Protein–interaction domains can create unique macromolecular complexes that drive evolutionary innovation. By combining bioinformatic and phylogenetic analyses with structural approaches, we have discovered that the docking and dimerization (D/D) domain of the PKA regulatory subunit is an ancient and conserved protein fold. An archetypal function of this module is to interact with A-kinase-anchoring proteins (AKAPs) that facilitate compartmentalization of this key cell-signaling enzyme. Homology searching reveals that D/D domain proteins comprise a superfamily with 18 members that function in a variety of molecular and cellular contexts. Further in silico analyses indicate that D/D domains segregate into subgroups on the basis of their similarity to type I or type II PKA regulatory subunits. The sperm autoantigenic protein 17 (SPA17) is a prototype of the type II or R2D2 subgroup that is conserved across metazoan phyla. We determined the crystal structure of an extended D/D domain from SPA17 (amino acids 1–75) at 1.72 Å resolution. This revealed a four-helix bundle-like configuration featuring terminal β-strands that can mediate higher order oligomerization. In solution, SPA17 forms both homodimers and tetramers and displays a weak affinity for AKAP18. Quantitative approaches reveal that AKAP18 binding occurs at nanomolar affinity when SPA17 heterodimerizes with the ropporin-1-like D/D protein. These findings expand the role of the D/D fold as a versatile protein-interaction element that maintains the integrity of macromolecular architectures within organelles such as motile cilia.

Protein–protein interactions constrain macromolecules to form molecular machines (1). A-kinase-anchoring proteins (AKAPs) confine PKA within ‘signaling islands’ to create highly organized signaling compartments (2–4). A defining attribute of AKAPs is an amphipathic α-helix that binds with high affinity to the docking and dimerization (D/D) domain of PKA regulatory (PKA-R) subunits (5, 6). The PKA holoenzyme is composed of two catalytic (C) subunits constrained by an R-subunit that mediates dimerization and AKAP binding (13). Subsequent NMR and crystallographic studies have characterized the structure of the RI and RII D/D domains in complex with AKAPs (14, 15). R-subunit protomers dimerize to form an X-type helix bundle in an antiparallel arrangement (Fig. 1A). A hydrophobic groove formed at the top of this substructure docks with an amphipathic α-helix on the surface of the AKAP (16–18). A helical segment at the amino terminus of RI subunits orients key determinants for AKAP binding (Fig. 1A). In RII subunits, the first five amino acids form β-strands that are essential for docking, with isoleucine’s 3 and 5 serving as key PKA-anchoring determinants (19) (Fig. 1A). About 60 AKAPs have been identified, each containing a PKA-anchoring helix that associates with D/D domains (Omar and Scott, 2020). These regions of secondary structure have degenerate sequences of 14 to 18 residues, but with a discernable pattern of hydrophobic amino acids critical for docking (5, 12). Peptide studies have uncovered primary structure determinants that influence AKAP binding to RI and to RII (6, 20–25).

While docking grooves were initially believed to be less modular than peptide motifs, the D/D domain is now designated as a bona fide modular unit (26–28). Studies on spermatozoa revealed that anchoring disruptor peptides, such as Ht31, impair flagellar motility (29). Flagellar motility was unaffected by the kinase inhibitor PKI, or the drug H-89 (29). This allowed these authors to conclude that anchored PKA was not involved in this process. Since then, D/D proteins such as sperm autoantigenic protein 17 (SPA17), ROPN1, ropporin-1-like protein (ROPNL), and CABYR have been recognized as nonkinase AKAP helix–binding partners (28, 30, 31). Functional studies report that genetic ablation of these D/D proteins or loss of association with AKAPs impairs motile ciliary action or flagellar motility (32, 33).

The bioinformatic and phylogenetic studies reported herein classifies the D/D domain superfamily. Eighteen superfamily

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members are subdivided into type I (R1D2) and type II (R2D2) lineages. Many of these proteins are ancient and present in diverse eukaryotic kingdoms. Others result from a gene expansion that took place at the advent of metazoan multicellularity. To gain further insight into the R2D2 lineage, we determined the crystal structure of apo SPA17 1 to 75 from Danio rerio. SPA17 can form homotetramers and displays a low affinity for AKAPs. AKAP binding is considerably enhanced when SPA17 heterodimerizes with another R2D2 protein ROPN1L. Thus, cross-member heterodimerization expands the repertoire and functionality of D/D domains.

Results

Annotation of the D/D domain superfamily

A combined strategy for data mining was utilized to generate an improved inventory and annotation of D/D domain–containing proteins. Three databases were interrogated to define relationship hierarchies (Fig. 1B). First, the NCBI Conserved Domain Database was searched for proteins within the “Dimerization/docking domain of the regulatory subunit of cAMP-dependent kinase and similar domains”. Second, the SuperFamily library was searched for proteins with the “dimerization-anchoring domain of cAMP-dependent PK regulatory subunit.” Third, protein BLAST analyses against metazoan and excluding metazoan species generated a comprehensive list of the RIIα clan across all taxa. Screening of the Pfam database refined the RIIα clan as consisting of Dumpy-30 (DPY-30) and PKA-R subunit superfamilies.

