Structural determinants for EB1-mediated recruitment of APC and spectraplakins to the microtubule plus end

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EB1 is a member of a conserved protein family that localizes to growing microtubule plus ends. EB1 proteins also recruit cell polarity and signaling molecules to microtubule tips. However, the mechanism by which EB1 recognizes cargo is unknown. Here, we have defined a repeat sequence in adenomatous polyposis coli (APC) that binds to EB1’s COOH-terminal domain and identified a similar sequence in members of the microtubule actin cross-linking factor (MACF) family of spectraplakins. We show that MACFs directly bind EB1 and exhibit EB1-dependent plus end tracking in vivo. To understand how EB1 recognizes APC and MACFs, we solved the crystal structure of the EB1 COOH-terminal domain. The structure reveals a novel homodimeric fold comprised of a coiled-coil and four-helix bundle motif. Mutational analysis reveals that the cargo binding site for MACFs maps to a cluster of conserved residues at the junction between the coiled coil and four-helix bundle. These results provide a structural understanding of how EB1 binds two regulators of microtubule-based cell polarity.

Introduction

The microtubule cytoskeleton, whose minus ends in most cells are located near the centrosome and plus ends extend toward the cell periphery, is involved in the long range transport of membrane organelles, mRNAs, protein complexes, and chromosomes (Karcher et al., 2002). To achieve these functions, microtubules serve as tracks for motor proteins that move along the microtubule surface. A second and more recently discovered type of intracellular transport is driven by the inherent polymerization of tubulin at the microtubule plus end (Desai and Mitchison, 1997). Several “plus end–tracking” proteins have been described that continually bind to the growing plus end of the microtubule and then dissociate, after a temporal delay, from the microtubule shaft. By tracking along the tips of microtubules, plus end–tracking proteins can be delivered to the cell cortex. Cortical polarity factors also can bind to plus end–tracking proteins, leading to the capture and selective stabilization of the microtubule and communication with the actin network (Korinek et al., 2000; L. Lee et al., 2000; Miller et al., 2000). Such interactions between microtubule plus ends and the cortex have been shown to be important for orienting the microtubule network, which can polarize the cell for migration or rotate the mitotic spindle during asymmetric cell divisions (L. Lee et al., 2000; Lu et al., 2001; Etienne-Manneville and Hall, 2003).

The two best characterized microtubule plus end binding proteins are Clip-170 and EB1, which are conserved from yeast to humans. The Clip-170 protein family is characterized by NH2-terminal microtubule-interacting Cap-Gly domains and a COOH-terminal region that recruits various cargos, including the dynein–dynactin complex (Lansbergen et al., 2004), and vesicles (Pierre et al., 1992). EB1 members have a similar bipartite composition: the NH2-terminal domain mediates microtubule plus end localization and a COOH-terminal cargo binding domain that captures cell polarity determinants. In Saccharomyces cerevisiae, an EB1 homologue, Bim1p, links microtubules to the cell cortex by interacting with the polarity determinant Kar9p (Korinek et al., 2000; L. Lee et al., 2000; Miller et al., 2000). In mammalian cells, the tumor suppressor adenomatous polyposis coli (APC) protein plays a
functionally homologous role (Su et al., 1995; Lu et al., 2001). In addition to its role as a scaffold for the regulated degradation of β-catenin (Su et al., 1993), APC plays a role in microtubule-based cell polarity (Rubinfeld et al., 1993; Lu et al., 2001). EB1 was originally identified as an APC binding protein, and this interaction appears to be important for the later pathway (Su et al., 1995; Lu et al., 2001). Several signaling pathways establish polarity via EB1–APC recruitment including: (a) the sequential activation of CDC42, Par6-associated atypical PKC, and GSK-3β, which in turn localizes APC to the leading edge of astrocytes in a wounded monolayer (Etienne-Manneville and Hall, 2003; Kodama et al., 2003), (b) an NGF-induced axon growth pathway (Zhou et al., 2004), (c) the Rho-activated mDia recruitment of EB1 and APC in fibroblast cell migration (Wen et al., 2004), and (d) a microtubule actin cross-linking factor (MACF)–based recruitment of EB1 and APC to the muscle–tendon junction in Drosophila melanogaster (Subramanian et al., 2003). The EB1–APC interaction also is likely to be critical for tumorigenesis/metastasis of intestinal epithelial, probably through a loss of the normal polarity of the epithelia. The majority of APC mutations in human tumors involve deletion of its EB1 binding domain (Powell et al., 1992; Su et al., 1993).

Previous biochemical studies have shown that the COOH-terminal half of EB1 is sufficient for binding APC (Askham et al., 2000; Bu and Su, 2003). Likewise, in yeast, the corresponding region of Bim1p is used for Kar9p binding (Miller et al., 2000). Overexpression of the EB1 COOH-terminal domain also produced a dominant-negative phenotype in cell polarization (Wen et al., 2004; Zhou et al., 2004). Collectively, these studies establish EB1’s COOH terminus as a cargo binding domain and a key determinant in search and capture processes. However, the structural mechanism by which EB1 recognizes its cargo is unknown. Here, we have identified a repeat motif in APC that binds EB1 and found a similar motif in the MACF family of spectraplakins, which allows these proteins to bind to EB1 and track along microtubule plus ends in vivo. We also solved the X-ray crystal structure of EB1’s cargo binding domain and mapped the residues involved in MACF binding by alanine scanning mutagenesis. These results provide a structural understanding of how EB1 recruits cargo to the microtubule plus end.

