Differential Recognition Preferences of the Three Src Homology 3 (SH3) Domains from the Adaptor CD2-associated Protein (CD2AP) and Direct Association with Ras and Rab Interactor 3 (RIN3)*

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Background: The CD2AP adaptor facilitates cell signaling using its in-built interaction modules (SH3 domains).
Results: RIN3 was characterized as a novel CD2AP SH3 binding protein by biophysical and biochemical methods.
Conclusion: CD2AP has many potential interactors and is probably a cellular hub.
Significance: We reveal how protein modules can interact with families of related recognition motifs rather than a single motif.

CD2AP is an adaptor protein involved in membrane trafficking, with essential roles in maintaining podocyte function within the kidney glomerulus. CD2AP contains three Src homology 3 (SH3) domains that mediate multiple protein-protein interactions. However, a detailed comparison of the molecular binding preferences of each SH3 remained unexplored, as well as the discovery of novel interactors. Thus, we studied the binding properties of each SH3 domain to the known interactor Casi tas B-lineage lymphoma protein (c-CBL), conducted a peptide array scan based on the recognition motif PxPxPR and identified 40 known or novel candidate binding proteins, such as RIN3, a RAB5-activating guanine nucleotide exchange factor.

CD2AP SH3 domains 1 and 2 generally bound with similar characteristics and specificities, whereas the SH3-3 domain bound more weakly to most peptide ligands tested yet recognized an unusually extended sequence in ALG-2-interacting protein X (ALIX). RIN3 peptide scanning arrays revealed two CD2AP binding sites, recognized by all three SH3 domains, but SH3-3 appeared non-functional in precipitation experiments. RIN3 recruited CD2AP to RAB5a-positive early endosomes via these interaction sites. Permutation arrays and isothermal titration calorimetry data showed that the preferred binding motif is Px(P/A)xPR. Two high-resolution crystal structures (1.65 and 1.11 Å) of CD2AP SH3-1 and SH3-2 solved in complex with RIN3 epitopes 1 and 2, respectively, indicated that another extended motif is relevant in epitope 2. In conclusion, we have discovered novel interaction candidates for CD2AP and characterized subtle yet significant differences in the recognition preferences of its three SH3 domains for c-CBL, ALIX, and RIN3.

CD2-associated protein (CD2AP)8, or CMS (Cas ligand with multiple Src homology 3 domains) is a multifunctional adaptor protein of 71 kDa, found ubiquitously in the body. First characterized as interacting with the T-cell surface antigen, CD2 (1), and the docking protein p130Cas (also known as BCAR1) (2), CD2AP is an essential protein for the proper functioning of the kidney glomerulus, playing a particularly important role in the
CD2AP SH3 Domain Recognition Preferences

development of the slit diaphragm. Homozygous CD2AP−/− mice develop progressive glomerular dysfunction caused by the loss of podocyte foot process integrity and die of renal failure within 6–7 weeks (3). Mutations identified in CD2AP (single nucleotide polymorphisms or deletions leading to protein sequence changes) result in abnormal kidney function and proteinuria with nephrotic syndrome caused by focal segmental glomerulosclerosis (4, 5).

Apart from podocytes, CD2AP has been studied in T-cells and dendritic cells among others. CD2AP plays an integral role in actin dynamics, endocytosis of cell surface receptors, and their degradation (e.g. binding to F-actin and promoting actin bundling (6); promoting internalization of EGF receptor (7) and VEGF receptor (8); and down-regulation of the T-cell receptor (9), the HER2 (ErbB2) receptor in a subset of breast cancers (10), and the RET (rearranged during transfection) receptor (11, 12)). Conversely, CD2AP positively regulates the plasma-cytoid dendritic cell receptor BDCA2; its degradation occurs only with a lack of CD2AP activity (13).

CD2AP can facilitate branched actin filament formation by direct interactions with actin capping protein and cortactin, giving rise to lamellipodia (14). A more direct activity in actin barbed-end capping at adherens junctions was recently reported (15). Various other CD2AP-interacting proteins have been identified, many of which are involved in vesicle trafficking and endocytosis, including the ubiquitin ligases c-CBL (herein termed CBL) (16) and CBL-b (17). Increasingly, endocytosis pathway genes (e.g. BIN1/amphiphysin II) are implicated in causal mechanisms underpinning late onset Alzheimer disease (LOAD) (18). Recently, CD2AP was strongly linked to LOAD through genome-wide association studies (19, 20).

CD2AP and its homologue Cbl-interacting protein of 85 kDa (CIN85) are composed of three Src homology 3 (SH3) domains in the N-terminal half followed by a proline-rich region (PRR) and a coiled-coil domain at the C terminus, interspersed with apparently intrinsically disordered regions. CD2AP differs from CIN85 by the presence of additional actin binding sites before the coiled-coil domain (Fig. 1A). SH3 domains are globular protein interaction modules, typically binding to proline-rich linear recognition motifs. A large subset of CD2AP interactions involves one or more of its SH3 domains. However, the protein sequences of human CD2AP SH3 domains are only moderately conserved: SH3-2 and SH3-3 share the highest pairwise global sequence identity of only 45%; SH3-1 and SH3-2 have 40% identity; and SH3-1 and SH3-3 have 38% (Fig. 1B).

Different biological processes regulated by CD2AP require specific binding partners, illustrating the versatility of CD2AP as an adaptor and hinting that each SH3 may have unique properties and requirements for binding. The key to understanding how specificity and selectivity are achieved to regulate multiple, heterogeneous binding events thus lies in combined knowledge of each molecular interaction. Most in vitro studies on the relationships between CD2AP SH3 domains and protein partners have tended to focus on a single domain or partner. None have systematically compared the binding modes of all three SH3 domains in comparative analyses with multiple interactors.

Therefore, we investigated the similarities and differences in the preferences governing protein interactions of each CD2AP SH3 domain with known and novel partners. Based on the PxxPxxR motif, which is characteristic for CBL peptide binding to SH3 domains, we used an in vitro peptide array screen to confirm interaction candidates, many with previously uncharacterized sequence requirements. Among 40 binders reported, including ALIX, BLNK, and zona occludens protein 2 (ZO-2), we followed up on the newly identified RIN3, a RAB5 and RAB31 guanine nucleotide exchange factor involved in endocytosis (21, 22), which we identified as a novel interactor in kidney and MCF10A breast cancer cells by several biochemical and biophysical methods.

Experimental Procedures

Generation of Expression Constructs

The sequences encoding the three human CD2AP SH3 domains, herein referred to as SH3-1, SH3-2, and SH3-3, were generated as follows: SH3-2, SH3-3, and SH3-3L (a longer form of SH3-3) were amplified by polymerase chain reaction (PCR) using Pfu polymerase; CD2AP SH3-2, -3, or -3L BamHI forward primer; CD2AP SH3-2, -3, or -3L EcoRI reverse primer; and full-length CD2AP cDNA (23) as template before subcloning into BamHI/EcoRI-digested pGEX-6P-1 vectors (24) as a template and subcloned into BamHI/EcoRI-digested pGEX-6P-1 (SH3-2 and -3) or pGEX-KN (SH3-3L) vectors. Primer sequences were as follows: CD2AP SH3-2-BamHI forward, 5′-CG GCCATC ATG GAT AAG GA-3′; CD2AP SH3-2- EcoRI reverse, 5′-ATA GAATTC TTA CTA GGT AAA-3′; CD2AP SH3-3-BamHI forward, 5′-CG GCCATC ATG GAT AAG GA-3′; CD2AP SH3-3L EcoRI reverse, 5′-ATA GAATTC TTA CTA GGT AAA-3′; CD2AP SH3-3L BamHI forward, 5′-ATA GAATTC TTA CTA GGT AAA-3′; CD2AP SH3-3L EcoRI reverse, 5′-ATA GAATTC TTA CTA GGT AAA-3′.

A thrombin site was inserted at 5′ of the SH3-3L sequence using the QuikChange™ XL site-directed mutagenesis kit (Agilent Technologies). SH3-1 was synthesized with BamHI and EcoRI restriction sites and cloned into pUC57 by GenScript Corp. The cDNA fragment was subcloned into BamHI/EcoRI-digested pUC57 by GenScript Corp. The cDNA was amplified by PCR using Pfu polymerase with RIN3-AAA78F-BamHI forward (5′-ATA GCC ATCC GCC ATC AAC CCT TT-3′) and RIN3-AAA467 R-EcoRI reverse primer (5′-ATA GAATTC TTA CTA GAT CCG TTT TTT CCT GG-3′) using full-length Myc-tagged RIN3 cDNA (kindly provided by Dr. T. Katada) (24) as a template and subcloned into BamHI/EcoRI-digested pGEX-6P-1. Myc- and GFP-tagged RIN3 single and/or double epitope PxxxR → AxxxxA mutants were generated using wild-type RIN3 cDNA and the QuikChange™ XL kit. All constructs were validated by DNA sequencing. A Qiagen Plasmid Maxi Kit was used to prepare DNA.