The output of our data mining strategy is diagrammatically presented in Figure 1B. We defined the RIIα clan as the group comprising the DPY-30 and PKA-R superfamilies. This has led to the identification of 18 PKA-R superfamily members based on sequence identity (Fig. 1C and Table 1). The group is further subdivided into type I and type II PKA-R subunit-like D/D proteins (Fig. 1, B and C).

Taxonomic distribution and phylogeny of the D/D domain superfamily

A total of 249 D/D domain–containing proteins across all taxa were selected for further analysis (Supplemental material). Metazoans have a full complement of PKA-R-like proteins (Fig. 2A). These include SPA17, ROPN1L, RIIAD1, CATIP, EFCAB10, TPGS1, AK5, AK8, VEST1, FBXL13, and TEX55. Taxa outside the metazoan kingdom contain the PKA-R-like proteins ROPN1L/RSP11, RSP7, TPGS1, EFCAB10, AK8, enolase, and RIIAD1 (Fig. 2A). Higher animals, including humans, additionally evolved the sperm fibrous sheath R2D2 proteins CABYR, ROPN1, and ROPN1B (Fig. 2A). The full gene name of each PKA-R superfamily member is listed in Table 1.

An evolutionary tree illustrates how D/D domains evolved across major eukaryotic clades (Fig. 2B). Metazoans lost the D/D domain on enolase despite the expansion of the domain to other proteins (Fig. 2, A and B; Fig. S1). Dendrograms displaying the phylogenetic topology of the D/D superfamily were generated using the RAxML and IQ-tree platforms. Virtually identical branch alignments were obtained on both platforms (Fig. 2C; Fig. S1). Interestingly, organisms which do not rely on flagella for reproduction experienced an evolutionary loss of PKA-R-like D/D proteins. For example, gymnosperms and
Defining the PKA docking and dimerization domain superfamily

Table 1

Human D/D domain of PKA-R superfamily members

| PKA-R DD superfamily | Chr | AA | Mouse KO phenotype |
|----------------------|-----|----|-------------------|
| Ropporin-1 (ROPN1)   | 3   | 73 | Sutherland        |
| Ropporin-1-Like (ROPNL1) | 5 | 73 | Ciliary dysmotility, Sutherland |
| Ropporin-1B (ROPN1B) | 3   | 73 | Not applicable    |
| Ca²⁺-binding Tyr-phosphorylation regulated (CABYR) | 18 | 75 | Fibrous sheath dysplasia, Sutherland |
| Sperm autoantigenic protein 17 (SPA17) | 11 | 75 | Not available |
| PKA type II regulatory subunit α (PKA-RII) | 3  | 45 | Reduced interaction with AKAPs |
| PKA type II regulatory subunit β (PKA-RII) | 7  | 45 | Body weight/fat (RIIβ) |
| Ciliogenesis-associated TTC17-interacting protein (CATIP) | 2  | 60 | Sutherland |
| RIα domain containing protein 1 (RIIAD1) | 1  | 85 | Absent whiskers, abnormal body wall, neonatal lethality |
| Tubulin polyglutamylation subunit 1 (TPGS1) | 19 | 45 | Sutherland |
| PKA type I regulatory subunit α (PKA-RI) | 17 | 50 | Carney complex (RIα) |
| PKA type I regulatory subunit β (PKA-RI) | 7  | 50 | LTD and LTP (RIβ) |
| Vestibulin-1 (VEST1) | 8  | 60 | Not available |
| Adenylate kinase 8 (AK8) | 9  | 60 | Hydrocephaly |
| Adenylate kinase 5 (AK5) | 1  | 65 | Not available |
| F-Box and Leu-rich repeat protein 13 (FBXL13) | 7  | 70 | Abnormal eye interior |
| Testis expressed 55 (TEX55) | 3  | 37 | Chamber depth |
| EF-Hand Ca²⁺ binding | 7  | 60 | Not available |
| Protein 10 (EFCAB10) |  |  | |

The gene name of each PKA-R superfamily member as listed in Figure 1C. The chromosomal location (Chr) and number of amino acids (AA) are indicated. Putative functions of each family member are inferred by listing the mouse KO phenotype obtained from the Mouse Genomics Data consortium. Not available denotes that a KO mouse has not been generated.

Abbreviations: LTP, long-term potentiation; LTD, long-term depression.

angiosperms use pollen to produce fertile seeds and do not have R2D2 proteins, but mosses and ferns which utilize sperm have R2D2 proteins (Fig. S2).

As previously mentioned, the Pfam algorithm assigns the DPY-30 and related proteins to the RIα clan. Our analyses designate the DPY-30 and related proteins as an outgroup that is equally related to RI and RII (Figs. 1C and 2C). The crystal structure of DPY-30 reveals a D/D fold similar to PKA-R domains (16, 34). Accordingly, the DPY-30 structure is superimposable over RII with an RMSD of 2.6 Å (Fig. S3A). Further delineation between DPY-30 and RII is evident from probabilistic hidden Markov modeling (Fig. S3B). This algorithm predicts that evolutionary changes have occurred at different positions within the D/D domains of both superfamilies.

