

ALIX-CHMP4 interactions in the human ESCRT pathway

John McCullough, Robert D. Fisher, Frank G. Whitby, Wesley I. Sundquist*, and Christopher P. Hill*

Department of Biochemistry, University of Utah, Salt Lake City, UT 84112-5650.

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The ESCRT pathway facilitates membrane fission events during enveloped virus budding, multivesicular body formation, and cytokinesis. To promote HIV budding and cytokinesis, the ALIX protein must bind and recruit CHMP4 subunits of the ESCRT-III complex, which in turn participate in essential membrane remodeling functions. Here, we report that the Bro1 domain of ALIX binds specifically to C-terminal residues of the human CHMP4 proteins (CHMP4A-C). Crystal structures of the complexes reveal that the CHMP4 C-terminal peptides form amphipathic helices that bind across the conserved concave surface of ALIX_{Bro1}. ALIX-dependent HIV-1 budding is blocked by mutations in exposed ALIX_{Bro1} residues that help contribute to the binding sites for three essential hydrophobic residues that are displayed on one side of the CHMP4 recognition helix (M/L/IxxLxxW). The homologous CHMP1–3 classes of ESCRT-III proteins also have C-terminal amphipathic helices, but, in those cases, the three hydrophobic residues are arrayed with L/I/MxxxLxxL spacing. Thus, the distinct patterns of hydrophobic residues provide a “code” that allows the different ESCRT-III subunits to bind different ESCRT pathway partners, with CHMP1–3 proteins binding MIT domain-containing proteins, such as VPS4 and Vta1/LIP5, and CHMP4 proteins binding Bro1 domain-containing proteins, such as ALIX.

cytokinesis | ESCRT-III | HIV | multivesicular body

The ESCRT pathway functions in the budding of HIV-1 and other lentiviruses (1), in the final abscission stage of cytokinesis (2), and in intraluminal vesicle formation at the late endosome or multivesicular body (MVB) (3, 4). Involvement in these seemingly diverse biological processes can be rationalized if the ESCRT machinery encodes membrane remodeling and fission activities that are required to resolve the thin membrane “necks” created during the final stages of virus budding, MVB vesicle formation, and cell division. Although mechanistic details are still lacking, there is increasing evidence that the ESCRT-III proteins may mediate such vesicle extrusion and/or membrane fission activities, possibly in conjunction with the AAA ATPase VPS4 (for example, see ref. 5). Humans express 11 related, but distinct ESCRT-III proteins (termed the CHMP proteins) that can be subdivided into seven different families (CHMP1–7) based on their similarities to one another and to the six ESCRT-III-like proteins in yeast. The different ESCRT-III proteins apparently adopt similar folds (6) and can copolymerize together on membranes, yet have evolved to interact differently with other ESCRT pathway components. For example, only the three human CHMP4 proteins (CHMP4A-C) can bind ALIX (yeast Bro1p), another protein in the ESCRT pathway (7–16).

The ESCRT machinery functions at different membranes, and ALIX plays important roles in targeting the pathway to function in retrovirus budding by binding directly to viral Gag proteins (16, 17) and to function in abscission by binding the midbody protein CEP55 (2). In both cases, ALIX must also bind the CHMP4 proteins, because ALIX point mutations that block CHMP4 binding inhibit HIV-1 budding (10, 11) and abscission (18). Thus, ALIX can serve as an adaptor that recruits CHMP4/ESCRT-III complexes to function at distinct biological membranes. Conversely, CHMP4 proteins can apparently recruit

ALIX to membranes because the membrane-bound Snf7p (yeast CHMP4) brings Bro1p/ALIX to the endosome to function in MVB vesicle formation (19).