**Results**

**The EB1 COOH-terminal domain binds to a repeat motif in APC and MACFs**

We first set out to define a minimal domain in human APC that interacts with the human EB1 COOH-terminal domain. Previous studies using a two-hybrid screen and in vitro binding revealed that the COOH-terminal 170 amino acids of APC can interact with EB1 (Su et al., 1995; Askham et al., 2000). To gain further insight on this APC region, we performed a detailed sequence analysis and identified four cryptic repeats of 21–25 amino acids (designated repeats I–IV), which are characterized by conserved prolines, serines, threonines, and positively charged residues (Fig. 1 B). The highest degree of homology was observed between repeats III and IV, which displayed 44% amino acid identity and 52% homology. To determine if all or a subset of these repeats constituted functional EB1 binding sites, we constructed several APC-GFP fusion constructs representing all contiguous combinations of the repeats and tested them for binding to a GST-EB1 column. APC constructs containing repeats III and IV were retained on this column, but proteins lacking the repeat III–IV pair failed to bind, as did the single III or IV repeat proteins (Fig. 1 A). Mapping the EB1 binding site to repeats III and IV (APC2781-2820) corroborates data from Bu and Su (2003), who delineated an overlapping APC region for EB1 binding (APC2781-2820). The minimal construct defined by

![Figure 1](image-url)

**Figure 1. Delineation of the EB1–APC and EB1–MACF2 interaction regions.** (A) Summary of EB1 binding assays. APC constructs are delineated in green, MACF2 in blue. (B) Alignment of APC repeat regions I–IV. Identity is indicated in red, homology in orange. (C) Alignment of the EB1 binding motif of human APC, human MACF2, murine MACF2, and Drosophila Shot, with conservation colored as in B. Asterisks denote tentative PKA (green) and CDC2 (blue) phosphorylation sites. (D) Coomassie-stained polyacrylamide gel showing MACF21595–1637 retention on a GST-EB1185–255 column. (E) GFP-APC2744–2843 and MACF21596–1637 compete for binding to GST-EB1185–255. 10 μl GST-EB1185–255–bound glutathione Sepharose beads were preincubated with 0, 138, 225, and 550 nmol MACF21596–1637 for 30 min before incubation with 550 nmol GFP-APC2744–2843 for an additional hour. Resin was washed three times in batch with wash buffer, and the amount of bound GFP-APC2744–2843 was analyzed by Coomassie-stained SDS-PAGE.
Bu and Su (2003) compromises the COOH-terminal 11 amino acids of repeat IV, which, although retaining EB1 binding, may result in decreased affinity. Additional support for EB1 binding to this region is derived from phosphorylation and phosphomimetic mutation studies of residues within repeats III and IV. In these studies, mutation of the CDC2 and PKA phosphorylation sites (Fig. 1 C) reduced or ablated EB1 binding (Askham et al., 2000; Nakamura et al., 2001; Bu and Su, 2003).

We investigated whether or not any other proteins have repeat motifs homologous to those in APC that bind EB1. After performing a BLAST search (Altschul et al., 1990) using APC repeats III and IV, we identified a similar sequence (38% identity and 57% homology; Fig. 1 C) in the human and mouse spectraplakin family member MACF2 (Sun et al., 2001). Previous work has shown that ACF7 (a mouse MACF) binds to the plus ends of microtubules in mammalian tissue culture cells, but an interaction with EB1 was not investigated (Kodama et al., 2003). Like APC, MACFs are large, multidomain proteins characterized by tandem, NH₂-terminal calponin homology domains that mediate binding to actin filaments, a plakin domain, a central rod-like region composed of dystrophin-like spectrin repeats that typically spans 1,000–4,000 residues, and a COOH-terminal region comprised of two EF-hand motifs, a GAS2/GAR2 domain that confers microtubule association (Sun et al., 2001), a series of quadruplet repeats typified by the sequence GSRA, and a final 50–100 residues of unknown function (Leung et al., 1999). The APC-like repeat motifs that we identified are found in this very COOH-terminal region of human MACF2, the murine MACF2, and the Drosophila MACF Shot (also known as Shortstop or Kakapo). Unlike APC, which has four repeat motifs, the MACFs only have two repeats that are more similar to APC repeats III and IV than to I and II.

We tested whether or not the dual MACF repeat is indeed capable of binding to EB1. We purified the human MACF2 repeat (MACF2₁₅₉₅-₁₆₃₇) and tested its binding to GST fusions of full-length EB1 or its COOH-terminal (EB1₁₈₅-₂₅₅) domain prebound to glutathione-Sepharose. Both full-length GST-EB1 (not depicted) and GST-EB1₁₈₅-₂₅₅ retained the MACF2₁₅₉₅-₁₆₃₇ peptide, whereas GST alone did not (Fig. 1 D). Thus, like APC, the MACF2 repeat motif is capable of specifically binding to the EB1 COOH-terminal domain in vitro. Additionally, competition experiments using GST-EB1₁₈₅-₂₅₅ prebound to glutathione-Sepharose, fixed levels of GFP-APC₁₇₄₄₋₂₈₄₁, and increasing level of MACF2₁₅₉₅-₁₆₃₇ revealed that MACF2 could compete with APC for binding to the EB1 COOH-terminal domain (Fig. 1 E).