Sequence determinants that delineate the RI and RII families predominantly lie in the amino-terminal flanking region and in the loop between helix I and helix II of the D/D domain (Fig. 1A). Phylogenetic and topological analyses have used this information to subdivide the PKA-R superfamily D/D domains into two distinct but overlapping groups (Fig. 1, B and C). A hallmark of RI subunits is the presence of two prolines on each end of the loop between helix I and helix II. Hence, enolase, EFCAB10, AK5, AK8, VEST1, FBXL13, and TEX55 are prototypic of the R1D2 clade (Fig. 1C, gold underlined). In contrast, a defining feature of the R2D2 clade is replacement of the second proline with a hydrophobic side chain (Ile, Leu, or Val, Fig. 1C, blue underlined). Five proteins, ROPN1, ROPNL1, SPA17, RIIAD1, and CATIP, follow this convention (Figs. 1C and 2C; Fig. S1).

Other determinants also contribute to the R1D2 or R2D2 designations. For example, TPGS1 is considered an R1D2 protein because it contains a predicted helical flanking amino-terminal motif and a second proline in the loop region. Yet, TPGS1 can also be considered an R2D2 protein because of features such as a glycine at the start of helix I and a conserved “YF” motif in helix II (Figs. 1C and 2C). Likewise, RIIAD1 and CATIP are intermediate to the R2D2 clade because they are predicted to have an amino-terminal helix rather than a β strand (Fig. 1C). Thus, our phylogenetic analyses have defined primary, secondary, and tertiary structure characteristics that are emblematic of the DPY-30, R1D2, and R2D2 subgroups of the RIα clan. All data have been deposited in the Dryad server.

Structure of the SPA17 D/D domain

We chose to focus our structural analyses on SPA17 because of its extended D/D domain. The zebrafish ortholog is 72% identical to the human ortholog and proved amenable to crystallization in multispecies trials (Fig. 3A). A construct spanning amino acids 1 to 75 of SPA17 from D. rerio was expressed in Escherichia coli, and the resultant protein was purified with a sequential three-step affinity, anion-exchange, and size-exclusion chromatography (SEC) protocol (Fig. S4). Crystals of SPA17 diffracted X-ray to 1.72 Å. The structure was determined by molecular replacement using the D/D domain of PKA-RIα as a search model (PDB ID: 2IZX) and subsequently refined to an R_work of 0.154 and R_free of 0.165 (Fig. 3B and Table 2).

Four copies of SPA17 are observed in the asymmetric unit of the crystal. The two central copies of SPA17 form the canonical four-helix bundle as previously observed in the homodimeric D/D domains of R1 and RII (Figs. 1A and 3B). The other two copies each form a similar homodimer with a symmetry related SPA17 chain. Interestingly, the two conserved sequence regions flanking the central helices of SPA17 both adopt a regular β-strand conformation. A four-stranded β-sheet is formed from the amino-terminal β-strand of two SPA17 molecules and the carboxyl-terminal β-strands of two other SPA17 chains (Fig. 3B). Because of the close involvement of these β-strands in crystal packing, the
Defining the PKA docking and dimerization domain superfamily

A. The table depicting evolutionary conservation of the PKA-R D/D domain superfamily. Each filled dot represents the presence of the D/D domain protein identified on the left in the genus listed on the top.

B. Evolutionary tree illustrates the prevalence of D/D domains across all major eukaryotic clades.

C. Dendrogram displaying the phylogenetic topology of the PKA-R D/D superfamily. 249 orthologs were used to build an estimated phylogeny by maximum likelihood with the RAxML algorithm. The dendrogram was drawn using MEGA-X and represents 2000 combined pseudoreplicates from runs with and without DPY-30 as a designated outgroup. Branch support values are indicated. D/D, docking and dimerization; PKA-R, PKA regulatory.

Figure 2. Taxonomic distribution and phylogenetic topology of the PKA-R D/D domain superfamily. A, the table depicting evolutionary conservation of the PKA-R D/D domain superfamily. Each filled dot represents the presence of the D/D domain protein identified on the left in the genus listed on the top. B, evolutionary tree illustrates the prevalence of D/D domains across all major eukaryotic clades. C, dendrogram displaying the phylogenetic topology of the PKA-R D/D superfamily. 249 orthologs were used to build an estimated phylogeny by maximum likelihood with the RAxML algorithm. The dendrogram was drawn using MEGA-X and represents 2000 combined pseudoreplicates from runs with and without DPY-30 as a designated outgroup. Branch support values are indicated. D/D, docking and dimerization; PKA-R, PKA regulatory.
Figure 3. The structure of SPA17 (1–75). A, sequence alignment of human (Hs); mouse (Mm); zebrafish (Dr), and Trichoplax adhaerens (Ta) SPA17 orthologs. Residues 1 to 75 are presented. Black arrows indicate conserved hydrophobic at positions flanking the helical sections. B, the first 75 amino acids of SPA17 from Danio rerio resolved in a crystal structure at 1.72 Å. Side view, four copies of SPA17 (indicated in green, yellow, purple, and magenta) are observed in the asymmetric unit of the crystal. Top view, extended regions observed in the sequence alignment appear to occlude the AKAP-binding site and facilitate tetramerization. C and D, the stoichiometry of homomeric SPA17 complexes was calculated by size-exclusion chromatography coupled to multiangle light scattering (SEC-MALS). SDS gels denote protein purity. Molecular weight markers are indicated. C, SPA17 1 to 75 with a predicted molecular weight (MW) of 8.8 kDa forms tetramers (35 kDa) and dimers (17 kDa). D, full-length SPA17 with a predicted MW of 17.4 kDa forms tetramers (72 kDa) and dimers (36 kDa). Representative gels from three experiments. AKAP, A-kinase-anchoring protein; SPA17, sperm autoantigenic protein 17.
Defining the PKA docking and dimerization domain superfamily