Protein Expression and Purification

GST fusion proteins were expressed in Escherichia coli BL21(DE3) and purified essentially as described (25). Eluted GST fusion proteins were dialyzed against the final buffer (5 mM Tris, pH 7.0, 2 mM β-mercaptoethanol), and their integrity was confirmed by SDS-PAGE and Coomassie Blue staining;
protein concentrations were assayed by the Bradford method. Purified proteins were snap-frozen in aliquots and stored at −80 °C until further use in peptide arrays or GST pull-downs.

For isothermal titration calorimetry (ITC) and crystallization, proteins were purified as described above, and the GST tags were cleaved by incubation with PreScission Protease (GE Healthcare) in 20 mM Tris, pH 7.5, 150 mM NaCl. The GST tag and cleaved protein were separated by size exclusion chromatography on a HiLoad 16/60 Superdex 75 column equilibrated with column buffer (20 mM Tris, pH 7.5, 150 mM NaCl for CD2AP SH3-1 or supplemented with 2 mM β-mercaptoethanol for CD2AP SH3-2 and -3). Purified protein was dialyzed against the final buffer (5 mM Tris, pH 7.5, for SH3-1; 5 mM Tris, pH 7.5, 2 mM β-mercaptoethanol for SH3-2 and SH3-3). Proteins were concentrated and quantified by UV absorption at 280 nm, and their integrity was checked by SDS-PAGE/Coomassie Blue staining.

**Synthesis of Soluble Peptides and Microarrays**

Apart from the RIN3 peptide permutation arrays, all arrays and soluble peptides were synthesized as described (25). RIN3 peptide permutation arrays were prepared on cellulose-(3-amino-2-hydroxypropyl)-ether membranes (26) using a SPOT robot (Intavis AG, Cologne, Germany) according to our standard spot synthesis protocol. The arrays were designed using in-house LISA software. Cellulose-(3-amino-2-hydroxypropyl)-ether membranes gave a better signal/noise ratio compared with standard β-alanine cellulose membranes (26, 27).

**Peptide Array Far-Western Blots**

**CBL**

**Scanning Array**—The entire human CBL protein sequence was spot-synthesized as 21-aa partially overlapping peptides, sliding 3 aa with each step. Membranes were prepared as described (28), briefly prewetted in ethanol, and then washed with TBST (20 mM Tris-HCl, pH 7.5, 100 mM NaCl, 0.1% (w/v) Tween 20) three times for 10 min and blocked in TBST with 5% (w/v) nonfat dry milk for 4 h at room temperature. Membranes were incubated with 0.56 μM (14 μg/ml) purified GST in blocking buffer overnight at 4 °C to detect any background signals elicited by the GST tag. After washing three times in TBST for 10 min, membranes were probed with an anti-GST mAb, washed again, and incubated with HRP-coupled secondary (anti-lg) antibody. Membranes were then reblocked for 1 h after washing and reprobed with 0.56 μM (20 μg/ml) GST-CD2AP SH3 domain overnight at 4 °C. Bound GST-CD2AP SH3 was detected as described for GST; signals were apparent after several seconds.

**Truncation Array**—N- and C-terminally truncated peptides containing the 21-aa CBL epitope (GSQVPERPPKPFPRRINSERK) were spot-synthesized in duplicate and probed as described above.

**Permutation Array**—21-aa peptides encompassing the CBL epitope GSQVPERPPKPFPRRINSERK were permuted in single positions (underlined) to all 20 common amino acid residues, in duplicate rows, and probed as described above.

**ALIX**

**Permutation Array**—Spot-synthesized 21-aa peptides containing the ALIX epitope AGGHAPTPPTPAPRTMPPTKP were permuted in single positions (underlined) to all 20 other residues, in duplicate rows. Membranes were probed as above for the CBL permutation array.

**Proteome-wide search for novel putative CD2AP SH3 binding partners**

The PxPxPR motif in CBL was used as a search sequence for scanning the human proteome, disregarding splice variants, with ScanProsite (29). From 695 hits, a subset of 74 potential binding sites in 62 proteins with known or possible roles in signaling was selected; preference was given to proteins involved in membrane trafficking and/or signaling. Selected sequences were spot-synthesized on a microarray as 21-aa peptides, in duplicate. The arrays were probed as described for the CBL scanning array.

**RIN3**

**Scanning Array**—The entire human RIN3 protein sequence was spot-synthesized as partially overlapping 27-aa peptides, sliding 3 aa with each step. Three array copies were probed as described above for the CBL arrays, except 0.1 μM (2.6 μg/ml) GST or 0.1 μM (3.6 μg/ml) GST-SH3 was used in the same blocking buffer (TBST, 5% (w/v) nonfat dry milk).

**Permutation Array**—Two 17-aa sequences from RIN3 (AKKNLPTAPPRRRVSE and TAKQPPVPPPRKRISR) that bind to CD2AP SH3 domains were permuted at 15 single positions (underlined) to 20 aa and spot-synthesized in single rows. Arrays were probed as for the RIN3 scanning arrays, except with a modified blocking buffer (2% (w/v) ovalbumin in RIPA-100, for GST-SH3-1 and -SH3-2, or supplemented with 2 mM DTT, for GST-SH3-3) partly due to the different membrane type. No background GST signal was detected after 2 min, whereas GST-SH3 signals appeared after 1 s.

**ITC**

ITC was performed at 25 °C with a VP-ITC microcalorimeter (MicroCal/Malvern Instruments) essentially as described (30), but peptides were dissolved at 0.75–1.5 mM, and protein solutions of CD2AP SH3-1, SH3-2, or SH3-3 domains were used at 0.05–0.1 mM in ITC buffer (25 mM HEPES, pH 7.5, 100 mM potassium acetate). An initial injection of 4 μl of peptide solution was followed by 18 15-μl injections into the sample cell containing protein solution. Background heats from injection of peptide into buffer solution without protein were subtracted from experimental heats and then analyzed in ORIGIN (version 5.0) software by fitting to a model for a single set of binding sites using χ² minimization.

**Protein Crystallography**

**CD2AP SH3-1/RIN3e1 complex**

The SH3-1 domain was mixed in a 1:3.5 ratio with a 16-mer RIN3e1 peptide (AKKNLPTAPRRVSE). Crystals were grown by vapor diffusion in a sitting nanodrop (total volume of 150 nl) at 20 °C from a 1:1 ratio of mother liquor (0.1 M calcium
acetate, pH 4.2, 0.2 M Li$_2$SO$_4$, and 25% PEG 10,000) to CD2AP SH3-1 protein solution (19 mg/ml). A cluster of needles (~160 μm long), which grew after 2 weeks, was transferred to cryoprotectant (mother liquor supplemented with 25% glycerol) before being snap-frozen in liquid nitrogen. Data were collected to 1.6 Å on beamline I04 at Diamond Light Source synchrotron (Harwell, UK). Data were reduced with the XDS package (31), converted to CCP4 mtz format, and scaled with SCALA (32); data to 1.65 Å were kept for refinement. Molecular replacement was done in PHASER (33) using as a search model the SH3 domain alone from a reported binary complex of CD2AP SH3-1 and CBL-b peptide (PDB code 2J6F). The single, correct solution in space group C2 was refined with REFMACS (34), and additional manual model building was done in COOT (35). TLS (torsion, libration, screw) restraints were included for six groups (i.e. each chain in the asymmetric unit) (three SH3-1 domains and three RIN3 peptides). The final model was generated by optimizing experimental data/geometry restraint weighting and refining the water structure in PHENIX (36) before deposition in the PDB under accession code 4WCI.

**CD2AP SH3-2-RIN3e2 complex**

The SH3-2 domain was mixed with a 16-mer RIN3e2 peptide (TAKQPPVPPRPKRIS) in a 1:3 ratio. Crystallizations were set up as for the SH3-1-RIN3e1 complex. After 3 weeks, a single, thin, square plate (~180 μm in length/width) grew from a 2:1 ratio of mother liquor (0.1 M HEPES, pH 7.5, 1.4 M trisodium citrate dehydrate) to CD2AP SH3-2 protein solution (16.7 mg/ml). This crystal was transferred to cryoprotectant (mother liquor supplemented with 10% glycerol) and then snap-frozen in liquid nitrogen. Data were collected to 1.1 Å on beamline I03 at the Diamond synchrotron. Data were reduced with the XDS package, proving incomplete in most resolution shells (space group C2), so an additional data set was later collected using the same crystal but reoriented in the beam (Diamond beamline I04-1). The two data sets were merged with XS SCALE before conversion to CCP4 mtz format. Molecular replacement with PHASER used an ensemble of SH3 models homologous to CD2AP SH3-2 (PDB codes 2G6F, 1OEB, 2AK5, 2FEI, and 3IQL). The single, correct solution was subjected to automatic model building with AutoBuild in the PHENIX suite and later REFMACS refinement. Additional manual model building was done with COOT. Anisotropic B-factors and several alternative conformations were included. The final model was deposited in the PDB with accession code 4WCI.