In addition to its involvement in HIV budding and cytokinesis, ALIX has been implicated in a variety of biological processes that may reflect other ESCRT pathway functions or possibly ESCRT-independent ALIX functions. These functions include lysobisphosphatidic acid (LBPA) binding (reviewed in ref. 20), endophilin binding (21), receptor trafficking (22–24), endosome distribution (25), cell motility/adhesion (26, 27), apoptosis (reviewed in ref. 28), actin and microtubule binding (26, 29) and regulation of JNK signaling (30). Thus, ALIX appears to play widespread roles in membrane biology and cell signaling. Similarly, ALIX has been implicated in the release of several other classes of enveloped viruses, including hepatitis B virus (31), human parainfluenza virus (32), and possibly Sendai virus (33) (however, see ref. 34). Thus, ALIX may play widespread roles in the release of highly divergent enveloped viruses.

ALIX has three distinct regions: an N-terminal Bro1 domain (residues 1–358), a central “V” domain (362–702), and a C-terminal proline-rich region (703–868). Crystal structures of different ALIX constructs (9, 10, 35) have revealed that the banana-shaped Bro1 domain is organized about a core of tetratricopeptide helical hairpins and that the V domain is composed of two extended helical arms that fold in the shape of the letter V. YP(X_n)L sequence motifs within retroviral Gag proteins bind on the inner face of the second arm of the V domain (10, 17, 35–37). The proline-rich region contains binding epitopes for a number of other cellular factors, including TSG101 (13, 15, 16), endophilins (21), and ALG-2 (38, 39). The Bro1 domain contains binding sites for both HIV-1 NC (40) and the CHMP4 proteins (7–11). Mutations that inhibit CHMP4 binding cluster within an exposed hydrophobic patch on the concave surface of the Bro1 domain, which is thought to be the CHMP4 binding site (9–11). This important interaction has yet to be characterized in molecular detail, however, and, we have therefore mapped the ALIX binding sites on the three human CHMP4 proteins and determined crystal structures of the relevant ALIX_{Bro1}-CHMP4 complexes.

Results

CHMP4-ALIX Interactions. Deletion analyses were used in conjunction with biosensor binding experiments to map the ALIX

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The authors declare no conflict of interest.

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Data deposition: The atomic coordinates and structure factors have been deposited in the Protein Data Bank, www.pdb.org [ID codes 3C3O (CHMP4A), 3C3Q (CHMP4B), and 3C3R (CHMP4C)].

*To whom correspondence may be addressed. E-mail: wes@biochem.utah.edu or chris@biochem.utah.edu.

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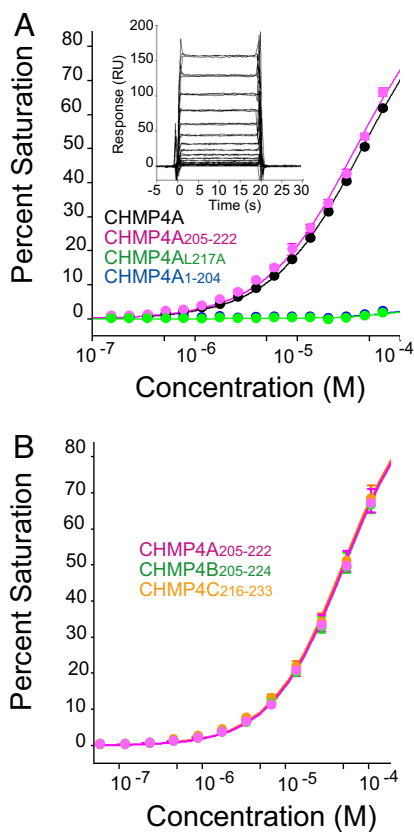