SPA17 coexists with ropporin-1-like proteins in the flagellum of mammalian sperm and motile cilia (31). Sequence similarities between these members of the R2D2 clade raised the possibility that SPA17 and its close relative ROPN1L may form heterodimers (Fig. 1C). In keeping with this notion, full-length SPA17 and ROPN1L comigrate as assessed by SEC-MALS analysis (Fig. 5A). Likewise, SPA17 1 to 75 and ROPN1L 1 to 75 multimerize when analyzed by SEC-MALS (Fig. 5B). Gel filtration traces further indicate that SPA17–ROPN1L complexes migrate with an apparent molecular weight that is consistent with a heterodimer with a minor tetrameric species (Fig. 5, A and B). Protein pulldowns verified interaction between glutathione-S-transferase (GST)-ROPN1L and SPA17 (Fig. 5C). Reciprocal pull-down experiments confirmed that GST-SPA17 binds ROPN1L (Fig. 5D). Thus, SPA17 has the capacity to form heterodimers with ROPN1L.

Homeric SPA17 exhibits limited interaction with AKAPs (31). However, AKAP interaction is more evident in cells that co-express SPA17 and ROPN1L (31). Therefore, it was imperative that we investigated the AKAP-binding properties of SPA17–ROPN1L heterodimers. SEC-MALS was used to evaluate full-length SPA17 and ROPN1L interaction with a GST-AKAP18 fusion (35). In solution, the AKAP18–SPA17–ROPN1L ternary complex elutes as a single peak at 10.4 ml that corresponds to a molecular weight ~330 kDa (Fig. 6: lanes 1 and Fig. 6B). This is equivalent to a tetrameric GST-AKAP18–SPA17–ROPN1L assembly. In contrast, SPA17–ROPN1L heterodimers (Fig. 6C) and GST-AKAP18 (Fig. 6D) elute with lower apparent molecular weights on the analytical size-exclusion column (Fig. 6A: lanes 2 and 3).

To quantify the AKAP18 interaction with SPA17 and ROPN1L, biolayer interferometry analysis was performed on the Octet system. A probe with anti-GST antibodies was loaded with GST-AKAP18, washed, and then incubated with either SPA17 alone or SPA17–ROPN1L over a range of concentrations. The steady state affinity is calculated as Response = (Rmax × concentration)÷(KD + concentration). Homeric SPA17 bound AKAP18 with an affinity of 9.6 ± 0.69 μM (Fig. 6F), supporting the notion that its AKAP-binding site is not blocked by the terminal β strands. Remarkably, the SPA17–ROPN1L heterodimer binds AKAP18 with an affinity of 82 ± 2.1 nM (Fig. 6G). For comparison, the KD for RIIα interaction with AKAP18 is 31 nM (25). Hence, SPA17–ROPN1L heterodimers exhibit a physiologically relevant binding affinity for AKAP18 and are likely to represent the preferred AKAP-binding module.

To further probe this phenomenon, we conducted in silico modeling of the AKAP18–SPA17–ROPN1L trimer (Fig. 5, A and B). Sequence alignments indicate that the AKAP18-anchoring helix is palindromic except for the central two residues (Fig. 5, C and D). Hence, the AKAP helix can be presented in two possible orientations (Fig. 5, A and B). Several factors could favor one of the orientations and lead to an enhanced affinity in comparison with the SPA17 homodimer. These include the asymmetrical nature of the SPA17–ROPN1L heterodimer.

### Table 2

| Property                              | Value                      |
|---------------------------------------|----------------------------|
| Space group                          | P 32                       |
| Cell constants a, b, c, α, β, γ       | 60.96 Å 60.96 Å 89.02 Å    |
| Resolution (Å)                       | 34.03–1.72                 |
| % Data completeness (in resolution range) | 98.5 (45.41–1.72)  |
| Wavelength                           | 0.99996 Å                  |
| Rmee                                 | 0.098                      |
| Rpm                                  | 0.035                      |
| Rs (1.72–1.75 Å)                     | 0.91 (0.269)               |
| Data redundancy (1.72–1.75 Å)        | 7.6 (5.6)                  |
| CC1/2                                | 0.993                      |
| R<sub>f</sub>(1)                     | 1.02 (at 1.72 Å)           |
| Refined program                      | phenix.refine             |
|                                     | 1.18.2 3874, PHENIX        |
|                                     | 1.18.2 3874                |
|                                     | 1.05, 0.165                |
|                                     | 0.154, 0.166               |
| Anisotropy                           | 100%                       |
| Bulk solvent ksoft/e(Å<sup>3</sup>)  | 0.42, 34.0                 |
| L-test for twinning                  | < |L| > = 0.51, < L<sup>2</sup> > = 0.34 |
| Estimated twinning fraction          | 0.479 for -h-k,l           |
|                                     | 0.480 for h,-k,-l          |
|                                     | 0.479 for -k,h,-l          |
| F0,Fc correlation                    | 0.96                       |
| Average B, all atoms (Å<sup>2</sup>)| 4799 (17.0)               |