**Cell Culture**

Conditionally immortalized human kidney podocytes were a kind gift from Prof. Moin A. Saleem (University of Bristol). Podocytes were maintained in RPMI 1640 medium supplemented with 10% FBS and 1% ITS (insulin-transferrin-selenium) and incubated at 33 °C with 5% CO$_2$. Differentiation was triggered by thermo-switching to 37 °C as described (37). HEK293 and HepG2 cell lines were grown in DMEM supplemented with 10% FBS and 1% penicillin/streptomycin and incubated at 37 °C with 5% CO$_2$. Jurkat cells were cultured in RPMI 1640, 10% FBS at 37 °C with 5% CO$_2$. Growth medium was changed every 2–3 days. Cells were split at a ratio of 1:3 to 1:5 when they reached 80–90% confluence by washing with PBS and incubating for <5 min with 1–2 ml of 0.25% trypsin, 0.1% EDTA in PBS solution until the cells detached before centrifuging at 200 × g for 5 min and resuspending in fresh medium. MCF10A cells were grown as described (38) without antibiotics.

**Western Blotting, Immunoprecipitations, and Pull-downs**

Total cell lysates were prepared and analyzed by Western blotting essentially as described (39). For immunoprecipitation of wild-type human RIN3 and CD2AP, HEK293T cells were transiently co-transfected with 0.5 μg each of FLAG-tagged CD2AP and Myc-tagged RIN3 per 60-mm dish using FuGENE-6 and then lysed in HNTG buffer (50 mM Heps, pH 7.4, 150 mM NaCl, 1% Triton X-100, 10 mM EDTA) with 1 mM PMSE, 1 mM aprotinin, and 1 mM leupeptin added. Washed GammaBind™ G-Sepharose™ beads (GE Healthcare) were then added to the samples for 90 min at 4 °C. Immunoprecipitates were washed four times with ice-cold HNTG buffer containing 0.1% Triton X-100 plus inhibitors as above and then subjected to Western blotting with specific antibodies (Abs). Monoclonal Abs were used against c-Myc (clone 9E10, sc-40, Santa Cruz Biotechnology, Inc.) and FLAG (M2, Sigma-Aldrich), and a polyclonal rabbit Ab was used against CD2AP (sc-9137, Santa Cruz Biotechnology).

For GST pull-downs, equal amounts of purified GST fusion proteins (GST-CD2AP SH3 domains; GST-RIN3(378–467)) were coupled to reduced glutathione-Sepharose beads (GE Healthcare) and incubated with 1 mg of total cell radiimmune precipitation assay protein extract for 16 h with rocking at 4 °C, followed by three 1-ml washes with chilled RIPA-100 buffer (20 mM Tris-HCl, pH 7.5, 1 mM EDTA, 1% Triton X-100, 0.5% sodium deoxycholate, 0.1% SDS, and 100 mM NaCl). Proteins were separated by SDS-PAGE followed by semidry transfer onto polyvinylidene difluoride (PVDF) membranes (Hybond-P, GE Healthcare) and then blocked, probed, and developed as described (40). The anti-CBL mouse monoclonal Ab (A-9) was from Santa Cruz Biotechnology (sc-1651). The anti-RIN3 antibody (41) was kindly donated by Prof. John J. Colicelli (UCLA, Los Angeles, CA).

**Immunofluorescence**

MCF10A cells plated on poly-L-lysine-coated coverslips were transfected with GFP-RAB5a, FLAG-CD2AP, RFP-RIN3, and RFP-RIN3mut. The next day, cells were washed, fixed with 2% formaldehyde-PBS for 10 min at room temperature, and permeabilized with 0.2% Triton X-100/PBS for 1 min. Cells were stained with monoclonal anti-FLAG antibody (M2, Sigma; 1:50,000) diluted in PBS-Triton (0.02%). After four washes, cells were incubated with Alexa647-goat anti-mouse IgG (Molecular Probes, Inc.; 1:500).

**Results**

Analysis of the Interaction between the First Two CD2AP SH3 Domains and CBL—CD2AP SH3-1 and SH3-2 bind to a known region in the C terminus of CBL (7, 16, 42) and participate in the linkage of EGF receptor internalization to the cytoskeleton. We
confirmed these interactions in Jurkat cells by showing that both CD2AP SH3-1 and SH3-2 could pull down CBL, but the third SH3 domain could not (Fig. 1C). Interestingly, SH3-2 was more effective than SH3-1, pulling down over twice the amount of CBL, suggesting a stronger interaction. The adaptor protein Grb2, cooperating with CBL, is indispensable for EGF receptor internalization (43) and may bind to CBL via both its N- and C-terminal SH3 domains (44), so we compared the latter interaction in our experiment. We found that Grb2 SH3C pulled down twice the amount of CBL compared with CD2AP SH3-2 (Fig. 1C). This suggested that the interaction of CBL with

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FIGURE 1. CD2AP SH3-1 and SH3-2, but not SH3-3, bind directly to CBL via a motif bounded by Pro-824 and Arg-829. A, schematic CD2AP domain composition. SH3, Src homology 3; proline-rich, proline-rich region; CC, coiled-coil domain; an actin binding region is shown with putative interaction sites. N and C represent the protein termini. B, sequence alignment of the human CD2AP SH3 domains. Identical residues down columns are shown as red boxes with white letters. Residues that have higher than 70% similarity (based on physico-chemical properties) in a column are shown as red letters in blue frames. The highest pairwise sequence identity is found between SH3-2 and SH3-3 (45%); SH3-1 and SH3-2 have less identity (40%); SH3-1 and SH3-3 have the lowest identity (38%). C, GST pull-downs. 1 mg of Jurkat total cell lysate was precipitated with 50 μg of bead-immobilized GST or GST-SH3 fusion proteins; precipitates were immunoblotted with an anti-CBL antibody. Values from densitometry below the blot show the CBL signal after pull-downs with each GST-SH3 relative to the intensity for CD2AP SH3-1. D, peptide scanning arrays. On cellulose membranes, peptides were spot-synthesized as overlapping 21 aa, sliding 3 aa with each step to cover the full human CBL sequence. Membranes were initially probed with 0.1 μM GST before immunodetection to confirm no nonspecific binding and then washed, reblocked, and probed again with 0.1 μM GST-CD2AP SH3-1, GST-CD2AP SH3-2, or GST-CD2AP SH3-3, as described under “Experimental Procedures.” The series of spots that gave the strongest signals are boxed in red. E, schematic domain composition of human CBL. The full sequence region spanned by the boxed series of spots in D (aa 811–843) is shown. The portion of this sequence common to each spot in the series, encompassing the CD2AP SH3 recognition sequence, is shown in red letters. TKB, tyrosine kinase binding domain; RING finger, really interesting new gene zinc finger domain; proline-rich, proline-rich region; UBA, ubiquitin-associated domain. F, truncation arrays. N- and C-terminally truncated peptides of the 21-aa CBL epitope were spot-synthesized in duplicate. Arrays were first probed with GST and then GST-CD2AP SH3-1 or GST-CD2AP SH3-2 as described in D. Truncation beyond Pro-824 and Arg-829 (colored in red) prevents binding, which identified the core motif boundaries.
CD2AP SH3 Domain Recognition Preferences

CD2AP is less dominant in EGF receptor endocytosis than the Grb2-mediated interaction, similar to the minor or redundant role reported for the CD2AP homologue CIN85 (43). We then validated in vitro the interactions of all three CD2AP SH3 domains with CBL (42) to confirm whether the C-terminal binding site was unique. Thus, with each SH3 domain as a GST fusion, we probed spot-synthesized CBL peptide scanning arrays of peptides that covered the entire human CBL sequence. We found that the C-terminal epitope is indeed the major specific binding site for SH3 domains 1 and 2 and, to a much lesser extent, also SH3-3 (Fig. 1, D and E). To determine the boundaries of the CBL epitope, we used arrayed CBL peptides (aa 816–836) sequentially truncated from either the N or C terminus and probed with the first two CD2AP SH3 domains (as GST fusions). The recognition motif extended from Pro-824 to Arg-829 in CBL (Fig. 1F). Membranes first probed with GST alone as a control gave no background signal in an equivalent exposure time (data not shown).