Fig. 1. Mapping the ALIX binding sites of CHMP4 proteins. (A) Binding isotherms and sensorgrams (*Inset*) showing ALIX_{Bro1-V} binding to GST-CHMP4A ($K_D = 30 \pm 11 \mu\text{M}$) (*Inset*), GST-CHMP4A₂₀₅₋₂₂₂ ($K_D = 44 \pm 6 \mu\text{M}$), GST-CHMP4A_{L217A} (binding not detectable), and GST-CHMP4A₁₋₂₀₄ (binding not detectable). The K_D estimates are averages from best fits to single-site binding models (mean \pm SD, $n \geq 3$). The shorter ALIX_{Bro1} construct also bound to CHMP4A₂₀₅₋₂₂₂ ($K_D = 40 \pm 0.6 \mu\text{M}$) and to the longer CHMP4A₁₉₅₋₂₂₂ ($K_D = 40.5 \pm 0.4 \mu\text{M}$) and CHMP4A₁₇₄₋₂₂₂ ($K_D = 36.5 \pm 0.4 \mu\text{M}$) C-terminal constructs with comparable affinities (dissociation constant and error were estimated from a statistical fit of a single binding isotherm derived from duplicate measurements at 10 different ALIX_{Bro1-V} concentrations; data not shown). Error bars are indicated on all biosensor figures, but are often too small to be readily visible. (B) ALIX_{Bro1} binds the C termini of CHMP4A, CHMP4B, and CHMP4C. Binding isotherms showing ALIX_{Bro1-V} binding to immobilized C-terminal peptides from CHMP4A-C. Estimated dissociation constants were: GST-CHMP4A₂₀₅₋₂₂₂, $44 \pm 6 \mu\text{M}$ (mean \pm SD, $n = 6$); GST-CHMP4B₂₀₅₋₂₂₄, $48 \pm 6 \mu\text{M}$ (mean \pm range, $n = 2$); GST-CHMP4C₂₁₆₋₂₃₃, $41 \pm 10 \mu\text{M}$ (mean \pm range, $n = 2$). Binding to a control GST surface was negligible (data not shown).

binding site on CHMP4A. As summarized in Fig. 1A, an ALIX construct spanning the Bro1 and V domains (ALIX_{Bro1-V}) bound both full-length CHMP4A and a minimal C-terminal CHMP4A₂₀₅₋₂₂₂ construct with similar affinities ($30 \pm 11 \mu\text{M}$ and $44 \pm 6 \mu\text{M}$), but did not bind detectably to a CHMP4A construct that lacked the final 18 residues (CHMP4A₁₋₂₀₄) or to a full-length CHMP4A protein that harbored a single point mutation within this region (CHMP4A_{L217A}) (15). Thus, the ALIX_{Bro1-V} binding site maps to the final 18 residues of CHMP4A.

The three human CHMP4 family members have similar but distinct C-terminal sequences, so we tested whether peptides corresponding to the termini of CHMP4B and CHMP4C also bound ALIX_{Bro1-V}. As shown in Fig. 1B, terminal fragments from all three CHMP4 proteins bound ALIX_{Bro1-V} with similar affinities (44, 48, and $41 \mu\text{M}$, respectively), demonstrating that all three CHMP4 proteins encode C-terminal ALIX binding sites. Similar binding data were also obtained for the shorter

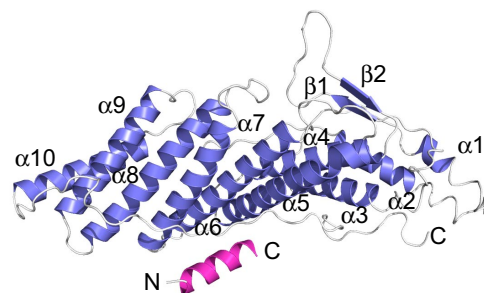


Fig. 2. Ribbon diagram showing the ALIX_{Bro1} domain (blue) in complex with the C-terminal helix from CHMP4A (purple).

ALIX_{Bro1} construct (see Fig. 1 legend), and the Bro1 domain of ALIX, therefore, binds the C termini of all three human CHMP4 proteins.

Crystal Structures of ALIX_{Bro1}-CHMP4 Complexes. Crystal structures of ALIX_{Bro1} in complex with peptides corresponding to the C-terminal binding sites from each of the three human CHMP4 proteins were determined in order to visualize the molecular basis for ALIX-CHMP4 interactions [Figs. 2 and 3 and [supporting information \(SI\) Figs. S1–S3](#)]. All three complexes crystallized isomorphously in space group C2 with a single ALIX_{Bro1}-CHMP4 complex in the asymmetric unit. The CHMP4A-C complexes were refined to resolutions of 2.15, 2.10, and 2.02 Å, respectively, with good geometries and R_{free} values $<30\%$ (Table S1).