formation of the four-stranded β-sheet is likely a crystallization artifact. Nonetheless, these β-strands might mediate SPA17 oligomerization, as biochemical studies indicate that SPA17 can exist in higher order configurations (Fig. 3, C and D). SEC coupled to multiangle light scattering (SEC-MALS) verifies that SPA17 tetramers and dimers exist in solution (Fig. 3, C and D). The SPA17 1 to 75 fragment (8.8-kDa monomer) has molecular masses of 35 and 17 kDa (Fig. 3C). Although less evident, multimerization of full-length SPA17 (17.4-kDa monomer) was also observed. The purified protein ensemble elutes with molecular masses of 72 and 36 kDa (Fig. 3D).

Collectively, the data in Figure 3 imply that, unlike RIIα, SPA17 can form higher order homo-oligomeric complexes.

Although SPA17 exhibits distinctive structural features, it still retains many hallmarks of a canonical R2D2 protein. Alignment of the SPA17 structure to the D/D domain of apo RIIα dimers results in an associated RMSD of 0.482 Å. Similarly, the apo structure of SPA17 superimposes over RIIα in complex with AKAP-in silico with an RMSD of 0.514 Å (Fig. 4A, left panel). The AKAP-binding site is, therefore, retained in the SPA17 homodimer, although it appears to be occluded by the β-sheet formed among the terminal strands of SPA17 protomers in the crystal (Fig. 4A, right panel). As expected, key hydrophobic residues necessary for dimerization (magenta squares) are strictly conserved, but only five of the docking determinants (purple dots) are invariant in the sequence alignment (Fig. 4B). Together, these features suggest that the extended D/D domain of SPA17 contains most necessary determinants for binding to AKAPs. Thus, the mode of SPA17 interaction with AKAPs might be slightly different than how RII interfaces with its anchoring proteins.
complex, the distinct pI values of both protomers and differential exposure of core DD domain residues. In addition, the unique N-terminal flanking sequences on SPA17 and ROPN1L might participate in customized protein–protein interactions that enhance affinity for the AKAP18 helix only in the context of the heterodimer.

Discussion

Our studies shed new light on the evolution of D/D domains. These regions were originally characterized as amino-terminal elements of the regulatory subunits of PKA that form homodimers to create an AKAP-binding surface (13, 16). Our subsequent identification of 18 family members presented in Figures 1 and 2 suggests that these four-helix bundle-like–forming proteins have a broader role in shaping subcellular architecture than previously appreciated. Another interesting outcome of our phylogenetic analysis has been the segregation of the D/D domain into the DPY-30 and the PKA-R superfamilies. DPY-30 proteins are universally found in all phyla and may be more closely related to the primordial D/D domain. In contrast, there are signature motifs within the PKA-R superfamily that subdivide this group into the R1D2 and R2D2 families. Each D/D domain class exhibits distinct features that contribute to their specialized roles in the coordination of organellar and subnuclear events.

R1D2 proteins are typified by a flanking N-terminal helix containing consecutive aromatic and hydrophobic amino acids and a loop with a PxxP motif. Conversely, R2D2 proteins have a β strand and a Pxx[L,I,V] motif in the corresponding regions. Our phylogenetic analyses in Figures 1C and 2A indicate that there was an expansion in both classes around the time of metazoan evolution. In contrast, fungi and plants have fewer D/D proteins. For example, BCY1, the PKA-R subunit gene in the budding yeast Saccharomyces cerevisiae contains a recognizable RII-like D/D domain that also forms crystallization-related oligomers (36–40). Structural evidence for the oligomerization of BCY1 calls into question the ability of this D/D domain to dock with fungal proteins (36). Parenthetically, there is scant evidence for AKAPs in the fungi kingdom (2–4). This may be because kinase anchoring is not a preferred mechanism for local signaling in these less morphologically developed organisms, and subcellular organelles including cilia are not prominent in these kingdoms. Taken together, this data mining venture indicates that metazoans express the most comprehensive set of D/D domain proteins, thereby lending further credence to the notion that compartmentalized signaling expanded at the base of animal evolution.

DPY-30 proteins are an outgroup to the D/D family that are present in all animal and plant phyla (41). The founding member DPY-30 encodes the core subunit of the SET1/MLL family of COMPASS histone methyltransferases (42). DPY-30 homodimers associate with a helical region on ASH2L, another core element of the COMPASS complex to stabilize intrinsic methyltransferase activity (43, 44). This protein–protein interaction bears a striking resemblance to the RI–AKAP and RII–AKAP interfaces as depicted in Fig. S6, A and B. In agreement with this concept, the DPY-30 four-helix bundle retains the capacity to interact with AKAPs and is thought to be a native ligand for AKAP95. This affords a means to incorporate AKAP95 into histone methyltransferase complexes (45). Previous studies have shown that AKAP95 is a predominantly nuclear protein that has the capacity to interact...
Defining the PKA docking and dimerization domain superfamily