Although Arg-829 has been identified as a critical residue for the interaction of CBL with CD2AP SH3-1 and SH3-2 (42), we aimed to map comprehensively the exact residues required for binding. First, we quantified the interactions of the three CD2AP SH3 domains with CBL by ITC (Table 1). We found that SH3-2 bound most strongly ($K_d = 6.7 \mu M$), whereas SH3-1 and SH3-3, respectively, bound 2.7- and 21-fold more weakly compared with SH3-2. These results reflected the differences observed in GST pull-downs (Fig. 1C) and the scanning array (Fig. 1D).

To explore the CBL motif in greater detail and to study any similarities and differences in the binding modes of the first two CD2AP SH3 domains, we probed spot-synthesized 20-aa permutation arrays of the CBL epitope (Fig. 2). Both CD2AP SH3-1 and SH3-2 recognized a core $P^xP^xP^{R^x}$ motif in CBL (superscript numbers refer to relative motif positions, and $x$ indicates that the residue position is unimportant for the interaction; this notation is used throughout). The array also detected some degree of selection for the interaction in motif positions 2 and 4, most notably for SH3-2, giving $P_xP^xP^x$ (or possibly $P_kP^xP^x$) as the full recognition sequence (lowercase letters denote less important residues for the interaction; Fig. 3). This sequence is, in part, similar to one previously reported highlighting a minor role for the equivalent Lys in CBL-b, which permitted binding to CD2AP SH3-1 in two orientations as heterotrimers (45). However, the peptide array revealed the sequence of $K_825A$ substitution reduced the binding signal, consistent with the reported effect of a CBL-b K907A mutation (45). However, binding between CBL and every CD2AP SH3 domain in ITC experiments had 1:1 stoichiometry, implying that each complex was heterodimeric (supplemental Table S1 and Fig. S1). Interestingly, the array revealed that an alanine in motif position 3 was equally acceptable in the case of both SH3-1 and SH3-2 binding, whereas SH3-2 tolerated additional residues in two other peptide positions. This suggested that CD2AP may also bind a $P_xA_xP^R$ motif.

Bioinformatics and in Vitro Screens Identified 40 Putative CD2AP SH3-interacting Proteins—We investigated how frequently the CBL $P_xP^xP^x$ motif featured in the human proteome in order to discover other CD2AP-interacting proteins. A proteome search using the ScanProsite Web tool (29) yielded 602 proteins (695 sequence hits in total) containing this motif. We filtered these hits to prioritize cytosolic signaling proteins,

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**TABLE 1**

| Protein | Peptide sequence | Length (aa) | $K_d$ ± SE (μM) |
|---------|-----------------|-------------|-----------------|
| CBL     | GSTDINVTEGQVPERPPKFPSTWYV | 28          | 17.5 ± 1.7 | 6.5 ± 0.5 | 137.6 ± 7.3 |
| ALIX    | AQGAVIPFTEAPTMHPE | 21          | 6.6 ± 0.002   | 4.2 ± 0.001 | 32.2 ± 1.9 |

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**FIGURE 2.** CD2AP SH3-1 and SH3-2 recognize a $P_xP^xP^x$ motif in CBL. 21-aa peptides of the CBL epitope GSQVPERPPKFPSTWYV were permuted in single positions (underlined) to the 20 common residues, in duplicate rows. Arrays were initially probed with GST and then with GST-CD2AP SH3-1 or GST-CD2AP SH3-2, as described for Fig. 1D. The CBL peptide sequence, and hence each residue position permuted, is displayed vertically down the left side of each array; the individual amino acid substitutions at these positions are listed across the top. Wild-type CBL peptide spots are boxed; those emphasized in red boxes show highly restricted positions (i.e. key motif determinants). Note that the amino acid preferences at these positions are further highlighted on the right side of each array. Lowercase red type indicates less critical positions; a black letter x marks unimportant positions within a motif. Red type, wild-type sequence; smaller black type (capital letters), amino acids giving a signal intensity similar to wild type.
ensuring the inclusion of those involved in membrane trafficking. Thus, 74 epitopes within 62 different proteins were selected for in vitro analysis by peptide microarray. These included a few reported partners (e.g. CBL and ALIX), useful as positive controls, but the majority were unknown as interactors. 21-residue peptides spanning the motif regions from these selected hits (supplemental Table S2) were spot-synthesized in duplicate on cellulose membranes, and three array copies were probed with one of each of the CD2AP SH3 domains (Fig. 3). Strikingly, the pattern of identified peptides binding to each SH3 was non-identical, with the proportion of hits as follows: 66% for SH3-1, 34% for SH3-2, and 9% for SH3-3. Note that in this analysis, we included all positive blotting reactions, however weak, because the intensity levels from probing immobilized peptides unoptimized for length and boundary positions are not always a reliable predictor of binding affinity in solution (46). Nevertheless, with the exception of CADH1, unique to SH3-2, the hits for SH3-2 formed an exact subset of those to which SH3-1 bound, and similarly for SH3-3, albeit a far smaller subset. Importantly, this third domain not only bound very few of the sequences, but also much more weakly, and thus appears poorly optimized for binding the core PxPxFPR motif. This implies that SH3-3 may have different and as yet unidentified binding preferences and a divergent functional role. In total, peptides from 40 potential or known CD2AP-binding proteins were identified in the screen (disregarding TAB3; see below).

**Differential Interactions of Each CD2AP SH3 Domain with an ALIX Peptide**—The signal intensity of the CBL peptide spot, the strongest binder, was normalized to 100%, and all others were adjusted relatively based on densitometric analysis. Thus, other strong binders (>75% of the CBL signal) included ALIX, BLNK, CBL, CEND3, DAB2, SYNJ2, and TAB3. Medium-strength binders (25–75% of the CBL control signal) comprised ANKS1, ARHGB, ATX7, BANK1, CSTD2, DAB2P, DDEF1, GA2L2, GATA6, MAP3K11, P85A, REP52, RHGO4, RHGO6, RIN2, RIN3, SAM68, SYNJ1, TAF6, and UBPS1. Low intensity binders (<25% of the control signal) were CYLC1, DOK2, DOK3, LAT51, NEDD4-like, PTN14, RGL2, RN123, SNIP, SPP3, TAF1A, WISP2, and ZO-2. Non-binders, taken to be those spots not visible to the naked eye, had a mean signal of <3% of CBL. Proteins that did not bind with a single epitope to any CD2AP SH3 were AKTS1, ARHGB, BORG3, CADH1, CLD23, ELK1, ESB1, FOXGA, GRASP, HNRPK, JIP1, P85B, PDPK1, PHAR1, PP1RA, PTN12, REP51, SGI1P, SHAN1, SOS2, STYK1, and SYGP1.
the ALIX epitope interacted with all three SH3 domains. ITC measurements with a synthesized 21-residue ALIX peptide and each SH3 revealed stronger binding to the first two SH3 domains ($K_d$ values of 6.6 and 4.2 μM, respectively) compared with 5–8-fold lower affinity for SH3-3 (Table 1).

To explore the ALIX interaction in greater detail, we synthesized 20-aa permutation arrays based on the ALIX epitope (Fig. 4). Broadly, the array blotting patterns with SH3 domains 1 and 2 were very similar, but domain 3 displayed significant variation, again highlighting its distinct properties. Interestingly, the array data revealed that the first two domains recognize the motif PtPxxR. This differs in two key aspects from the CBL epitope: (i) Pro in position 5 of the CBL motif $P^{3}tP^R$ was surprisingly not a strict requirement according to the array yet was present in the ALIX motif; (ii) the second position is more restricted than in the CBL motif, mainly preferring a Thr as observed for the CBL epitope in immobilized form. Thus, the two RIN (Ras and Rab interactor) family proteins, RIN2 and RIN3, were analyzed further. Note that low intensity binders (<25% of the control signal) may also be relevant but were not pursued further. The RIN proteins were of special interest due to their link to endocytic processes, as guanine nucleotide exchange factors for the small GTPase RAB5, and hence potential candidates of physiological interplay with CD2AP; also, relatively little was known about their biological roles. Both RIN2 and RIN3 displayed noticeable binding to SH3-1, very weak binding to SH3-2, and yet no discernible binding to SH3-3 (Fig. 3). Therefore, to test the relevance of these interactions with full-length proteins, we used a co-immunoprecipitation system between CD2AP and RIN2 (data not shown), whereas a CD2AP-RIN3 complex could be co-immunoprecipitated (Fig. 5A).

RIN3 Is a Novel Interaction Partner for the CD2AP SH3 Domains in Kidney Cells—We also considered medium-intensity spots from the peptide array screen as useful hits because boundary positions may have influenced the signal strengths observed for the array peptides in immobilized form. Thus, the CD2AP SH3-3 recognizes an extended epitope in ALIX unlike domains 1 and 2. 21-aa peptides of the ALIX epitope $AGGHPATP^R$ were permuted in single positions (underlined). The arrays were probed as described for Fig. 1D using GST-CD2AP SH3-1 (left), GST-CD2AP SH3-2 (middle), or GST-CD2AP SH3-3 (right). Wild-type ALIX peptide spots are boxed, and all other figure labeling is the same as for the CBL arrays (Fig. 3) except that blue type and boxes in the array probed with SH3-3 indicate additional restricted residue positions in an extended ALIX motif, not seen with SH3-1 or -2.