As shown in Fig. 2, CHMP4A₂₀₅₋₂₂₂ forms an amphipathic helix that binds across the concave surface of ALIX_{Bro1}, con-

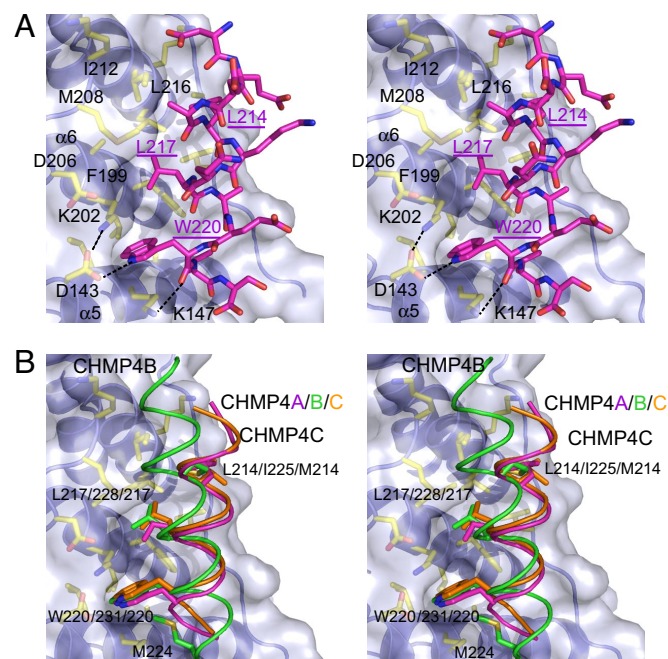


Fig. 3. ALIX_{Bro1}-CHMP4 interfaces. (A) Stereoview of the ALIX_{Bro1}-CHMP4A interface. The CHMP4A helix is oriented N to C from top to bottom, ALIX residues within the binding interface are shown explicitly, dashed lines indicate hydrogen bonds or salt bridges, and key hydrophobic CHMP4 residues are underlined. (B) Stereo-view showing an overlay of the bound CHMP4A-C helices. The orientation is the same as in A, and the three key hydrophobic CHMP4 binding residues are shown in sticks. Note that the CHMP4A (purple) and CHMP4C (orange) helices overlay well, whereas the CHMP4B helix (green) is rotated by $\approx 20^\circ$.

tacting helices 5–7 and the extended C-terminal strand that traverses the domain. ALIX_{Bro1} does not change conformation notably upon CHMP4 binding, although several sidechains in the binding site shift slightly, with the largest adjustment ($\approx 1.5\text{\AA}$) being made by the ALIX Phe-199 ring. The CHMP4A₂₀₅₋₂₂₂ and CHMP4C₂₁₆₋₂₃₃ peptides have similar lengths and sequences and bind ALIX_{Bro1} in very similar fashions, whereas the longer CHMP4B₂₀₅₋₂₂₄ peptide binds at the same site, but in a slightly different fashion (see below).

In all three complexes, important interactions are made by hydrophobic residues located on three successive turns of the CHMP4 recognition helix (CHMP4A residues Leu-214, Leu-217, and Trp-220). The most distinctive contact is made by the indole ring of Trp-220, which binds in a hydrophobic pocket located between helices 5 and 6 (Fig. 3A). The indole nitrogen also forms a hydrogen bond with the conserved ALIX Asp-143 carboxylate, which in turn is buttressed by a salt bridge with the ALIX Lys-202 side chain. Leu-217 binds in an adjacent hydrophobic pocket located between ALIX helices 6 and 7, whereas Leu-214 binds on a more open hydrophobic surface of ALIX helix 6. No other CHMP4A side chains make substantial contacts, and the structures therefore indicate that the three hydrophobic residues of the CHMP4A recognition helices are the primary determinants of ALIX binding and specificity. The terminal carboxylates of CHMP4A and CHMP4C probably also contribute to ALIX recognition, because they make water-mediated interactions with the ALIX Lys-151 side chain.