**DmEB1 associates with the Drosophila spectraplakin Shot**

Next, we wished to determine if the MACF interaction with EB1 also occurs in vivo. We pursued this question using Drosophila S2 cells because DmEB1 can be very efficiently depleted in these cells by RNAi (Rogers et al., 2002). The solitary MACF homologue in Drosophila is Shot, which is expressed as many different isoforms. Shot was initially isolated in a screen for proteins involved in integrin-mediated adhesion and has subsequently been found to play a critical role during axon extension as a cytoskeletal cross-linker (Gregory and Brown, 1998; Lee and Kolodziej, 2002). Additional work demonstrated that Shot recruits DmEB1 and APC1 to the muscle–ten don junction to promote microtubule assembly, but it was not investigated whether the Shot–DmEB1 association was direct or indirect (Subramanian et al., 2003). In S2 cells, DmEB1 localizes to the plus ends of microtubules in a “comet-tail” pattern throughout the cell cycle as well as to centrosomes and spindle poles (Rogers et al., 2002). When cells were double stained for DmEB1 and Shot, we found that the distribution of the proteins overlapped but were not identical (Fig. 2, A–C). Shot antibodies labeled microtubules along their entire length but were enriched at their tips and also stained small punctae throughout the cell. However, close examination of the microtubules with double staining of Shot and DmEB1 revealed a terminal “cap” of DmEB1 (~1 μm long) that lacked Shot followed by a region of overlap of both proteins in 106 out of 120 microtubules examined (Fig. 1 C). These results indicate that DmEB1 and Shot colocalize on microtubules but also maintain individual zones of localization, which is similar to observations made for EB1 and APC in mammalian cells (Juwana et al., 1999; Askham et al., 2000; Mimori-Kiyosue et al., 2000; Nakagawa et al., 2000; Barth et al., 2002).

We expressed Shot (B isoform) fused to a COOH-terminal GFP and observed its localization in living S2 cells using spinning disk confocal microscopy. As predicted by immunofluorescence, Shot-GFP associated with microtubules, but exhibited two types of dynamic behaviors. On dynamic microtubules, Shot-GFP tracked along with growing microtubule plus ends at a velocity of 4.0 ± 0.8 μm/min (the rate of microtubule elongation in GFP-tubulin–expressing cells [Rogers et al., 2002]). This tip-associated Shot-GFP exhibited a comet-tail distribution, but unlike the short (~2 μm) comet tails of EB1, the length of the microtubule decorated by Shot-GFP was often >5 μm (Video 1, available at http://www.jcb.org/cgi/content/full/jcb.200410114/DC1). Association of Shot-GFP with microtubule plus ends was also observed in mitosis, particularly on the halo of astral microtubules that emanated toward the cortex. A second population of microtubules associated with Shot-GFP along their length and were stable over extended (>20 min) periods of observation, exhibiting neither growth nor shrinkage typified by dynamic instability. Such highly stable microtubules were never observed in S2 cells expressing GFP-tubulin alone (unpublished data). Thus, overexpression of Shot-GFP compared with wild-type cells is likely to account for this suppression of microtubule dynamics. In support of this idea, when Shot-GFP was expressed at very high levels (2 d after transfection), exaggerated microtubule bundling and a complete suppression of microtubule dynamics was observed (Lee and Kolodziej, 2002; unpublished data).

We tested if EB1 was required for the plus end tracking of Shot-GFP by depleting EB1 (<1% of wild type) using RNAi (Rogers et al., 2002). RNAi-treated cells expressing GFP-tubulin under a constitutive actin promoter were double labeled for Shot and DmEB1 to identify unambiguously the DmEB1-depleted cells. Control cells showed colocalization of Shot at microtubule plus ends. In EB1 RNAi-treated cells, Shot never associated with microtubules (n = 900 cells), either at...
their tips or along their length (Fig. 2 E). Instead, Shot immuno-
fluorescence staining revealed a punctuate pattern with no
discernible microtubule association. In contrast, RNAi deple-
tion of D-Clip-190 (another plus end–tracking protein) did not
alter Shot’s microtubule localization (unpublished data). These
results indicate that association of Shot with microtubules re-
quires DmEB1; however, albeit less likely, it cannot be ruled
out that Shot mislocalization in DmEB1 RNAi-treated cells
could be due to the relative undynamic nature of microtubules
after DmEB1 depletion (Rogers et al., 2002).

The crystal structure of EB1’s COOH-
terminal cargo binding domain
To better understand the structural basis of cargo recognition
by EB1, we crystallized its conserved COOH-terminal cargo
binding domain (residues 185–255). Two crystal forms were
obtained: P2₁ (two molecules per asymmetric unit, 2.0 Å reso-
lution) and C222₁ (one molecule per asymmetric unit, 1.8 Å
resolution) (see Materials and methods). Phase information
was obtained for both crystal forms by multiwavelength anom-
alous diffraction (MAD) using the selenomethionine (SeMet)-
derivatized construct V243M, in which a methionine was in-
corporated at the homologous site of a methionine in the EB1
homologue RP1 (Fig. 3 D) to enhance MAD phasing power.
The refinement of the structures yielded the following statistics:
P2₁ crystal form, R = 21.9, R_free = 25.1; C222₁ crystal
form, R = 22.1, R_free = 25.9. Further details on the diffraction
data, phasing, refinement, and model statistics are presented in
Table I. Coordinates have been deposited in the Protein Data
Bank under accession codes 1YIG (P2₁ crystal form) and 1YIB
(C222₁ crystal form).