According to the array is $P^{3}tP^R$ (Fig. 4, right). This is an unusually long recognition sequence compared with known CD2AP SH3 interaction motifs. Similar to domains 1 and 2, SH3-3 strictly required Pro in position 1, Arg in position 5, and Pro in position 3 (with Ala demonstrably possible). However, we observed a marked difference with SH3-3 in position 10, where proline was necessary for a strong interaction, and in position 12, a basic residue was essential (Lys in ALIX). This indicates that a patch beyond the canonical binding groove may be accessible for full interaction with SH3-3. However, SH3-3 still bound with lower affinity than SH3-1 and -2 to the ALIX peptide ($K_d = 32 \mu M$; Table 1), as observed with the CBL peptide (and RIN3e1; see below). The arrays gave no background signal when first probed with GST alone (data not shown).

RIN3 Is a Novel Interaction Partner for the CD2AP SH3 Domains in Kidney Cells—We also considered medium-intensity spots from the peptide array screen as useful hits because boundary positions may have influenced the signal strengths observed for the array peptides in immobilized form. Thus, the two RIN (Ras and Rab interactor) family proteins, RIN2 and RIN3, were analyzed further. Note that low intensity binders (<25% of the control signal) may also be relevant but were not pursued further. The RIN proteins were of special interest due to their link to endocytic processes, as guanine nucleotide exchange factors for the small GTPase RAB5, and hence potential candidates of physiological interplay with CD2AP; also, relatively little was known about their biological roles. Both RIN2 and RIN3 displayed noticeable binding to SH3-1, very weak binding to SH3-2, and yet no discernible binding to SH3-3 (Fig. 3). Therefore, to test the relevance of these interactions with full-length proteins, we used a co-immunoprecipitation system in HEK293 kidney cells with Myc-RIN2 or -RIN3 and FLAG-CD2AP. In contrast, there was barely a detectable interaction between CD2AP and RIN2 (data not shown), whereas a CD2AP-RIN3 complex could be co-immunoprecipitated (Fig. 5A).

To determine the contribution of each SH3 domain to the RIN3 interaction, we carried out pull-down experiments using GST fusions of each individual SH3 domain and lysates from immortalized kidney podocytes. Only very low levels of endogenous RIN3 were detectable in undifferentiated cells (33 °C); increasing the temperature to 37 °C to trigger differentiation increased RIN3 amounts in pull-down experiments with undifferentiated podocytes, but after cell differentiation, substantially more RIN3 was pulled down (Fig. 5B). SH3 domains 1 and 2 bound...
equal amounts of RIN3, but SH3-3 failed to elicit a positive pull-down, as did the GST-only control. Recently, the SH3-3 domain from the homologue CIN85 showed greater thermal stability and expression yield compared with a shorter construct when its N and C termini were extended (48). Therefore, in case the negative SH3-3 pull-down result was somehow related to suboptimal boundaries of our GST-SH3-3 domain construct (should protein stability affect ligand binding), we generated a longer SH3-3 construct. This was extended at both termini in total by 10 residues, equivalent to the region elongated in CIN85 SH3-3 (48). However, the longer construct did not pull down RIN3 either (labeled 3L in Fig. 5B), confirming the different behavior of SH3-3 relative to SH3-1 and -2.

**RIN3 Harbors Two CD2AP Interaction Motifs**—RIN3 has an extended PRR in its N-terminal half. Given the potential for
CD2AP SH3 Domain Recognition Preferences

FIGURE 6. Each CD2AP SH3 domain recognizes the same two RIN3 epitopes. A, RIN3 scanning arrays. Overlapping 27-aa human RIN3 peptides were spot-synthesized on cellulose membranes sliding 3 aa with each step to cover the full protein sequence. Arrays were initially probed with GST and then with GST-CD2AP SH3-1 (left), GST-CD2AP SH3-2 (middle), or GST-CD2AP SH3-3 (right), as described for Fig. 1D. Two principal binding regions are evident (ringed in green and red). The exact sequence of each spot is found below the arrays (same color coding). Note that, unlike results obtained with CBL (Fig. 1, D and F), the actual RIN3 peptide boundaries appear to influence probe binding onto the arrayed peptides much more strongly. B, schematic domain composition of RIN3. The boxed sequences represent the minimal common segment found in all binding spots for that region (again, green for epitope 1, aa 367–390; red for epitope 2, aa 445–462). SH2, Src homology 2; RH, RIN homology; VPS9, vacuolar sorting protein 9; RA, Ras association. Note that only PRR-2 and -3 have affinity for CD2AP.

variant motifs suggested by differences in the binding preferences observed in the ALIX and CBL permutation arrays, we tested whether RIN3 contained additional CD2AP binding sites (Fig. 6). Spot-synthesized scanning arrays of peptides covering the entire human RIN3 sequence were probed with the three individual CD2AP SH3 domains (Fig. 6A). Interestingly, each SH3 domain showed some binding to the originally screened motif region (Fig. 6B, red box), but an additional RIN3 epitope recognized by all three SH3 domains was also identified (Fig. 6B, green box). It differs from the screened sequence by not containing an exact PxPxPR motif; rather, Ala is present in motif position 3. The lack of immunoreaction to a longer contiguous series of array peptides harboring the motif indicated that the peptide boundaries were indeed critical for binding to these epitopes in short peptide form (cf. CBL scanning array; Fig. 1D) and explains why the initial screening array peptides sometimes gave weaker signals for this hit (Fig. 3).

Having mapped these binding epitopes, herein referred to as RIN3e1 and RIN3e2, we created a GST-fused RIN3 construct that spanned both epitopes in order to perform the reciprocal pull-down of CD2AP from cell lysates. GST-RIN3(378–467) was able to pull down endogenous CD2AP, whereas no CD2AP band was present with GST alone (Fig. 5C). Note that extra blot bands were visible due to cross-reactivity of the anti-CD2AP antibody with species in the recombinant protein sample, as determined by a lysate-free control. We proceeded to characterize the CD2AP-RIN3 interaction with comprehensive biochemical and biophysical approaches.

Substitution Analyses Revealed Px(P/A)xPr in RIN3 as the Preferred Recognition Motif—First, to delineate the recognition motifs as for CBL and ALIX, we synthesized 20-aa permutation arrays of RIN3e1 and RIN3e2 peptides (Fig. 7) and probed them with each CD2AP SH3 domain. These arrays were synthesized on a different carrier material, and although the initial GST-only control probe gave no appreciable signal (data not shown), a higher background signal was observed from nonspecific membrane binding by the SH3 domains, particularly the RIN3e2 arrays. However, the results still allowed clear detection of specific interactions (Fig. 7). In general, the signal pattern was strikingly similar between all three SH3 domains (in contrast to the ALIX peptide arrays; Fig. 4). Indeed, the residues PxPxPR formed the general recognition motif in RIN3e2, with Thr in position 2 of some importance. In RIN3e1, the second proline (motif position 3) was replaced by alanine, thereby forming the basis for recognition of the related motif: PxAPxP. In fact, the peptide arrays revealed that in the sequence contexts of either RIN3 epitope, Ala or Pro are equally strong binders in position 3, as seen for ALIX and CBL. Val in this position is also tolerated, only slightly weakening the interaction. In contrast, in position 1, only Pro appears possible for a strong interaction with every SH3 domain because any other residue prevented or weakened binding. The effect was more pronounced with RIN3e1 than with RIN3e2. Arg in position 6 was strictly required for strong binding in all cases. Pro in position 5 of RIN3e1 was essential for binding to SH3-2 and SH3-3 although not SH3-1. The equivalent Pro, however, was generally preferred for the strong interaction of all SH3 domains with RIN3e2, although greater tolerances exist in this position.

Both RIN3 Motifs Are Necessary for Full CD2AP Interaction and Its Recruitment by RIN3 to RAB5a-positive Endosomes—The ability of each RIN3 motif in the context of the full-length protein to interact with CD2AP was next assessed using the HEK293T transfection system, with mutated RIN3 variants. Motifs in epitopes 1 or 2 were rendered non-functional by dual

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Ala substitutions of the critical first and last motif positions (PxxxxR to AxxxxA); a third variant with both epitopes simultaneously mutated in this manner was also tested. Mirroring the previous result with endogenous RIN3 in podocytes (Fig. 5B), wild-type RIN3 (W) bound to SH3-1 and -2, but not SH3-3 or GST alone (Fig. 5D). Furthermore, whereas dual Ala substitutions in RIN3-e1 or -e2 individually still allowed binding to the first two SH3 domains, indicating the independent functionality of each motif, only when both epitopes were mutated was binding prevented (double epitope mutant (D); Fig. 5D). The two binding sites may thus work in tandem for proper CD2AP interaction in the cell, engaging with the first two SH3 domains simultaneously.