The ALIX_{Bro1}-CHMP4B₂₀₅₋₂₂₄ Complex. The longer CHMP4B₂₀₅₋₂₂₄ peptide binds ALIX_{Bro1} at the same site and forms an amphipathic helix that places the same three hydrophobic side chains in analogous binding sites. In this case, however, the helix is rotated by $\approx 20^\circ$ relative to the CHMP4A/C helices, which displaces the N and C termini of the CHMP4B helix by $\approx 5\text{\AA}$ and 3\AA , respectively. (Fig. 3B and Figs. S2B and S3). The C-terminal displacement allows the final two CHMP4B residues (not present in the shorter CHMP4A/C helices) to make unique interactions that appear to dictate the helix orientation. Specifically, the CHMP4B Ser-223 hydroxyl caps the helix and hydrogen bonds with the ALIX Lys-147 side chain (whereas ALIX Lys-147 hydrogen bonds with the Trp-220 main-chain carbonyl in the CHMP4A and 4C complexes). The terminal CHMP4B Met-224 residue, in turn, contacts a hydrophobic patch between ALIX helices 5 and 6 (Fig. 3B and Fig. S3B). Thus, the C-terminal helices of all three human CHMP4 proteins bind the same site on ALIX, although the detailed interactions can differ depending on the length of the recognition helix.

Mutational Analyses of the ALIX_{Bro1}-CHMP4A Interaction. Biosensor binding assays were also performed to test the importance of the three conserved hydrophobic residues of the CHMP4 recognition helix. As shown in Fig. 4A, single alanine substitutions of CHMP4A residues Leu-214, Leu-217, and Trp-220 abolished ALIX_{Bro1-V} binding, confirming the energetic importance of all three residues. A cluster of acidic residues is also conserved at the N-terminal end of the CHMP4 recognition helix (Fig. 4B). These residues do not contact ALIX directly in the crystal structures, but do approach basic surface residues Lys-209 and Lys-215, and could therefore also contribute to binding. Mutation of the glutamate residue present in all three human CHMP4 proteins (CHMP4A Glu-209) reduced ALIX_{Bro1-V} binding affinity by twofold, indicating that hydrophilic flanking residues can also contribute to ALIX_{Bro1} binding, albeit modestly.

Discussion

Our biochemical and structural analyses show that ALIX binds amphipathic C-terminal helices on all three human CHMP4 proteins. Three hydrophobic residues on the CHMP4 recognition