The crystal structure of the EB1 COOH-terminal domain
reveals a unique homodimeric structure formed by a coiled coil
that folds back at the COOH-terminal region to form a four-
helix bundle (Fig. 3 A). A search of structural domains in the Pro-
tein Data Bank failed to identify a similar homodimeric fold.
The homodimer engages both protomers in the P2₁ asymmetric
unit, whereas a crystallographic two-fold in the C222₁ form
delineates the molecule’s endogenous symmetry. Entirely helical
in structure, the COOH-terminal domain is defined by helices
A and A that span residues 190–230, B and B that span residues
238–248, and a solitary helix C that spans residues 251–254 found in only
one protomer of the P2₁ crystal form. Helices A and A engage
in a coiled coil over residues 190–218, burying conserved
hydrophobic side chains and the aliphatic region of a conserved
lysine (K204). No density was observed for residues 185–189
in either crystal form, thus delineating the start of the dimeriza-
tion domain at residue 190. The remaining portion of A and
A cooperate with B and B’ to form a four-helix bundle
that contributes 1,540 Å² to the total 3,340 Å² of buried surface
area formed in homodimerization. Superimposition of the P2₁
and C222₁ crystal forms along A reveals a plasticity in the rel-
ative twist of the coiled coil and four-helix bundle (Fig. 3 C).
Differences in the organization of the loop bridging helices A and
B are apparent, correlating with this region’s low homol-
ogy and exemplified by a five amino acid insert located within the corresponding region of *Schizosaccharomyces pombe* *Mal3* (Fig. 3 D). The final helix, αC, is ordered in one protomer of the P2 crystal form, most likely due to crystal packing contacts. Due to its disorder in the C222 crystal form and in one protomer of the P2 crystal form, we suspect αC is flexible in solution, but due to its proximity to the MACF binding region described in the following section, it may transit to an ordered state upon cargo binding.

To elucidate tentative sites responsible for cargo interaction, we examined the spatial positions of conserved residues (defined by invariance, 85% identity and 85% homology) across 13 representative EB1 family members from yeast to vertebrates (Fig. 3 D). Under this classification system, conserved residues within the NH₂-terminal region of the coiled coil (αA and αA') are predominantly buried and constitute the hydrophobic core of the coiled coil (Figs. 3 D and 4 B). The highest near-continuous stretch of conservation occurs along the second half of αA, spanning residues 211–229 at the coiled coil–four-helix bundle junction (Fig. 3 D). Amidst this stretch of conservation are the invariant residues F216, Y217, and F218 (Fig. 3, B and D) that form symmetrical, stacked arrays of solvent-exposed hydrophobic side chains that flank each side of the coiled coil. The register of the coiled coil dictates that the spatial FYF motif be comprised of F216 and Y217 from one chain and F218 from the homodimeric mate (Fig. 3 B). Additional conservation at this junction is provided by two solvent-exposed residues of the αB helix: I245 and Y247. This conserved surface is characterized by both a hydrophobic character and a net negative charge contributed by the acidic residues E211, E213, D215, and E225 (Fig. 4 D). The composite architecture of a conserved solvent-exposed region with hydrophobic and charged character implicated this zone as the tentative cargo binding site.

Conserved residues constitute the APC-MACF binding site

To identify the residues in EB1 that contact cargo, we developed a surface plasmon resonance binding assay. We prepared a construct of the MACF21595-1637 EB1 binding region with an
NH$_2$-terminal His-tag and a COOH-terminal motif (GGGLNDIFEAQQKIEWHE) that becomes biotinylated at the lysine residue during expression in *Escherichia coli* (Beckett et al., 1999). MACF2$_{1595-1637}$ peptide was immobilized on a streptavidin sensor chip, increasing concentrations of NH$_2$-terminally His-tagged full-length EB1 were injected, and binding was detected by the increased mass density at the sensor chip surface. Full-length EB1, rather than EB1$_{185-255}$, was used to maximize the binding signal. Analysis revealed that the on-rate and maximal signal increased with increasing concentrations of EB1; however, a very slow off-rate was observed, which is most likely due to bivalent complex formations of two peptides with EB1, which retarded dissociation from the surface. The best fit of these curves was also achieved assuming a bivalent binding reaction (unpublished data). However, the very slow off-rate made it difficult to accurately determine $k_{	ext{on}}$ and the equilibrium binding constant.