Immunofluorescence microscopy was performed to investigate the association of RIN3 and CD2AP in transiently transfected cells in culture. In CD2AP-expressing cells, CD2AP is mainly localized in the cytoplasm and shows no obvious accumulation on RAB5a-labeled early endosomes (Fig. 5, G and H). As shown previously, RIN3 localizes to RAB5a-positive endosomes (Fig. 5H) (21). In addition, a RIN3 variant with the CD2AP binding sites mutated also targets to RAB5a-positive endosomes. Importantly, when RIN3 and CD2AP were co-expressed CD2AP became co-localized with RIN3 on early endosomes (Fig. 5, E and I). The recruitment of CD2AP to early endosomes was mediated by RIN3 because the RIN3 variant mutated for CD2AP interaction failed to recruit CD2AP to early endosomes (Fig. 5, F and I). These data revealed the importance of both RIN3 binding sites for interaction with CD2AP in cells.

**ITC Data Confirmed Differences in CD2AP SH3 Interactions with Free RIN3 Peptides**—To quantify binding, ITC experiments were performed between individual CD2AP SH3 domains (with GST tags cleaved off) and synthetic 16-mer RIN3-e1 and RIN3-e2 peptides (Table 2). In general, RIN3-e1 binding affinities were comparable with those of ALIX peptide, but RIN3-e2 displayed tighter binding to all SH3 domains. Specifically, RIN3-e1 bound to SH3-1 and SH3-2 with similar binding affinities (Kd values of 8.9 and 10.0 μM, respectively) but unexpectedly bound with 4-fold weaker binding than SH3-1 to RIN3-e1 only (Kd ~40 μM) as shown in Table 2.

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**FIGURE 7.** All three CD2AP SH3 domains recognize variants of a Pt(P/A)xPXR motif in RIN3. The two RIN3 binding regions identified in the scanning array were spot-synthesized as 17-aa peptides on cellulose-(3-amino-2-hydroxypropyl)-ether membranes (RIN3-e1, AKKNLPTAPRRRVR; RIN3-e2, TAKQPPVPVRKRISR) and permuted in 15 single positions (underlined) against all 20 natural residues, in single rows. Three copies of the RIN3-e1 permutation array (all left panels) or the RIN3-e2 array (all right panels) were initially probed with GST as described for Fig. 1D and then with GST-CD2AP SH3-1 (A), GST-CD2AP SH3-2 (B), or GST-CD2AP SH3-3 (C). See also legend to Fig. 2 for an explanation of the array annotations.
more similar affinity to RIN3e2 (6.7 μM), the tightest binding so far observed for this domain. Shorter RIN3 peptide lengths, such as 10- or 11-mers, reduced binding affinity relative to 16-mers, an effect that was more pronounced for epitope 2 than epitope 1. Thus, additional contributions to the binding energy may derive from residues beyond the canonical motif.

Next, to confirm the importance of the three principal motif determinants, focusing on the tighter binder, RIN3e2, we synthesized mutant peptides and measured their interactions with each CD2AP SH3 domain by ITC (Table 2). Thus, we introduced alanine point mutations in place of Pro-457, Pro-459, and Arg-462 (i.e., motif positions 1, 3, and 6). The P457A mutant peptide greatly weakened binding, most notably in the case of SH3-1, resulting in a >30-fold reduction in affinity relative to the wild-type sequence (and nearly 10-fold with SH3-2; 16-fold with SH3-3). The R462A mutant abolished the interaction with each SH3 to unquantifiable levels, indicating its key role in recognition. However, the P459A peptide maintained binding in the low micromolar range with all SH3 domains, not too dissimilar from the wild-type peptide. This was expected, reflecting the functional interchangeability of Ala for Pro in motif position 3 observed in the peptide array. Finally, ITC data showed that each SH3-peptide interaction was predominantly enthalpically driven, as is common for SH3s, but the tighter binding RIN3e2 peptide had a more favorable entropic contribution (supplemental Table S1).

**High-resolution Crystal Structures of Heterodimers Revealed a More Extended Class II Binding Mode with a CD2AP SH3-2–RIN3e2 Complex Compared with CD2AP SH3-1–RIN3e1**—In parallel, we attempted co-crystallization of each SH3 domain in complex with peptides encompassing each wild-type RIN3 motif. We solved high-resolution structures of both CD2AP SH3-1 in complex with RIN3e1 to 1.65 Å (PDB code 4WCI), and SH3-2 in complex with RIN3e2 to 1.11 Å (PDB code 3U23) (Fig. 8 and Table 3). However, crystallization attempts with SH3-3 did not yield crystals in apo form or as a complex. Both CD2AP SH3 domains 1 and 2 display the familiar SH3 fold of a six-stranded β-barrel. In the SH3-1–RIN3e1 structure, three copies of the complex form the asymmetric unit in a triangular arrangement. Each complex is a heterodimer (i.e., 1:1 stoichiometry), consistent with the ITC result (supplemental Table S1). Differences between monomers include the positions of the N termini (specifically, the residues are vector artifacts conjoined to the SH3), which extend out from the domain proper and provide various crystal-packing contacts. In each complex, the RIN3e1 peptide forms a left-handed polyproline type II helix and binds to the SH3 in class II orientation. Pro-382, Ala-384, and Arg-387 from the RIN3e1 peptide (i.e., the essential residues of the PxApRxP motif) selectively dock into the two hydrophobic pockets and the negatively charged specificity pocket of SH3-1, respectively (Fig. 8, A and C, left panels). Furthermore, Pro-386 makes no contacts with the SH3 at all, as inferred previously by the array, although Pro in this motif position 5 within a CBL epitope is essential for binding (Fig. 2). Pro-386 therefore presumably serves an essential structural role within the polyproline type II helix.

The structure of the SH3-2–RIN3e2 complex was solved with a single binary complex in the asymmetric unit (i.e., 1:1 stoichiometry), again consistent with our ITC data. The heterodimer reveals a similar complementary fit with Pro-457, Pro-459, and Arg-462 from RIN3e2, in bonding distance to the SH3-2 domain and in class II orientation (Fig. 8, A and C, right panels). As with RIN3e1, the RIN3e2 peptide adopts a polyproline type II helical conformation, but four extra residues beyond the C-terminal end of the canonical motif were visible in the electron density maps. Ile-466 in motif position 10 lies within a distal hydrophobic patch, flanked by Val-141, Leu-154, and Leu-156 on SH3-2, implicating Ile-466 in the interaction (Fig. 8, A–C, right panels). This is consistent with the reduced affinity observed for an 11-mer RIN3 peptide truncated at Lys-464 (motif position 8) compared with the 16-mer co-crystallized (ITC data; Table 2). Thus, the RIN3 recognition motif for SH3-2 may be more extended (i.e., P1xPxpRxxrl10), so we analyzed this further by ITC. An I464A mutant peptide curiously had no major effect on the binding affinity for SH3 domain 1 or 2 (Table 2), a result that mirrored our peptide array data, but

**CD2AP SH3 Domain Recognition Preferences**

TABLE 2

| Binding affinities of wild-type and mutant RIN3 peptides to each CD2AP SH3 domain |
|---|
| **Data were obtained by ITC. Two wild-type RIN3 epitopes (e1 and e2) were tested, and mutant versions of RIN3e2, the tighter binder. Affinities are grouped by increasing red colour intensity; the stronger the shade of red, the higher the affinity (the lower the Kd value). Color groups represent Kd ranges in half-log (μM) increments (e.g., strongest red: >0–0.5 log Kd (μM) = Kd range of >1–3.16 μM). TLQ, too low for quantification (of accurate affinities); wt, wild-type sequence; point mutants of RIN3e2 are also shown.** |
| **TABLE 2** Binding affinities of wild-type and mutant RIN3 peptides to each CD2AP SH3 domain |
| **Data were obtained by ITC. Two wild-type RIN3 epitopes (e1 and e2) were tested, and mutant versions of RIN3e2, the tighter binder. Affinities are grouped by increasing red colour intensity; the stronger the shade of red, the higher the affinity (the lower the Kd value). Color groups represent Kd ranges in half-log (μM) increments (e.g., strongest red: >0–0.5 log Kd (μM) = Kd range of >1–3.16 μM). TLQ, too low for quantification (of accurate affinities); wt, wild-type sequence; point mutants of RIN3e2 are also shown.** |
| **RIN3 epitopes** | **Peptide sequence** | **Length (a.a.)** | **Kd ± SE (μM)** |
|---|---|---|---|
| **| | **| **SH3-1** | **SH3-2** | **SH3-3** |
| e1 | wt | AKKNLPTAPPPQVSKQ | 16 | 8.9 ± 0.2 | 10.0 ± 0.2 | 39.8 ± 0.1 |
| | | NLTPAPPERRKK | 10 | 13.5 ± 0.1 | 10.7 ± 0.3 | 84.8 ± 0.3 |
| | P457 → A | TAKQQVAPPSPPKRK | 16 | 16.1 ± 0.1 | 2.1 ± 0.8 | 6.7 ± 0.3 |
| | | P459 → A | 11 | 10.4 ± 1.3 | 15.1 ± 0.9 | 57.5 ± 0.06 |
| | R462 → A | TAKQPPVPPPPKKRIS | 16 | 126.0 ± 1.7 | 19.9 ± 2.6 | 68.0 ± 0.24 |
| | | R462 → L | 16 | 22.2 ± 0.3 | 3.1 ± 0.4 | 6.9 ± 0.02 |
| | | R462 → V | 16 | TLQ | TLQ | TLQ |
| | | R462 → I | 16 | TLQ | TLQ | TLQ |
| | I464 → A | TAKQPPVPPPKKRAS | 16 | 1.9 ± 0.1 | 1.6 ± 0.2 | 11.8 ± 2.0 |