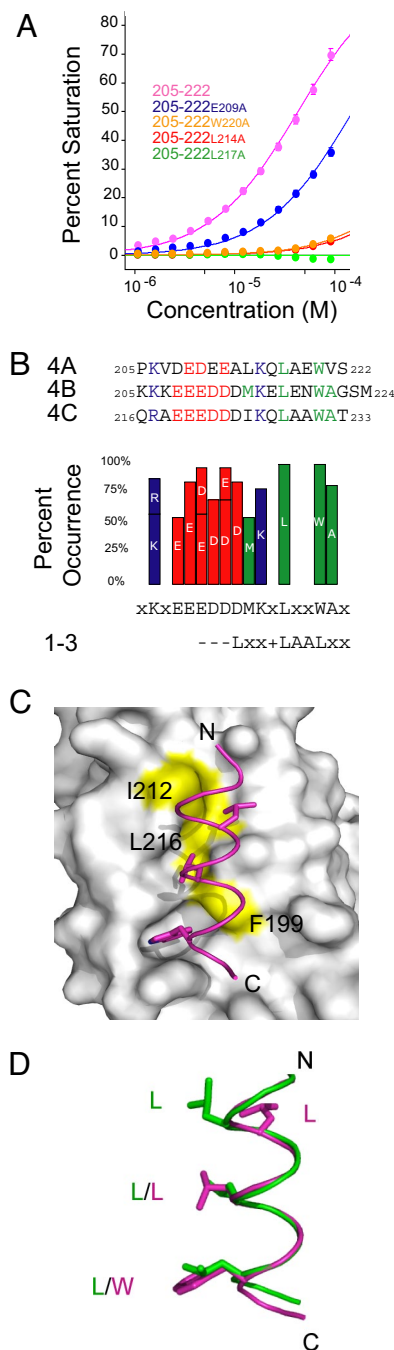


Fig. 4. Molecular recognition in ALIX-CHMP4 complexes. (A) Isotherms showing ALIX_{Bro1-V} binding to immobilized WT CHMP4A₂₀₅₋₂₂₂ and to CHMP4A₂₀₅₋₂₂₂ constructs with the following mutations: W220A, L217A, L214A, and E209A. For the E209A mutant, $K_D = 95 \pm 2 \mu\text{M}$ (dissociation constant and error were estimated from a statistical fit of a single binding isotherm derived from duplicate measurements at 10 different ALIX_{Bro1-V} concentrations.). (B) Sequences of the C termini of human CHMP4 proteins are shown (Upper), together with a bar graph showing sites of >50% identity across metazoan CHMP4 proteins, the resulting CHMP4 consensus sequence (see also Table S2) and the distinct consensus sequence of the C-terminal amphipathic helices of CHMP1–3 proteins (Lower) (47). (C) Model of the ALIX_{Bro1}-CHMP4A interaction, with mutation sites that block ALIX binding and ALIX-dependent HIV-1 budding highlighted in yellow on the ALIX surface. (D) Overlay of the C-terminal recognition helices from CHMP4A (magenta) and CHMP1A (green, PDB 2jq9), extracted from the bound ALIX_{Bro1}-CHMP4A₂₀₅₋₂₂₂ and VPS4A MIT-CHMP1A₁₈₀₋₁₉₆ complexes. The three key hydrophobic residues from each recognition helix are shown explicitly, and the figure emphasizes that the first hydrophobic residue is positioned differently in the two helices.

helices contact ALIX extensively, and their energetic importance was confirmed by mutagenesis. The C-terminal Leu and Trp residues are invariant in metazoan CHMP4 proteins, whereas the first hydrophobic position of the helix can vary between Met, Leu, Ile, and Phe (Fig. 4B and Table S2). This pattern of conservation is nicely explained by the ALIX_{Bro1}-CHMP4 structures, which show that the terminal Leu and Trp residues of the CHMP4 recognition helix bind in well defined pockets, whereas the first hydrophobic residue in the helix binds against a flat hydrophobic surface that can presumably tolerate greater side chain variability. Several sets of flanking hydrophilic residues are also conserved in the recognition helix, including an upstream acidic cluster (residues 208–212 in CHMP4A) and two basic residues (CHMP4A Lys-206 and 215). These side chains are solvent exposed and do not contact ALIX extensively, but a mutation within the CHMP4A acidic cluster did reduce ALIX_{Bro1} binding slightly, indicating that the cluster may interact favorably with the basic ALIX_{Bro1} surface. It is alternatively possible, however, that flanking hydrophilic residues are conserved primarily because they make important contacts when CHMP4 proteins adopt alternative conformations or bind other partners.

CHMP4 binding appears to be a conserved function of Bro1 domains, because three other Bro1 domain-containing proteins, Rim20p, HD-PTP, and Brox, also bind CHMP4 proteins (41, 42). Alignment of the Bro1 domains from metazoan ALIX proteins, human Brox and HD-PTP, and yeast Rim20p and Bro1p reveal strong, although not absolute conservation of CHMP4 contact residues, suggesting that most Bro1 domains will bind CHMP4 proteins in a similar fashion (Fig. S4). For example, the ALIX_{Bro1} Asp-143 and Lys-202 residues, which form a salt bridge that buttresses the ALIX_{Bro1} Asp-143-CHMP4A Trp-220 hydrogen bond, are absolutely conserved as an acidic/basic residue pair in all of the aligned Bro1 domains.