Figure 5. SPA17 heterodimerizes with ROPN1L. A, size-exclusion chromatography evaluated complex formation between SPA17 and ROPN1L. Left panel, Coomassie-stained SDS gel of peak fractions (indicated below each lane). Right trace, SPA17–ROPN1L multimerization assayed by SEC. Apparent molecular weights of heterotetramer and heterodimer are indicated. B, size-exclusion chromatography coupled to multiangle light scattering (SEC-MALS) evaluated SPA17 1 to 75 complex formation with ROPN1L 1 to 75. Left panel, Coomassie-stained SDS gel of peak SPA17 1 to 75–ROPN1L 1 to 75 fraction. Right trace, SPA17: ROPN1L multimerization assayed by SEC-MALS. Apparent molecular weights of heterotetramer and heterodimer are indicated. C and D, reciprocal GST pull-down experiments demonstrate interaction between GST-ROPN1L (pink) and SPA17 (purple) and SPA17 1 to 75 (lavender). C, Coomassie-stained SDS gel shows GST-ROPN1L (lane 1). Pull down of full-length SPA17 (lane 2), human SPA17 1 to 75 (lane 3), zebrafish SPA17 1 to 75 (lane 4), and beads only (lane 5). D, Coomassie-stained SDS gel of reciprocal GST-pull-down binding experiments. Protein GST-SPA17 (lane 1). ROPN1L binding to GST-SPA17 (lane 2). Molecular weight markers are indicated. Representative gels of three experiments. SPA17, sperm autoantigenic protein 17.

with PKA but only when the nuclear envelope is dissolved during mitosis (46, 47). Nonetheless, AKAP95–PKA holoenzyme complexes may participate in different signaling events during cell division, as a significant proportion of the anchoring protein remains associated with DYP-30 during mitosis (45, 48). Thus, differential association with RII or DYP-30 may determine whether AKAP95 functions as a kinase-anchoring protein or an ancillary component of histone methyltransferases. The latter function may also be relevant to AKAP95 nuclear role in interphase cells as this anchoring protein has been implicated as a positive regulator of pre-mRNA splicing and gene expression during tumorigenesis (49, 50). Overall, these observations imply that subtle but conserved differences in the D/D domain not only influence...
Figure 6. The SPA17–ROPN1L heterodimer is the functional AKAP-binding module. A, Coomassie-stained gel showing the peak fractions from the gel filtration experiment to the right. Electrophoretic mobility of GST-AKAP18 (gray), ROPN1L (magenta), and SPA17 (purple) and molecular weight markers are indicated. B, GST–AKAP18–SPA17–ROPN1L ternary complex (lane 1 in panel A) elutes in a single peak in 10.4 ml. C, SPA17–ROPN1L heterodimer (lane 2 in panel A) elutes in 13.1 ml. D, GST-AKAP18 (lane 3 in panel A) elutes in 13.4 ml. E, diagram of the Octet BLI system. In the first step, a probe with anti-GST antibodies is washed with a buffer. GST protein is applied to the probe, and the detector shows a shift in the refractive index corresponding to increased

- **Baseline Loading**
- **Association**
- **Dissociation**

F, Ligand: GST-AKAP18

- **Analyte:** SPA17

G, Analyte: R2D2 Heterodimer

- **Steady State**

R² = 0.9993
R_max = 0.38 +/- 0.002
K_D = 82nM +/- 2.1nM

R² = 0.9978
R_max = 0.38 +/- 0.002
K_D = 9.6μM +/- 0.69μM.
Defining the PKA docking and dimerization domain superfamily

which binding partners this protein module interacts with but
also have marked effects on the differential functionality of the
resulting macromolecular complexes.

Although SPA17 and RIIa are homologous, our sequence
analysis identified unique residues in the helical flanking re-

gions of SPA17. For this reason, we determined the crystal
structure of the extended D/D domain of SPA17. Interestingly,
sequences at either end of the helical segments form a four-
stranded β sheet that occludes the AKAP interface. Although
AKAP-binding determinants are conserved in the core of
SPA17, our in vitro–binding studies presented in Figure 6
reveal that the anchoring protein AKAP18 has a surprisingly
low affinity toward this homomeric D/D domain. Although
interference from flanking regions could explain poor AKAP
binding, it is currently unclear whether or not this molecular
mechanism is operational in the context of the native protein.
Likewise, participation of these flanking β strands in the crystal
packing that lead to the formation of a tetramers could be
construed as a consequence of the unnatural physiochemical
conditions imposed by protein crystallization. Yet, our SEC
and MALS experiments presented in Figure 3, C and D indi-
cate that soluble SPA17 exists in both dimeric and tetrameric
configurations. Thus, an equally plausible explanation is that
the conserved flanking regions facilitate supplementary oligo-
merization of SPA17.