**Color key**

- **Kd range (in log μM)**
- **Colour key**
- >0 to 0.5
- >0.5 to 1.0
- >1.0 to 1.5
- >1.5 to 2.0
- >2.0 to 2.5
- >2.5 (TLQ)
weakened SH3-3 binding nearly 2-fold. However, inspection of the crystal structure indicated that a water molecule modeled with high confidence mediates hydrogen bonding between the main-chain nitrogen of Ile-466 in RIN3 and SH3-2 (Fig. 8D). In fact, it appears to stabilize Asp-125 in the SH3 domain, which participates in the key salt bridge to Arg-462 of the RIN3 motif residue. This backbone hydrogen bond probably drives binding rather than the side chain, explaining the apparent discrepancy. Moreover, the extended interaction involving Ile-466 may contribute to the generally higher affinity of RIN3e2 to CD2AP SH3 domains than RIN3e1. Although the Ile side chain docked well on the hydrophobic patch, we acknowledge that crystal packing could have influenced its positioning.

In the SH3-2-RIN3e2 complex, SH3-2 Phe-117 and Phe-161 in the binding groove make hydrophobic contacts with RIN3 Pro-457, whereas Pro-459 interacts with SH3-2 Phe-161 and Tyr-119. RIN3 Arg-462 forms the characteristic salt bridge to SH3-2 Asp-125 and hydrophobic contacts with the aliphatic portion of its side chain along Trp-145 while also in polar bonding distance to Glu-126 and a backbone main-chain carbonyl oxygen in SH3-2. To evaluate the importance to SH3-2 Trp-145 binding of the hydrophobic portion of the RIN3 Arg-462 side chain, rather than electrostatic interaction with its guanidinium group, as shown previously for a Grb2 SH3 interaction (25), additional Arg-462 substitutions were tested by ITC. Arg was replaced by Ala, Ile, or Val, and the effects on binding were measured (Table 2). Typically, removal of the crucial salt bridge

FIGURE 8. Crystal structures of CD2AP SH3-1 and SH3-2 domains in binary complexes with RIN3 peptides. CD2AP SH3-1 in complex with RIN3e1 (left panels) and CD2AP SH3-2 in complex with RIN3e2 (right panels) are depicted as follows. A and C, electrostatic potential surface representation; RIN3 peptides are shown in stick representation and colored by element (white, carbon; blue, nitrogen; red, oxygen). C, close-up views of RIN3 peptide docking. These two views were generated by rotating the view in A by −30° around the y axis, and 20° around the x axis. B, refined 2Fo − Fc electron density maps (green mesh) of docked RIN3 peptides (SH3 surface omitted) contoured at 1.0 electron/Å³ illustrate the data quality. RIN3e1 (orange sticks) was modeled at 1.65 Å resolution (left), and RIN3e2 (blue sticks) was modeled at 1.11 Å resolution (right). RIN3 residues critical for the interaction are labeled. D, further detail of RIN3e2 (pink sticks) binding to SH3-2 (yellow sticks) within 2Fo − Fc electron density, a snapshot from the molecular graphics program COOT (35). Ile-466 docks onto a distal hydrophobic patch flanked by Val-141, Leu-154, and Leu-156 on the SH3 domain. A water oxygen atom (H₂O) stabilizes the interaction between the backbone amide nitrogen of Ile-466 in the RIN3e2 peptide and the SH3-2 residue Asp-125, which bonds to the key RIN3 motif residue Arg-462. The two alternative partial side chain conformations of this Arg are visible. E, Web logos of strong and medium binders from the PxPxPR array screen compared with non-binders (Fig. 3), created using the online tool (67).
CD2AP SH3 Domain Recognition Preferences

Table 3: X-ray data collection, processing, and refinement parameters for two CD2AP SH3-RIN3 structural complexes

| Parameters                                      | Complex 1  | Complex 2  |
|------------------------------------------------|------------|------------|
| Structures deposited                            | CD2AP SH3-1/RIN3e1 | CD2AP SH3-2/RIN3e2 |
| Protein/peptide PDB code                        | 4WCI       | 3U23       |
| Data collection and processing                  |            |            |
| Diamond Light Source beamline                   | I04        | I03        |
| Detector                                        | Pilatus 2M  | Pilatus 6M  |
| Wavelength (Å)                                  | 0.9173     | 0.97625    |
| Space group                                     | C2         | C2         |
| Unit cell parameters                            |            |            |
| \(a, b, c\) (Å)                                 | 95.53, 55.36, 64.56 | 53.13, 32.04, 39.18 |
| \(\alpha, \beta, \gamma\) (degrees)            | 90.00, 131.7, 90.00 | 90.00, 92.56, 90.00 |
| Resolution range (Å)                            | 48.19–1.65  | 27.43–1.11  |
| Outer shell (Å)                                 | 1.74–1.65   | 1.2–1.11    |
| Mean I/\(\sigma\)                               | 6.6 (2.4)   | 15.0 (2.9)  |
| Completeness (%)                                | 97.7 (93.0) | 92.8 (74.5) |
| \(R_{free}\) (%)                               | 13.2 (47.9) | 5.2 (34.1)  |
| Refinement and validation                      |            |            |
| \(R_{free}\) (%)                               | 19.2 (27.8) | 15.1 (33.7) |
| Highest resolution bin (Å)                      | 21.9 (32.1) | 16.7 (29.1) |
| Root mean square deviation                      | 1.70–1.65   | 1.14–1.11   |
| Bond angles (degrees)                           | 1.01        | 2.27        |
| Bond lengths (Å)                                | 0.007       | 0.025       |
| Ramachandran plot (%)                           |            |            |
| Favored regions                                 | 98.61       | 98.31       |
| Allowed regions                                 | 1.39        | 1.69        |
| Outliers                                        | 0           | 0           |
| Molprobity score; percentile                    | 1.03; 100th | 1.06; 97th  |

Additional data

No. of binary complexes in a.u. | 3 | 1

Discussion

The multifunctional adaptor protein CD2AP binds multiple signaling partners. Through motif screening, we have identified further potential binders and characterized RIN3 as a novel, functional CD2AP interactor, co-recruited to Rab5a-positive endosomes. Although we report the first direct interaction, since this study began, both RIN3 and CD2AP have been implicated in the same “biological space” in several independent settings. First, two recent genome-wide association study reports based on single nucleotide polymorphism (SNP) data uncovered CD2AP as a novel highly associated risk factor gene in LOAD (19, 20), and later RIN3 from a meta-analysis (49), thus potentially connecting them functionally in LOAD. To date, 21 genes have been strongly associated with LOAD, including endocytosis genes, such as amphiphysin II (BIN1). An amphiphysin II-RIN3 interaction is involved in vesicle trafficking in the early endocytic pathway (21), and it is undoubtedly relevant that amphiphysin II can also bind to CD2AP (50).

Second, both RIN3 and CD2AP were among 21 proteins pulled down from kidney cells by the SH3-5/6 domains of the large adaptor protein TUBA (51). TUBA contains six SH3 domains and has guanine nucleotide exchange factor activity toward CDC42. It binds dynamin, providing a link in endocytosis to actin and actin regulatory proteins (e.g. Ena/VASP and N-WASP) as CD2AP binds, bundles, and caps actin (6, 15, 52). N-WASP was also present in the TUBA complex(es) along with CIN85, not surprising given its structural and functional overlap with CD2AP, as well as tight junction proteins ZO-1 and ZO-2. Interestingly, ZO-2 emerged as a hit in our peptide array screen (Fig. 3) and thus can bind to CD2AP SH3 domains directly. The proteomics study (51) did not ascertain whether or not TUBA interacted directly or was merely present in associated complexes. Therefore, a strong possibility remains that a CD2AP-RIN3 and/or a CD2AP-ZO-2 co-complex bind to TUBA, or indeed a multiprotein complex forms between several of these newly identified interacting proteins. Alongside ZO-2 and RIN3, other physiological interaction partners may feature in the array screen hits (Fig. 3), but these require further functional exploration. For instance, SYNJ2 but not SYNJ1, both of which were screen hits, was reported to interact functionally with the CD2AP homologue CIN85 (53).