To test the individual binding contribution of residues within the 211–229 conserved span, we prepared single alanine mutations, with the exception of Y217 for which both alanine (Y217A) and phenylalanine (Y217F) mutants were generated. Mutant EB1 proteins were analyzed for MACF2$_{1595-1637}$ binding using the surface plasmon resonance assay at a fixed concentration of 2.5 μM and compared with the behavior of wild-type EB1 at the same concentration (Fig. 5 B). A spectrum of association rates was obtained for the various mutants and classified according to effect (altered association, yellow; no binding, red). Only one mutant of a nonconserved glycine (G219A) displayed wild-type association kinetics (Fig. 3 D). Several alanine mutations of charged/polar residues (E211A, K212A, E213A, R214A, D215A, R222A, N223A, and Q229A) displayed wild-type association kinetics. No binding was obtained for the F216A, Y217A, F218A, K220A, F224A, and E225A mutants. All mutants that showed no MACF2 binding were analyzed by gel filtration to determine if homodimerization had been compromised. All mutants eluted at the position of a monomer (Fig. 5 C). I224 is the only residue mutated that is completely buried, situated at the core of the four-helix bundle. These re-
results indicate that destabilization of the four-helix bundle compromises dimerization and that dimerization is essential for MACF2 binding.

Mapping of mutant association results on the crystal structure reveals a cluster of key binding residues along helix αA, the dominant contributors to which are E211, D215, F216, Y217, F218, R222, E225, and L226 (Fig. 6). The FYF motif provides a dramatic contribution to the binding interaction as mutation of any of the three residues to alanine ablates binding. In contrast, the Y217F mutation produces only a modest decrease in binding. Thus, the tyrosine’s hydrophobic Van der Waals contacts, rather than hydrogen bonding by the hydroxyl, makes a dominant contribution, either by contributing direct binding energy to the protein–protein interface or by positioning the spatially flanking phenylalanine residues 216 and 218’, which in turn confer crucial contacts for MACF2 association.

**Discussion**

In this work, we have characterized the binding of EB1 to APC and have identified a new EB1 binding partner, the spectraplakin MACF. We show using X-ray crystallography and structure-function studies that a repeat region shared by APC and MACF binds at an interface created by a four-helix bundle motif in the COOH-terminal domain of EB1. We discuss the implication of these findings for understanding how EB1 molecules attract cargo binding partners to microtubule tips.

**Diverse cell polarity determinants share a common EB1 binding motif**

We have elucidated a motif shared by mammalian APC and several mammalian spectraplakins MACFs. Although these two families bear no apparent sequence homology outside the identified EB1 binding site, they do share a similar composite domain architecture and cellular functions. Both proteins are extremely large (1,000–5,000 amino acids) due to repeated domains in their central regions (β-catenin and axin binding sites in APC [Su et al., 1993; Rubinfeld et al., 1993] and dystrophin-like spectrin repeats in MACFs [Leung et al., 1999]). Both proteins also are characterized by microtubule binding motifs (Smith et al., 1994; Deka et al., 1998; Sun et al., 2001) followed immediately after by the COOH-terminal EB1 recognition motif. In recent studies, both APC and the spectraplakins ACF7 and Shot have been shown to play roles in axon growth, cell polarity, and migration (S. Lee et al., 2000; Lu et al., 2001; Etienne-Manneville and Hall, 2003; Kodama et al., 2003; Subramanian et al., 2003; Wen et al., 2004; Zhou et al., 2004). In these studies, overexpression of the EB1 COOH-terminal domain resulted in a dominant-negative disruption of polarity signaling, indicative that an interaction with full-length EB1 is required for proper binding to microtubule plus ends.
signal transduction/cytoskeletal stabilization by mammalian APC and spectraplakins. GSK-3β, a kinase implicated in cell polarity signal transduction pathways, also has been implicated in the regulation of APC and MACFs (Etienne-Manneville and Hall, 2003; Kodama et al., 2003; Zhou et al., 2004). It is possible that this kinase might, at least in part, control polarity by regulating EB1 interaction because in vitro studies have shown that phosphorylation of APC abrogates binding to EB1 (Askham et al., 2000; Nakamura et al., 2001; Bu and Su, 2003).

Interestingly, the two APC genes in Drosophila have the β-catenin/axin binding domains, but lack the COOH-terminal region in mammalian APC that contains the EB1 binding motif. To date, Drosophila APCs have not been observed tracking along microtubule tips, although DmAPC2 has been found to tether mitotic spindles to cortical actin (McCartney et al., 2001). Furthermore, the spectraplakin interaction with EB1 and microtubule tips appears to be widely conserved in all metazoans. Previous work has implicated the MACF Shot as an EB1-associated factor based on colocalization and coimmunoprecipitation, but a direct interaction was not shown (Subramanian et al., 2003). Here, we have definitively demonstrated that the in vivo association of Shot with the tips of microtubules is mediated by EB1. Thus, we speculate that the spectraplakins may have represented the original metazoan invention of a microtubule tracking, cell polarity factor and that vertebrates subsequently incorporated this feature into APC by duplication and gene integration. Consistent with this idea, the EB1 binding motifs are more similar between vertebrate MACF and APC than they are between MACFs from vertebrates and invertebrates.

In addition to the EB1 interaction, our localization studies suggest other features about the mechanism by which spectraplakins and APC interact with microtubules. Shot colocalizes only with the third of EB1 furthest from the microtubule plus end (Fig. 2 C). This pattern, which is not observed for other proteins such as RhoGEFs (Rogers et al., 2004), suggests that Shot does not coassemble with EB1 on the growing end of the microtubule but recognizes EB1 after a temporal delay, perhaps also suggesting some additional mechanism of regulation.