Analogous flanking regions are detected in other R2D2
family members. In addition, proteomic screens often report
that clusters of D/D domain proteins exist within the same
macromolecular complexes (51). This raises the intriguing
possibility that certain D/D domain proteins have the capacity
to heterodimerize. Consistent with this notion, our biochem-
ical studies presented in Figures 5 and 6 show that SPA17 can
dimerize with the ROPN1L. This latter D/D protein also fea-
tures extended flanking regions that are highly conserved in all
orthologs. A particularly fascinating aspect of this observation
is that SPA17–ROPN1L heterodimers display a higher affinity
for AKAPs as assessed by our quantitative binding measure-
ments presented in Figure 6G. Heterodimerization may be a
result of the complementary isoelectric point values of
ROPN1L and SPA17 N-terminal domains as indicated in
Fig. S5E. The increased affinity for AKAP binding is likely due
to central asymmetric residues in the context of a largely
palindromic AKAP helix binding to distinct residues exposed
within the binding groove formed by the SPA17–ROPN1L
heterodimer. Indirect support for this concept is provided by
analysis of the closely related R2D2 proteins, RSP7 and RSP11
(52). In this context, the N-terminal flanking regions are
observed to form interactions with distal portions of amphi-
pathic helix on their binding partner RSP3 (as depicted in
Fig. S5, A and B). Thus, the formation of mixed four-helix
bundles could expose additional AKAP-binding determinants
that are masked in homomeric conformations. Alternatively,
flanking regions of the SPA17–ROPN1L heterodimers might
not be able to interfere with the AKAP interface. Irrespective
of which explanation is correct, our discovery of D/D domain
heterodimers expands the repertoire of D/D modules that may
be operational within the intracellular environments.

A recurring theme throughout this study is evidence that D/
D domain proteins participate in the organization and struc-
tural integrity of flagella and motile cilia (51). The rhythmic
beating of these motile appendages enhances microorganism
motility and the propulsion of sperm. Interestingly, the sperm
fibrous sheath and ciliary radial spoke complexes are organized
by AKAPs and D/D proteins (53, 54). Moreover, our explo-
dation of D/D heterodimerization presented in Figures 5 and 6
is reminiscent of the recent cryo-EM analyses of the axonemal
radial spoke complex (52, 55). The radial spoke is a complex
of 23 proteins that functions as a mechanochemical transducer
that modulates activity of dynein motors to promote flagellar
motility. Eukaryotic flagella and motile cilia share a common
“9 + 2” structure, in which nine peripheral microtubule dou-
blets surround the central pair of microtubules. These sub-
structures are connected by radial spokes. The radial spoke is a
T-shaped macromolecular assembly that anchors peripheral
microtubules to the central pair. The stalk of the radial spoke
is organized by an anchoring protein called RSP3 that co-
dinates microtubule sliding (52). The radial spoke was
originally thought to contain PKA by virtue of evidence that
RII binds RSP3 in a far-Western overlay assay (56, 57).
However, proteomic screens have never detected PKA as a
component of this complex (53). Likewise, the Chlamydo-
monas reinhardtii protein RSP3, which originally was designated
as an AKAP, is now recognized to anchor RSP7–RSP11 het-
erodimers. This heterodimeric complex of D/D proteins might
provide rigidity to this flagellar substructure (52). The archi-
tectural integrity imparted by RSP7–RSP11 heterodimers in-
volves two classes of protein–protein interaction. The D/D
domain tethers to RSP3, whereas binding motifs in each pro-
tomer cross-link with other elements of the radial spoke.
Interestingly, algal RSP11 is an ortholog of metazoan ROPN1L.
Likewise, RSP7 and SPA17 share a conserved extended D/D
domain and related calcium-binding motifs. In total, these
observations infer that this D/D–AKAP interface is an
adaptable architectural element, rather than just a platform for
kinase signaling.

In conclusion, this study highlights the discovery and clas-
sification of interactive motifs patterned after the D/D do-
main of the PKA-R subunits. We show that this emergent
protein module functions as an AKAP-interaction domain.
Undoubtedly, the R1D2 and R2D2 proteins described herein
create platforms for intracellular signaling. However, these D/
D domains may equally serve as architectural components of

mass on the surface of the probe. Excess protein is washed away with the buffer to record a baseline. Finally, the ligand of interest is loaded and washed to calculate the binding kinetics. Each color track represents an increasing concentration of the ligand (as indicated). F, SPA17 alone (top subpanel) has micromolar affinity for AKAP18. G, SPA17 1 to 75–ROPN1L 1 to 75 heterodimer has nanomolar affinity for AKAP18. F and G, left subpanels show raw data with fitted curves to calculate $k_\text{on}$ and $k_{off}$. Right subpanel graphs are calculated as follows: $R = (R_{\text{max}} \times \text{concentration})+(K_\text{D} + \text{concentration})$. Amalgamated data from five experiments derived dissociation constants. AKAP, A-kinase-anchoring protein; BLI, bilayer interferometry analysis; SPA17, sperm autoantigenic protein 17.
motile cilia and integration points for processing cellular cues including calcium and histone methylation. Moreover, cross-member heterodimerization may represent a previously unconsidered mechanism to expand the repertoire and functionality of D/D domains well beyond the compartmentalization of PKA. Future studies are necessary to elucidate roles of the RIIα clan as multipurpose adapters that are utilized in expanding number of molecular and cellular contexts.