ALIX can be recruited to facilitate enveloped virus budding and release, and this function has been studied most extensively for HIV-1 (10, 11). These studies have identified three different ALIX point mutations (F199D, I212D, and L216D) that block both CHMP4 binding and ALIX-mediated release of HIV-1 constructs that cannot recruit TSG101/ESCRT-I. All three of these residues map to the CHMP4 binding interface, and the crystal structures are consistent with the observed loss of CHMP4 binding for Asp mutations at these positions (Fig. 4C). The ALIX I212D mutation has also been shown to block the ALIX-dependent abscission step of cytokinesis (18). Thus, the crystallographic ALIX-CHMP4 interface visualized here is essential for ALIX-dependent steps in HIV-1 budding and cytokinesis. This interaction recruits CHMP4 proteins, which in turn presumably recruit additional ESCRT-III subunits and VPS4 complexes to function in membrane remodeling and fission.

The recruiting order appears to be reversed in the case of MVB vesicle formation, where copolymerization of the different ESCRT-III subunits on endosomal membranes creates a surface that recruits other ESCRT pathway proteins, including ALIX, VPS4, Doa4p, IST1, and Vta1p/LIP5. Like the CHMP4 subunits, the CHMP1–3 subunits of ESCRT-III also have C-terminal amphipathic helices [MIT interacting motifs (MIM)], but in these cases the helices bind the MIT domains of VPS4, Vta1p/LIP5, and AMSH and thereby recruit ATPase and deubiquitylating activities to the membrane (43–48). Hence, the terminal helices of different ESCRT-III subunits must display distinct binding surfaces to ensure specificity in protein recruitment. This is borne out by the observation that the C-terminal CHMP4A helix binds ALIX_{Bro1} but not the VPS4A MIT domain, whereas the C-terminal CHMP1B helix binds the VPS4A MIT domain but not ALIX_{Bro1} (Fig. S5). As shown in Fig. 4D, the MIM helices of the CHMP1–3 proteins display three key leucine/hydrophobic residues that make important MIT domain contacts (47, 48). In these cases, the three hydrophobic residues are spaced by three and two intervening residues, whereas the

three hydrophobic residues of the CHMP4 recognition helix are each separated by two intervening residues. As a result, the initial hydrophobic residues of the CHMP1–3 and CHMP4 recognition helices occupy different relative positions. Furthermore, the terminal hydrophobic CHMP4 residue is a Trp, whereas the terminal CHMP1–3 hydrophobic residue is a Leu. Thus, the different identities and positions of the hydrophobic residues in their terminal recognition helices help ensure that each ESCRT-III subunit recruits its proper binding partner(s).

Methods

Expression Constructs and Plasmids. Expression constructs for GST-CHMP4A (WISP06–197) and GST-CHMP4A L217A (WISP06–61) are described in ref. 15. Quikchange mutagenesis (Stratagene) was used to create expression constructs for GST-CHMP4A_{1–204} (WISP06–198), GST-CHMP4A_{205–222} (WISP06–60), GST-CHMP4B_{205–224} (WISP06–201), GST-CHMP4C_{216–233} (WISP06–202), GST-CHMP4A_{205–222W220A} (WISP06–203), GST-CHMP4A_{205–222L217A} (WISP06–204), GST-CHMP4A_{205–222L214A} (WISP06–205), and GST-CHMP4A_{205–222E209A} (WISP06–206), using expression constructs for GST-CHMP4A, GST-CHMP4B (WISP06–199), and GST-CHMP4C (WISP06–200) as templates.

ALIX and CHMP4 Protein Expression and Purification. CHMP4 peptides used for crystallization studies were synthesized with an N-terminal acetyl capping group: ₂₀₅PKVDEEALKQLAEWVS₂₂₂ (CHMP4A), ₂₀₅KKKEEEDDDMKELLENWAGSM₂₂₄ (CHMP4B), and ₂₁₆QRAEEEDDDIKQLAAWAT₂₃₃ (CHMP4C). ALIX_{Bro1} (residues 1–359) and ALIX_{Bro1-V} (residues 1–702) proteins used in biosensor and crystallization studies were expressed and purified as described in ref. 10. GST-CHMP4 proteins used in biosensor experiments were expressed in BL21(DE3) Codon⁺ (RIPL) *Escherichia coli* cells grown in autoinduction media, ZYP-50502 (49). Cells were grown at 37°C for 4 h, then transferred to 23°C or 17°C for growth to saturation. Subsequent purification steps were performed at 4°. Cells from 100 ml of cultures were harvested, resuspended in 4 ml of 20 mM sodium phosphate (pH 7.2), 150 mM NaCl, 5 mM β-mercaptoethanol, and protease inhibitors (Roche Diagnostics) and lysed by the addition of 1 mg/ml lysozyme followed by sonication. The lysate was clarified by centrifugation, and GST-CHMP4 proteins were affinity purified by incubation with 500 μl of glutathione beads in biosensor running buffer [20 mM sodium phosphate (pH 7.2), 150 mM NaCl, 0.01% P20, 0.2 mg/ml BSA, 5 mM β-mercaptoethanol] for 30 min, washed extensively, and eluted with 20 mM reduced glutathione in running buffer.