The EB1 COOH-terminal domain is a structural scaffold for dimerization and cargo binding

Our X-ray crystallographic studies show that the conserved EB1 COOH-terminal domain constitutes a unique coiled coil–four-helix bundle motif that confers both dimerization and cargo binding. Although EB1 and Bim1p were suggested to have a coiled coil domain based on sequence analysis (Rehberg and Graf, 2002), this is the first physical evidence that EB1 proteins are indeed dimers. Dimerization appears to rely not only on the coiled coil but also on the four-helix bundle motif, which provides nearly half of the buried surface area formed in dimerization. Consistent with this idea, destabilization of the four-helix bundle with the I224A mutation abolishes dimerization.

We also used alanine scanning mutagenesis, based on the crystal structure, to uncover a bivalent cargo binding site
in the EB1 COOH-terminal domain. Both subunits of the homodimer appear to contribute residues to the cargo binding site, which is also supported by the finding that abrogating dimerization prevents cargo binding. Although the EB1 COOH-terminal domain is highly negatively charged, many of the key determinants in MACF2 binding are hydrophobic and form a discrete exposed hydrophobic pocket in the center of the domain. In comparison to the high conservation of EB1’s COOH-terminal domain (from yeast to man), the repeat binding motifs found in APC and MACF2 are less well conserved, especially if one compares Drosophila to human. However, general features are the presence of several hydrophobic residues (mainly prolines) and a net positive charge. We speculate that these prolines and other hydrophobics dock into EB1’s hydrophobic pocket, whereas the charged residues may augment the binding interaction by providing a peripheral gasket.

The conservation of solvent-exposed residues in EB1’s COOH-terminal domain extends from vertebrates down to the yeast homologues Bim1 and Mal3. Kar9 is functionally homologous to APC and Shot in its general role of microtubule search and capture (Miller et al., 2000); however, Kar9 does not have a repeat motif architecture similar to APC/spectraplakins (Miller and Rose, 1998). A weak segment of homology with hydrophobic character was reported between regions of Kar9 and the EB1 binding motif (Bienz, 2001), but this site has not been experimentally confirmed. It will be interesting to compare and contrast the molecular basis for the EB1–APC interaction and the Bim1–Kar9 interaction to determine if a similar or unique set of determinants govern these binding interactions.

Our results also help to explain previous mutagenesis results of EB1 that were performed before the availability of a crystal structure. Wen et al. (2004) mutated conserved hydrophilic residues within aa 211–229 in EB1 and found that the mutations E211A/E213A, E211A/D215A, K220A/R222A, and E211A/E213A/K220A/R222A inhibited EB1–APC interaction in vitro and yielded supporting phenotypic behavior in vivo in an mDia-mediated cell polarity pathway. Our results predict the strongest effect for constructs containing the K220A mutation and milder effects for the one with the E213A mutation, which agrees with the observations by Wen et al. (2004). This correlation between EB1 mutants binding to MACF21595-1637 and APC also substantiates the likelihood of a similar binding architecture between these two classes of proteins and the EB1 COOH-terminal domain.

Implications of the EB1 COOH-terminal domain structure for other cargo binding interactions

Several EB1 family members exist in metazoan genomes (four in Drosophila [Rogers et al., 2002] and three in humans [Su and Qi, 2001]). Although the crystal structures reported in this paper describe a homodimeric structure, we cannot rule out the possibility that EB1 heterodimers exist and are used for diverse cargo recognition. The length of the coiled-coil domains and the residues in the COOH-terminal domain that constitute the hydrophobic core and mediate dimer contacts are highly conserved across family members, especially within species (Fig. 3 D). Because each bivalent cargo binding site is formed from residues contributed by each of the EB1 molecules, heterodimers could confer complex, pair-wise cargo recognition motifs. Thus, the possibility of whether or not heterodimers of EB1 can form in vitro and within cells merits investigation.

The EB1 COOH-terminal domain also may contain cargo binding sites other than the one that we defined for APC and MACF at the coiled-coil–four-helix bundle junction. Several proteins, including the dynein–dynactin complex (Valetti et al., 1999; Vaughan et al., 2002), Ncd (unpublished data; Ghoshima, G, personal communication), the microtubule-destabilizing kinesin Klp10A (Mennella et al., 2005), and RhoGEF2 (Rogers et al., 2004), have been reported to track along microtubule tips and associate with EB1. However, only in the cases of the p150Glued dynactin subunit (Askham et al., 2002; Tirnauer et al., 2002; Bu and Su, 2003; Wen et al., 2004) and Klp10A (Mennella et al., 2005) has a direct interaction with EB1 been shown. In the case of p150Glued, deletion studies have suggested that p150Glued and APC use overlapping yet unique binding sites on EB1’s COOH-terminal domain (Askham et al., 2002; Bu and Su, 2003; Wen et al., 2004). The p150Glued binding site maps to EB1’s coiled-coil–four-helix bundle but also requires EB1’s COOH terminus (13 residues not included in our crystallization construct). Thus, it is possible that EB1 contains multiple cargo binding sites, some of which might be independent and others which might be mutually exclusive. The crystal structure reported here should provide a valuable tool for probing these future questions concerning EB1–cargo interactions.
Materials and methods