**Experimental procedures**

*Phylogenetic analysis*

A combined approach was used to inventory PKA-R-like D/D domain–containing proteins. In an attempt to identify all putative R1D2 and R2D2 proteins, the NCBI Conserved Domain Database was searched for proteins within the “D/D-R-PKA: Dimerization/Docking domain of the Regulatory subunit of cAMP-dependent kinase and similar domains” superfamily and the superfamily Hidden Markov models, and genome assignment library was searched for proteins within the expanding number of molecular and cellular contexts.

**Molecular biology and protein purification**

Human ROPN1L, ROPN1L (1–75), SPA17, SPA17 NTD (1–75), SPA17 NTD (1–80), AKAP18α, and zebrafish SPA17 (1–75) were cloned into pGEX-6P-1 using the BamHI and XhoI restriction sites. These plasmids were then transformed into BL21 cells from Invitrogen for protein expression. Induction was performed at 16 °C overnight using 1 mM IPTG starting at an absorbance at 600 nm of about 0.8. Cells were lysed in a buffer containing 50 mM Tris, pH 8, 200 mM NaCl, and 5 mM DTT. Lysates were clarified via centrifugation at 40,000g. The clarified lysate was first purified using a glutathione affinity column (GE Healthcare) pre-equilibrated in the lysis buffer. Protein was eluted from the column by overnight on-column cleavage of the GST tag by 3C-HRV protease produced in house at 4 °C. Eluted material was further purified using the AKTA system with a 5-ml HiTrap Q-HP column (GE Healthcare), followed by a Superdex 200 gel filtration column (GE Healthcare) in a final buffer containing 10 mM Tris, pH 8, 200 mM NaCl, 5 mM DTT (Fig. S5). Proteins were concentrated to 15 to 20 mg/ml, flash-frozen in liquid nitrogen, and stored in −80 °C.

**Crystallization**

Extensive crystallization screening was conducted to obtain high-quality crystals. Purified *D. rerio* SPA17 NTD (1–75) protein was mixed with mother liquor in a 1:1 ratio using 150 nl of each component and spotted onto hanging drop seals on a Mosquito nanodrop robot. The crystallization trays were then placed at 4 °C. The crystals used for analysis grew in 0.1 M Hepes, pH 7.5, and 0.5 M magnesium formate dihydrate from the Index Screen by Hampton Research. Crystals appeared after 2 to 3 days and reached their final size in 1 to 2 weeks. Crystals were directly transferred to a cryoprotectant buffer containing 50% glycerol and 50% mother liquor and frozen in liquid nitrogen for synchrotron data collection.

**X-ray data collection and structure determination**

X-ray diffraction data were collected at the Advanced Light Source at Lawrence Berkeley National Laboratory. Diffraction data were indexed, integrated, and scaled with the HKL-2000 package (67). Resolution cutoffs were determined using completeness (>80%) and I/σ >1 as primary criteria. Initial phases were determined by the molecular replacement method using PKA RIα (PDB ID: 2IZX) as a search model. The

XSEDE with 1000 replicate trees for calculation of transfer bootstrap expectation support metrics. This analysis was conducted twice, once with DPY-30 as an outgroup and once without an outgroup. The resulting dendrograms were identical, so the branch support values were listed for 2000 replicate trees. A second maximum likelihood analysis was conducted for comparison using IQ-Tree (65, 66) on XSEDE with 2000 replicates trees for calculation of UFBoot support metrics. Dendrograms were drawn and visualized in MEGA-X.
Defining the PKA docking and dimerization domain superfamily

majority of the model was subsequently build using the autobook function in PHENIX (68). Minor rebuilding was done manually in Coot (69). Refinement was also conducted in PHENIX, and the final model had an Rwork/Rfree of 0.154/0.165. All structural figures used for data analysis were visualized in PyMOL.

Protein pull-down

GST-AKAP18α and GST-RIIα were transformed into BL21 cells and purified using glutathione-affinity purification and anion-exchange chromatography. About 0.3 mg of purified GST proteins, GST-AKAP18α, GST-SPA17, GST-ROPN1L, and GST-RIIα was loaded onto a GST purification column (GE Healthcare) in a bed volume of 200 μl. Solutions containing 0.3 mg of SPA17 NTD, SPA17, or ROPN1L were then applied to the column, which was washed three times with ten column volumes of the lysis buffer (50 mM Tris, pH 8, 200 mM NaCl, 5 mM DTT). Samples were eluted with 10 mM glutathione in the lysis buffer, mixed with SDS-PAGE loading buffer, analyzed by SDS-PAGE, and visualized by Coomassie staining.

Protein quantitation

Protein quantitation was conducted using a NanoDrop with A280 extinction coefficients calculated for each protein using ExPASy ProtParam.

Data availability

All data are contained within the article. Coordinates for the crystal structure of SPA17 Docking and Dimerization Domain from Danio rerio have been deposited in the PDB database. Accession Number: PDB ID: 7MY4.

Supporting information—This article contains supporting information.

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Abbreviations—The abbreviations used are: AKAPs, A-kinase-anchoring proteins; D/D, docking and dimerization; DPY-30, Dumpy-30; MALS, multiangle light scattering; PKA-R, PKA regulatory; SEC, size-exclusion chromatography; SEC-MALS, size-exclusion chromatography coupled to multiangle light scattering; SPA17, sperm autoantigenic protein 17.

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