Biosensor Binding Studies. Binding experiments used Biacore 2000 and T100 optical biosensor instruments. Research-grade CM5 sensor chips were derivatized with anti-GST antibody, using amine coupling, and were used to capture affinity purified GST-CHMP4 proteins or GST alone (reference) at surface densities of 300–3030 response units (RU). Similar binding data were obtained for both ALIX_{Bro1} and ALIX_{Bro1-V} constructs, but the ALIX_{Bro1-V} construct was more soluble and therefore allowed more complete sampling of the binding isotherms. Pure ALIX_{Bro1-V} (diluted in running buffer to the designated concentrations) was injected in duplicate (50 μl/min, 20°C) and binding data were collected at 2 Hz during the 30-s association and dissociation phases. All interactions reached equilibrium rapidly and dissociated within seconds during the dissociation phase, and all were studied at more than one surface density to rule out crowding and mass transport effects. Binding responses at 10–20 sec were fit to simple 1:1 binding isotherms to obtain equilibrium constants.

Crystallization. ALIX_{Bro1}-CHMP4 crystals were grown by sitting drop vapor diffusion at 4°C at protein:peptide molar ratios of 1:1.2 at final ALIX_{Bro1} concentrations of 25 mg/ml (CHMP4A) or 23 mg/ml (CHMP4B/C). Crystals grew from drops containing 0.5 μl of protein solution mixed with 0.5 μl of reservoir solution: 15% PEG 8000, 100 mM Na MES (pH 6.5), and 200 mM Na Acetate (CHMP4A) from drops containing 1.2 μl of protein solution mixed with 0.7 μl of reservoir solution [15% PEG 8000, 100 mM Na MES (pH 6.5), 200 mM Na Acetate (CHMP4B)] or from drops containing 1.2 μl of protein solution mixed with 0.7 μl of reservoir solution [10% PEG 2000, 100 mM Na MES (pH 6.5) (CHMP4C)].

Data Collection and Structure Refinement. Crystals were cryoprotected in reservoir solutions made up with 20% glycerol, suspended in nylon loops, and flash frozen in liquid nitrogen. Data were collected on a copper rotating anode generator with confocal optics (Rigaku; MicroMax 007HF) and a Rigaku R-Axis IV image plate (CHMP4B and CHMP4C) and at beam lines 11-1 (MAR325, 0.97607 Å wavelength) and 7-1 (Quantum 315, 0.98397 Å wavelength) of the Stanford Synchrotron Radiation Laboratory (CHMP4A). Crystals were maintained at 100 K during data collection. Data were integrated and scaled with

Denzo and Scafeck, using the HKL2000 suite (50). ALIX_{Bro1}-CHMP4 crystals were isomorphous with crystals of unliganded ALIX_{Bro1} (PDB entry 2OEW) (10). The unliganded ALIX_{Bro1} structure was used as a starting model for all three refinements, which were performed with REFMAC, using the maximum likelihood target function and incorporating TLS analysis (51, 52) as implemented in the CCP4 package (53). Models were built in O (54) and COOT (55), geometry was analyzed with PROCHECK (56), and figures were generated in PyMOL (57). The following residues were modeled in the different CHMP4 complexes: CHMP4A, 210–222; CHMP4B, 207–224; and CHMP4C, 221–233. Crystallographic statistics are presented in Table S1.

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