DNA manipulation, protein preparation, and immunoblot

Human cDNA encoding EB1 was obtained from the American Type Culture Collection and subcloned into the E. coli expression vectors pGEX-6P2 (Amersham Biosciences) and pET28a (Novagen). The EB1 conserved NH2-terminal domain (residues 12–133) and the conserved COOH-terminal domain (residues 185–255) were independently subcloned into pGEX-6P2. A COOH-terminal mutant, V243M, in EB1158525S was created using the PCR sewing method and inserted into pGEX-6P2. Site-directed mutagenesis was conducted on the full-length EB1-pET28a construct using the Quickchange method (Stratagene) to create the point mutants described in Figs. 4–6. APC constructs (residues 2739–2806, 2739–2781, 2739–2759, 2760–2831, 2760–2806, 2760–2809, 2760–2778, 2760–2781, 2760–2806, 2760–2809, 2760–2778, 2760–2781, 2760–2806, and 2760–2809) and EB1 constructs (full length and residues 185–255) were subcloned into a modified pET15 (Novagen) vector to yield an NH2-terminal hepta-histidine tag followed by GFP and the specified region of APC. APC cDNA was provided for recombinant expression (excluding a SeMet growth of the E. coli expression vectors pGEX-6P2) EB1185-255 V243M mutant in which the construct was transformed into B212 (DE3). E. coli for recombinant expression (excluding a SeMet growth of the pGEX-6P2 EB1 constructs) was resuspended in lysis buffer (25 mM Tris, pH 8.0, 200 mM NaCl, and 0.1% β-mercaptoethanol (β-ME)) and lysed using a microfluidizer (Microfluidics, Inc.). 2 mM PMSF was added, and the lysate was clarified by centrifugation at 36,000 g for 10 min, loaded onto a column, washed with 200 column volumes of lysis buffer, and then protein was applied to the column to test for retention. Additional purification of GST-EB1158525S and GST-EB1158525V243M proceeded as follows. GST fusion protein was step eluted using 5 column volumes of lysis buffer supplemented with 50 mM glutathione, pH 8.0. 30 µl of PreScission protease (Amersham Biosciences; 2 mg/ml stock) was added and incubated at 4°C for 12 h, which yielded near complete cleavage. Cleaved fusion protein was concentrated in a Centriprep YM3 (Millipore) and diluted 100-fold into 25 mM Tris, pH 7.0, 0.1% β-ME, and loaded onto SP Sepharose Fast Flow resin (Amersham Biosciences) and eluted using a NaCl gradient of 0–1 M over 20 column volumes. Protein fractions were pooled, concentrated in a Centriprep YM3, and exchanged into storage buffer (10 mM Tris, pH 7.0, 25 mM NaCl, and 0.1% β-ME). All buffers used for the EB1158525S V243M SeMet purification also contained 5 mM methionine as a supplemental reducing agent. Additional purification of GST1H-MACF2159-163-Biotinylation Tag entailed PreScission protease cleavage, followed by binding to Ni2+-nitrilotriacetic acid resin, washing in 25 mM Tris, pH 8.0, 15 mM imidazole, 300 mM NaCl, 0.1% β-ME, and step elution using washing buffer supplemented with 285 mM imidazole. Eluted protein was concentrated and exchanged into 50 mM Tris, pH 8.0, 150 mM NaCl, and 0.1% β-ME using a Centricon YM3. All H2-GFP fusion proteins were purified as follows. Bacterial cell pellets were thawed and resuspended in lysis buffer (25 mM Tris, pH 8.0, 300 mM NaCl, 15 mM imidazole, and 0.1% β-ME), lysed in a microfluidizer, supplemented with 2 mM PMSF, and centrifuged at 36,000 g for 45 min at 4°C. Supernatant was incubated in batch with Ni2+-nitrilotriacetic acid resin for 1 h. Protein was harvested by centrifugation at 200 g at 4°C, loaded into a column, and washed with 100 column volumes of lysis buffer. EB1 mutants were washed with 15 column volumes of lysis buffer supplemented with 15 mM imidazole, and step eluted with lysis buffer supplemented with 285 mM imidazole. All other H2-GFP fusion proteins were eluted using a gradient of 15–300 mM imidazole over 20 column volumes. Eluted H2-GFP fusion proteins were concentrated in a Centricon YM3, exchanged into 10 mM Tris, pH 8.0, 150 mM NaCl, and 0.1% β-ME, frozen in liquid nitrogen, and maintained at –80°C until used.

Drosophila tissue culture and immunofluorescence

Drosophila S2 cells were cultured, treated for RNAi, and processed for immunofluorescence as described previously (Rogers et al., 2002). Primary antibodies used for these studies were murine monoclonal anti-Shot (raised against the antigen Shot LIA) 1450–1909, which was prepared as a PreScission protease cleavage product from a GST fusion (Amersham Biosciences) rabbit anti–Clip-190 (Rogers et al., 1998), and rabbit anti–DmEB1 (Rogers et al., 2002). Secondary antibodies (TMR anti-rabbit, Cy5 anti-mouse, Cy2 anti-rabbit, and Cy3 anti-guinea pig; Jackson ImmunoResearch Laboratories) were used at a dilution of 1:300.