Third, RIN3 was identified as a risk factor for Paget disease of bone (54), with a P386S polymorphism in RIN3e1, one of sev-
eral in the proline-rich region suggested to be functional (55). Furthermore, most recently, RIN3 has been linked to modulation of bone mineral density of the lower limbs (56). CD2AP has also been associated with the bone disorder Kashin-Beck disease (57). Thus, it is tempting to speculate that, aside from the specific environmental factors contributing to each, there may be some overlapping etiologies between these diseases.

CD2AP SH3-1 was previously crystallized with peptides derived from CBL-b and CD2 (45) as heterotrimers, with two SH3-1 domains clamped around a single CBL-b or CD2 peptide. However, crystals containing CD2 heterodimers in class II orientation were also reported. The SH3-1 domain from CIN85 and β-PIX, each co-crystallized with the same CBL-b peptide, also formed heterotrimers (58). Subsequently, solution data were inconsistent with heterotrimer formation; rather, CIN85 SH3-1 most likely supported either class I or II binding in independent heterodimers (59). Most recently, a detailed biophysical study of the solution state revealed important contrasts between CIN85 and CD2AP (60). CD2AP appeared only to form heterodimers with CD2 or CBL-b peptides. Whereas a CD2 peptide bound to both CD2AP and CIN85 SH3-1 domains purely in class II orientation, CBL-b peptide could adopt class I or II arrangements with both. However, heterotrimer formation was consistent only with the CIN85 data and not CD2AP. In our study, peptides (from CBL, ALIX, and RIN3) bound with 1:1 stoichiometry, and the crystal structures confirmed heterodimers with class II orientation for both RIN3 peptides, consistent with ITC data. This is explainable by the lack of Lys or Leu in position 2 (i.e., P_K[L]PxPR), both of which have been proven to enable distinct partner peptides to bind in the reverse class I orientation under certain conditions (45). In addition, Arg in motif position −2, which could promote class I binding (58, 60), is absent.

The CD2AP SH3-2 structure presented herein is the first crystal structure of this domain and consequently the first high-resolution structural view of it in complex with a partner peptide; the NMR solution structure of SH3-2 alone has previously been reported (61, 62). Our crystal structure was recently used in the reconstruction of an elegant, three-dimensional model of tetrameric full-length recombinant CD2AP in density maps generated by electron microscopy (EM) (63). The EM model suggests a conformational arrangement in which all SH3 domains present their binding sites accessibly. This would potentially enable SH3 interactions of individual monomers with their multiple, varied partners simultaneously, as discussed similarly in the context of CIN85 clustering (53). In short, CD2AP probably acts as a major hub in cell signaling. Consequently, how cooperativity or synergy of binding and avidity effects between the three SH3s might influence their physiological affinities and interaction dynamics is an important area of further study.

The inherent plasticity of the CD2AP SH3 domains, which is possibly a general characteristic of protein-binding modules (64), implies that there is no single, exact motif for each SH3 but rather a family of related motifs. These typically share a core of critical residues (i.e., Px(PxA)xxR) but diverge in the type of residue in flanking positions and their strengths of interaction. Thus, the sequence context of the binding partner dictates exactly which residues will fully determine its interaction potential. A corollary of this is the difficulty of predicting precisely which potential motif-containing proteins bind to a given SH3. On a related note, differential residue tolerances with each CD2AP SH3 were revealed by ALIX peptide permutation arrays (e.g., FxPxXR bound only to SH3-2 but not SH3-1 or SH3-3; Fig. 4), illustrating that selectivity can be achieved between similar SH3 domains.

Our study highlighted the usefulness of (20-aa) permutation arrays for correctly delineating all recognition determinants, because Pro in motif position 3 is interchangeable with Ala with respect to CD2AP SH3 binding. This vital information is undetectable in a simple Ala scan and hence explains why, historically, the importance of motif position 3 has often been overlooked by P → A mutagenesis studies in various experimental contexts. The permutation arrays also indicated that motifs lacking Pro or Ala in this position generally should not bind to CD2AP. This was indeed shown for the non-binding proline-rich region 1, PRR-1, in B cell linker protein, BLNK (or SLP-65), also a hit in our array screen, unlike its PRR-2 and -3, which both contain PxPxPR and were necessary and sufficient for an in vivo interaction (65). CIN85 has also been reported as a BLNK interactor (66).

Exactly why the SH3-1 domain bound to more partner peptides than SH3-2, such as those from CLD23, UBP51, and CSTF2, based on the same motif is not clear, given that their interactions with CBL-derived peptides exhibited a similar pattern (Fig. 2). However, the underlying reasons point toward subtle differences in physicochemical properties of the two SH3 binding surfaces governing specificity. It is noteworthy that −34% of the screened peptides failed to bind to any CD2AP SH3 domain, although each contained a PxPxPR motif. Multiple factors may contribute to such “negative selection.” For instance, Weblog (67) analysis revealed a higher Gly and Pro content around the motif residues in non-binders compared with the medium/strong SH3-1 binders (Fig. 8E). This is rather paradoxical but implies that too little or too much rigidity could hamper adoption of the conformational state(s) optimal for binding. In line with this, the population size of preformed peptide conformers competent for binding to a partner domain surface would appear to play a decisive role in the potency of protein-peptide interactions, as demonstrated very recently for a Gab peptide interaction with Grb2 SH3-C (68). Furthermore, in the non-binding group, negatively charged side chains (Asp and Glu) are prevalent close to the critical Arg of the recognition motif but absent in binders, implying that local charge repulsion negates this positive charge required for tight binding.

The third SH3 domain is a curiosity because it bound very few of the screened peptides compared with SH3-1 and SH3-2, which displayed greater cross-similarity; its exact binding preferences are the least understood. However, SH3-3 is unique in being linked to a disease-associated mutation; a polymorphism (K301M) in SH3-3 may play a role in hereditary focal segmental glomerulosclerosis (5). Lys-301 lies outside of the canonical binding groove and so would not be expected to interfere directly with peptide ligand docking, raising questions as to its exact role. We found that SH3-3 had generally lower affinity for ARALIX peptide binding required a very extended sequence (Fig. 5), implicating additional contact points on the SH3 domain, yet
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despite multiple attempts, we could produce no crystal structure to visualize this. An SH3-3 structure has been solved by NMR in apo form (69) and as a complex bound to ubiquitin ($K_d \sim 130 \mu M$) (70, 71). Ubiquitin docks partially across the P(A)xPR peptide-binding groove in SH3-3 but also extends to the distal hydrophobic pocket (equivalent to that observed in the SH3-2-RIN3e2 complex); however, no prolines are present. This is probably the same site occupied by the extended ALIX motif. Subtle plasticity in the SH3 domain enabling it to accommodate aliphatic and other side chains, a general property discerned from our array data, is thus exploited in the ubiquitin interaction.

In summary, we have identified putative CD2AP SH3 binders with an in vitro screen using prior knowledge of recognition requirements and elucidated varied and distinct binding preferences of the three CD2AP SH3 domains with CBL, ALIX, and the novel partner RIN3. Finally, we have characterized by detailed biochemical and biophysical methods CD2AP-RIN3 interactions, revealing that RIN3 recruits CD2AP to early endosomes via two binding sites. Details of their biological interplay as well as the significance and relative importance of each SH3 domain to this and other interactions awaits further investigation.

Author Contributions—P. C. S. and S. M. F. designed the experiments, analyzed the results, and wrote the paper, with substantial contributions from K. H. K. and S. K. E., M. J., T. K., and P. C. S. performed peptide microarray and ITC experiments. P. C. S., E. R., M. J., J. R. C. M., S. K., and F. v. D. performed peptide microarray and ITC experiments. P. C. S., E. R., M. J., T. K., and P. C. S. contributed. B. R. and K. H. K. designed, performed peptide microarray and ITC experiments. P. C. S., E. R., M. J., T. K., and P. C. S. contributed. B. R. and K. H. K. designed, performed, and analyzed the results of immunofluorescence experiments, with P. C. S. contributing. B. R. and K. H. K. designed, performed, and analyzed the results of immunofluorescence experiments. All authors reviewed the results and approved the final version of the manuscript.

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