. H., Blacklow, S. C., Boggon, T. J., et al. (2016) Data publication with the structural biology data grid supports live analysis. Nat. Commun. 7, 10882
41. Robert, X., and Gouet, P. (2014) Deciphering key features in protein structures with the new ENDscript server. Nucleic Acids Res. 42, W320–W324