Time-lapse observation of Shot-GFP in living cells was performed by transfection of S2 cells with pUAS-Shot (L) B-GFP (Lee and Koh, 2002) and pMT-GAL4 (Rogers et al., 2004) using the GAL4 transfection reagent (Invitrogen) following the manufacturer’s protocols. GAL4-mediated Shot-GFP was induced with a 2 h induction using 1 µM copper sulfate and imaged using a spinning disk confocal microscope (Solamere Technology Group) mounted on an Axiovert 200 (Carl Zeiss Microimaging, Inc.) equipped with an X/Mega10 intensified charge coupled device camera (Stanford Photonics) driven by QED In Vivo software (QED Imaging).

Crystallization, data collection, data processing, model building, and refinement

The EB1 COOH-terminal domain (residues 185–255) was crystallized using the hanging drop vapor diffusion method at 20°C. 2 µl EB1 COOH-terminal domain at 15 mg/ml was added to 2 µl of mother liquor containing 22% PEG 200, 100 mM sodium citrate, pH 4.6, and equilibrated over 1 M mother liquor. Native crystals were crushed and used to streak-seeds of the EB1 COOH-terminal domain mutant V243M SeMet. The V243M SeMet protein crystallized in the space group P21 with two molecules in the asymmetric unit. Attempts to cocrystallize a complex of EB1158525S with MACF2159-163 using similar crystallization conditions yielded native and SeMet crystals of EB1 (without MACF2159-163 present) in the space group C2221 with one molecule in the asymmetric unit. Crystals were transferred to Paratone-N and flash frozen in liquid nitrogen.

Diffraction data were measured at the Lawrence Berkeley National Laboratory Advanced Light Source beamline 8.3.1 using an ADSC 210 charge coupled device detector and maintained at 100 K using a nitrogen cryo stream. Data were processed and scaled using DENZO and SCALEPACK (Otwinowski, 1993). Heavy atom searches were performed using the CNS package (Brunger et al., 1998). Two selenium sites were found in each crystal form using an automated Patterson heavy atom search method and peak (P2; crystal form) or inflection (C222; crystal form) anomalous wavelength data. The additional two selenium sites for the P2; crystal form were found using an anomalous difference Fourier map. Experimental phase calculations used phases from the two selenium sites found in the initial automated search. The coordinates for these two additional selenium sites were refined using the CNS package and used for subsequent phase determination and map building. MAD phasing used the selenium coordinates and the low energy remote wavelength as reference. Phases were improved by solvent flipping and histogram matching. Model building was done with O (Jones et al., 1991) and refinement was performed with the CNS package and monitored with a random 10% of the data used for cross-validation. Model refinement used the MLH target function, torsion angle molecular dynamics simulated annealing, B-factor refinement, and rebuilding in O with eA-weighted difference Fourier maps. The C222; crystal form structure refinement was extended to the high resolution of the native data set using the MLF target function. Figures were prepared with PyMOL (DeLano Scientific) and GRASP (Nicholls et al., 1991). Coordinates have been deposited in the Protein Data Bank under accession codes 1YIG (P21 crystal form) and 1YIB (C222; crystal form).

Surface plasmon resonance studies

Surface plasmon resonance experiments were conducted using a Biacore 1000 and Streptavidin sensor chip (Biacore). Flow cells were pretreated with 2 x 1 min injections of 1 M NaCl, 50 mM NaOH, and 3 x 1 min injections of regeneration solution (0.25% SDS). Flow cells were normalized and one flow cell was injected with 50 µl of 10 µM H2-MACF2159-163-Biotinylation Tag protein at a 5 µl/min flow rate to yield an increase of 600 response units. The second flow cell was used as a control surface over which identical analyte injections were performed in triplicate. All runs were conducted at a flow rate of 20 µl/min in running buffer (10 mM Tris, pH 8.0, 200 mM NaCl, 0.1% β-ME, and 0.05% Tween-20) at 25°C. Sensor chips were prestablized for 2 min with running buffer. Analyte was injected for 2.5 min followed by 6 min of running buffer for the anal-
ysis of dissociation. The sensor chip was regenerated by injection of 0.25% SDS for 1 min followed by a 5.5-min stabilization of running buffer before the start of the next sensorgram. For full-length EB1 analysis, eight different concentrations of H6-EB1 were injected onto the sensor chip in triplicate (2.4 nM, 10 nM, 39 nM, 160 nM, 630 nM, 2.5 μM, 10 μM, and 40 μM). Binding studies using wild-type and mutant EB1 proteins were conducted at an analyte concentration of 2.5 μM and conducted in triplicate. Data was processed using BIAevaluation 3.2 R1C1 software. Control analyte runs were subtracted from corresponding analyte plus MACF2 peptide runs and calibrated for comparison via baseline adjustment before analyte injection.

**Gel filtration assay**

Full-length human EB1 and EB1 point mutants (F216A, Y217A, F218A, K220A, I224A, and E225A) were diluted in running buffer (25 mM Tris, pH 8.0, 200 mM NaCl, and 0.1% β-ME) to a final concentration of 32 μM. 100 μl of each protein was injected onto a Superdex200 Tricorn analytical gel filtration column (Amersham Biosciences) equilibrated in running buffer and analyzed at a flow rate of 0.3 ml/min, measuring the absorbance at 280 nm.

**Online supplemental material**

GFP-Shot in interphase S2 cells shows microtubule plus end localization (Video 1). Frames for Video 1 were acquired at a rate of one frame per 5 s. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200410114/DC1.

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